

INTEGRAL Science Data Centre

IBIS ANALYSIS USER MANUAL

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INTEGRAL Science Data Centre Chemin d'Écogia 16 CH–1290 Versoix Switzerland

http://isdc.unige.ch

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Prepared by :	M. Chernyakova
Agreed by :	R. Walter
Approved by :	T. Courvoisier

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Acronyms and Abbreviations

AD	Architectural Design	HEPI	Hardware Event Processor	
ADD	Architectural Design Document	HV	High Voltage	
A/D	Analog-Digital	IC	Instrument Characteristics	
AFEE	Analog Front End Electronics	IJD	Integral Julian Day	
ASIC	Application Specific Integrated Circuits	ISDC	Integral Science Data Center	
BGO	Bismuth Germanate	ISOC	Integral Science Operations Centre	
CdTe	Cadmium-Telluride	MCE	Module Control Electronics	
CsI	Caesium-Iodide	MDU	Modular Detection Units	
DBB	Detector Bias Box	OBT	On-Board Time	
DFEE	Digital Front End Electronics	OG	Observation Group	
DOL	Data Object Locator	PCFOV	Partially Coded Field of View	
DPE	Data Processing Electronics	PEB	PICsIT Electronic Box	
DS	Data Structure	PIF	Pixel Illuminated Factor	
FCFOV	Fully Coded Field of View	PMT	Photomultiplier Tube	
FIFO	First-In, First-Out	PLM	Payload Module	
FOV	Field of View	RMF	Redistribution Matrix Files	
FWHM	Full Width at Half Maximum	ScW	Science Window	
GPS	Galactic Plane Scan	SWG	Science Window Group	
GTI	Good Time Interval	TBW	To be written	
GUI	Graphical User Interface	TM	Telemetry	

Glossary of Terms

- *ISDC system:* the complete ground software system devoted to the processing of the *INTEGRAL* data and running at the ISDC. It includes contributions from the ISDC and from the *INTEGRAL* instrument teams.
- Science Window (ScW): For the operations, ISDC defines atomic bits of INTEGRAL operations as either a pointing or a slew, and calls them ScWs. A set of data produced during a ScW is a basic piece of INTEGRAL data in the ISDC system.
- Observation: Any group of ScW used in the data analysis. The observation defined from ISOC in relation with the proposal is only one example of possible *ISDC observations*. Other combinations of Science Windows, *i.e.*, of observations, are used for example for the Quick-Look Analysis, or for Off-Line Scientific Analysis.
- *Pointing:* Period during which the spacecraft axis pointing direction remains stable. Because of the *INTEGRAL* dithering strategy, the nominal pointing duration is of order of 20 minutes.
- *Slew:* Period during which the spacecraft is manoeuvred from one stable position to another, i.e., from one pointing to another.

1 Introduction

The 'IBIS Analysis User Manual', *i.e.*, this document, was edited to help you with the IBIS specific part of the *INTEGRAL* Data Anaysis.

A more general overview on the *INTEGRAL* Data Analysis can be found in the 'Introduction to the *INTEGRAL* Data Analysis' [1]. For the ISGRI and PICsIT analysis scientific validation reports see [3] and [4].

The 'IBIS Analysis User Manual' is divided into two major parts:

- Description of the Instrument This part, based to some extent on the ISOC AO-2 document [2], introduces the *INTEGRAL* on-board Imager (IBIS).
- Description of the Data Analysis This part starts with an overview describing the different steps of the analysis. Then, in the Cookbook Section, several examples of analysis and their results and the description of the parameters are given.

Finally, the used algorithms are described. A list of the known limitations of the current release is also provided.

In the Appendix of this document you find the description of the Raw and Prepared Data and also the description of the Scientific Products. If you are interested in Data Structures not described in the Appendix go to the ISDC web-page http://isdc.unige.ch/index.cgi?Data+templates

Part I Instrument Definition

2 Scientific Performances Summary

IBIS is a gamma-ray telescope observing celestial objects of all classes ranging from the most compact galactic systems to extragalactic objects, with powerful diagnostic capabilities of fine imaging, source identification and spectral sensitivity in both continuum and lines. It is able to localize weak sources at low energy to better than a few arcminutes accuracy, covering the entire energy range from a few tens of keV to several MeV. Table 1 gives an overview of the scientific capabilities of IBIS. The effective area curves are given on the Figure 1.

Table 1: Scientific Parameters of IBIS.			
Operating energy range	15 keV - 10 MeV		
Energy resolution (FWHM)	7% @ 100 keV		
	9% @ 1 MeV		
Effective Area	ISGRI: 960 cm^2 at 50 keV		
	PICsIT: 870 cm^2 at 300 keV (single events)		
	PICsIT: 275 cm^2 at 1 MeV (multiple events)		
Field of view	$9^{\circ} \times 9^{\circ}$ (fully coded)		
	$19^{\circ} \times 19^{\circ}$ (partially coded, 50%)		
Angular resolution (FWHM)	12'		
Point source location accuracy	30'' @100 keV		
(90% error radius)	<5' @1 MeV		
Continuum sensitivity,	3.8×10^{-7} @100 keV		
photons $\mathrm{cm}^{-2} \mathrm{s}^{-1} \mathrm{keV}^{-1}$	$1 - 2 \times 10^{-7} @ 1 \text{ MeV}$		
$(3\sigma \text{ detection}, \Delta E = E/2, 10^6 \text{ s integration})$			
Narrow line sensitivity,	1.3×10^{-5} @100 keV		
photons $\rm cm^{-2} \ s^{-1}$	$4 \times 10^{-5} @ 1 \text{ MeV}$		
$(3\sigma, 10^6 \text{ s integration})$			
Absolute timing accuracy (3σ)	ISGRI: 61 μ s		
	PICsIT: 0.976 – 500ms (selected from ground)		



Figure 1: IBIS effective area

3 Instrument Description

3.1 The Overall Design

IBIS is a gamma-ray imager operating in the energy range 20 keV to 10 MeV, with two simultaneously operating detectors covering the full energy range, located behind a Tungsten mask which provides the encoding.

The coded mask is optimized for high angular resolution. As diffraction is negligible at gamma-ray wavelengths, the angular resolution of a coded-mask telescope is limited by the spatial resolution of the detector array. The angular resolution of a coded mask telescope $d\theta$ is defined by the ratio between the mask element size C (11.2 mm) and the mask-to-detection plane distance H (3133 mm).

$$d\theta = \arctan\left(\frac{C}{H}\right) = 12'$$

IBIS is made of a large number of small, fully independent pixels.

The detector features two layers, ISGRI and PICsIT: the first is made of Cadmium-Telluride (CdTe) solidstate detectors and the second of Caesium-Iodide (CsI) scintillator crystals. This configuration ensures a good broad line and continuum sensitivity over the wide spectral range covered by IBIS. The double-layer discrete-element design of IBIS allows the paths of interacting photons to be tracked in 3D if the event involves detection units of both ISGRI and PICsIT. The application of Compton reconstruction algorithms to these types of events (between few hundred keV and few MeV) allows an increase in signal to noise ratio attainable by rejecting those events unlikely to correspond to source photons inside the field of view.

The detector aperture is restricted, in the hard X-ray part of the spectrum, by passive shielding covering the distance between mask and detection plane. An active BGO scintillator VETO system shields the detector bottom as well as the four sides up to the bottom of ISGRI.

Figure 2 shows a cut-away drawing of the various components of IBIS (except the mask and tube). Figure 3 shows the distances between the different parts of the detector assembly. Figure 4 shows the spacecraft & instruments coordinate systems.



Figure 2: Cutaway drawing of the IBIS detector assembly, together with the lower part of the collimator (Hopper).



Figure 3: IBIS detector assembly in numbers.



Spacecraft & Instrument Coordinate Systems

Figure 4: Spacecraft & Instrument Coordinate Systems. Note that the X-axis of the spacecraft is defined by the pointing direction.

3.2 The Subsystems

3.2.1 The Mask

The IBIS Mask Assembly is rectangular with external dimensions of $1180 \times 1142 \times 114 \text{ mm}^3$, and consists of three main subsystems: the Coded Pattern, the Support Panel and the Peripheral Frame with the necessary interface provisions.

The Coded Pattern is a square array of size $1064 \times 1064 \times 16 \text{ mm}^3$, made up of 95×95 individual square cells of size $11.2 \times 11.2 \text{ mm}^2$. The mask chosen for IBIS is based on a cyclic replication of MURA (Modified Uniformly Redundant Array) of order 53. The properties of the MURA patterns are described e.g. in [11] and [12].

Approximately half of the mask cells are opaque to photons in the operational energy range of the IBIS instrument, offering a 70% opacity at 1.5 MeV. The other 50% of cells are open, *i.e.*, with an off-axis transparency of 60% at 20 keV. Figure 5 shows the mask pattern.



Figure 5: The IBIS coded mask pattern.

The Support Panel includes additional elements to support the code mask pixels, providing the necessary stiffness and strength to overcome the launch environment and the in-orbit operational temperatures. This panel is done from the material known as "nomex". Its transparency should be taken into account in the data analysis, as it absorbs part of the flux.

Figure 6 shows the cross section of the support panel.

The Peripheral Frame reinforces the sandwich panel.



Figure 6: The cross section of the support panel.

The mechanical interfaces with the INTEGRAL payload module also provide extra Tungsten shielding to the diffuse background through the gap between the mask edges and the payload vertical walls.

3.2.2 The Collimator

In order to maintain the low energy response of IBIS despite the dithering needed for SPI, the collimation baseline consists of a passive lateral shield that limits the solid angle (and therefore the cosmic gamma-ray background) viewed directly by the IBIS detector in the full field of view up to a few hundreds of keV. The tube collimation system is implemented with three different devices:

- *The Hopper*: four inclined walls starting from the detector unit with a direct interface to the IBIS detector mechanical structure. The hopper is not physically connected to the payload module structure.
- The Tube: The Tube is formed by four payload module walls shielded with glued Lead foils.
- *The additional side shielding on the mask.* Four strips of 1 mm thick Tungsten provide shielding from the diffuse background in the gaps between the mask edges and the top of the tube walls.

3.2.3 Detector

The ISGRI CdTe and PICsIT CsI(Tl) detectors are layered with respect to each other, with PICsIT below ISGRI with respect to the coded mask (and hence the astronomical source).

• Upper Detector Layer: ISGRI

Cadmium Telluride (CdTe) is a semiconductor operating at ambient temperature. $0^{\circ} \pm 20^{\circ}$ C is the optimum range. With their small area, the CdTe detectors are ideally suited to build an image with good spatial resolution.



Figure 7: ISGRI and PICsIT division in modules and submodules

The CdTe layer is made of 8 identical Modular Detection Units each having 32×64 pixels (see Figure 7). Total sensitive area of the detector is 2621 cm².

• Lower Detector Layer: PICsIT

Caesium Iodide is a scintillation crystal. The CsI(Tl) layer is divided into eight rectangular modules of 16×32 detector elements (see Figure 7). In each module there are two independent semi-module each one with its independent Front End Electronics. Total sensitive area of the detector is 2994 cm².

• Noisy Pixels

It is possible that with the time some of the pixels of the detector may become out of order and start to produce outputs not triggered by an income photon, *i.e.*, to become "noisy". If the particular pixel countrate is too high relatively to the module countrate, then the on-board electronics switch it off. In ISGRI case the noisy pixels can recover after being switched off for some time and disabled pixels are periodically reset to check their status.

In PICsIT case, pixels cannot be recovered that easily. PICsIT pixel will remain off once killed. Only if half of the detector (or so) will be off, an attempt will be made to turn pixels on. The current situation is shown on Figure 8. Overall the killed pixels are less than 1%.

3.2.4 On-board Calibration Unit

IBIS contains an on-board collimated radioactive ²²Na source. This allows regular calibration of PICsIT at both the 511 keV line (calibration to better than 1% in 4 hours) and 1275 keV (1% in 8 hours). ISGRI can also use the 511 keV line, albeit at lower efficiency. Any energy deposits from untagged photons will have an impact of < 1% on the overall continuum sensitivity between 100 keV and 2 MeV.

3.2.5 Veto Shield

The Veto shield is crucial to the operation of IBIS. IBIS uses anticoincidence logic to accept or reject detected events as real photons in the field of view, or background particles or photons propagating through,



Figure 8:

gure 8: The schematic view of PICsIT layer. Each module number is indicated. The dotted lines represents the division in semimodules whose number is indicated at the top. The black pixel are the killed ones. The (Y,Z) coordinates are the IBIS ones both ranging from 0 to 63. X-axis is directed toward the source located above the page. The Z-axis is pointing positively to the sun.

or induced in, the spacecraft.

The sides, up to the ISGRI bottom level, and rear of the stack of detector planes are surrounded by an active Bismuth Germanate (BGO) veto shield. Like the detector array, the Veto shield is modular in construction. There are 8 lateral shields, *i.e.*, 2 modules per side, and 8 bottom modules.

The high density and mean Z of BGO ensures that a thickness of 20 mm is sufficient to reduce the detector background due to leakage through the shielding of cosmic diffuse gamma-ray background and gamma-rays produced in the spacecraft, to less than the sum of all other background components.

4 How the Instrument works

4.1 Event Types

The photon entering the telescope can be detected due to its interaction with the absorbing material of the detector. Three major types of interactions play a dominant role: photoelectric absorption, Compton scattering and pair production. In the photoelectric absorption process a photon undergoes an interaction with an absorber atom in which the photon completely disappears. In its place an energetic photoelectron is ejected by the atom, carrying away most of the original photon energy. The Compton scattering takes place between the incident gamma-ray photon and an electron in the absorbing material. The incoming photon is deflected and it transfers a portion of its energy to the electron. The energy transferred to the electron can vary from zero to a large fraction of the initial gamma-ray energy. In the pair production process the gamma-ray photon disappears and is replaced by an electron-positron pair. The positron will annihilate in the absorbing medium and two annihilation photons are normally produced as secondary products of the interaction. Depending on the size of the detector and on the energy of the incoming photon, a photon scattered in a Compton interaction can escape the detector, or undergo a second interaction. The pairs of 511 keV photons, produced by the annihilation of the positrons resulting from pair creation, can also produce other interactions or escape the detector.

Both ISGRI and PICsIT record the coordinates of each event registered in the corresponding layer, to build up an image. The anticoincidence VETO is used to reject background events.

The coded mask produces a shadowgram. Photons from the source and the background are distributed across the entire field of view, but cross-correlation techniques allow the full image to be reconstituted for the fully coded field of view ($9^{\circ} \times 9^{\circ}$) at each pointing. For the partially coded field of view (out to $29^{\circ} \times 29^{\circ}$), special cleaning techniques must be applied to the data to properly reconstruct the image. The actual sky coverage in an observation of course depends on the dither pattern.

The on-board electronics classify registered events according to the activated layer and the number of events detected by a submodule practically simultaneously. Events detected by different submodules are treated as independent ones. There are five main events type:

• ISGRI single event

Photon is stopped in a single pixel of the ISGRI layer, generating an electric pulse.

In principle, the amplitude of the pulse yields the energy of the incident photon. However, above 50 keV the energy is a function of not just the pulse height but also the pulse rise time, so both are used to determine the energy of the incident photon. In addition the resulting line profile (energy resolution) is no longer Gaussian, but more similar to a Lorentzian. The energy resolution depends on the operating temperature, and also on the bias voltage; the bias voltage has to be optimized as a trade-off between high resolution but more noise (high voltage) or lower noise but lower resolution (low voltage).

All cases of multiple ISGRI detection units excitation (in one module) are rejected. In case of the excitation of the detection units in different modules, such events are treated as independent single events.

