

OGIP Calibration Memo CAL/GEN/92-015

How to Manage a Calibration Database

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SUMMARY

This document describes how to include new files into an existing Calibration Database (CALDB). Documentation on how to install a local Caldb is covered in OGIP Calibration Memo “How to Install a Calibration Database” CAL/GEN/94-004.

Log of Significant Changes to this document

Release Date	Sections Changed	Brief Notes
1997 Jun 20		First Draft by Lorraine Breedon
2005 Nov 16	All	Completely re-written by MFC
2006 Feb 08	4.2.1, 4.2.2	added example showing the explicit dependence of <code>udcif</code> conflict checking on the order of the CBD keywords
2012 Nov 26	all	minor corrections

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1 Introduction

This document describes the steps needed to include new files in the calibration database (CALDB) for a given mission and instrument. In order to update the CALDB to include a new calibration file, these steps must be followed:

1. The file must be updated with the mandatory CALDB FITS keywords;
2. The file must be moved to the appropriate directory in the CALDB for this mission and instrument;
3. The caldb.indx file for this mission and instrument must be updated to include the new file (perhaps to replace existing files in the caldb.indx)
4. The update to the CALDB should be announced and documented in a publicly accessible way; at the HEASARC this is done via the CALDB RSS feed and the CALDB News Archive page, among other locations

Typically, missions will control the structure and content of their CALDBs and will be responsible for keeping their CALDBs current and correct. Usually missions will communicate with the HEASARC CALDB manager that a CALDB update has been performed; in such cases the HEASARC CALDB manager will incorporate and make the update publicly available from the HEASARC CALDB (see “Automated Delivery of Calibration Data to the CALDB”, Cal/gen/03-001, for a discussion of data delivery by a mission to the HEASARC CALDB).

2 Including CALDB Keywords in a Calibration File

Calibration files which are listed in the CALDB are identified by a set of “calibration parameters”. These parameter are:

1. the mission (given by the FITS TELESCOP header keyword);
2. the the instrument (given by the FITS INSTRUME header keyword);
3. (optionally) a detector on the instrument (given by the FITS DETNAM header keyword);
4. (optionally) a filter which may be inserted into the light path before the instrument or detector (given by the FITS FILTER header keyword);
5. A date (given by the FITS CVSD0001 header keyword) and time (given by the FITS CVST0001 header keyword) after which the calibration file is considered appropriate or “valid”;
6. the “type” of calibration file (given by the file’s “CALDB name”, or “CAL_CNAM”; see “Required and Recommended FITS keywords for Calibration Files”, Section 3A, Cal/gen/92-011);

7. (optionally) a set of “calibration parameters” (or “calibration boundaries”) which can specify the dependence of the calibration on non-standard parameters (temperature, SAA, earth elevation, or other variables), given by the FITS CBDN0001 header keywords.

The implementation of these parameters in the CALDB is summarized in Table 1.

Table 1: The CALDB Calibration File Parameter Set

Parameter Description	CALDB File HEADER Keyword	caldb.indx File Column Name	Required or Optional
The mission	TELESCOP	TELESCOP	Required
The instrument	INSTRUME	INSTRUME	Required
The detector	DETNAM	DETNAM	Optional
The filter	FILTER	FILTER	Optional
Validity Start Date	CVSD0001	REF_TIME ^a	Required
Validity Start Time	CVST0001	REF_TIME	Required
Calibration Date Type	CCNM0001	CAL_CNAM	Required
Calibration Boundaries	CBDn0001	CAL_CBD	Optional

^aREF_TIME is the MJD corresponding to CVSD0001 and CVST0001.

These parameters are specified in FITS calibration files by header keywords for each FITS extension. The full list of CALDB header keywords is given in cal/gen/92-011, “Required and Recommended FITS keywords for Calibration Files”. If there are questions about use of these keywords please contact the HEASARC Caldb Manager for assistance. For the rest of this document we’ll assume that all required keywords are present in the new CALDB file.

