reduce Users Manual Release X1

Kennneth Anderson

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INTRODUCTION

This document is version 1.0 of the reduce Users Manual. This manual will describe the usage of reduce as an application provided by the Gemini Observatory Astrodata package suite. reduce is an application that allows users to invoke the Gemini Recipe System to perform data processing and reduction on one or more astronomical datasets.

This document presents details on applying reduce to astronomical datasets, currently defined as multi-extension FITS (MEF) files, both through the application's command line interface and the application programming interface (API). Details and information about the astrodata package, the Recipe System, and/or the data processing involved in data reduction are beyond the scope of this document and will only be engaged when directly pertinent to the operations of reduce.

1.1 Reference Documents

- The Gemini Recipe System: a dynamic workflow for automated data reduction, K. Labrie et al, SPIE, 2010.
- Developing for Gemini's extensible pipeline environment, K. Labrie, C. Allen, P. Hirst, ADASS, 2011
- Gemini's Recipe System; A publicly available instrument-agnostic pipeline infrastructure, K. Labrie et al, ADASS 2013.

1.2 Overview

As an application, reduce provides interfaces to configure and launch the Gemini Recipe System, a framework for developing and running configurable data processing pipelines and which can accommodate processing pipelines for arbitrary dataset types. In conjunction with the development of astrodata, Gemini Observatory has also developed the compatible astrodata_Gemini package, the code base currently providing abstraction of, and processing for, Gemini Observatory astronomical observations.

In Gemini Observatory's operational environment "on summit," reduce, astrodata, and the astrodata_Gemini packages provide a currently defined, near-realtime, quality assurance pipeline, the so-called QAP. reduce is used to launch this pipeline on newly acquired data and provide image quality metrics to observers, who then assess the metrics and apply observational decisions on telescope operations.

Users unfamiliar with terms and concepts heretofore presented should consult documentation cited in the previous sections (working on the Recipe System User Manual).

1.3 Glossary

adcc – Automatated Data Communication Center. Provides XML-RPC and HTTP services for pipeline operations. Can be run externally to reduce. Users need not know about or invoke the adcc for reduce operations. reduce will launch an adcc instance if one is not available. See Sec. *The adcc* for further discussion on adcc.

astrodata (or Astrodata) – part of the **gemini_python** package suite that defines the dataset abstraction layer for the Recipe System.

AstroData – not to be confused with **astrodata**, this is the main class of the astrodata package, and the one most users and developers will interact with at a programmatic level.

AstroDataType – Represents a data classification. A dataset will be classified by a number of types that describe both the data and its processing state. The AstroDataTypes are hierarchical, from generic to specific. For example, a typical GMOS image might have a set of types like

'GMOS_S', 'GMOS_IMAGE', 'GEMINI', 'SIDEREAL', 'IMAGE', 'GMOS', 'GEMINI_SOUTH', 'GMOS_RAW', 'UNPREPARED', 'RAW' (see **types** below).

astrodata_Gemini – the **gemini_python** package that provides all observatory specific definitions of data types, **recipes**, and associated **primitives** for Gemini Observatory data.

astrodata_X – conceivably a data reduction package that could reduce other observatory and telescope data. Under the Astrodata system, it is entirely possible for the Recipe System to process HST or Keck data, given the development of an associated package, astrodata_HST or astrodata_Keck. Pipelines and processing functions are defined for the particulars of each telescope and its various instruments.

Descriptor – Represents a high-level metadata name. Descriptors allow access to essential information about the data through a uniform, instrument-agnostic interface to the FITS headers.

gemini_python – A suite of packages comprising **astrodata**, **astrodata_Gemini**, **astrodata_FITS**, and **gempy**, all of which provide the full functionality needed to run **Recipe System** pipelines on observational datasets.

gempy – a **gemini_python** package comprising functional utilities to the **astrodata_Gemini** package.

MEF – Multiple Extension FITS, the standard data format not only for Gemini Observatory but many observatories.

primitive – A function defined within an **astrodata_[X]** package that performs actual work on the passed dataset. Primitives observe tightly controlled interfaces in support of re-use of primitives and recipes for different types of data, when possible. For example, all primitives called flatCorrect must apply the flat field correction appropriate for the data's current AstroDataType, and must have the same set of input parameters.

recipe – Represents the sequence of transformations. A recipe is a simple text file that enumerates the set and order of **primitives** that will process the passed dataset. A **recipe** is the high-level pipeline definition. Users can pass recipe names directly to reduce. Essentially, a recipe is a pipeline.