• PICsIT single event

Photon passes through ISGRI and is stopped in a single pixel of the PICsIT layer, generating one scintillation flash.

The energy of the incident photon is derived, in each crystal bar, from the intensity of the flash recorded in the photodiode. The energy resolution of PICsIT is a function of the signal-to-noise of the event, which in turn is governed by factors operating conditions and PIN capacity.

• PICsIT multiple event

Several PICsIT detection units in one submodule were excited during one event, generating several scintillation flashes. The energy of the primary photon is determined from the sum of the energies of all detected events. The position of the incoming photon is attributed to the position of the most energetic event.

- Compton single event Photons arriving in either ISGRI or PICsIT produce secondary photon via Compton scattering, detected in another layer. The position of the incoming photon is attributed to the position of the most energetic event, and the energy is determined as the sum of the detected events energies.
- Compton multiple event

One ISGRI detection unit and several PICsIT detection units in one submodule were excited. As in previous cases the position of the incoming photon is attributed to the position of the most energetic event, and the energy is determined as the sum of the detected events energies.

In Fig.9 the efficiencies of the various detection techniques is shown.



Figure 9: IBIS sensitivities for the various detection techniques.

4.2 IBIS observing modes

IBIS has several observing modes, for engineering and calibration purposes. However, for scientific use there is only one operating mode, Science Mode.

In Science Mode, ISGRI registers and transmits events on a photon-by-photon basis, *i.e.*, every event is tagged with (X,Y) position on the detector plane, event energy (from the pulse height and rise time) and event time.

PICsIT in principle can also operate in photon-by-photon mode. However, with the higher background compared to ISGRI, there would be unacceptable data loss. Therefore, the standard mode for PICsIT is histogram. Images and spectra (full spatial resolution, 256 energy channels) are accumulated for about 30 minutes before transmission to ground. There is no time-tagging internal to the histogram, *i.e.*, spectral imaging has time resolution of 30 minutes.

In addition, coarse spectra, without imaging information, are accumulated by PICsIT and transmitted with far higher time resolution, but without imaging information. Thus their usefulness is limited to observations of very strong sources where the source countrate dominates the background. The time resolution, and the number of energy channels, for this spectral timing data can be commanded from ground. The time resolution can take values between 1 and 500 ms; the current default is 500 ms and two energy channels, but the values to be used for routine observations will be decided when the in-flight background of PICsIT is measured (and compared with the available telemetry rate), during the commissioning phase.

In Table ${\bf 2}$ the properties of the all modes are summarized.

	Detector Image	Timing	Spectral	
Observing Mode	Resolution	Resolution	Resolution	
	(pixels)		(channels)	
ISGRI				
photon-by-photon	128×128	$61.035 \mu s$	2048	
PICsIT				
Photon-by-Photon	64×64	$64 \mu s$	1024	
Spectral-Imaging	64×64	$\leq \sim 30 \min$	256	
Spectral-Timing	None	$1-500\mathrm{ms}$	2-8	

 Table 2: Characteristics of the IBIS Telemetry Formats

Part II Cookbook

This Part was completely rewritten by

M.Chernyakova A.Paizis I. Lecoeur-Taibi

5 Overview

In this Section an overview of the analysis of IBIS data is given.

Each photon detected by IBIS is analyzed with the on-board electronics and tagged with the arrival time, type (ISGRI, PICsIT, Compton¹ etc.), energy, position etc. according to the operation mode (i.e. photon-by-photon, standard, calibration etc.). These data are then sent to ground in telemetry (TM) packets.

During Pre-Processing the TM packet information is deciphered and rewritten into the set of FITS files (RAW data). Then the local on-board time is converted into the common on-board time (OBT) and the House Keeping (HK) parameters into physical units (PRP data).

These steps are done at ISDC and you do not have to redo them. In the Appendix you will find the description of the raw and prepared data and also the description of the instrument characteristic files that are used in the Scientific Analysis.

INTEGRAL data is organized into the so-called Science Windows (see Introduction to the *INTEGRAL* Data Analysis [1] for more explanations). During the scientific analysis, all the Science Windows belonging to the same observation are grouped together to form the "Observation Group".

Figure 10 shows in details the different steps performed by the scientific analysis script, *ibis_science_analysis*. This high level script consists of three smaller ones: *ibis_scw1_analysis*, *ibis_obs1_analysis* and *ibis_scw2_analysis*. *ibis_scw1_analysis* and *ibis_scw2_analysis* work on a Science Window basis while *ibis_obs1_analysis* works on the Observation Group basis. Each subscript performs the tasks shown in Figure 10, explained in more details in the text below.



Figure 10: Science Analysis Overview

¹For the time being, Compton analysis is not available

• The first script *ibis_scw1_analysis* performs the following tasks:

<u>COR – Data Correction</u>

Tags noisy pixels, corrects energy of the photons for rise time and temporal variations of the gain, transforms channels to energy. See Section 12.1 for more details.

GTI – Good Time Handling

Generates, selects, and merges Good Time Intervals (GTI) to produce a unique GTI that is then used by the software to select good events. See Section 12.2 for more details.

DEAD – Dead Time Calculation

Calculates the total dead time during which the incoming photons may be lost due to the processing of the previous events. Also veto strobe signals generated by BGO (Bismuth Germanate) shield, calibration source and Compton events are taken into account. See Section 12.3 for more details.

BIN_I – Event Binning for Imaging

Sorts data into energy bins. For each energy range, the intensity shadowgram and a corresponding efficiency map are created. See Section 12.4 for more details.

BKG_I – Background Correction

Creates rebinned maps for background and absorption of support mask (see Section 3.2.1) corrections. Corrects for efficiency and subtracts background. See Section 12.6 for more details.

After these steps the high-level analysis is performed.

• The second script *ibis_obs1_analysis* takes the whole Observation Group previously created as input and performs the following tasks:

CAT_I - Catalog Source Selection for Imaging

Selects from the given catalog a list of sources in the Field of View matching the criteria defined by script parameters, and creates an output list with location and expected flux values of the selected sources. See Section 12.7 for more details.

IMA - ISGRI and PICsIT (staring) Image Reconstruction

In the case of ISGRI, shadowgrams are deconvolved, source search is performed in the single images as well as in the mosaic (combination of different images) and a list of detected sources is created.

If *INTEGRAL* was stable during the whole period of interest, then, at your request, all PICsIT shadowgrams are combined into one and then are deconvolved into a single image. See Section 12.8 for more details.

• The third script *ibis_scw2_analysis* again works Science Window by Science Window and performs the following tasks:

IMA2 - PICsIT Image Reconstruction

PICsIT shadowgram deconvolution is done at this step, creating a separate image for each science window. See Section 12.8.3 for more details. Nothing is done at this step for ISGRI.

BIN_S – Event Binning for Spectra

Creates rebinned maps for background and absorption of support mask (see Section 3.2.1) corrections. Sorts data into energy bins. For each energy range the shadowgram and a corresponding efficiency shadowgram is created. See Sections 12.4, 12.6 for more details.

SPE - ISGRI spectra extraction

For each source of interest, one PIF² is produced. ISGRI spectral extraction is done for all catalog

 $^{^{2}}$ PIF is a number between 0 and 1, which expresses the theoretical degree of illumination of each detector pixel for a given source in the sky.

sources with the use of these PIFs. See Section 12.9 for more details.

LCR - PICsIT Detector Light Curve Creation and ISGRI source lightcurve extraction

At this step, PICsIT Detector light curves are built from the spectral timing data. For all sources from the input ISGRI catalog light curves are extracted. See Section 12.10 for more details.

CLEAN – Last step

At this step PICsIT mosaic is created. See Section 12.11 for more details.

As of October 18, 2004, all public INTEGRAL data are available in two formats: revision 1 and revision 2. In revision 2 data, the correction of all JD time stamps for the offsets between the OBT of each instrument is done, and as all data are available in revision 2 format, data in revision 1 format become obsolete.

For revision 2 data, the data correction step (COR), as well as the instrumental GTI and deadtime handling (DEAD) steps have already been performed using version 4.2 of OSA at the science window level. However, the data correction implemented in OSA 5.0 is much better and it is highly recommended to rerun these steps, starting from the COR level.

6 Getting started

This chapter describes how to set up the the environment and the analysis data and how to analyse data from the two instruments that are part of IBIS: ISGRI and PICsIT. These two instruments are quite different in energy range (ISGRI starts from 15 keV and PICsIT starts from 200 keV) and sensitivity, and are optimised for different targets. This is why we have decided to guide you through the analysis of the crowded Galactic Centre around 4U 1700-377 for ISGRI and of the bright Crab for PICsIT.

Here we assume that you have already successfully installed ISDC Off-line Scientific Analysis (OSA) Software version 5.0 (the directory in which OSA is installed is later on referred to as the ISDC_ENV directory). If not, then look at the "Installation Guide for the INTEGRAL Data Analysis System" [5], for detailed help.

6.1 Setting up the analysis data

In order to set up a proper environment, you first have to create an analysis directory (e.g ibis_data_rep) and "cd" into it:

mkdir ibis_data_rep
cd ibis_data_rep

This working directory ibis_data_rep will be referred to as the "REP_BASE_PROD" directory in the following. All the data required in your analysis should then be available from this "top" directory, and they should be organized as follows

- scw/ : data produced by the instruments (e.g., event tables) cut and stored by ScWs;
- **aux**/ : auxiliary data provided by the ground segment (e.g., time correlations);
- **cat**/ : ISDC reference catalogue (OSA_CAT package);
- ic/ : Instrument Characteristics (IC), such as calibration data and instrument responses (OSA_IC package);
- idx/ : set of indices used by the software to select approriate IC data (OSA_IC package).

The cat/, ic/ and idx/ directories are part of the OSA software distribution and should be installed following the "Installation Guide for the *INTEGRAL* Data Analysis System" [5]. The actual data along with the auxiliary files (scw/ and aux/) are sent to the Principal Investigators of the observation. Alternatively, the public data can be downloaded from the archive (see Section 6.1.1). In case the data are already available on your system you can either copy these data to the relevant working directory, or better, create soft links as shown below. Alternatively, if you do not have any of the above data on your local system, or if you do not have a local archive with the scw/ and the aux/ branch available, follow the instructions in the next section to download data from the ISDC WWW site.

```
ln -s directory_of_ic_files_installation__/ic ic
ln -s directory_of_ic_files_installation__/idx idx
ln -s directory_of_cat_installation__/cat cat
ln -s directory_of_local_archive__/scw scw
ln -s directory_of_local_archive__/aux aux
```

Then, just create a file 'isgri_gc.lst' containing the 5 lines:

```
scw/0051/005100410010.001/swg.fits[1]
scw/0051/005100420010.001/swg.fits[1]
```

```
scw/0051/005100430010.001/swg.fits[1]
scw/0051/005100440010.001/swg.fits[1]
scw/0051/005100450010.001/swg.fits[1]
```

and a file 'picsit.lst' containing:

scw/0039/003900020020.001/swg.fits[1] scw/0039/003900020030.001/swg.fits[1] scw/0039/003900020040.001/swg.fits[1] scw/0039/003900020050.001/swg.fits[1] scw/0039/003900020060.001/swg.fits[1]

The created files contain the list of Scws you want to analyze (technically, we call them DOLs -Data Object Locators-, i.e. a specified extension in a given FITS file 3).

These file names 'isgri_gc.lst' and 'picsit.lst' will be used later as an argument for the og_create program (see Sections 7, 10).

6.1.1 Downloading data from the archive

• To retrieve the required ISGRI analysis data from the archive, go to the following URL:

http://isdc.unige.ch/index.cgi?Data+browse

You will reach the W3Browse web page which will allow you to build a list of Science Windows (SCWs) that you will analyse with OSA.

- Type the name of the object (4U 1700-377) in the 'Object Name Or Coordinates:' field

- Do not forget to change 'Search Radius' if you are interested in science windows where your source is in the partially coded field of view. Set it to e.g. 10 degrees

-Click on 'More Options' button at the top or at the bottom of the web page

- Deselect the 'All' checkbox at the top of the Catalog table, and select the 'SCW - Science Window Data' one

- Press the 'Specify Additional Parameters' button at the bottom of the web page

Introduce values in the fields of interest. For instance:

- Sort output by $\mathbf{scw_id},$ for that check the 'Sort' column.

- Put pointing in the field 'scw_type' (to specify that only pointings should be returned and not e. g. slews)

- Put >=2003-03-15 23:00:00 in the field 'start_date' and put <= 2003-03-16 02:30:00 in the field 'end_date'

- Put public in the field 'ps' (to specify that only public ScWs should be returned)

- Put >100 in the field 'good_isgri' (to select Science Windows with good ISGRI time higher than 100 seconds)

- Press the 'Start Search' button at the bottom of the web page. In our case, a table with 5 ScWs will be displayed.

- Select the SCWs of interest. To follow the example in the Cookbook click on All (for all SCWs).

- Press the 'Save SCW list for the creation of Observation Groups' button at the bottom of that table and save the file with the name 'isgri_gc.lst'. This file 'isgri_gc.lst' will be used later as input for the og_create program (see Section 7). From this file you need the 5 lines:

scw/0051/005100410010.001/swg.fits[1] scw/0051/005100420010.001/swg.fits[1]

 $^{^{3}}$ When an analysis script asks you to specify the DOL, you should specify the path of the corresponding FITS file, and the corresponding name or number of the data structure in square brackets (do not forget that numbering starts with 0!). See more details in the Introduction to the INTEGRAL Data Analysis [1]. Please note that the naming scheme is different for revision 1 and revision 2 data. For the revision 1 data, the name of the prepared Science Window Group is swg_prp.fits instead of swg_fits.

scw/0051/005100430010.001/swg.fits[1] scw/0051/005100440010.001/swg.fits[1] scw/0051/005100450010.001/swg.fits[1]

You should then download them pressing the '**Request** data products for selected rows' button. In the 'Public Data Distribution Form', provide your e-mail address and press the 'Submit Request' button. You will be e-mailed the required script to get your data and the instructions for the settings of the IC files and the reference catalogue. Just follow these instructions.

- To retrieve the required <u>PICsIT analysis data</u> from the archive proceed in the same manner with the following parameters:
 - 'Object Name Or Coordinates:' Crab
 - 'Search Radius' : default value
 - 'scw_id' : 0039% (To select only science windows starting with 0039)
 - 'start_date' : >=2003-02-07 06:44:19 and 'end_date': <=2003-02-07 12:44:05

and save your results in a file called 'picsit.lst' which should contain:

scw/0039/003900020020.001/swg.fits[1] scw/0039/003900020030.001/swg.fits[1] scw/0039/003900020040.001/swg.fits[1] scw/0039/003900020050.001/swg.fits[1] scw/0039/003900020060.001/swg.fits[1]

6.2 Setting the environment

Before you run any OSA software, you must also set your environment correctly.

The commands below apply to the csh family of shells (i.e csh and tcsh) and should be adapted for other families of shells⁴.

In all cases, you have to set the REP_BASE_PROD variable to the location where you perform your analysis (e.g the directory ibis_data_rep). Thus, type:

setenv REP_BASE_PROD \$PWD

Then, if not already set by default by your system administrator, you should set some environment variables:

```
setenv ISDC_ENV directory_of_OSA_sw_installation
setenv ISDC_REF_CAT "$REP_BASE_PROD/cat/hec/gnrl_refr_cat_0023.fits[1]"
source $ISDC_ENV/bin/isdc_init_env.csh
```

The latter command executes the OSA set-up script (isdc_init_env.csh) which initialises further environment variables relative to ISDC_ENV. Ignore all warnings mentioning ROOTSYS.