3 Placing the File in the CALDB Directory Structure

The CALDB software relies on a specific directory structure to enable users to find calibration data appropriate to their observing setup. The directory structure has the form:

```
$CALDB/data/<mission>/<instrument>/<class>
```

where \$CALDB is an environment variable which defines the top-level location of the CALDB, <mission> is the name of the mission, and <instrument> is the name of the instrument. For largely historical reasons, class is either `cpf`, `bcf`, or `pcf` which distinguishes “calibration product files” from “basic calibration files” from “primary calibration files”, respectively. This progression is supposed to represent distance from the raw calibration data: Calibration products are derived from basic calibration files, which in turn are derived from primary calibration data. In practice CALDB access software will

only be able to retrieve and use `cpf` or `bcf` files; `pcf` files, as raw calibration files, do not necessarily even need to be in FITS format, while `cpf` and `bcf` files do. For more information on the differences between classes see CAL/GEN/92-003.

For example, the file `pspcb_gain2_256.rmf`, which contains the spectral response file (usually considered a “calibration product”) for the ROSAT PSPC-B instrument is located in

```
$CALDB/data/rosat/pspcb/cpf/matrices/pspcb_gain2_256.rmf
```

in the HEASARC CALDB. Missions should define the appropriate directory structure for their CALDBs within these conventions.

4 Including the Calibration File in the `caldb.indx`

Access to calibration files in the CALDB is accomplished through the `caldb.indx` file (which is located at `$CALDB/<mission>/<instrument>`), which contains a summary of files in the CALDB, along with their calibration parameter sets. In order for a file to be retrieved by the standard CALDB access software¹, the file must be included in the `caldb.indx` file.

A file may be included in the `caldb.indx` file by using the `udcif` tool included with the `caltools` distribution. For a given calibration file, `udcif` reads information from the file (specifically the mandatory CALDB keywords from the file FITS extension headers) then, for each valid FITS extension, creates an entry in the `caldb.indx` table.

4.1 An Example

As an example, suppose the file `$CALDB/data/suzaku/hxd/cpf/ae_hxd_pinxinom_20051104.rsp` needs to be added to the `caldb.indx` file for the SUZAKU HXD instrument. This file contains 2 extensions, containing the response matrix for the HXD (CCNM0001= 'SPECRESP MATRIX') along with an extension containing the energy boundaries for the HXD (CCNM0001= 'EBOUNDS '), as described in “The Calibration Requirements for Spectral Analysis”². Both extensions are to be considered “good” (CAL_QUAL=0) for their specified calibration parameter set.

```
% cd $CALDB/data/suzaku/hxd/cpf
% udcif
Name of file containing dataset[-] ae_hxd_pinhxnom_20051104.rsp
```

¹a subroutine called `gtcalf`, included as part of the `callib` software library distributed in the standard FTTOOLS release, is usually used to retrieve files from the CALDB.

²cal/gen/92-002; see also the addendum, cal/gen/92-002a.

```

Name of Calibration Index File[../caldb.indx] ../caldb.indx
Dataset:      SPECRESP MATRIX
Quality value for the dataset being entered[0] 0
Dataset:      EBOUNDS
Quality value for the dataset being entered[0] 0

```

In general it's better to run `udcif` in the directory in which the calibration file is located, and to specify the path to the `caldb.indx` file; this has the benefit of including the full path to the file (relative to `$CALDB`) in the `caldb.indx` column `CAL_DIR`, and the filename (without any directory path) in the `CAL_FILE` column. An alternative is to run `udcif` in the directory where the `caldb.indx` file is located (namely `$CALDB/data/<mission>/<instrument>`); in this case the resulting `CAL_FILE` column will contain the directory path (relative to `$CALDB/data/<mission>/<instrument>`) in addition to the file name. In practice the CALDB access software will transparently handle either case.

4.2 Understanding and Handling Conflicts

The CALDB works best when there's a one-to-one correspondence between calibration files and calibration parameter sets (mission, instrument, detector, filter, validity date and time, and calibration boundary parameters), i.e. when one particular parameter set corresponds to one and only one file (really, an extension in a FITS calibration file). In many cases this simple situation holds, but other more complex cases may arise.

A *conflict* is a situation in which two or more calibration file extensions (hereafter called “*calibration units*”) overlap for a given set of calibration parameters. Table 2 lists the criteria for which `udcif` considers a file to be ingested in conflict with a file already listed in the `caldb.indx` file.

The `udcif` tool tries to avoid situations where 2 or more calibrations may conflict, since in such cases ambiguities may arise in determining which of the calibrations should be used for the observation in question. Avoiding such conflicts is especially important for the CALDB, since the assumption which underpins the CALDB is that the end user should not have to know the details of which instrument calibrations are needed to properly interpret an observation. In other words, situations in which the end user may need to choose between two or more calibration files should be avoided as much as possible.