Recipe System – The gemin_python framework that accommodates an arbitrary number of defined recipes and the primitives

reduce – The user/caller interface to the Recipe System and its associated recipes/pipelines.

type or **typeset** – Not to be confused with language primitive or programmatic data types, these are data types defined within an **astrodata_[X]** package used to describe the kind of observational data that has been passed to the Recipe System., Eg., GMOS_IMAGE, NIRI. In this document, these terms are synonymous with **AstroDataType** unless otherwise indicated.

CHAPTER

TWO

USER ENVIRONMENT

Once a user has has retrieved the gemini_python package, available as a tarfile from the Gemini website (http://gemini.edu), and untarred only minor adjustments need to be made to the user environment in order to make astrodata importable and allow reduce to work properly.

2.1 Installation

Download the gemini_python X1 distribution (tar archive), place the tarfile as desired, and extract the archive:

```
$ tar -xvf gemini_python_X1.tar.gz
```

Next, invoke the usual Distutils command for a standard python module installation:

```
python setup.py install --prefix=/somewhere/
```

This will place executables in /somewhere/bin and the package modules in /somewhere/lib/python2.7/site-packages/.

Users will then need to have /somewhere/bin in \$PATH and /somewhere/lib/python2.7/site-packages in either \$PYTHONPATH or add the site-packages to sys.path.

reduce is made available on the command line once the installation is complete.

2.2 Test the installation

Start up the python interpreter and import astrodata:

```
$ python
>>> import astrodata
```

Next, return to the command line and test that reduce is reachable and runs. There may some delay as package modules are byte compiled:

```
$ reduce -h [--help]
```

This will print the reduce help to the screen.

If users have Gemini fits files available, they can test that the Recipe System is functioning as expected with a test recipe provided by the astrodata_Gemini package:

```
$ reduce --recipe test_one /path/to/gemini_data.fits
```

If all is well, users will see something like:

```
Resetting logger for application: reduce
Logging configured for application: reduce
                    --- reduce, v4890 ---
            Running under astrodata Version GP-X1
All submitted files appear valid
Starting Reduction on set #1 of 1
 Processing dataset(s):
       gemini_data.fits
______
RECIPE: test_one
______
PRIMITIVE: showParameters
 ______
rtf = False
suffix = '_scafaasled'
otherTest = False
logindent = 3
logfile = 'reduce.log'
reducecache = '.reducecache'
 storedcals = 'calibrations/storedcals'
 index = 1
retrievedcals = 'calibrations/retrievedcals'
cachedict = {'storedcals': 'calibrations/storedcals', 'retrievedcals':
            'calibrations/retrievedcals', 'calibrations': 'calibrations',
            'reducecache': '.reducecache'}
 loglevel = 'stdinfo'
 calurl_dict = {'CALMGR': 'http://fits/calmgr',
              'UPLOADPROCCAL': 'http://fits/upload_processed_cal',
              'QAMETRICURL': 'http://fits/qareport',
              'QAQUERYURL': 'http://fits/qaforgui',
              'LOCALCALMGR': 'http://localhost:%(httpport)d/calmgr/%(caltype)s'}
 logmode = 'standard'
 test = True
writeInt = False
calibrations = 'calibrations'
Wrote gemini_data.fits in output directory
Shutting down proxy servers ...
ADCC is running externally. No proxies to close
reduce exited
               on status: 0
```

Exit status 0 indicates nominal operations.

INTERFACES

3.1 Introduction

The reduce application provides a command line interface and an API, both of which can configure and launch a Recipe System processing pipeline (a 'recipe') on the input dataset. Control of reduce and the Recipe System is provided by a variety of options and switches. Of course, all options and switches can be accessed and controlled through the API.

3.2 Command line interface

We begin with the command line help provided by reduce --help, followed by further description and discussion of certain non-trivial options that require detailed explanation.

```
usage: reduce [options] fitsfile [fitsfile ...]
positional arguments:
fitsfile [fitsfile ...]
```

The [options] are described in the following sections.