Besides these mandatory settings, the optional environment variable COMMONLOGFILE can also be useful. By default, the software logs messages to the screen (STDOUT). To have these messages in a file (i.e common_log.txt), and make the output chattier⁵ use the command:

setenv COMMONLOGFILE +common_log.txt

⁵For example, the exit status of the program will now appear.

 $^{^{4}}$ If the setenv command fails with a message like: 'setenv: command not found' or 'setenv: not found', then you are probably using the sh family. In that case, please replace the command 'setenv my_variable my_value' by the following command sequence 'my_variable=my_value; export my_variable'

In the same manner, replace the command 'source a_given_script.csh' by the following command '. a_given_script.sh' (the '.' is not a typo!).

6.3 Two ways of launching the analysis

6.3.1 Graphical User Interface (GUI)

When you launch the analysis the Graphical User Interface (GUI) is launched, providing an opportunity to set the values of all desired parameters, see Figure 12. On the right side of the panel you see the following buttons:

- Save as With the "Save As" button a file is created. This file stores all parameters as they are currently defined in the GUI as a command line script. This file is an executable one and calling it from the command line will launch the instrument analysis program with the parameters as they were defined in the GUI.
- Load With the "Load" button a previously saved file (see "Save As") can be read and the GUI will update all parameters with the values as they are defined in the loaded file.
- ResetWith the "Reset" button the parameters in the GUI will be reset to the default values as they are defined in the parameter file of the instrument analysis program and stored in the \$ISDC_ENV/pfiles directory.
- Run With "Run" button the analysis is launched.
- Quit With "Quit" button you quit the program without analysis launch.
- Help With "Help" button the help file of the main script is opened in a separate window.
- hidden With the "hidden" button you have an access to the hidden parameters with values defined by the instrumental teams. Change them with care!

6.3.2 Launching scripts without GUI

Alternatively, parameters can be specified on the command line typing 'name = value' after the script name.

If you are running your own scripts that call OSA many times you don't want GUI to pop up each time. In such a case set **COMMONSCRIPT** variable to "1" with:

setenv COMMONSCRIPT 1

This is automatically done within the file created with the help of "Save as" - button, see above.

To have the GUI back again, unset the variable:

unsetenv COMMONSCRIPT

6.4 Useful to know!

- How do I get some help with the executables? All the available help files are stored under \$ISDC_ENV/help. To visualize a help file interactively type tool_name --h once your environment is set (i.e. *which tool_name* returns the path to it).
- Where are the parameter files and how can I modify them? All the available executables for the analysis of *INTEGRAL* data are under \$ISDC_ENV/bin. The corresponding parameter files are stored under \$ISDC_ENV/pfiles/*.par. The first time you launch a script, the system will copy the specific tool.par from \$ISDC_ENV/pfiles/ to a local directory (/user_name/pfiles/). The parameter file in the local directory is the one used for the analysis and is

the one you can modify. If this parameter file is missing (e.g. you have deleted it), the system will just re-copy it from \$ISDC_ENV/pfiles/ as soon as you launch the script again. The system knows what to copy from where thanks to the \$PFILES environment variable that is also used in FTOOLS (http://heasarc.gsfc.nasa.gov/ftools/). Each parameter is characterized with a letter that specifies the parameter type, i.e:

"q" (query) parameters are always asked to the user

"h" (hidden) parameters are not asked to the user and the indicated value is used

"l" (learned) parameters are updated with the user's value during the use of the program. The GUI is a fast and easy way to change the parameters, see section 6.3.1 for details.

• What are groups and indices?

The ISDC software makes extensive use of groups and indices. While it is not necessary to grasp all the details of these concepts, a basic understanding is certainly quite useful.

As implied by their names, "groups" make possible the grouping of data that are logically connected. Groups can be seen as a kind of data container, not completely unlike standard directories. At ISDC, we create separate groups for each pointing, in which we store the many different data types produced by Integral and its instruments. The user then only has to care about one file, the group, many tens of files being silently included. Several pointings (the "Science Window Groups") can be arbitrarily grouped into bigger groups (the "Observation Group") to select data very efficiently according to the user's needs.

Indices are a special kind of groups, which differ only in the fact that all the the data sets they contain are similar and that the indices know the properties of the data sets they contain. Indices are a kind of poor man's database. For example, an imaging program creates several images of different types (flux map, significance map,...) in different energy bands. These images are stored in an index, in which the image type and energy band information is replicated. ISDC software is then able to select very efficiently the needed images. The user can also make use of the indices; just by looking at the index (for instance using "fv"), the user can identify immediately the content of each image.

• Why do I need "[1]" after a FITS file name?

A FITS file can have many extensions and sometimes it is necessary to specify as input to a given parameter not the file name alone (file.fits) but the extension too (file.fits[1], or file.fits[2], etc). The file name with a specified data structure (extension) is called DOL (Data Object Locator). When you modify the parameter file itself (see above) or use the GUI, the extension will be correctly interpreted in the file.fits[1] case. On the command line though, the normal CFITSIO and FTOOLS rules apply, i.e. you have to specify it as one of the following file.fits+1 "file.fits=1".

Note that if no extension is specified explicitly then the first one ([1]) will be used by default.
7 A Walk through ISGRI Analysis

After setting up as described in the previous section, you are ready to analyse the data.

Please do remember that you are dealing with a coded mask instrument not with a focussing telescope and a CCD. It is not possible to deal with one source at a time: each source is background for the others, the whole field of view - and not just the few pixels around your source - matters!

In this Section, we guide you through your first IBIS analysis, but please read also Section 8, where more details on the main parameters are given. You could end up with fake sources that are created by a blind use of parameters! More tips and tricks are given in Section 9 for advanced users.

In the example below we analyze observations of the Galactic Center, using data we have downloaded and installed as it is described in Section 6.1.

Create the Observation Group with the og_{create} program (see its description in the Toolbox section of [1]):

cd \$REP_BASE_PROD

og_create idxSwg=isgri_gc.lst ogid=isgri_gc baseDir="./" instrument=IBIS

As a result of the og_create command, the directory $REP_BASE_PROD/obs/isgri_gc$ is created. In this directory you find all you need for the analysis, its structure is illustrated in Figure 11.



Figure 11: Structure of the directory created with *og_create*

7.1 Image Reconstruction

The first thing to do when you are looking for the first time at your data is to create an image in order to know how the portion of sky you are interested in looks like, whether your source is detected, and what other sources you should take into account to do spectral and lightcurve analysis in a proper way. To start the analysis, move to the working directory $REP_BASE_PROD/obs/isgri_gc$ and call the *ibis_science_analysis* script:

cd obs/isgri_gc ibis_science_analysis

After a few seconds the main page of the IBIS Graphical User Interface (GUI) appears, as shown in Figure 12.

		ibis_science_analysis		
Main-				<u>S</u> ave As
				Load
	startLevel:	COR 💌		<u>R</u> eset
	endLevel:	IMA2		Run
	GENERAL_levelList:	COR,GTI,DEAD,BIN_I,BKG_I,CAT_I,IMA,IMA2,BIN_S,SPE,LCR,COMP,CLEAN		Quit
				<u>H</u> elp
	CAT_refCat:	/isdc/arc/rev_2/cat/hec/gnrl_refr_cat_0020.fits[1][ISGRI_FLAG==1]	browse	hidden
	SWITCH_disableIsgri:	Checked: yes		
	SWITCH_disablePICsIT:	Checked: yes		
	SCW1_GTI_gtiUserI:	<u></u>	browse	
	SCW1_GTI_TimeFormat:			
-	ISGRI IMA	ISGRI SPE and LCR	PICsI	T analysis

Figure 12: Main page of the IBIS GUI

Keeping all the default values, you will make an analysis starting from the energy correction level (startLevel = COR) until the image reconstruction level (endLevel = IMA2⁶). The default input catalog (CAT_refCat = $SISDC_REF_CAT[ISGRI_FLAG==1]$) will be used: it contains the sources that were detected by ISGRI in the public data up to April 2005. ISGRI data alone will be processed (SWITCH_disablePICsIT=yes) through all the available levels shown in GENERAL_levelList within COR and IMA2. No additional good time interval will be applied to the data: SCW1_GTI_gtiUserI field is empty, which also implies that the last parameter SCW1_GTI_TimeFormat is not effective.

You are ready to set the parameters specific for imaging. Press the ISGRI_IMA button at the bottom of the GUI. Another box, shown in Figure 13 appears.

With the parameters displayed in the GUI, you will create 4 different images of the sky, corresponding to 4 energy bands (IBIS_II_ChanNum=4) in the ranges 20–40, 40–60, 60-100 and 100-200 keV. You will let the software look for the 15 brightest sources in the field of view (OBS1_Searchmode=2 and OBS1_ToSearch=15), with detection significance higher than 6 for catalog sources (OBS1_MinCatSouSnr=6), and 7 for new sources (OBS1_MinNewSouSnr=7). Given that you are analysing the crowded Galactic center, it might be better to increase the number of sources you are looking for. Change manually OBS1_ToSearch to 25. The position of all the catalog sources will be fitted (OBS1_SouFit=0). Resulting images will be cleaned with the available background maps provided by the IBIS team (empty value of SCW1_BKG_LisgrBkgDol means usage of the default map

\$REP_BASE_PROD/ic/ibis/bkg/isgr_back_bkg_0005.fits). After creation of all the individual science window images the mosaic image will be created (OBS1_DoPart2=1). In this mosaic image the photons belonging to a source are spread around the single central peak (OBS1_PixSpread=1), resulting in better source location.

Now that you have checked all the parameters, press **Ok**, the Imaging window disappears and you are back to the main GUI page. Press **Save**, and then **Run**.

 $^{^{6}\}mathrm{IMA2}$ level is for PICsIT analysis only, and there is no difference for ISGRI analysis whether endLevel is set to IMA, or IMA2.

	ISGRI IMA	· [
- ISGRI IMAGING		<u>k</u> <u>H</u> elp
IBIS_II_ChanNum:	4	
IBIS_II_E_band_min: 20 40 6	0 100	Energy
IBIS_II_E_band_max: 40 60 10	00 200	Roundaries
IBIS_II_inEnergyValues:		browse
OBS1_SearchMode:		
OBS1_ToSearch:	15	
OBS1_MinCatSouSnr:	6 ∉ Source Search	
OBS1_MinNewSouSnr:	7	
OBS1_DoPart2:	1 Ansaic	
OBS1_PixSpread:		
OBS1_SouFit:	🛛 🛃 Source Position	
SCW1_BKG_L_isgrBkgDol:		browse Background ma
SCW1_BKG_I_isgrBkgDol:		browse Background r

Figure 13: Imaging page of IBIS GUI

7.1.1 Results from the Image Step

When the analysis is finished, you will find that new files have appeared in your working directory. In Figure 14 an overview of the files related to the image reconstruction is given. The full description of all files produced at different levels is given in Appendix C.

As it is shown in Figure 14, there are results for each science window as well as for the overall group.

Science window results include:

- Images: scw/0051004X0010.001/isgri_sky_ima.fits (X=1..5) for each energy range four images are produced: intensity, variance, significance⁷ and residual⁸. To know which type of image and which energy band correspond to a given extension, you can either
- check in the header of the isgri_sky_ima.fits files, or check the first extension of the file an index, summarizing the file content, see Figure 15. More details on index concept are in [1].
 Detected source list: scw/0051004X0010.001/isgr_sky_res.fits (X=1..5)
- list of the sources detected in each energy range with reconstructed RA, DEC, flux, error and significance.

Observation group results include:

- Images: isgri_mosa_ima.fits The structure of this file with the mosaic images is similar to the one of isgri_sky._ima.fits, but instead of residual maps you have the exposure ones.
- Detected source list: isgri_mosa_res.fits list of the sources detected in the mosaic image in each energy range with reconstructed RA, DEC, flux, error and significance.

The single science window and mosaic results are merged in the file isgri_srcl_res.fits. This file contains all the sources from the isgri_catalog.fits plus all the new sources, with the information on their fitted position, fluxes, and detection level.

⁷Note that systematic errors are not included in the calculation of the detection significance (DETSIG).

⁸The intensity image from which all photons attributed to the detected sources were removed



Figure 14: Overview of the IMA level products

You find more details on the structure of the output files in Section C.7.1.

7.1.2 Displaying the Results from the Image Step

It is convenient to look at the images with the help of the ds9 program. First create a region file from the catalog using the **cat2ds9** program. In the example below we create the region file **found.reg** with all the sources found in the mosaic image, **isgri_mosa_res.fits**, for the first energy band (extension number 2), and a region file **cat.reg** with all sources that were in the input catalog **isgri_catalog.fits**.

```
cd $REP_BASE_PROD/obs/isgri_gc
cat2ds9 isgri_mosa_res.fits\[2] found.reg symbol=box color=green
cat2ds9 isgri_catalog.fits\[1] cat.reg symbol=box color=white
```

To see the resulting images:

```
ds9 $REP_BASE_PROD/obs/isgri_gc/isgri_mosa_ima.fits\[2] \
    -region $REP_BASE_PROD/obs/isgri_gc/cat.reg\
    -cmap b -scale sqrt -scale limits 0 60 -zoom 2\
    $REP_BASE_PROD/obs/isgri_gc/isgri_mosa_ima.fits\[4] \
    -region $REP_BASE_PROD/obs/isgri_gc/found.reg\
    -cmap b -scale sqrt -scale limits 0 60 -zoom 2
```

In Figure 16 you see the intensity \$REP_BASE_PROD/obs/isgri_gc/isgri_mosa_ima.fits[2] (left) and significance \$REP_BASE_PROD/obs/isgri_gc/isgri_mosa_ima.fits[4] (right) mosaic images in the 20 - 40 keV energy range. In the left image we have shown all catalog sources (white boxes), and in the right one only the detected ones (green boxes). As you can see, there are three sources labelled as NEW. This example



Figure 15: Sketch of the isgri_sky_ima.fits structure (left), and extract from the Index table (right).

shows that you should be careful with the new sources found by the software, and always make a detailed analysis to verify whether they are real or spurious ones. In a coded mask instrument a true point source will cause secondary lobes, 8 main "ghosts" aligned with the detector edges, at a distance that is a multiple of the mask basic pattern, 10.7 degrees in IBIS/ISGRI case. If the source is detected, its ghosts will be cleaned and will not be included in the mosaic. However, if a source is too weak to be automatically detected in a single Science Window, its ghosts are not cleaned, they can appear in the mosaic image and even be found by the software as new sources.

In the present case, three sources were recognized by the software as new ones. These three sources, NEW_1, NEW_2, and NEW_3, are the ghosts of 4U 1722-30.

There is an easy way to collect from different science windows all the information related to a given source and energy band. In the example below we create a file 4U1700-377_scwlc.fits with all the information on $4U \ 1700-377$ in 20 - 40 keV energy band. The structure of this file is explained in Appendix C.7, Table 53.

In Figure 17 the 20 - 40 keV science window per science window 4U 1700-377 lightcurve is shown.

7.2 Spectral Extraction

It is not possible to extract the spectrum of only the source you are interested in. All sources brighter or compatible with the one you are interested in should be taken into account too. Thus it is strongly recommended, when you deal for the first time with your data, to run the analysis until the IMA2 level, as described in Section 7.1, check the results, and call *ibis_science_analysis* once again to run the spectral extraction part. The description of the algorithm used for spectral extraction is given is Section 12.9.1.

Launch *ibis_science_analysis*, and on the main GUI page change Start Level to **BIN_S**, and End Level to **SPE**. After that press the **ISGRI_SPE_and_LCR** button.