4.2.1 Understanding Boundary Value Conflicts

After determining that the file to be added to the `caldb.indx` file matches criteria 1–6 in table 2, the `udcif` tool then looks at the boundary keywords to determine if a calibration boundary keyword conflict exists. Recall that calibration boundaries are specified by the syntax

PARAM(VALUE)UNIT

Table 2: Conflict Criteria Used by UDCIF

A Conflict exists if All these Conditions are True		
Criteria #	File Value	Value from CALDB.INDX file Entry
1	TELESCOP	matches TELESCOP
2	INSTRUME	matches INSTRUME
3	DETNAM	matches DETNAM
4	FILTER	matches FILTER
5	CCNM0001	matches CAL_CNAM
6	REF_TIME ^a	matches REF_TIME
7	And if a calibration boundary keyword conflict exists	

^aREF_TIME for the file is calculated from the CVSD0001 and CVST0001 header keywords

(see Section B.3 of CAL/GEN/92-011 for more information about this syntax). PARAM is a parameter name (for example, TEMPERATURE), VALUE is the value of the parameter (for example (100)), or an allowed parameter range (for example (100:110)), and UNIT³ is the unit of the VALUE (for example, 'DEGREES').

The `udcif` tool looks for boundary conflicts by examining the first PARAMETER entry in the CAL_CBD array for the first row in the `caldb.indx` file that satisfies criteria 1 – 6. The tool then compares this CAL_CBD parameter name to the names of the parameters in the CBDN0001 keywords in the file to be ingested. If the names match, then `udcif` checks the value of the parameter. If the parameter values are the same (or the range overlaps) then `udcif` considers that these two files are in conflict. If the parameter names are the same but the values are different (or the ranges do not overlap) then the files are considered to be not in conflict.

Table 3 shows some permutations of the calibration boundaries, and the resulting determination (CONFLICT or NO CONFLICT) of `udcif`, for an example where the file to be ingested into the `caldb.indx` file meets criteria 1 – 6 for some row in the `caldb.indx` file, and the file has 2 calibration boundary keywords. Table 3 shows the first two CAL_CBD keywords for the matching row in the `caldb.indx` file, and the rest of the nine CAL_CBD values are 'NONE'.

It's important to note the first and second cases in table 3. These cases cover exactly the same calibration boundary parameter space, but the conflict checking used by `udcif` yields different results (in the first case `udcif` doesn't consider the two files to be in conflict; in the second case it does). This is because, once `udcif` finds a calibration boundary keyword parameter name that matches, it stops checking the rest of the calibration boundary parameters. Thus it's important to realize that the conflict checking performed by `udcif` is **sensitive to the order** of the CBD keywords.

³Note that, currently, `udcif` does not check to confirm that the UNITS of matching PARAMS are the same!

Table 3: Permutations of Boundary Values

caldb.indx Entry		CBDs in File for Ingest		Conflict?	Explanation
CAL_CBD1	CAL_CBD2	CBD1	CBD2		
SAA(NO)	TEMP(100)	SAA(YES)	TEMP(100)	No conflict	CAL_CBD1's have same parameter but different values
TEMP(100)	SAA(NO)	TEMP(100)	SAA(YES)	conflict	CAL_CBD1's have same parameter and same values
SAA(YES)	DAY(YES)	SAA(YES)	DAY(NO)	Conflict	Even though the DAY values are different, the SAA match takes precedence
SAA(NO)	NONE	SAA(YES)	TEMP(100)	No conflict	CAL_CBD1 and CBD1 have same parameter name but different values
NONE	NONE	SAA(NO)	NONE	Conflict	If caldb.indx entry has first CAL_CBD value equal to NONE, then it's assumed all the values are NONE; udcif regards this as a conflict regardless of the CBD values in the file to be ingested
SAA(NO)	NONE	NONE	NONE	Conflict	If CBD1 is NONE in the file to be ingested, then it's assumed all the values are NONE; udcif regards this as a conflict regardless of the CAL_CBD values of the matching caldb.indx entry
SAA(YES)	DAY(YES)	TEMP(100)	DAY(YES)	Conflict	No match for SAA in the CBDs of the file to be ingested, so the DAY parameter is tested, and found to conflict
THETA(100)	TEMP(105)	DETCANS(256)	PROCV(1)	Conflict	If there are no explicit boundary matches between the two arrays, both arrays claim validity over the same parameter space and they are considered to be in conflict.