3.2.1 Informational switches

- -h, -help show the help message and exit
- -v, -version show program's version number and exit
- -d, -displayflags Display all parsed option flags and exit.

When specified, this switch will present the user with a table of all parsed arguments and then exit without running. This allows the user to check that the configuration is as intended. The table provides a convenient view of all passed and default values. Unless a user has specified a recipe (-r, -recipe), 'recipename' indicates 'None' because at this point, the Recipe System has not yet been engaged and a default recipe not yet determined.

Eg.,:

```
['--invoked'] :: invoked :: False
['--addprimset'] :: primsetname :: None
['-d', '--displayflags'] :: displayflags :: True
['-p', '--param'] :: userparam :: None
['--logmode'] :: logmode :: ['console']
['-r', '--recipe'] :: recipename :: None
['--throw_descriptor_exceptions'] :: throwDescriptorExceptions :: False
['--logfile'] :: logfile :: reduce.log
['-t', '--astrotype'] :: astrotype :: None
['--override_cal'] :: user_cals :: None
['--context'] :: running_contexts :: None
['--calmgr'] :: cal_mgr :: None
['--suffix'] :: suffix :: None
['--loglevel'] :: loglevel :: stdinfo
Input fits file(s): fitsfile.fits
```

3.2.2 Configuration Switches, Options

- -addprimset <PRIMSETNAME> Add this path to user-supplied primitives for reduction. eg., path to a primitives module.
- **-calmgr <CAL_MGR>** This is a URL specifying a calibration manager service. A calibration manager overides Recipe System table. Not available outside Gemini operations.
- **-context <RUNNING_CONTEXTS>** Use <RUNNING_CONTEXTS> for primitives sensitive to context. Eg., context QA When not specified, the context defaults to 'QA'.
- **-invoked** Boolean indicating that reduce was invoked by the control center.
- **-logmode <LOGMODE>** Set logging mode. One of 'standard', 'console', 'quiet', 'debug', or 'null', where 'console' writes only to screen and 'quiet' writes only to the log file. Default is 'standard'.
- -logfile <LOGFILE> Set the log file name. Default is 'reduce.log' in the current directory.
- **-loglevel <LOGLEVEL>** Set the verbose level for console logging. One of 'critical', 'error', 'warning', 'status', 'stdinfo', 'fullinfo', 'debug'. Default is 'stdinfo'.
- **-override_cal <USER_CALS [USER_CALS ...]>** The option allows users to provide their own calibrations to reduce. Add a calibration to User Calibration Service. '-override_cal CALTYPE:CAL_PATH' Eg.,:
 - -override_cal processed_arc:wcal/gsTest_arc.fits
- -p <USERPARAM [USERPARAM ...]>, -param <USERPARAM [USERPARAM ...]> Set a primitive parameter from the command line. The form '-p par=val' sets the parameter in the reduction context such that all primitives will 'see' it. The form
 - -p ASTROTYPE:primitivename:par=val

sets the parameter such that it applies only when the current reduction type (type of current reference image) is 'ASTROTYPE' and the primitive is 'primitivename'. Separate parameter-value pairs by whitespace: (eg. '-p par1=val1 par2=val2')

See Sec. Overriding Primitive Parameters, for more information on these values.

-r <RECIPENAME>, -recipe <RECIPENAME> Specify an explicit recipe to be used rather than internally determined by a dataset's <ASTROTYPE>. Default is None and later determined by the Recipe System based on the AstroDataType.

- -t <ASTROTYPE>, -astrotype <ASTROTYPE> Run a recipe based on this AstroDataType, which overrides default type or begins without initial input. Eg., recipes that begin with primitives that acquire data. reduce default is None and determined internally.
- -suffix <SUFFIX> Add 'suffix' to output filenames at end of reduction.
- -throw_descriptor_exceptions Boolean indicating descriptor exceptions are to be raised. This is a development switch.

3.2.3 Nominal Usage

The minimal call for reduce can be

```
$ reduce <dataset.fits>
```

While this minimal call is available at the Gemini Observatory, if a calibration service is unavailable to the user – likely true for most users – users should call reduce on a specified dataset by providing calibration files with the –overrride_cal option. For example:

```
$ reduce --override_cal processed_arc:wcal/gsTest_arc.fits <dataset.fits>
```

Such a command for complex processing of data is possible because AstroData and the Recipe System do all the necessary work in determining how the data are to be processed, which is critically based upon the determination of the *typeset* that applies to that data.