On the screen that appears (see Figure 18), you can specify:

Spectral energy binning:



Figure 16: Intensity (left) and significance (right) mosaic in the 20 – 40 keV energy band

With the help of the parameter IBIS_SI_inEnergyValues you can specify the file (and its extension) describing the desired binning of the response matrix. By default (set with empty line), the latest available file with the rebinned response matrix is used. Currently it is

\$REP_BASE_PROD/ic/ibis/rsp/isgr_rmf_grp_0017.fits[3] with 16 channels in it.

See sections 8.3 and 9.4 if you want to define your own spectral binning.

Background maps:

With the **SCW2_BKG_LisgrBkgDol** parameter you can specify the background map to be used in the spectral and timing analysis. It is recommended to use the same background map as in the imaging case. Default empty value corresponds to the default map (\$REP_BASE_PROD/ic/ibis/bkg/isgr_back_bkg_0005.fits).

Input catalog for Spectral and Lightcurve extraction:

By default a spectrum will be created for all sources listed in the isgri_srcl_res.fits catalog created during the IMA step. The position of the source will be taken from RA_FIN, DEC_FIN columns. Note however that spectral extraction is time consuming, and it is not recommended to include sources that are faint in comparison with the source of interest. It is not recommended to have more than 30 sources in the input catalog, as the software might crash otherwise. Copy isgri_srcl_res.fits to another file, say specat.fits as shown below, leaving only sources with significance higher than 6:

```
cd $REP_BASE_PROD/obs/isgri_gc
fcopy "isgri_srcl_res.fits[ISGR-SRCL-RES][DETSIG >= 6.0]" specat.fits
```

(Note, that there should be spaces around ">=").

As we have discussed in Section 7.1.2, all found new sources are fake ones, so remove them with the help of the fv command, and make the resulting catalog read only (this is important because the file could otherwise get corrupted during the analysis):

chmod -w specat.fits

The resulting specat.fits catalog is shown in Figure 19 (only a few selected columns are shown). The same



Figure 17: 4U 1700-377 science window per science window lightcurve in the 20 - 40 keV energy band

	ISGRI_SPE_and_LCR	
RI Spectral extraction and light curve		<u>_</u>
IBIS_SI_inEnergyValues:	Spectral Energy Binning	browse
SCW2_cat_for_extra.ct:	Input Catalog	browse
SCW2_BKG_I_isgrBkgDol:	Background Map	browse
ILCR_num_e: 4	Number of Energy Bins —	
ILCR_e_min: 20 40 60 100 ILCR_e_max: 40 60 100 200	Energy Boundaries	Parameters for
ILCR_delta_t: 100 🗲	Time Resolution	LightCurve Extraction

Figure 18: Page of IBIS GUI for Spectral and Ligtcurve extraction

		RA_OBJ	DEC_OBJ	DETSIG
elect	ZUA	1E dou	1E dou	1E
nvert		ueg	ueg	
1	0A0 1657-415	2.551996E+02	-4.167306E+01	4.683989E+01
2	4U 1700-377	2.559865E+02	-3.784414E+01	1.708829E+02
3	GX 349+2	2.564354E+02	-3.642278E+01	3.195103E+01
4	н 1705-440	2.572275E+02	-4.410056E+01	1.421409E+01
5	IGR J17252-3616	2.613000E+02	-3.626667E+01	1.358841E+01
6	4U 1722-30	2.618883E+02	-3.080194E+01	9.551276E+00
7	GX 354-0	2.629892E+02	-3.383472E+01	2.222851E+01
8	4U 1735-444	2.647429E+02	-4.445000E+01	6.183074E+00
9	1E 1740.7-2942	2.659779E+02	-2.974528E+01	8.209898E+00

Figure 19: List of sources used for spectral analysis

catalog will be used later for the lightcurve extraction.

In the GUI, set the **SCW2_cat_for_extract** parameter to point to **specat.fits[1]** and press **Ok**, the window disappears and you are back to the main GUI page. Press **Save**, and then **Run**.

7.2.1 Results of the Spectral Extraction

Spectra are produced for each science window:

```
scw/0051004X0010.001/isgri_spectrum.fits (X=1..5)
```

In these files you find the spectra of the desired sources plus the background spectrum. In the header of each data structure, the Name and ID of each source are given, and the first extension of the file $scw/0051004X0010.001/isgri_spectrum.fits[1]$ is an index, summarising the content of all the extensions. With the help of this index you can easily see which extension contains the spectrum of the source you are interested in.

To achieve a better signal to noise ratio, it can be convenient to sum up spectra of a source from different science windows. This can be done with the *spe_pick* tool. In the following example we create a joint spectrum of 4U 1700-377 from all the available science windows (included in the og_ibis.fits file).

```
cd $REP_BASE_PROD/obs/isgri_gc
spe_pick group="og_ibis.fits[1]" source="4U 1700-377" rootname=4U1700
```

As a result two files with spectra of 4U 1700-377 will be created: 4U1700_sum_pha.fits contains the final average spectrum of 4U 1700-377, while 4U1700_single_pha2.fits stores all the original spectra of 4U 1700-377 that were used to create the average one. It is forseen that in the nearest future there will be several ARFs, depending on the data time. In order to deal with the data from different time periods *spe_pick* was updated and creates now a resulting ARF for your particular dataset. This ARF is written to 4U1700_sum_arf.fits and 4U1700_single_arf.fits files. At the moment only one ARF is present and is copied as it is to the files mentioned above.

7.2.2 Displaying the Results of the Spectral Extraction

To analyze the average spectrum with XSPEC do

```
cd $REP_BASE_PROD/obs/isgri_gc
fparkey 0.02 4U1700_sum_pha.fits SYS_ERR add=yes
xspec
    cpd /xw
    data 4U1700_sum_pha.fits
    setpl energy
    ign **-22.5
    ign 150.-**
    plot ldata
    model cutoffpl
    1.7
    33
    3
    fit 100
    plot ldata del
```

The resulted 4U 1700-377 total spectrum is shown in Figure 20. Please note that in OSA 5.0 reconstructed ISGRI spectra are not accurate below 17 keV. ISGRI systematic is believed to be equal to 1%.

In the very same manner, you can sum up the spectra for all the sources of the input catalog, specified in the **SCW2_cat_for_extract** parameter⁹.



Figure 20: Total 4U 1700-377 spectrum.

Usage of *spe_pick* for spectra created in different observation groups is explained in Section 9.

7.3 Lightcurve Extraction

After you have done the spectral extraction, you may want to produce a lightcurve for the same sources. Stay in the working directory **\$REP_BASE_PROD/obs/isgri_gc** and call the *ibis_science_analysis* script again. On the main GUI window (see Figure 12), change both Start Level and End Level to **LCR**, and press the **ISGRI_SPE_and_LCR** button. The GUI shown in Figure 18 will appear.

Note that to extract the lightcurve for a source you need the Pixel Illuminated Fraction (PIF¹⁰) map. Such a map is created during the spectral step that we have just run. This means that at this stage you can extract the lightcurve only of the sources for which a spectrum has been created already. So leave SCW2_cat_for_extract to point to specat.fits.

By default, for each source from the input catalog (specat.fits in our example), four lightcurves (in 20 - 40, 40 - 60, 60 - 100, and 100 - 200 keV energy bands) with 100 sec time bins will be created.

The lightcurve extraction is performed by building shadowgrams for each time and energy bin. Hence this step is time and space consuming. Note that due to CFITSIO limitations, the product of number of energy bins by number of time bins in a science window should be less than 250.

Press Ok, the window will disappear and you are back to the main GUI page. Press Save, and then Run.

⁹To read a given extension in XSPEC you have to specify it in curly brackets: "data pha.fits{3}"

 $^{^{10}}$ PIF is a number between 0 and 1, which expresses the theoretical degree of illumination of each detector pixel for a given source in the sky.

7.3.1 Results of the Lightcurve Extraction

Lightcurves are produced for each science window:

scw/0051004X0010.001/isgri_lcr.fits (X=1..5)

To create a merged lightcurve in the 20 - 40 keV energy range (i.e. to store in one file all the available lightcurves of a given source) use lc_pick :

lc_pick source='4U 1700-377' attach=n \
 group=og_ibis.fits+1 lc=4U1700-377.fits emin="20" lcselect='E_MAX==40'

The result is written to **4U1700-377.fits**. It is possible to create more than one merged lightcurve at a time if you specify a list of desired minimum energies with the parameter **emin**. If you have more than one energy range with the same minimal energy, then you should use the parameter **lcselect** to define the unique energy band, otherwise the **lcselect** parameter can be omitted.

7.3.2 Displaying the Results of the Lightcurve Extraction

To see the source lightcurve, you should plot the column RATE with error ERROR versus column TIME, whereas to see the background lightcurve, you should plot column BACKV with error BACKE versus column TIME.

To display the resulting lightcurve it is convenient to use the *lcurve* program from the FTOOLS package:

As a result, the 4U1700-377_lc.ps file was produced and is shown in Figure 21.



Figure 21: 4U1700-377.fits lightcurve in 20 – 40 keV energy range with 100 sec binning.

8 More on ISGRI relevant parameters

In the first part of the Cookbook (Section 7), the default values of the parameters were used. Now, it is time to discuss in more details the usage of the main parameters. The full list of parameters along with short explanations is given in Table 61, Appendix D.

8.1 How to choose the start and end level for the analysis

When you launch an analysis you have to decide which levels to perform (see Section 5 for a description of the different levels). This choice is done with the parameters startLevel and endLevel. We recommend to use startLevel="COR" in any case, even if you are using revision 2 data: in OSA 5, you have more accurate energy correction than was available at the time of the revision 2 data archiving. Therefore, it is worth redoing this step.

Once you have specified startLevel and endLevel, all the steps listed in the GENERAL_levelList parameters between these two levels (both included) will be performed. For instance, in section 7.1 you used:

```
startLevel="COR" endLevel="IMA2"
GENERAL_levelList="COR,GTI,DEAD,BIN_I,BKG_I,CAT_I,IMA,IMA2,BIN_S,SPE,LCR,COMP,CLEAN"
```

The levels from COR to IMA2 (included) listed in GENERAL_levelList were performed.

You can choose what to run according to what kind of output you need. Basically:

• If you are interested only in imaging results set

```
startLevel="COR" endLevel="IMA2"
```

with the complete list of GENERAL_levelList given above, as done in section 7.1.

• If you have already a catalog of sources for which to extract the spectra (e.g.specat.fits¹¹) and you are not interested in the imaging results anymore you can skip all the imaging related levels and do:

¹¹specat.fits has to be of the ISGR-SRCL-RES type, meaning that it has to be created by the imaging step (with OBS1_DoPart2=1, since this file is created at mosaic stage). See section 7.2 to learn how to go from the imaging result file isgri_srcl_res.fits to your specat.fits, to avoid software crash.

startLevel="COR" endLevel="SPE" GENERAL_levelList="COR,GTI,DEAD,BIN_S,SPE"

Remember that if you do not do the imaging part then isgri_srcl_res.fits is not created! This file is the default one that is used for spectral extraction (7.2). Do not forget to specify the catalog you have built for this purpose setting the SCW2_cat_for_extract equal to specat.fits[1].

• Likewise, if you are interested only in lightcurve results you should create PIFs for all source of interest with *ii_pif* program (see Section 9.10 for an example), and afterwards you may use:

startLevel="COR" endLevel="LCR" GENERAL_levelList="COR,GTI,DEAD,BIN_S,LCR"

again with SCW2_cat_for_extract equal to specat.fits[1].

• If you are familiar with the sources of your field (i.e. you have already run the imaging part, for instance) and you want to extract images, spectra and lighturves in one go, set SCW2_cat_for_extract equal to specat.fits[1] and run your analysis with

startLevel="COR" endLevel="LCR"

and the complete list of levels in GENERAL_levelList.

8.2 Imaging

Source detection is an important and delicate process as the background and all the sources (especially the bright ones) in the field of view will have an impact on it. Because of this intrinsic complexity of the instrument, there are different options as far as source search and background correction are concerned. The choice of the best method to use for a given case may require a bit of iteration on your side.

In the next two paragraphs we describe the parameters involved in the source search, along with some practical advices. The single Science Window case is treated in Section 8.2.1, while the mosaic image case in Section 8.2.2. Comments on the background subtraction issue are given in Section 8.2.3 while a set of miscellanea on imaging parameters is given in Section 8.2.4.

8.2.1 How to choose the source search method in the Science Window analysis

Basically the idea is that you can either let the software look for the excesses in the field or you can tell the software to look for sources from a given catalog. This catalog can be either the one provided by ISDC (CAT_refCat="\$ISDC_REF_CAT"), from which you can extract e.g. only the sources that have been detected by ISGRI ("\$ISDC_REF_CAT[ISGRI_FLAG==1]"), or a catalog made on your own (see section 9.6 to create a catalog on your own). The number of sources that are detected in an image is important because detecting a source means also removing its ghosts from the images ¹² (see section 7.1).

The source search method is defined with the parameters OBS1_SearchMode and OBS1_ToSearch. These parameters allow you the following possibilities:

• $OBS1_SearchMode = 0$

This mode was developed for testing purposes and is not recommended for scientific use.

• OBS1_SearchMode = 1

the software will look for all the sources of the input catalog that are in the field of view (regardless of their detection level). If the source gives a positive detection, then the relevant results are saved in the isgri_sky_res.fits file, but see also discussion on OBS1_NegModels in Section 8.2.4.

This mode can be useful in case your Science Window images are noisy and many fake new sources are detected. Searching only the sources from a given catalog will avoid having ghost cleaning for fake sources.

 $^{^{12}}$ Note that the run time is proportional to the number of sources detected.

• OBS1_SearchMode = 2

The software will look for the K brightest sources in the field, where K = OBS1_ToSearch. They can be either known sources (more significant than OBS1_MinCatSouSnr) or new sources (more significant than OBS1_MinNewSouSnr).

This mode is useful if you are interested in having a first glance at your field, if you are interested in bright sources only, or if you run your analysis on large data sets. In fact, in these cases you would not like to spend most of your time collecting information on weak sources that are not detected in a single Science Window. The drawback of this method is that these undetected weak sources can produce ghosts in the mosaic image. We recommend you to start the analysis with OBS1_SearchMode = 2 with a high OBS1_ToSearch value (around 20, especially for the Galactic plane or Centre crowded regions), see the sources that are detected and prepare your own catalog of detected sources (see section 9.6). Then launch again the analysis with OBS1_SearchMode = 3 - described below - and give your catalog as input for the analysis (CAT_refCat=''usrcat.fits[1]'').

• OBS1_SearchMode = 3

The software will look for all the catalog sources (regardless of their detection level) *plus* for K significant excesses more significant than <code>OBS1_MinNewSouSnr</code>, where <code>K = OBS1_ToSearch</code>. All these sources will be saved in the final <code>isgri_sky_res.fits</code> file if their detection significance is positive, but see also discussion on OBS1_NegModels in Section 8.2.4. This mode is very efficient in cleaning the ghosts of all the sources in the field of view. It is of course more time consuming since a model is created and deconvolved for each given source.

We suggest that you should use this mode either with your own catalog (built from the previous run with

OBS1_SearchMode = 2, see above) or, if no personalised catalog is available, using the ISDC catalog with only the sources that have been detected by ISGRI (\$ISDC_REF_CAT[ISGRI_FLAG==1], see above). Any bright source out of the catalog (new or forgotten!) will be detected as one of the K new sources.