4.2.2 Avoiding or Overriding Conflicts

Conflicts can be avoided if files which satisfy criteria 1 – 6 have the same PARAMETER NAMES in their calibration boundary array (i.e. the values of the CBD keywords in the extension header) with well-defined boundary constraints, but this may not always be practical or even possible. Furthermore, it's necessary to put the boundary parameters **in the same order** in the file as in the matching entry in the `caldb.indx` file for accurate conflict checking.

If a conflict is identified, `udcif` presents the user with the choice of changing the “quality” of the conflicting dataset (i.e. the `CAL_QUAL` value of the file that is already tabulated in the `caldb.indx` file) to a value of 5 (“bad”). This would make the file inaccessible via normal CALDB access methods (since by default the CALDB access routine returns only “good” files, with `CAL_QUAL=0`). This would be appropriate if the file that's being ingested into the CALDB is meant to replace the existing file; the new file would then be considered the “good” version and the old file the “bad” version.

There are times when `udcif` might identify a potential conflict between two files which in practice is not a conflict, and the retrieval software is written such that by specifying the appropriate parameter a unique file name (and FITS extension number) can be retrieved. If it is certain that there is no conflict even though `udcif` claims that there is, the user has the option of overriding `udcif` and including the file in the `caldb.indx` file, without flagging the matching `udcif` entry as bad, as shown in the example below:

```
% udcif
Name of file containing dataset[-] ae_hxd_pinxinom_20051104.rsp
Name of Calibration Index File[../caldb.indx]
Dataset:      SPECRESP MATRIX
Quality value for the dataset being entered[0]
```

```
Another dataset has been found which is valid for the same
conditions as the dataset being indexed
```

```
The conflicting dataset:
```

```
-----
Instrument: HXD
Code Name: SPECRESP MATRIX
File: ae_hxd_pinhxnom_20051104.rsp[ 1]
Description:
RESPONSE MATRIX
-----
```

```
Change conflicting dataset's Quality value to 5 (Y/N)?[N] n
Include new data file in CALDB.INDX (Y/N)> Y
```

```
Dataset:      EBOUNDS
```

Quality value for the dataset being entered[0]

Another dataset has been found which is valid for the same conditions as the dataset being indexed

The conflicting dataset:

```
-----
Instrument: HXD
Code Name: EBOUNDS
File: ae_hxd_pinhxnom_20051104.rsp[ 2]
Description:
EBOUNDS
-----
```

```
Change conflicting dataset's Quality value to 5 (Y/N)?[no] N
Include new data file in CALDB.INDX (Y/N)> Y
```

and extensions 1 & 2 for both the new file (ae_hxd_pinxinom_20051104.rsp) and the old file (ae_hxd_pinhxnom_20051104.rsp) are retained in the caldb.indx file with CAL_QUAL=0.

Including such “conflicts” in the caldb.indx file might pose difficulties in the future for users who might want to unambiguously retrieve a specific file from the CALDB and so should be used with caution. It is often the case that judicious choice of CBD keyword values can help avoid such conflicts. The burden is on the instrument team to adequately describe the calibration files using appropriate calibration boundary parameters and values in each calibration file of a give calibration type.

5 Verifying the Update

After a file is ingested into the CALDB, the user should try to verify that the file has been included properly in the caldb.indx, file. Probably the easiest way to do this is to use a FITS table viewer like fv. After opening the caldb.indx in the FITS viewer, look to see if all the entries for the new file appear correct in the EXTNAME= 'CIF' binary extension table.