Without any user-specified recipe (-r -recipe), the default recipe is qaReduce, which is defined for various Astro-DataTypes and currently used at the summit. For example, the qaReduce recipe for a GMOS_IMAGE specifies that the following primitives are called on the data:

```
prepare
addDQ
addVAR
detectSources
measureIQ
measureBG
measureCCAndAstrometry
overscanCorrect
biasCorrect
ADUToElectrons
addVAR
flatCorrect
mosaicDetectors
makeFringe
fringeCorrect
detectSources
measureIQ
measureBG
measureCCAndAstrometry
addToList
```

The point here is not to overwhelm readers with a stack of primitive names, but to present both the default pipeline processing that the above simple reduce command invokes and to demonstrate how much the reduce interface abstracts away the complexity of the processing that is engaged with the simplist of commands.

3.2.4 Overriding Primitive Parameters

In some cases, users may wish to change the functional behaviour of certain processing steps, i.e. change default behaviour of primitive functions.

Each primitive has a set of pre-defined parameters, which are used to control functional behaviour of the primitive. Each defined parameter has a "user override" token, which indicates that a particular parameter may be overridden by the user. Users can adjust parameter values from the reduce command line with the option,

-p, -param

If permitted by the "user override" token, parameters and values specified through the **-p, -param** option will *override* the defined parameter default value and may alter default behaviour of the primitive accessing this parameter. A user may pass several parameter-value pairs with this option.

Eg.:

```
$ reduce -p par1=val1 par2=val2 [par3=val3 ... ] <fitsfile1.fits>
```

For example, some photometry primitives perform source detection on an image. The 'detection threshold' has a defined default, but a user may alter this parameter default to change the source detection behaviour:

```
$ reduce -p threshold=4.5 <fitsfile.fits>
```

3.2.5 The @file facility

The reduce command line interface supports what might be called an 'at-file' facility (users and readers familiar with IRAF will recognize this facility). This facility allows users to provide any and all command line options and flags to reduce via in a single acsii text file.

By passing an @file to reduce on the command line, users can encapsulate all the options and positional arguments they might wish to specify in a single @file. It is possible to use multiple @files and even to embed one or more @files in another. The parser opens all files sequentially and parses all arguments in the same manner as if they were specified on the command line. Essentially, an @file is some or all of the command line and parsed identically.

To illustrate the convenience provided by an '@file', let us begin with an example *reduce* command line that has a number of arguments:

```
$ reduce -p GMOS_IMAGE:contextReport:tpar=100 GMOS_IMAGE:contextReport:report_inputs=True
-r recipe.ArgsTest --context qa S20130616S0019.fits N20100311S0090.fits
```

Ungainly, to be sure. Here, two (2) user parameters are being specified with **-p**, a recipe with **-r**, and a context argument is specified to be **qa**. This can be wrapped in a plain text @file called reduce args.par:

```
S20130616S0019.fits
N20100311S0090.fits
--param
GMOS_IMAGE:contextReport:tpar=100
GMOS_IMAGE:contextReport:report_inputs=True
-r recipe.ArgsTests
--context qa
```

This then turns the previous reduce command line into something a little more keyboard friendly:

```
$ reduce @reduce_args.par
```

The order of these arguments is irrelevant. The parser will figure out what is what. The above file could be thus written like:

```
-r recipe.ArgsTests
--param
GMOS_IMAGE:contextReport:tpar=100
GMOS_IMAGE:contextReport:report_inputs=True
--context qa
```

```
S20130616S0019.fits
N20100311S0090.fits
```

Note: Comments are accommodated, both line and in-line. '=' signs *may* be used but this has meaning only for arguments that expect unitary values. The '=' is entirely unnecessary.

White space is the only significant separator of arguments: spaces, tabs, newlines are all equivalent when argument parsing. This means the user can 'arrange' their @file for clarity.