8.2.2 Parameters related to the mosaic step

The OBS1_SearchMode parameter concerns only images at a Science Window level. In the mosaic image the software always looks for K=OBS1_ToSearch sources (more significant than OBS1_MinCatSouSnr for catalog sources, or OBS1_MinNewSouSnr for the new sources), regardless the real number of sources in the field of view. Thus, if you are working with a crowded part of the sky (mainly Galactic plane and Centre) you should use a high OBS1_ToSearch number (at least 20).

There are two ways to calculate the source flux in the mosaic given the original Science Window information:

• OBS1_PixSpread = 0

The whole input (Science Window) pixel count is put into one output (mosaic) pixel, no "spread" is done. This method optimises the flux and signal-to-noise ratio (SNR) evaluation but can sometimes give some undesired effects such as double source peaks.

• OBS1_PixSpread = 1 (default value)

The input pixel count is spread between different output map pixels. This method is better for the source position estimate but is less accurate than the previous method for source flux estimation. SNR in the new spread mosaic will be lower than in the previous case.

It is possible to decide whether you want the mosaic to be created along with the Science Window image reconstruction, or if you prefer it to be created later on. The mosaic production is controlled by the OBS1_DoPart2 parameter:

- OBS1_DoPart2 = 0 The mosaic step is not performed and the imaging part ends after the Science Window based part.
- OBS1_DoPart2 = 1 (default value) The mosaic is done after the Science Window image reconstruction, in the same run.

• OBS1_DoPart2 = 2

The mosaic is performed for images obtained from different, already existing, runs (see 9.3.1).

8.2.3 Background Subtraction

In the current OSA you have background maps provided by the ISGRI team. These maps are created on the base of observations containing only faint sources in their fields of view. Currently, a single set of maps for the complete mission is available, meaning that background variations are not taken into account. In most cases background subtraction improves the result, nevertheless we cannot state *a priori* in which cases the background subtraction will improve (or worsen) the results. The best approach is to verify in the images if the background subtraction is working correctly (giving a better image) before deciding what results to use and before applying it also to the spectral and lightcurve extraction levels.

Background subtraction can be tuned with the help of the SCW1_BKG_I_isgrBkgDol parameter. If it is left empty then the background map is taken automatically from the IC files. To have no background subtraction, set SCW1_BKG_I_isgrBkgDol=''-''.

As a general guideline, for short timescale (e.g. 1 Science Window), the background features should be less than the statistical uncertainties so that background removal is not expected to change the result by much. On longer time scales (mosaic image), things can be different, especially in the case of a Staring observation in which background features can accumulate and become important.

8.2.4 Miscellanea on Imaging

- With the parameter OBS1_ExtenType (to find it on the GUI pages press the hidden button, and go to the IMA page) you can choose whether to keep the <u>Residual and/or the true Exposure</u> image in the isgri_sky_ima.fits file. With the default, "0", 4 maps will be created per energy range (Intensity, Variance, Significance and Residuals). With OBS1_ExtenType = 1 true exposure maps will be written instead of residual images at the 4*n extension of isgri_sky_ima.fits. OBS1_ExtenType = 2 will add the true Exposure maps in an additional extension. OBS1_ExtenType = 3 will add single true Exposure map as the last extension.
- If you are interested in running the analysis in more than 10 energy bands please see Section 9.4.
- You can decide if you want to let the software find the best fit position of a source (in the vicinity of the one in the catalog, default OBS1_SouFit=0), or if you would prefer the catalog position (for all sources from the input catalog!) to be used for the flux determination (OBS1_SouFit=1). If OBS1_SouFit=1, then the position of a new source is always attributed to the pixel center, and for catalog sources the catalog position is used for SearchMode=1,3, or the pixel center for SearchMode=2. Note that OBS1_SouFit=1 fixes all the source positions only in the individual Science Window analysis. In the mosaic, all source positions are always fitted regardless of the value of OBS1_SouFit. This fitted values are written to RA_FIN and DEC_FIN columns of isgri-srcl-res.fits, and are later used in spectral and lightcurve extraction steps as a source position. To fix the positions of only several sources, set for them ISGRI_FLAG=2 in the input catalog.
- For OBS1_SearchMode = 1 or 3 (where you are forcing a catalog extraction) ghosts can be cleaned even from the known sources that happen to have a negative peak. This is done using OBS1_NegModels = 1 while with OBS1_NegModels empty or equal to "0" (the default) the deconvolved images will be cleaned from positive source ghosts only.

8.3 Spectral and Timing Analysis

8.3.1 Spectral Energy Binning

With the help of the IBIS_SI_inEnergyValues parameter you can specify the desired binning of the response matrix. See Section 9.4 for the detailed discussion on how to rebin the original response matrix.

In case you using a response matrix rebinned on your own you should set

SCW2_ISPE_idx_isgrResp="your_rebinned_matrix.fits". Warning: This parameter is a hidden one in OSA 5.1. Thus you should either provide the value of this parameter during the script launch

```
ibis_science_analysis SCW2_ISPE_idx_isgrResp="your_rebinned_matrix.fits"
```

or press "hidden"-button and set this parameter on the SPE page.

8.3.2 Background Subtraction

In spectral and timing analysis you can choose (with the help of the parameter SCW2_BKG_I_isgrBkgDol) whether you want to subtract the instrumental background or not, similarly to the imaging case. We recommend you to use the option that gives the best results at the imaging level (smoother image, no clear patterns etc). See 8.2.3 for more information.

8.3.3 Input catalog

The single Science Window and mosaic imaging results are merged in the file isgri_srcl_res.fits. By default a spectrum and a lightcurve will be created for all sources listed in this file. Spectral/lightcurve extraction is time consuming and we recommend you to create an input catalog for spectral and timing extraction, as discussed in Section 7.2. The spectrum and lightcurve extraction tools use the source position saved in the RA_FIN and DEC_FIN columns. These are the values computed during the imaging step, thus in case you want to use catalog source positions you have to modify them manually. In case you prefer the catalog positions for all catalog sources, you can remove RA_FIN and DEC_FIN columns (RA_OBJ and DEC_OBJ are used then) with the following commands:

```
chmod +w specat.fits
fdelcol infile=specat.fits+1 colname="RA_FIN" confirm=no proceed=yes
fdelcol infile=specat.fits+1 colname="DEC_FIN" confirm=no proceed=yes
chmod -w specat.fits
```

Important: make the file with the catalog **read only**, otherwise it may be corrupted in the course of the analysis!

9 Useful recipes for the ISGRI data analysis

In this Section we give a number of recipes that can be useful in the analysis of ISGRI data.

9.1 Rerunning the Analysis

Read this if you would like to redo part of your analysis, e.g. if your run has crashed, or if you want to change some parameters.

In case you want to re-run the analysis with different parameters, run og_create but this time with a different "ogid" parameter. This will create a new tree under obs/ogid where all the new results will be stored. If the pipeline has crashed¹³, in general it is safer to restart your analysis from scratch removing the obs/ogid directory and restarting from the og_create step.

In any case, we give below a set of recipes that can be useful.

Because of the group concept you cannot just delete the result you do not like and restart the pipeline. All results that were produced in the course of the analysis are linked to the group, and should be detached before you relaunch the script. To do this you can use the *og_clean* program, that will clean an Observation Group up to the level specified with parameter **endLeve1**. All data structures with a level equal or prior to **endLeve1** will be kept, while the data structure with a later level will be erased. For example, to run the spectral extraction (SPE level) you should clean from group whatever comes after the BIN_S level, as this is the level immediately preceeding the spectral one (see Fig.12).

```
og_clean ogDOL="og_ibis.fits[1]" endLevel="BIN_S"
```

If og_clean fails it could be due to the fact that the group was corrupted. You should try to fix it with *dal_clean* program

dal_clean inDOL="og_ibis.fits[1]" checkExt="1" backPtrs="1" checkSum="1"

and launch *og_clean* only afterwards.

Unfortunately the current version of og_{clean} is very slow. In some cases, if you know exactly which data structure has to be detached it would be much faster to launch *dal_detach*. For example if you run your analysis till SPE, or LCR level, and would like to produce a mosaic image afterwards, you do not have to clean the group, deleting all your results, but just have to detach ISGRI-SRCL-RES data structure (note that with the option "delete=y" all files with this data structure will be deleted):

dal_detach og_ibis.fits\[1] pattern=ISGR-SRCL-RES delete=y

9.2 Make your own Good Time Intervals

Read this if you are interested to select photons arrived at a particular time period (e.g. in analysis of flares, or for phase resolved spectroscopy).

You should define Good Time Intervals (GTIs) with the help of the *gti_user* program. To create a GTI for IBIS starting on IJD 1322.68 and lasting 1 minute (do not forget to convert it into days!) give the command:

 $^{^{13}{\}rm If}$ any executable crashes then it terminates with non zero status. The meaning of the status value can be found at <code>http://isdc.unige.ch/index.cgi?Soft+errors</code>

Then in the main page of the GUI, Fig. 12, you should set SCW1_GTI_gtiUserI="user_gti.fits[1]", and specify the time format that was used in this file, in the example above SCW1_GTI_TimeFormat="IJD". More details on *gti_user* are in the Data Analysis section of [1].

9.3 Combining results from different observation groups

Read this if you have a set of science windows belonging to different runs for which you have already built images, spectra or lightcurves and want to combine the results.

Section 9.3.1 explains how to combine all the existing images in a final mosaic, while Section 9.3.2 shows how to merge different lightcurves and spectra.

9.3.1 Creating a mosaic from different observation groups

Suppose you want to analyse 5 Science Windows and are not interested in the final mosaic. You create the group with og_create and then you launch the analysis till the imaging step but without the mosaic step (i.e. from COR till IMA with OBS1_DoPart2 = 0, see 8.2.2). An image is created per Science Window but you do not have the overall final mosaic. If you then change your mind and decide that you want the mosaic, all you have to do is move in the working directory \$REP_BASE_PROD/obs/xxx and relaunch the ibis_science_analysis command with startLevel="IMA", endLevel="IMA" and OBS1_DoPart2=1.

This is relatively simple because all the Science Windows belong to the same group, so combining the results is trivial. But if this is not the case, i.e. if you have run different sets of analysis, each one with its own og_create command, then you need to make some intermediate steps ¹⁴. You basically need to create a file that points to all the Science Windows you want to co-add. We call this file an "index". To create an index make a list of the Science Window groups you want to combine and save it as, e.g., dols.txt. To ensure a proper work of the software give the full path, i.e. your file should look like

```
/WORKING-DIR/obs/GROUP1/scw/011901070010.001/swg_ibis.fits[1]
/WORKING-DIR/obs/GRPUP1/scw/011901080010.001/swg_ibis.fits[1]
/WORKING-DIR/obs/GROUP2/scw/012000360010.001/swg_ibis.fits[1]
/WORKING-DIR/obs/GROUP2/scw/012000370010.001/swg_ibis.fits[1]
```

The first 2 files belong to a run with the og_create parameter "ogid" equal to "GROUP1" while the latter two to a run with "ogid" equal to "GROUP2" (remember the "[1]"!!!). /WORKING-DIR/ has to be the extensive name of \$REP_BASE_PROD. WARNING: make sure that obs/*/scw/*.001/isgri_sky_ima.fits files exist in all the Science Windows you mention, otherwise the merging will not work. Then give the command that actually builds the fits file from the ASCII file:

```
cp dols.txt $REP_BASE_PROD/obs/GROUP1/
cd $REP_BASE_PROD/obs/GROUP1/
txt2idx element="dols.txt" index="index_comb.fits[1]"
```

The file "index_comb.fits" is created and you can look at it with **fv**. In the first extension you have 4 rows, each row has the link to a given swg_ibis.fits file. What you need to do now it to let the software know that it has to use this particular set of Science Windows for the analysis. You do this by pointing **og_ibis.fits** to this file. This has to be done because no matter what level of the scientific analysis you are performing, the software will analyse the Science Windows pointed to by **og_ibis.fits**. Create an ASCII file **\$REP_BASE_PROD/obs/GROUP1/modfile** with one line in it:

 $^{^{14}}$ Please note that the images that you intend to merge MUST have the same energy boundaries i.e. all the ScW maps must have been analysed in the same way. You cannot merge a 20–40 keV map with a 30–50 keV one.

1 index_comb.fits

and then replace the first row of og_ibis.fits[1] automatically with:

```
cd $REP_BASE_PROD/obs/GROUP1/
fmodtab og_ibis.fits+1 MEMBER_LOCATION modfile
fdelrow og_ibis.fits+1 2 1 N Y
rm isgri_catalog.fits
```

With the fdelrow command you detach from the first extension of og_ibis.fits row number "2" (= the first row to be deleted, number of rows to be deleted = "1"), you were shown no keyword values, "N", and you agreed to proceed, "Y". This second row contained the catalog that was created in previous run during CAT_I step, and contain only sources that were in the field of view of science windows from GROUP1. To have a common catalog for GROUP1 and GROUP2 it would be necessary to rerun CAT_I step for the group referring to all science windows of interest.

You may check with **fv** that actually the first row of og_ibis.fits is indeed "index_comb.fits".

Before running the mosaic step please be aware that:

• If you had previously run the mosaic step you will see that **og_ibis.fits** points to the mosaic output (after you have deleted a reference to isgri_catalog.fits rows 2 to 4 of the first extension are isgri_srcl_res.fits, isgri_mosa_ima.fits and isgri_mosa_res.fits). This will interfere with the new mosaic you are about to launch, thus you have to detach these former mosaic results from the group and delete (or rename) them:

cd \$REP_BASE_PROD/obs/GROUP1
fdelrow og_ibis.fits+1 2 3 N Y
rm isgri_mosa_ima.fits
rm isgri_mosa_res.fits
rm isgri_srcl_res.fits

• If you had previously run the spectral and lightcurve steps you have to detach the ISGRI-SRCL-RES data structure from all your Science Window groups (swg_ibis.fits[1]). That is to each Science Window (\$scw) group located in the \$REP_BASE_PROD/obs/\$dir directory, you should do the following command:

cd \$REP_BASE_PROD/obs dal_detach object="\$dir/scw/\$scw.001/swg_ibis.fits[1]" pattern=ISGR-SRCL-RES delete=n

At this point you are ready to launch the analysis and to create a joint mosaic:

```
cd $REP_BASE_PROD/obs/GROUP1
ibis_science_analysis ogDOL="og_ibis.fits[1]" \
startLevel="CAT_I" endLevel="IMA" OBS1_DoPart2=2
```

See 8.2.2 for a description of the main parameters of the mosaic step. The above command is valid in case you have just run the ScW analysis part so that you indeed keep the same energy boundaries. Otherwise, if you parameter file has changes in between, you should add in the above the definition of IBIS_II_ChanNum, IBIS_II_E_band_min and IBIS_II_E_band_max according to the boundaries of the single ScW maps you want to merge.

9.3.2 Combining spectra and lightcurves from different observation groups

In Sections 7.2 and 7.3 you have seen how to merge lightcurves and spectra from different Science Windows belonging to the *same* group. In that case the file **og_ibis.fits** points to all the Science Windows and it is possible to launch the collecting/merging tools lc_{pick} and spe_{pick} directly on the group (group=og_ibis.fits+1).

If you have Science Windows belonging to *different* groups, you need an intermediate step. Basically you need to create a file (an index) that points to all the Science Windows you want to co-add, similarly to the case seen in section 9.3.1. Then this file will be given as input via the "group" parameter at the place of **og_ibis.fits**.