Users should also verify that the correct file is returned from the CALDB by using the FTOOL quzcif. For example, SUZAKU response matrices are stored in the first extension of the files ae_hxd_pinhxnom_20051104.rsp and ae_hxd_pinxinom_20051104.rsp. These matrices are used for simulations of the HXD instrument response. They differ because ae_hxd_pinhxnom_20051104.rsp is used when the SUZAKU pointing is “nominal” for the HXD instrument, while ae_hxd_pinxinom_20051104.rsp is used for simulations of the HXD response for nominal pointings of the XIS instrument. They can be distinguished by their CDB keywords: ae_hxd_pinhxnom_20051104.rsp has CBD30001= 'NOMINAL("HXD")' while ae_hxd_pinxinom_20051104.rsp has CBD30001= 'NOMINAL("XIS")'. Some sample calls to quzcif are shown below:

NO BOUNDARY CONDITIONS SPECIFIED:

```
=====
[corcoran:data/suzaku/hxd] corcoran% quzCIF
Name of Mission[-] suzaku
Name of Instrument[-] hxd
Name of Detector (- if not required)[-]
Name of Filter (- if not required)[-]
Calibration Dataset Codename[-] SPECRESP MATRIX
Requested Date in yyyy-mm-dd format[-] NOW
Requested Time in hh:mm:ss format[-] NOW
Boolean selection expression for Boundary params(- if not required)[-]
/caldb/test/staging/data/suzaku/hxd/cpf/ae_hxd_pinhxnom_20051104.rsp 1
/caldb/test/staging/data/suzaku/hxd/cpf/ae_hxd_pinxinom_20051104.rsp 1
```

Extensions in both files are identified by quzCIF since these particular extensions are distinguished by the values in their calibration boundary arrays, and no boundary condition was specified.

BOUNDARY = DETCHANS.EQ.256

```
=====
[corcoran:data/suzaku/hxd] corcoran% quzCIF
Name of Mission[suzaku]
Name of Instrument[hxd]
Name of Detector (- if not required)[-]
Name of Filter (- if not required)[-]
Calibration Dataset Codename[SPECRESP MATRIX]
Requested Date in yyyy-mm-dd format[NOW]
Requested Time in hh:mm:ss format[NOW]
Boolean selection expression for Boundary params(- if not required)[-] DETCHANS.EQ.256
/caldb/test/staging/data/suzaku/hxd/cpf/ae_hxd_pinhxnom_20051104.rsp 1
/caldb/test/staging/data/suzaku/hxd/cpf/ae_hxd_pinxinom_20051104.rsp 1
```

Extensions in both files are identified by quzCIF since these particular extensions are valid for the same values of DETCHANS in their calibration boundary arrays.

BOUNDARY= NOMINAL.EQ.HXD

```
[corcoran:data/suzaku/hxd] corcoran% quzCIF
Name of Mission[-]suzaku
Name of Instrument[-]hxd
Name of Detector (- if not required)[-]
Name of Filter (- if not required)[-]
Calibration Dataset Codename[-]SPECRESP MATRIX
Requested Date in yyyy-mm-dd format[-] NOW
Requested Time in hh:mm:ss format[-] NOW
```

```
Boolean selection expression for Boundary params(- if not required)[-] NOMINAL.EQ.HXD
/caldb/test/staging/data/suzaku/hxd/cpf/ae_hxd_pinhxnom_20051104.rsp 1
```

```
BOUNDARY= NOMINAL.EQ.XIS
```

```
=====
```

```
[corcoran:data/suzaku/hxd] corcoran% quzcif
Name of Mission[-] suzaku
Name of Instrument[-] hxd
Name of Detector (- if not required)[-]
Name of Filter (- if not required)[-]
Calibration Dataset Codename[-]SPECRESP MATRIX
Requested Date in yyyy-mm-dd format[-] NOW
Requested Time in hh:mm:ss format[-] NOW
Boolean selection expression for Boundary params(- if not required)[-] NOMINAL.EQ.XIS
/caldb/test/staging/data/suzaku/hxd/cpf/ae_hxd_pinxinom_20051104.rsp 1
```

One extension is selected since the files differ in the value of the “NOMINAL” boundary condition.

6 Notification

Once files have been successfully added to the mission’s CALDB, users should be notified. If the files are to be included in the HEASARC CALDB, the HEASARC CALDB manager should be notified. Instructions concerning the inclusion of calibration files in the HEASARC CALDB are available from CAL/GEN/2003-001, “Automated Delivery of Calibration Data to the CALDB”.

7 Error Messages

Sometimes the following errors are encountered as given in table 4:

Table 4: Error Messages

Error	Meaning
Cannot get directory from CGDR	This usually means the \$CALDB is undefined.
Error calculating the REF_TIME value	One of the REF_TIME values can’t be calculated because of trouble parsing the CVSD or CVST values.
Error searching alias config file	the \$CALDBALIAS file can’t be found or read