Eg., a more readable version of the above file might be written as:

```
# reduce parameter file
# yyyy-mm-dd
# GDPSG
# Spec the recipe
-r
   recipe.ArgsTests # test recipe
# primitive parameters here
# These are 'untyped', i.e. global
--param
   tpar=100
   report_inputs=True
--context
   qa
                      # QA context
S20130616S0019.fits
N20100311S0090.fits
```

All the above examples of reduce_args.par are equivalently parsed. Which, of course, users may check by adding the -d flag:

```
$ reduce -d @redpars.par
----- switches, vars, vals -----
                                   var 'dest'
                                                                 Value
['--invoked']
['--addprimset']
                                                                 :: False
                                   :: invoked
['-addprimset'] :: primsetname :: None
['-d', '--displayflags'] :: displayflags :: True
['-p', '--param'] :: userparam :: ['tpar=100', 'report | '-logmode'] :: logmode :: standard
['-r', '--recipe'] :: recipename :: ['recipe.ArgTests']
                                                                  :: ['tpar=100', 'report_inputs=True']
['--throw_descriptor_exceptions'] :: throwDescriptorExceptions :: False
['--logfile'] :: logfile :: reduce.log
['-t', '--astrotype'] :: astrotype :: None
['--override_cal'] :: user_cals :: None
['--context'] :: running_contexts :: ['QA']
['--calmgr']
                                   :: cal_mgr
                                                                  :: None
['--suffix']
                                   :: suffix
                                                                  :: None
['--loglevel'] :: loglevel
                                                                 :: stdinfo
Input fits file(s): S20130616S0019.fits
Input fits file(s): N20100311S0090.fits
```

3.2.6 Recursive @file processing

As implemented, the @file facility will recursively handle, and process correctly, other @file specifications that appear in a passed @file or on the command line. For example, we may have another file containing a list of fits files, separating the command line flags from the positional arguments.

We have a plain text 'fitsfiles' containing the line:

```
test_data/S20130616S0019.fits
```

We can indicate that this file is to be consumed with the prefix character "@" as well. In this case, the 'reduce_args.par' file could thus appear:

```
# reduce test parameter file

@fitsfiles  # file with fits files

# AstroDataType
-t GMOS_IMAGE

# primitive parameters.
--param
    report_inputs=True
    tpar=99
    FOO=BAR

# Spec the recipe
-r recipe.ArgTests
```

The parser will open and read the @fitsfiles, consuming those lines in the same way as any other command line arguments. Indeed, such a file need not only contain fits files (positional arguments), but other arguments as well. This is recursive. That is, the @fitsfiles can contain other at-files", which can contain other "at-files", which can contain ..., ad infinitum. These will be processed serially.

As stipulated earlier, because the @file facility provides arguments equivalent to those that appear on the command line, employment of this facility means that a reduce command line could assume the form:

```
$ reduce @parfile @fitsfiles
or equally:
$ reduce @fitsfiles @parfile
```

where 'parfile' could contain the flags and user parameters, and 'fitsfiles' could contain a list of datasets.

Eg., fitsfiles comprises the one line:

```
test_data/N20100311S0090.fits
```

while parfile holds all other specifications:

```
# reduce test parameter file
# GDPSG

# AstroDataType
-t GMOS_IMAGE

# primitive parameters.
--param
    report_inputs=True
    tpar=99  # This is a test parameter
```

```
# Spec the recipe
-r recipe.ArgTests
```

3.2.7 Overriding @file values

The reduce application employs a customized command line parser such that the command line option

-p or -param

will accumulate a set of parameters *or* override a particular parameter. This may be seen when a parameter is specified in a user @file and then specified on the command line. For unitary value arguments, the command line value will *override* the @file value.

It is further specified that if one or more datasets (i.e. positional arguments) are passed on the command line, all fits files appearing as positional arguments in the parameter file will be replaced by the command line arguments.

Using the parfile above,

Eg. 1) Accumulate a new parameter:

Eg. 2) Override a parameter in the @file:

Eg. 3) Override the recipe:

Eg. 4) Override a recipe and specify another fits file

```
$ reduce @parfile -r=recipe.FOO test_data/N20100311S0090_1.fits
parsed options:
```

```
AstroDataType: GMOS_IMAGE

FITS files: ['test_data/N20100311S0090_1.fits']

Parameters: tpar=100, report_inputs=True

RECIPE: recipe.FOO
```

3.3 Application Programming Interface (API)

Note: The following sections discuss and describe programming interfaces available on reduce and the underlying class Reduce.