To create the index make a list of the Science Window groups you want to combine and save it as, e.g., **dols.txt** under \$REP_BASE_PROD/obs/. To ensure a proper work of the software give the full path, i.e. your file should look like

```
/WORKING-DIR/obs/GROUP1/scw/011901070010.001/swg_ibis.fits[1]
/WORKING-DIR/obs/GRPUP1/scw/011901080010.001/swg_ibis.fits[1]
/WORKING-DIR/obs/GROUP2/scw/012000360010.001/swg_ibis.fits[1]
/WORKING-DIR/obs/GROUP2/scw/012000370010.001/swg_ibis.fits[1]
```

The first 2 files belong to a run with the og_create parameter "ogid" equal to "GROUP1" while the latter two to a run with "ogid" equal to "GROUP2" (remember the "[1]"!!!). /WORKING-DIR/ has to be the extensive name of \$REP_BASE_PROD. WARNING: make sure that the lightcurve and spectra result files exist for each Science Window you want to co-add (isgri_lcr.fits and isgri_spectrum.fits files).

Then give the command that actually builds the fits file from the ASCII file:

```
cd $REP_BASE_PROD/obs/
txt2idx element="dols.txt" index="index_comb_2.fits[1]"
```

Then run *lc_pick* and *spe_pick* putting the parameter "group" equal to "index_comb_2.fits[1]":

```
cd $REP_BASE_PROD/obs/
lc_pick source='GRS 1758-258' attach=n \
group=index_comb_2.fits+1 lc=GRS1758.lc.fits emin="20" \
lcselect='E_MAX==40' instrument="ISGRI"
```

GRS1758.lc.fits contains the merged lightcurve of GRS 1758-258 in the 20–40 keV band (energy range that of course has to exist in the isgri_lcr.fits original files!).

GRS1758_sum_pha.fits (the combined spectrum of GRS 1758-258) and GRS1758_single_pha2.fits (a file with the four spectra of the initial four Science Windows collected together) are created. It is forseen that in the nearest future there will be several ARFs, depending on the data time. In order to deal with the data from different time periods *spe_pick* was updated and creates now a resulted ARF for your particular dataset. This ARF is written to GRS1758_sum_arf.fits and GRS1758_single_pha2.fits files. The names of the response and ancrfile are inserted in the keyword of the final files, so that they are automatically recognised by XSPEC.

9.4 Rebinning the Response Matrix

Read this if you want to use a spectral binning different from the default one. You will also learn how to rebin the response matrix to extract images in more than 10 energy ranges.

The file **\$REP_BASE_PROD/ic/ibis/rsp/isgr_rmf_grp_0016.fits** contains the latest full response matrix with 2048 channels. With the help of the FTOOLS 5.3 program **rbnrmf**¹⁵ you can rebin this matrix according to your needs. The default rebinned matrix is **\$REP_BASE_PROD/ic/ibis/rsp/isgr_rmf_grp_0017.fits**

¹⁵please note that it will not work with the earlier versions of the program

and has 16 energy bins in its third extension. The optimum way to rebin the matrix depends on the source and on what you are interested to study, so there is not a general best way to rebin it. Nevertheless, it is reasonable not to spend time on a fine energy binning where the source is too weak and background contaminated as well as it is better to avoid too narrow energy bins for weak sources: extracting a spectrum in wider energy bins from the very beginning is better than extracting it in too fine bins, being background contaminated, and then rebinning the spectrum afterwards.

To use **rbnrmf** you should create an ASCII file with three columns, representing minimal and maximal channels, and the compression factor. In the example below we create the file **new_bin.txt** in order to apply a compression factor of 16 to channels 16 - 415 (20.6592 - 212.1392 keV), and ignore all the others. This will lead to [(maximal_channel-minimal_channel+1)/compression factor] number of final bins, i.e. in this example to 25 bins.

0 15 -1 16 415 16 416 2047 -1

To rebin the matrix give the command:

```
rbnrmf infile="$REP_BASE_PROD/ic/ibis/rsp/isgr_rmf_grp_0016.fits" \
outfile="new-rmf.fits" binfile="new_bin.txt"
```

As a result of this, you now have the rebinned response matrix new-rmf.fits. With the help of fv you can see that the rebinned matrix is saved in the third extension of the file. For the successful analysis you should make new rebinned matrix read-only:

chmod -w new-rmf.fits

To tell the software to use this newly created matrix set in your analysis IBIS_SI_inEnergyValues="/PATH-TO-THE-FILE/new-rmf.fits[3]" otherwise the default 16 channels one will be used (corresponding to an empty value of IBIS_SI_inEnergyValues).

You should also set SCW2_ISPE_idx_isgrResp="new-rmf.fits". Warning: This parameter is a hidden one in OSA 5.1. Thus you should either provide the value of this parameter during the script launch

ibis_science_analysis SCW2_ISPE_idx_isgrResp="new-rmf.fits"

or press "hidden"-button and set this parameter on the SPE page.

9.4.1 Extracting images in more than 10 energy ranges

If you want to create images in more than ten energy bands, you have to create a rebinned matrix **new-rmf.fits** as shown in the text just above. Then, to create images in the energy ranges defined in the third extension of this file set

IBIS_II_ChanNum=-1

IBIS_II_inEnergyValues="/PATH-TO-THE-FILE/new-rmf.fits[3]".

The first parameter instructs the program to use the energy boundaries specified in the second parameter.

9.5 Some tricks on saving disk space and CPU time.

To gain time and space it is possible to create the rebinned correction and background maps rebinned_*.fits only once and then give them as an input for the rest.

Analyse one science window from COR to SPE, copy the maps somewhere and make them read only. Then launch the analysis for other science windows specifying the following parameters in your *ibis_science_analysis* call:

rebinned_corrDol_ima="/your_path/rebinned_corr_ima.fits[1]"
rebinned_corrDol_spe="/your_path/rebinned_corr_spe.fits[1]"
rebinned_backDol_ima="/your_path/rebinned_back_ima.fits[1]"
rebinned_backDol_spe="/your_path/rebinned_back_spe.fits[1]"

IMPORTANT: rebinned maps are created for a given set of energies only, so these have to be kept the same.

9.6 Create your own catalog

Read this if you are familiar with the sources in your field of view and you want to build your own catalog for the Imaging step.

As already discussed in Section 8.2.1, during the imaging step (IMA) it can be very useful to use your own input catalog. The easiest way to build it is to modify the general catalog with the help of the fcopy program of FTOOLS. For example, if you want to create a catalog (usrcat.fits) with only three sources, Crab, 3C111, and XPer do the following:

cd \$REP_BASE_PROD
fcopy "\$ISDC_REF_CAT[NAME == 'Crab'|| NAME == '3C 111'|| NAME == 'X Per']" usrcat.fits
chmod -w usrcat.fits

If you want to add a new source, not specified in the general reference catalog, add a line to usrcat.fits with the help of the fv tool, and fill in the Source_ID, NAME, RA_OBJ and DEC_OBJ columns. As Source_ID you can choose whatever you like provided that this identifier is unique.

NOTE: usrcat.fits keeps the same format of the general reference catalog from which it was built, i.e. it has the extension of the type "GNRL-REFR-CAT". This is the correct extension to be used as input for the Imaging step. In Section 7.2 you have been shown how to build a catalogue for the lightcurve (LCR) and spectral (SPE) step. In that case the catalogue you obtained, specat.fits, was of the type of "ISGR-SRCL-RES" which is the correct format for the LCR and SPE steps. The two formats cannot be switched, meaning that you cannot use a "GNRL-REFR-CAT" type catalog for SPE and LCR or an "ISGR-SRCL-RES" one for IMA.

9.7 Alternative Spectral Extraction from the Mosaic

Read this if you want to extract the average spectrum of a source from a set of mosaic images in different energy bands.

It is possible to extract the spectrum of a source from a set of mosaic images (in several large energy bands). In general this method gives results similar to the Science Window average spectrum you obtain from *spe_pick* (see Sections 7.2 and 9.3.2). Once you have the spectrum, you will need the corresponding rebinned matrix to be able to analyse it in XSPEC. We suggest to build such a matrix before launching OSA as shown in Section 9.4 and then extract images with the use of this same matrix as shown in Section 9.4.1. See Sections 7.1 and 8.2.2 to learn how to create mosaic images (for a better flux estimate use OBS1_PixSpread=0).

Once you have the set of mosaic images in the desired energy bands, to extract the spectrum from the mosaics (attached to the og_ibis.fits file!) at the position of 4U 1700-377 (ra=255.9865, dec=-37.84414) use *mosaic_spec*:

```
cd $REP_BASE_PROD/obs/xxx
mosaic_spec DOL_inp="og_ibis.fits[1]" DOL_out="og_ibis.fits[1]"\
EXTNAME="ISGR-MOSA-IMA" DOL_spec="4U1700_mosa_pha.fits(ISGR-PHA1-SPE.tpl)"\
ra=255.9865 dec=-37.84414 size=4
```

Note that if the significance of the point is less than five, then *mosaic_spec* will assign a non-zero value to its quality, and the point will be not used by XSPEC. If you are interested to see this point do change its quality value to zero manually. You find more details on *mosaic_spec* in Section 12.12.1.

9.8 Barycentrisation

Read this if you want to apply barycentrisation to an extracted lightcurve.

The tool making such a correction is called *barycent*. Note that it re-writes the input file, so it may be worth to copy the original file first. Below we show an example for 4U 1700-377, Science Window number 011800900010 and observation group identification ("ogid" parameter of og_create) equal to "OGID":

```
cd $REP_BASE_PROD/obs/OGID
cp scw/011800900010.001/isgri_lcr.fits isgri_lcr_011800900010_bar.fits
barycent inCOL=TIME outCOL=TIME \
outDOL="isgri_lcr_011800900010_bar.fits[2]" \
inDOL=', auxDOL="../../aux/adp/0118.001/orbit_historic.fits[1]" \
raOBJ=255.9865 decOBJ=-37.84414
```

In the example we have applied barycentrisation to the first energy range of 4U 1700-377 that happens to be in the extension number 2 ("[2]"). Thus the extension you give as "outDOL" depends on the source and energy range you need to correct. The overview of the content of each extension is in the input file isgri_lcr.fits[1], i.e. the first extension (GROUPING).

Note that *barycent* tool does not use group concept, and is used directly with the lightcurve. But for it successful work **the REF_BASE_PROD** variable should be set and point to the place in which there is an ic directory, e.g. directory_of_ic_files_installation__/ic.

See more details on *barycent* tool in the Data Analysis section of [1].

9.9 Alternative Timing Analysis

Read this if you are interested in extracting lightcurves on smaller time bins (i.e. up to about 0.1 sec) than the ones allowed by the standard tool.

9.9.1 ii_light

The standard lightcurve extraction tool $(ii_lc_extract)$ called within the general analysis script builds shadowgrams for each requested time and energy bin. Thus this program is quite time consuming and it is not recommended to use it with time bin less than about 60 seconds. Besides, be aware that the higher the number of total bins is, the higher is the time and space needed.

The ii_light tool (not called in the analysis script but available as a stand alone tool) uses a different algorithm: the lightcurve is extracted not via the shadowgram creation but with the use of the Pixel Illuminated Fraction (PIF). This allows to extract lightcurves up to a time bin of about 0.1 sec. This is not the official lightcurve extraction tool and should be used mainly to check relative variability of bright sources within a given Science Window, rather than for a long term absolute flux estimate. Note that to be able to run ii_light you need to have ROOT available.

9.9.2 Run ii_light

To be able to use *ii_light* you have to run the standard analysis until the SPE level (startLevel="COR" endLevel="SPE"). You need to run the IMA level with the *same* energy ranges in which you are going to extract the lightcurves. This is due to the fact that the IMA step will produce correction and background maps (rebinned_corr_ima.fits and rebinned_back_ima.fits) that are energy dependent and that will be put as input to *ii_light*. Alternatively you can use *ii_map_rebin* to produce these maps inthe desired energy bands. You need to run the SPE level because during this level the PIF, needed as input to *ii_light*, is created. Alternatively you can create PIF with *ii_pif* program, as described in Section 9.10.

Once you have run the script until the SPE level (included) you are ready to launch the stand alone lightcurve extraction tool. *ii_light* works on the Science Window level only, meaning that you have to launch it once per Science Window. It does not change the structure of the Science Window group, so if you would like to compare the results of *ii_light* and *ii_lc_extract* it is worth to run the analysis until SPE, use *ii_light* as shown below, then go back to \$REP_BASE_PROD/obs/OGID and relaunch the analysis from LCR to LCR in the standard way. The output lightcurve files of the standard software are automatically called isgri_lcr.fits.

Alternatively, if you have first run the standard lightcurve extraction (i.e. the LCR level) you can still run ii_light following the instructions below.

We give an example of the shell script that launches *ii_light* for all Science Windows of your group OGID from a particular revolution, (0051 in example below).

```
cd $REP_BASE_PROD/obs/OGID
setenv run 0051
foreach file (scw/$run*/swg_ibis.fits)
echo $file
ii_light \
inSwg=$file \
num_e=4 e_min="20 40 60 100" e_max="40 60 100 200" delta_t=10 \
outLC="$file:h/lcr.fits(ISGR-SRC.-LCR-IDX.tpl)" \
GTIname="MERGED_ISGRI" \
corrDol="rebinned_corr_ima.fits" \
context="../../scw/$run/rev.001/idx/isgri_context_index.fits[1]" \
backDol="rebinned_back_ima.fits"
| tee out.log
end
```

As a result of this script you will have in each scw/ run^* directory a file lcr.fits with 4 lightcurves (in 20 – 40, 40 – 60, 60 – 100, 100 – 200 keV energy bands) with 10 seconds binning (delta_t=10). The sources for which a lightcurve is extracted are the ones for which a PIF was created at the SPE level, i.e. specified in **SCW2_cat_for_extract** (see Section 7.2.)

9.9.3 Merge the ii_light results from different Science Windows

If you want to merge in one file all the lightcurves provided by *ii_light* for a given source, you can use the *lc_pick* tool as shown in Section 9.3.2.

 ii_light is a stand alone tool and does not update the proper files with its results. Even if you have run both the lightcurve tools (in whichever order), only the standard one called within the script (LCR step, $ii_lc_extract$) will have updated the proper files with its results (i.e. the swg_ibis.fits file). Thus if you run lc_pick to collect the lightcurves, only the standard ones will be selected. If you want to collect the results of ii_light, then you have to create an index file that points to all the ii_light produced lightcurves and then give this file as input to lc_pick, as shown below:

```
cd $REP_BASE_PROD/obs/
setenv run 0051
dal_create obj_name=all_lcr.fits template=ISGR-SRC.-LCR-IDX.tpl
foreach file (OGID/scw/$run*/lcr.fits)
idx_collect index="all_lcr.fits[1]" template="" element="$file[1]"
end
lc_pick source='GRS 1758-258' attach=n \
group=all_lcr.fits+1 lc=GRS1758.iilight.lc.fits emin="20" \
lcselect='E_MAX==40' instrument="ISGRI"
```

GRS1758.iilight.lc.fits contains the merged lightcurve of GRS 1758-258 in the 20–40 keV band (that of course has to exist in the lcr.fits original files!). To display it is is convenient to use the *lcurve* program of the FTOOLS package, as shownin Section 7.3.2.

9.10 Timing Analysis without the deconvolution

Read this if you are interested in fast variability studies (up to milliseconds).

In this section we describe a way of doing timing analysis in a non binning way, i.e. starting from the single events. This way is suitable for very short time scales (up to milliseconds) and is less recommended for long time bins for which the binning tools *ii_light* and *ii_lc_extract* are suitable.

In the text we will use one of the Science Windows with Crab data you have downloaded to run the PICsIT analysis (e.g. 003900020020).