The reduce application is essentially a skeleton script providing the described command line interface. After parsing the command line, the script then passes the parsed arguments to its main() function, which in turn calls the Reduce() class constructor with "args". Class Reduce() is defined in the module coreReduce.py. reduce and class Reduce are both scriptable, as the following discussion will illustrate.

3.3.1 reduce.main()

The main() function of reduce receives one (1) parameter that is a Namespace object as returned by a call on ArgumentParser.parse_args(). Specific to reduce, the caller can supply this object by a call on the parseUtils.buildParser() function, which returns a fully defined reduce parser. As usual, the parser object should then be called with the parse_args() method to return a valid reduce parser Namespace. Since there is no interaction with sys.argv, as in a command line call, all Namespace attributes have only their defined default values. It is for the caller to set these values as needed.

As the example below demonstrates, once the "args" Namespace object is instantiated, a caller can set any arguments as needed. Bu they must be set to the correct type. The caller should examine the various "args" types to determine how to set values. For example, args.files is type list, whereas args.recipename is type string.

Eg.,

```
>>> from astrodata.adutils.reduceutils import reduce
>>> from astrodata.adutils.reduceutils import parseUtils
>>> args = parseUtils.buildParser("Reduce, v2.0").parse_args()
>>> args.files
[]
>>> args.files.append('S20130616S0019.fits')
>>> args.recipename = "recipe.FOO"
>>> reduce.main(args)
--- reduce, v2.0 ---
Starting Reduction on set #1 of 1
Processing dataset(s):
S20130616S0019.fits
```

Processing will proceed as usual.

3.3.2 Class Reduce and the runr() method

Class Reduce is defined in astrodata.adutils.reduceutils module, coreReduce.py.

The reduce.main() function serves mainly as a callable for the command line interface. While main() is callable by users supplying the correct "args" parameter (See *reduce.main()*), the Reduce() class is also callable and can be used

directly, and more appropriately. Callers need not supply an "args" parameter to the class constructor. The instance of Reduce will have all the same arguments as in a command line scenario, available as attributes on the instance. Once an instance of Reduce() is instantiated and instance attributes set as needed, there is one (1) method to call, **runr**(). This is the only public method on the class.

Note: When using Reduce() directly, callers must configure their own logger. Reduce() does not configure logutils prior to using a logger as returned by logutils.get_logger(). The following example will illustrate how this is easily done. It is *highly recommended* that callers configure the logger.

Eg.,

```
>>> from astrodata.adutils.reduceutils.coreReduce import Reduce
>>> reduce = Reduce()
>>> reduce.files
[]
>>> reduce.files.append('S20130616S0019.fits')
>>> reduce.files
['S20130616S0019.fits']
```

Once an instance of Reduce has been made, callers can then configure logutils with the appropriate settings supplied on the instance. This is precisely what reduce does when it configures logutils.

At this point, the caller is able to call the runr() method on the "reduce" instance.

```
>>> reduce.runr()
All submitted files appear valid
Starting Reduction on set #1 of 1
Processing dataset(s):
S20130616S0019.fits
```

Processing will then proceed in the usual manner.

SUPPLEMENTAL TOOLS

The astrodata package provides a number of command line driven tools, which users may find helpful in executing reduce on their data.

With the installation and configuration of astrodata and reduce comes some supplemental tools to help users discover information, not only about their own data, but about the Recipe System, such as available recipes, primitives, and defined AstroDataTypes.

If the user environment has been configured correctly these applications will work directly.

4.1 listprimitives

The application listprimitives is available as a command line executable. This tool displays available primitives for all AstroDataTypes, their parameters, and defaults. These are the parameters discussed in Sec. *Overriding Primitive Parameters* that can be changed by the user with the **-p**, **-param** option on reduce. under the AstroDataTypes. The help describes more options:

```
$ listprimitives -h
Usage: listprimitives [options]
Gemini Observatory Primitive Inspection Tool, v1.0 2011
Options:
-h, --help
                   show this help message and exit
                   apply color output scheme
-c, --use-color
-i, --info
                   show more information
                   show parameters
-p, --parameters
-r, --recipes
                   list top recipes
-s, --primitive-set show primitive sets (Astrodata types)
-v, --verbose
                   set verbose mode
--view-recipe=VIEW_RECIPE
                     display the recipe
```

4.1.1 listprimitives information

The following section presents examples of the kind of information that listprimitives may provide. Show available recipes:

\$ listprimitives -r

RECIPES_Gemini

- 1. basicOA
- 2. checkOA
- makeProcessedArc.GMOS_SPECT
- 4. makeProcessedBias
- 5. makeProcessedDark
- 6. makeProcessedFlat
- 7. makeProcessedFlat.GMOS_IMAGE
- 8. makeProcessedFlat.GMOS_SPECT
- 9. makeProcessedFlat.NIRI_IMAGE
- 10. makeProcessedFringe
- 11. qaReduce.GMOS_IMAGE
- 12. gaReduce.GMOS_SPECT
- 13. qaReduce.NIRI_IMAGE
- 14. qaReduceAndStack.GMOS_IMAGE
- 15. qaStack.GMOS_IMAGE
- 16. reduce.F2_IMAGE
- 17. reduce.GMOS_IMAGE

Subrecipes

- 1. biasCorrect
- correctWCSToReferenceCatalog
- 3. darkCorrect
- 4. flatCorrect
- 5. lampOnLampOff
- 6. makeSky
- 7. overscanCorrect
- 8. prepare
- 9. skyCorrect
- 10. standardizeHeaders
- 11. thermalEmissionCorrect

Users can request display the contents of a particular recipe, as listed above. This will present the 'stack' of primitives that will be called by the Recipe System when the particular recipe is either specified through the reduce command line by the user, or selected internally by the Recipe System itself.

For example, a user may like to see the primitive stack called by the default 'QA' recipe for GMOS_IMAGE data. As seen in the above example, these 'qa' recipes are defined for several AstroDataTypes.

Show the primitive stack for the 'qa' GMOS_IMAGE type:

\$ listprimitives --view-recipe qaReduce.GMOS_IMAGE

RECIPE: qaReduce.GMOS_IMAGE

 \sharp This recipe performs the standardization and corrections needed to convert

the raw input science images into a single stacked science image

prepare addDQ

```
addVAR(read_noise=True)
detectSources
measureIQ(display=True)
measureBG
measureCCAndAstrometry
overscanCorrect
biasCorrect
ADUToElectrons
addVAR(poisson_noise=True)
flatCorrect
mosaicDetectors
makeFringe
fringeCorrect
detectSources
measureIQ(display=True)
measureBG
measureCCAndAstrometry
addToList(purpose=forStack)
```

parameters in a more focused way, i.e., report only those primitives and parameters relevant to a given dataset. As it currently stands, users can request that <code>listprimitives</code> display primitive parameters (as may be passed to reduce through the **-p** or **-param** option, Sec. *Overriding Primitive Parameters*), but this results in a list of all AstroDataTypes, their primitives and associated parameters. Admittedly, this list is rather ungainly, but users may see, for example, that the primitive <code>detectSources</code> has several user-tunable parameters:

```
detectSources
    suffix: '_sourcesDetected'
    centroid_function: 'moffat'
    threshold: 3.0
    sigma: None
    fwhm: None
    method: 'sextractor'
    max_sources: 50
```

See the discussion in Sec. *Overriding Primitive Parameters* on command line override of primitive parameters, and where overriding the 'threshold' parameter is discussed specifically.

4.2 typewalk

As with listprimitives and reduce, the command line application typewalk becomes available once the user has configured astrodata correctly. typewalk examines files in a directory or directory tree and reports the types and status values through the AstroDataType classification scheme. Running typewalk on a directory containing some Gemini datasets will demonstrate what users can expect to see. If a user has downloaded gemini_python X1 package with the 'test_data', the user can move to this directory and run typewalk on that extensive set of Gemini datasets.