In general the table with the events is very big, so if you are interested in only part of the Science Window (e.g. in the case of a burst) it is better to define a user good time interval (see Section 9) and work within it .

To select the photons that come from a given source in the field of view, you need to have the corresponding PIF. PIF is automatically created during the SPE step, but can also be created with a standalone tool *ii_pif*.

Create with og_create observational group \$REP_BASE_PROD/obs/crab/og_ibis.fits, and run analysis from COR to DEAD level, prepare the catalog, with Crab only and run *ii_pif*.

Now you are ready to create the lists of photons

```
cd $REP_BASE_PROD/obs/crab
evts_extract group="og_ibis.fits[1]" \
events="crabevts.fits" instrument=IBIS \
sources="crab_specat.fits[1]" gtiname="MERGED_ISGRI" \
pif=yes deadc=yes attach=no barycenter=1 timeformat=0 instmod=""
```

To increase signal-to-noise ratio select only events with PIF=1:

```
fcopy "crabevts.fits[2][PIF_1==1]" crab_pif1.fits
chmod -w crab_pif1.fits
```

Now you can produce the Crab power spectrum:

```
powspec
Ser. 1 filename +options (or @file of filenames +options)[] crab_pif1.fits
Name of the window file ('-' for default window)[] -
Newbin Time or negative rebinning[] 0.001
Number of Newbins/Interval[] INDEF
Number of Intervals/Frame[] INDEF
Rebin results? (>1 const rebin, <-1 geom. rebin, 0 none)[] 0
Name of output file[default]
Do you want to plot your results?[] yes
Enter PGPLOT device[] /XW
hardcopy crab_powerspec.ps/PS
```

As a result, the crab_powerspec.ps plot, shown in Figure 22, was produced. The 33 millisecond pulsation of the Crab is visible. For the details on *INTEGRAL* absolute timing see Walter et al. 2003 [13].

If your data have many short GTIs (e.g. in the case of telemetry saturation due to a solar flare or when PICsIT is in non standard mode) you can obtain spurious results. A typical case is finding an 8 sec period in your data due to the fact that the telemetry restart is synchronized with an 8 sec frame! When possible, compare your results with ii.light that is immune of this problem and can reach about 0.1 sec binning.

10 PICsIT data analysis

Unfortunately PICsIT sensitivity is not high enough to create a good image for a single Science Window, thus you should expect good results only if *INTEGRAL* is in staring mode (and the source is really bright!). In the example below we use Crab observations done in staring mode.

Download and install, as described in Section 6.1, the following Science Windows, whose DOLs are assumed to be written to the file **picsit.lst**:

scw/0039/003900020020.001/swg.fits[1] scw/0039/003900020030.001/swg.fits[1] scw/0039/003900020040.001/swg.fits[1] scw/0039/003900020050.001/swg.fits[1] scw/0039/003900020060.001/swg.fits[1]

Create the working directory picsit_ima with the *og_create* program:

```
cd ibis_data_rep
setenv REP_BASE_PROD $PWD
og_create idxSwg=staring.lst ogid=picsit_ima baseDir="./" instrument=IBIS
cp -r obs/picsit_ima obs/picsit_spe
```



Figure 22: Crab power spectrum.

10.1 PICsIT Image Reconstruction

To start the analysis, go to the working directory **\$REP_BASE_PROD/obs/picsit_ima** and call the *ibis_science_analysis* script:

```
cd obs/picsit_ima
ibis_science_analysis
```

After a few seconds, the main page of the IBIS Graphical User Interface (GUI) appears, as shown in Figure 23. Press "Reset"-button to be sure that you have the default ISDC_ENV parameters.

For PICsIT there were no major changes since the creation of revision 2 data, so you can start directly from the BIN_I level (startLevel=BIN_I). The end level for PICsIT image creation should be the default endLevel=IMA2. Disable ISGRI analysis by checking SWITCH_disableIsgri button, and enable PICsIT by unchecking SWITCH_disablePICsIT button.

PICsIT operates in an energy range (0.175-10 MeV) where background subtraction plays a very important role. The OSA for PICsIT provides a default set of maps for the background subtraction in certain energy bands, selected to optimize the instruments performance. By default, the software will automatically take subtraction with one set of maps (with about 1.7 Ms of integration time). The default energy bands are (in keV) for single events: 203 - 252, 252 - 336, 336 - 448, 448 - 672, 672 - 1036, 1036 - 1848, 1848 - 3584, 3584 - 6720. For multiple events: 336 - 448, 448 - 672, 672 - 1036, 1036 - 1848, 1848 - 3584, 3520 - 9072 - 13440.

-		ibis_science_analysis	· []
F	Main		<u>S</u> ave As
			Load
	startLevel:	COR 💌	<u>R</u> eset
	endLevel:	IMA2 💌	<u>R</u> un
	GENERAL_levelList:	COR,GTI,DEAD,BIN_I,BKG_I,CAT_I,IMA,IMA2,BIN_S,SPE,LCR,COMP,CLEAN	Quit
			Help
	CAT_refCat:	/isdc/arc/rev_2/cat/hec/gnrl_refr_cat_0020.fits[1][ISGRI_FLAG==1]	hidden
	SWITCH_disableIsgri:	Checked: yes	
	SWITCH_disablePICsIT:	Checked: yes	
			1
	SCW1_GTI_gtiUseri:	browse	J
	SCW1_GTI_TimeFormat:		
	ISGRI IMA	ISGRI SPE and LCR	PICsIT analysis
	1		

Figure 23: Main page of the IBIS GUI

This works for standard mode (mode 41) 16 only. For photon-by-photon mode (mode 43) the default values are¹⁷:

Number of Energy bands IBIS_IP_ChanNum=3 List of low energy boundaries, keV (single events) IBIS_IP_E_band_min_s="175 600 1000" List of high energy boundaries, keV (single events) IBIS_IP_E_band_max_s="600 1000 10000" List of low energy boundaries, keV (multiple events) IBIS_IP_E_band_min_m="350 600 1000" List of high energy boundaries, keV (multiple events) IBIS_IP_E_band_max_m="600 1000 13500"

To select parameters specific for PICsIT analysis press the PICsIT button. The PICsIT GUI page is shown in Figure 24.

Depending on the observation mode (staring or dithering), there are two ways to analyze the PICsIT data. The first, by using the parameter staring=yes, can be used only for staring observation, where the difference between all the pointings is less than the value defined by the tolerance parameter. In this case, the corrected shadowgrams are integrated before the deconvolution. If staring=no, then deconvolution will be performed for each science window individually.

The science windows we use in the current example were observed in staring mode, so check the staring button.

 $^{^{16}}$ The IBIS data mode can be checked in the archive by selecting IBIS mode column on the page with additional parameters 17 It is also possible to select any other energy band, but in this case it is up to you to generate and use the proper background maps. If you really want to change the energy bands press the hidden button, and go to BIN_I page.

-		PICsIT analysis 🛛 🕢 📃
	PICsIT analysis staring: tolerance: OBS1_ScwType: SCW1_BKG_P_method: PICSIT_inCorVar: PICSIT_outVarian: PICSIT_source_name: PICSIT_source_RA: PICSIT_source_DEC:	□k □k Help □.0001 □ <tr< th=""></tr<>
	r.	

Figure 24: PICsIT page of the IBIS GUI

OBS1_ScwType=POINTING means that only pointings (not slews) will be treated.

SCW1_BKG_P_method defines the way to normalize the background maps to the observed shadowgram before performing the background subtraction. 0 means scaling with exposure, and 1 means scaling with the average counts value over the whole detector. Both generally provide the same results, though the second method, which serves as default, is sometimes a bit better.

PICSIT_inCorVar defines whether you would like the software to partially correct the variance shadowgram for systematic effects due to the background (value 1), or not, in order to estimate by yourself the degree of systematics present in the deconvolved maps (value 0).

PICSIT_outVarian defines whether you would like to have the variance maps in output (value 1), or not (value 0). Note that the mosaic executable of the pipeline (*ip_skymosaic*) works only in case PICSIT_outVarian=0.

Return to the main GUI window by clicking on OK. Without changing other default parameters, you are now ready to run the analysis; do so by clicking on Run.

10.1.1 Results of PICsIT image analysis

In case of a staring the output images are in file

```
./picsit_ima.fits
```

For each energy range and each type of event (single, multiple), 2 types of images, INTENSITY and SIG-NIFICANCE, are produced. In case PICSIT_inCorVar=1, a variance map is also produced.

The list of found sources is in the file

./picsit_sky_res.fits

In our example the Crab is the only source found.

In Figure 25 you see the resulting image in the 252 - 336 keV energy range. The Crab is clearly seen in the center. The figure was produced with the help of ds9:

In the case of observations with dithering pattern (staring=no), the pipeline performs the shadowgram deconvolution in every Science Window and the output files are under:

./scw/RRRPPPPSSSF.001/picsit_ima.fits
./scw/RRRPPPPSSSF.001/picsit_sky_res.fits

and the integration of all these images (mosaic) is in:

./picsit_ima.fits

10.2 PICsIT spectral extraction from the mosaic image

It is also possible produce the source spectrum from the mosaic image.



Figure 25: Crab significance image in the 252 - 336 keV energy band as seen by PICsIT.

At http://isdc.unige.ch/index.cgi?Soft+scripts, you find an example of such a of perl script *spextract_pics.pl*. To use it, create the picsit_default_bins.txt file with the grouping of channels corresponding to the energy boundaries of the images we have

0	4	5
5	11	7
12	24	13
25	40	16
41	74	34
75	128	54
129	250	122
251	509	259
510	977	468
978	1023	46

and then rebin the response matrix with the **rbnrmf** from the FTOOLS package.

rbnrmf infile="\$REP_BASE_PROD/ic/ibis/rsp/pics_srmf_grp_0002.fits" \
outfile="pics-rmf.fits" binfile="picsit_default_bins.txt"

Create also a list of files you want to use for spectral extraction. In the case of staring, file picsit.lst will contain a single line:

picsit_ima.fits

Now you are ready to extract the spectrum. To do this you can either fill the values manually from the picsit_sky_res file, or extract the fluxes from the brightest pixel around the catalog position with the help of the *spextract_pics.pl*:

spextract_pics.pl -i picsit.lst -r 83.605 -d 21.95\

```
-n 8 -o crabspe_pics.fits -m pics-rmf.fits\
-a $REP_BASE_PROD/ic/ibis/rsp/pics_sarf_rsp_0002.fits
```

(In this command, switches -i and -o define the input and output files, -r and -d define the RA and DEC of the desired pixel, -n defines the number of energy ranges, -m and -a define the locations of rmf and arf matrices).

First and last channels of the resulting spectrum crabspe_pics.pha are fictive. Do not forget to ignore them.

The result is shown in Figure 26.



Figure 26: PICsIT Crab spectrum extracted from the mosaic.

10.3 PICsIT pipeline spectral extraction

An improvement in the PICsIT spectrum extraction executable with the PIF method has been implemented in the OSA 5.1 and is described below. Despite the improvement, under certain conditions (e.g. staring observations), the algorithm is unstable. We suggest that the user adopts the method of extracting spectra from images, as explained in Section 10.2.

At the moment it is not possible to run the pipeline spectral analysis if you have already executed the BIN_I and IMA2 steps in the group. Thus to start spectral analysis, move to a new working directory \$REP_BASE_PROD/obs/picsit_spe and call *ibis_science_analysis*:

```
cd obs/picsit_ima
ibis_science_analysis
```

After a few seconds, the main page of the IBIS Graphical User Interface (GUI) appears, as shown in Figure 23.

Change start (startLevel=BIN_S) and end (endLevel=SPE) levels. Disable ISGRI analysis by the checking SWITCH_disableIsgri button, and enable PICsIT by unchecking the SWITCH_disablePICsIT button.

To select parameters specific for the PICsIT analysis press the PICsIT button. PICsIT GUI page is shown in Figure 24.

Set the name of the source, and its coordinates (note that below we give the syntax for the command line case, while in the GUI there should be no quotes):

PICsIT_source_name="Crab" PICsIT_source_ra=83.63 PICsIT_source_dec=22.01

Click on Ok and return to the main GUI window. Press Run.

10.3.1 Displaying the results of PICsIT spectral analysis

The output spectra are created for each science window, even in staring mode. They are located at $scw/0039000200X0.001/picsit_spec.fits$. To sum them up, you can use *mathpha* FTOOL. Below an example of a c shell script to sum up the spectra is given.

```
cd $REP_BASE_PROD/obs/picsit_spe/scw
mkdir pha
foreach dir ( *.001 )
 set fn='echo $dir| cut -c 1-12'
set fb = 'echo $fn.pha'
fextract "$dir/picsit_spec.fits[2]" "pha/$fb"
end
cd pha
mv 003900020020.pha sum.pha
set npha=1
foreach pha (0*.pha)
 @ npha++
mathpha units=R errmeth='gaus' expr="sum.pha+$pha"\
         outfil=tmp.pha exposure=CALC ncomments=0 areascal='-'
mv tmp.pha sum.pha
echo $npha
end
mathpha units=R errmeth='gaus' expr=sum.pha/$npha\
        outfil=tmp.pha exposure=sum.pha ncomments=0 areascal='-'
mv tmp.pha sum.pha
fparkey value=../../../ic/ibis/rsp/pics_srmf_grp_0003.fits\
       fitsfile=sum.pha keyword=RESPFILE
fparkey value=../../../ic/ibis/rsp/pics_sarf_rsp_0002.fits\
        fitsfile=sum.pha keyword=ANCRFILE
```

Now you are ready to launch xspec and check the resulted spectrum

xspec XSPEC> data sum.pha XSPEC> setplot energy XSPEC> ign 1000.-** XSPEC> plot ldata

The result is shown in Figure 27.



Figure 27: PICsIT Crab spectrum created as described in Section 10.3.

10.4 PICsIT Timing Analysis

It is possible to produce a lightcurve of the PICsIT detector from the spectral-timing data. To build it, it is enough to run the analysis from the COR to DEAD level and then from LCR to LCR (of course, it is not a problem if you have already done the analysis up to the IMA2 level). Do not forget to disable ISGRI and enable PICsIT for the analysis.

The results are in the following FITS files:

./scw/RRRRPPPPSSSF.001/picsit_lcr.fits

11 Known Limitations

Please note, that with the time between the releases this list can change. For the most update version see

http://isdc.unige.ch/Soft/download/osa/osa_sw/osa_sw-5.0/osa_issues.txt

11.1 ISGRI

- 1. The SPSF (System Point Spread Function) fit may fail (does not converge) during the sky image generation when there are close sources (within 15-20 pixels from the target source).
- 2. In the mosaic build with the option spread=1 the source flux is slightly reduced (10 the fluxes measured in the Science Window.
- 3. Secondary lobes of strong very-off-axis sources are sometimes not fully corrected/cleaned in reconstructed images. This is possibly due to the approximation in the source modelling.
- 4. The maximum number of sources handled by *ii_spectra_extract* is 100 but it is strongly recommended to only fit spectra of the sources that are effectively active (visible, detectable) during the Science Window.
- 5. In general, a safe lower limit for the response is 22.5 keV.
- 6. A problem on-board IBIS causes event times to be shifted by 2 seconds under some circumstances. The software corrects for it. The keyword TIMECORR found in *-*-ALL or *-*PRP structures of a Scw group, indicates whether the correction was done somewhere in this Scw. The algorithm is the following: (1) detect if there could be a possible 2 sec jump (2) if (1) then look if there was indeed a jump. (3) if (2) then correct, if not (2) then don't correct (4) if we cannot determine either (2) or not (2) then flag. As a result, the TIMECORR keyword can have the following values: 0: no possible jump
 - 1: possible jump, jump indeed exists, corrected
 - 2: possible jump, jump did not take place, not corrected
 - 3: possible jump, cannot decide if jump took place, not corrected
 - 4: possible jump, algorithm has a problem, to be investigated

If you are doing absolute timing and your data contains TIMECORR>0 take great care. If TIMECORR=1 or 2 it should be OK. If TIMECORR=3 you should better not use it. If TIMECORR=4 contact ISDC

7. The lightcurve extraction (*ii_lc_extract* program) is performed by building shadowgrams for each time and energy bin. It potentially takes a large amount of CPU time and moreover there is a limit on the minimum duration of the time bins for a given number of energy bins and the duration of the given Science Window.