By default, typewalk will recurse all subdirectories under the current directory. Users may specify an explicit directory with the **-d** or **-dir** option.

typewalk provides the following options [-h, -help]:

4.2. typewalk

```
-h, --help
                      show this help message and exit
-b BATCHNUM, --batch BATCHNUM
                     In shallow walk mode, number of files to process at a
                      time in the current directory. Controls behavior in
                      large data directories. Default = 100.
--calibrations
                     Show local calibrations (NOT IMPLEMENTED).
-c, --color
                     Colorize display
-d TWDIR, --dir TWDIR
                     Walk this directory and report types. default is cwd.
-f FILEMASK, --filemask FILEMASK
                     Show files matching regex <FILEMASK>. Default is all
                     .fits and .FITS files.
-i, --info
                     Show file meta information.
--keys KEY [KEY ...] Print keyword values for reported files.Eg., --keys
                     TELESCOP OBJECT
-n, --norecurse
                     Do not recurse subdirectories.
--or
                     Use OR logic on 'types' criteria. If not specified,
                     matching logic is AND (See --types). Eg., --or --types
                      GEMINI_SOUTH GMOS_IMAGE will report datasets that are
                      either GEMINI_SOUTH *OR* GMOS_IMAGE.
-o OUTFILE, --out OUTFILE
                     Write reported files to this file. Effective only with
                      --types option.
--raise
                     Raise descriptor exceptions.
--types TYPES [TYPES ...]
                     Find datasets that match only these type criteria.
                      Eq., --types GEMINI_SOUTH GMOS_IMAGE will report
                      datasets that are both GEMINI_SOUTH *and* GMOS_IMAGE.
--status
                      Report data processing status only.
--typology
                      Report data typologies only.
```

Files are selected and reported through a regular expression mask which, by default, finds all ".fits" and ".FITS" files. Users can change this mask with the **-f**, **-filemask** option.

As the **-types** option indicates, typewalk can find and report data that match specific type criteria. For example, a user might want to find all GMOS image flats under a certain directory. typewalk will locate and report all datasets that would match the AstroDataType, GMOS_IMAGE_FLAT.

A user may request that a file be written containing all datasets matching AstroDataType qualifiers passed by the **types** option. An output file is specified through the **-o, -out** option. Output files are formatted so they may be passed *directly to the reduce command line* via that applications 'at-file' (@file) facility. See *The @file facility* or the reduce help for more on 'at-files'.

Users may select type matching logic with the **–or** switch. By default, qualifying logic is AND, i.e. the logic specifies that *all* types must be present (x AND y); **–or** specifies that ANY types, enumerated with **–types**, may be present (x OR y). **–or** is only effective when the **–types** option is specified with more than one type.

For example, find all GMOS images from Cerro Pachon in the top level directory and write out the matching files, then run reduce on them (**-n** is 'norecurse'):

```
$ typewalk -n --types GEMINI_SOUTH GMOS_IMAGE --out gmos_images_south
$ reduce @gmos_images_south
```

Find all F2_SPECT and GMOS_SPECT datasets in a directory tree:

```
$ typewalk --or --types GMOS_SPECT F2_SPECT
```

This will also report match results to stdout, colourized if requested (-c).

CHAPTER

FIVE

DISCUSSION

5.1 Fits Storage

blah blah Fits Storage blah blah ...

5.2 The adcc

As a matter of operations, reduce and the Recipe System depend upon the services of what is called the adcc, the Automated Data Communication Center. The adcc provides services to pipeline operations through two proxy servers, an XML-RPC server and an HTTP server. The XML_RPC server serves calibration requests made on it, and retrieves calibrations that satisfy those requests from the Gemini FITS Store, a service that provides automated calibration lookup and retrieval.

The adcc can be run externally and will run continuously until it is shutdown. Any instances of reduce (and the Recipe System) will employ this external instance of the adcc to service a pipeline's calibration requests. However, a user of reduce need not start an instance of the adcc nor, indeed, know anything about the adcc per se. If one is not available, an instance of the adcc will be started by reduce itself, and will serve that particular reduce process and then terminate.

This note is provided should users notice an adcc process and wonder what it is.

5.3 Future Enhancements

5.3.1 Intelligence

One enhancement long imagined is what has been generally termed 'intelligence'. That is, an ability for either reduce or some utility to automatically do AstroDataType classification of a set of data, group them appropriately, and then pass these grouped data to the Recipe System.

As things stand now, it is up to the user to pass commonly typed data to reduce. As shown in the previous section, *typewalk*, typewalk can help a user perform this task and create a 'ready-to-run' @file that can be passed directly to reduce. Properly implemented 'intelligence' will *not* require the user to determine the AstroDataTypes of datasets.

5.3.2 Local Calibration Service

blahbadeeblah about a local calibration service ...

CHAPTER

SIX

6. ACKNOWLEDGMENTS

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APPENDIX

A

REDUCE DEMO