The time bin for the light curve must be such that the total number of maps in the file isgr-corr-shad does not exceed 2 Gig worth of disk space. Since 3 maps (intensity, variance, efficiency) are created per image and each image takes 130*134*4 bytes or 24 fits blocks, plus one fits block for the header, for a total of 3*25*2880=216000 bytes, the product of the number of time bins, per each science window, and the number of energy bands must be less than about 9942, assuming that no images are left from image and spectral analysis.

Example: assuming Nim+Nsp=30 shadowgrams already created for images and spectra, for a Tscw=3000 s science window duration, 4 light curves in Ne=4 energy bands can be created (for a total of Nlci=9942-(Nim+Nsp)=9912 light curve images) with the minimum time bin Tbin given by Tbin = Ne x (Tscw/Nlci) = $4 \times 3000/9912 = 1.2 \text{ s}$
- 8. Version 12 of XSPEC cannot read the ISGRI spectra extracted per ScW in "isgri_spectrum.fits" beyond the first extension. In case you want to study the ScW by ScW spectra of a given source, you may either use XSPEC 11 or use *spe_pick* to extract the spectra in PHA2 format ("single=y" option), which can be read by XSPEC 12.
- 9. *ii_pif* crashes if the input catalog inCat contains more than 500 sources.

11.2 PICsIT

1. An improvement in the PICsIT spectrum extraction executable with the PIF method has been implemented in the OSA 5.1. Despite the improvement under certain conditions (e.g. staring observations), the algorithm is still unstable. We still suggest the user to continue adopting the spectra extraction from images, as explained in OSA4.2. A RMF already rebinned for the channels of the imaging pipeline is available in the IC files. Part III Data Analysis in Details

12 Science Analysis

In the Cookbook you have seen that in order to run the Scientific Analysis you should just launch the main script *ibis_science_analysis* with a desired set of parameters. As discussed in the Overview (Section 5) the main script consists of smaller scripts, which in turn unify executables with similar tasks for different types of events (see Figure 28). In this chapter we describe these small scripts in more detail in order to explain how the main script works and what parameters you have to enter for a proper analysis.

While describing the executables we describe those parameters which were included into the main script. You find in the Appendix the detailed description of the results produced at each step . In the Appendix you also find the description of raw and prepared data with which you start the analysis.

12.1 *ibis_correction*

This script produces corrected Data Structures for all types of the events. This script combines the following executables:

- *ibis_isgr_evts_tag*
- *ibis_isgr_energy*
- *ip_ev_correction*

Using the information about the noisy pixels (see Section 3.2.3) the script flags all the noisy events. Then the script performs the energy correction of all the events for which correction is possible (no correction can be done for the spectral-timing mode). The output Data Structures contain the list of the photon energy in keV (see Appendix C.1).

$12.1.1 \quad ibis_isgr_evts_tag$

ibis_isgr_evts_tag tags, as noisy, all photons from a pixel if this pixel had a switch off during the Scw. It also looks at the distribution of the time between events in each given pixel. If this distribution is abnormal, all photons from this pixel are flagged as noisy (within this Scw). Noisy events are ignored in the subsequent analysis.

Table 3:	ibis isar	evts taa pa	rameters	included	into th	ne main	script.
rabie o.	1010_1091_	cous_oug pu	amouto	merudea	11100 01	ic main	beript.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_ICOR_idxSwitch	idxSwitch	string	DOL of the index of pixels switch list.
			default: ""
SCW1_ICOR_probShot	probShot	real	Probability of shot time decay.
			default: 0.01

12.1.2 *ibis_isgr_energy*

Due the charge loss in the CdTe crystal, for a given energy deposit, events which have interacted at different depths in the crystal get different measured amplitudes (PHA). The rise-time of the signal induced by these events is also different, and a clear quasi-linear correlation between the charge loss and the rise-time variation is observed (a deeper interaction induces a larger charge loss, giving the longer rise-time). It is then possible to correct this charge loss effect, by taking into account the rise-time information of the signal.



ibis_science_analysis

Figure 28:

Composition of the main script *ibis_science_analysis*. For further descriptions of the BIN BKG steps for the DEAD, IMA and BIN_S levels, see Figures 29, 30 and 31 respectively.



Binning - backgroud step for [IMA]

Figure 29: Overview of the binning - background step for Imaging.

ibis binning $\xrightarrow{19}{9} og_jibis \longrightarrow -ii_shadow_build \\ -ip_(*)_shadow_build (*)=ev,si} \longrightarrow og_jibis$ $EFFI IC file \longrightarrow -ii_map_rebin \longrightarrow rebinned maps$ $ibis background_cor$ $og_jibis \longrightarrow -ii_shadow_ubc \\ -ip_shadow_ubc \longrightarrow og_jibis$ $\xrightarrow{19}{9}$



Figure 30: Overview of the binning - background step for Spectra.





Figure 31: Overview of the binning - background step for Lightcurves.

The executable *ibis_isgr_energy* performs a rise-time correction for each raw ISGRI event, using the ISGRI rise-time correction table (Section B.2). The corrected energy is given in keV (ISGRI_ENERGY), and the corrected rise-time (ISGRI_PI) is the row number used in the IC file. The correction is done by rescaling the measured spectra in accordance with the observed rise-time rt, so that the corrected event corresponds to the deposited energy.

Also the temperature and bias corrections are done at this step.

Table 4: $ibis_isgr_energy$ parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_ICOR_GODOL	GODOL	string	DOL of the Gain-Offset correction table
			default: ""
SCW1_ICOR_riseDOL	riseDOL	string	DOL of the rise-time correction table
			default: ""
SCW1_ICOR_icDOL	icDOL	string	DOL of the IC file for calibration parameters
			default: ""

$12.1.3 \quad ip_ev_correction$

The executable $ip_ev_correction$ performs energy correction of the events received by PICsIT in photon-byphoton mode. Single events are corrected for each pixel separately with the use of the pixel-dependent gain and offset factors, and the pixel-independent channel-to-keV conversion factors avgain and avoffset, see Section B.2 for more details.

 $energy[keV] = avgain \cdot gain \cdot PICSIT_PHA + avoffset \cdot offset$

Multiple events are corrected for gain and offset on-board, and thus they are simply transferred from channels to keV:

$energy[keV] = avgain \cdot PICSIT_PHA$

Table 5: *ip_ev_correction* parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_PCOR_enerDOL	enerDOL	string	DOL of the energy correction table (PICsIT).
			default: ""

12.2 *ibis_gti*

This script builds Good Time Interval (GTI) information from housekeeping data, information about the satellite stability and data gaps. It calls the following executables to obtain the GTIs:

- gti_create
- gti_attitude
- gti_data_gaps
- gti_import
- gti_merge

For the definition of GTI see Introduction to the INTEGRAL Data Analysis [1]

$12.2.1 \quad gti_create$

This program generates all GTIs for one instrument that depend on HK and other parameters and are defined by a limit in a limit table, see details in Section B.3. In ISGRI case it creates **VETO** GTIs during which VETO was switched on, and **ISGRI_MCE7X** (where X varies from 0 to 7) GTIs during which the Xth module was switched on. **VETO** GTIs are taken into account by *gti_merge* executable (see Section 12.2.5), and **ISGRI_MCE7X** GTIs are taken into account later during BIN_I, BIN_S and BIN_T levels.

Table 6:	gti_create	parameters	included	into	the	main	script.
----------	---------------	------------	----------	------	-----	-----------------------	---------

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_GTI_LimitTable	LimitTable	string	The DOL of the GTI limit table.
			default: ""

12.2.2 gti_attitude

A GTI is defined for each period of time where the pointing stability is better than the accepted tolerance (parameter AttStability). This GTI is named "ATTITUDE". For slews this GTI is always set to be good independently of any input data. If the necessary data are not available the GTI is set to be "not good" for the whole Science Window.

Table 7:	$gti_{-}attitude$	parameters	included	into	$_{\rm the}$	main	script.
----------	-------------------	------------	----------	------	--------------	-----------------------	---------

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_GTI_attTolerance	AttStability	real	Defines the accepted attitude stability tolerance in units of arc minutes. A GTI is created if the stability is better than this tolerance
			default: "0.5"

$12.2.3 \quad gti_data_gaps$

This program generates GTIs for ISGRI and PICsIT. The GTIs depend on the presence of the science data. A time is defined as bad if a science packet of the instrument is missing. The expected science packets depend on the actual mode of the instrument. This is taken into account while the GTI is being created. The program does not distinguish between a pointing, a slew and an engineering window. The time is also set to be "bad" if there are scientific data that do not agree with the reported mode of the instrument.

Time is also defined bad if the telemetry indicates that there was an interruption of the on-board processing due to a telemetry saturation. In normal operation this case is rather seldom.

The final GTI is written into the IBIS index group and has the name "ISGRI_DATA_GAPS" and "PIC-SIT_DATA_GAPS" for ISGRI and PICsIT correspondingly.

12.2.4 gti_import

The *gti_import* reads the user GTI table and converts it into a table in ISDC format. The user GTI can be defined either in units of OBT, IJD, or UTC. The output is always in OBT. The user table can define either bad or good time intervals. The output time intervals are always good ones. See more details in the Introduction to the *INTEGRAL* Data Analysis [1].

Table 8: The *gti_import* parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_GTI_gtiUser	InGti	string	DOL of the user GTI table
			" "=there is no user GTI.
			default: ""
SCW1_GTI_TimeFormat	TimeFormat	string	Time format in which the user GTI is given.
			possible values: "IJD", "UTC", "OBT"
			default: "OBT"
SCW1_GTI_Accuracy	Accuracy	string	Used accuracy for OBT to IJD conversion and vice
			versa.
			possible values: "any", "inaccurate", "accurate"
			default: "any"

12.2.5 *gti_merge*

This program merges input GTIs to form a new GTI. It is an AND operation: a time in the resulting GTI is defined to be "good" if the time is "good" in every input GTI. The names of the GTIs and the instrument to which the GTI belongs have to be defined as program parameters.

	<i>j</i>	<i>J I i i</i>	I I I I I I I I I I I I I I I I I I I
Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_GTI_ISGRI	MergedName	string	GTIs to be merged for ISGRI.
			default: ''VETO ATTITUDE ISGRI_DATA_GAPS''
SCW1_GTI_PICsIT	MergedName	string	GTIs to be merged for PICsIT.
			default: ''VETO ATTITUDE P_SGLE_DATA_GAPS
			P_MULE_DATA_GAPS''
SCW1_GTI_SCI	SC_Names	string	Names of spacecraft GTIs to be merged for ISGRI

string

SC_Names

default: '' ''

default: '' ',

Names of spacecraft GTIs to be merged for PICsIT

Table 9: gti_merge parameters included into the main script.

12.3 *ibis_dead*

SCW1_GTL_SCP

This script combine two executables calculating the dead time for ISGRI, PICsIT and Compton cases.

- *ibis_isgr_deadtime*
- *ibis_pics_deadtime*

The detector efficiency is divided into the real efficiency and the dead time D which is due to the delay following a photon detection during which another event can not be recorded.

The observed count rate $C_{\rm o}$ is connected with the real count rate C as

$$C_{\rm o} = C(1-D)$$

The presence of the BGO shielding (see Section 3.2.5), calibration source, and the Compton coding induces an efficiency loss due to good events which by chance fall in their respective time windows.

For IBIS the term "efficiency" is reserved for the real physical detection efficiency. The effects induced by Veto, calibration source, and Compton tagging are considered as additional dead times to be added to instrumental dead time.

Numerically, dead time is calculated as the product of the count rate and the time window (e.g. the dead time due to the time of the photon detection is equal to the product of the module count rate and the coding time. The dead time due to the veto is equal to the product of the veto count rate and the veto time window, etc.) The resulting dead time is taken as a sum of all the dead times connected with different effects. Simple summing of dead times can be done since the estimations show that the coincidence probability between different types of events is low. The values of the count rates is taken from the HK Data Structures **IBIS-DPE.-HRW** and **IBIS-DPE.-CNV** and the corresponding time windows are either measured (instrumental) or read into HK or IC (all others: VETO, Compton).

12.3.1 *ibis_isgr_deadtime*

The executable $ibis_isgr_deadtime$ calculates dead times resulting from count rates in ISGRI modules, taking into account the instrument configuration.

ISGRI dead time is due to the combined effect of:

- Instrument count rate,
- Veto count rate (from the lateral, or lateral+bottom (according to IC file) shielding),
- Calibration count rate,
- Compton coincidences, if Compton mode is on.

Note that the problem of the VETO swapping on-board is corrected almost every time through IC file: **IBIS-VETO-MOD**. VETO also sometimes has a wrong level; this will be corrected in the future. The 3 cases when it happened are in the IC file **GNRL-INTL-BTI** with the BTI_TYPE: **ISGRI_VETO_LEVEL**.

Table 10: *ibis_isgr_deadtime* parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_veto_mod	icDOL	string	DOL of the IC file for VETO model and width of
			Compton window
			default ""

$12.3.2 \quad ibis_pics_deadtime$

The executable *ibis_pics_deadtime* extracts from the housekeeping data the PICsIT intrinsic dead time of and dead time induced by the fortuitous coincidence with VETO and calibration system.

Contrary to the ISGRI case, the dead time is measured by the on-board electronics and we have only to decipher it from the telemetry data.

12.4 *ibis_binning*

The main function of *ibis_binning* is to split the data into energy and time bins. The number of the time and energy bins depend on the purpose of the analysis. The user can choose if the binning is done for imaging, spectral extraction, or light curve production. Typically just one bin in time and few bins in energy are chosen for the image production; one time bin and many energy bins for spectral extraction; many time bins and just few energy bins for timing analysis.

This step is repeated for each Science Window in the observational group. The shadowgram is created for each energy interval. Script *ibis_binning* combines the following executables:

- *ii_shadow_build*
- $\bullet ~ip_ev_shadow_build$
- *ip_si_shadow_build*

12.4.1 *ii_shadow_build*

The executable ii_shadow_build creates shadowgrams of ISGRI events in the given time (Tbin) and energy (Ebin) ranges. For every shadowgram a corresponding efficiency shadowgram is created. For pixels active during the revolution (switch status ON), efficiency falls into two components and is null for the others: the first factor takes into account the dead time D of the corresponding module MDU; the second factor reflects the efficiency energy dependence, LT, for the lower energy bins due to the low threshold operation limit, such that: Eff[Tbin,Ebin,y,z]= (1-D[Tbin,mdu]) * LT[Ebin,y,z]. It is this value that is given in the shadowgram efficiency maps.

This executable also applies two different noisy pixel detection techniques. The first one is time-based; it discriminates every pixel for which the scientific SELECT_FLAG equals one (even only once over all the Science Window events). Note: this flag was filled by the executable *ibis_isgr_evts_flag*. The second method is spectral-based; it discriminates pixels by estimating their spectral deviation from the average ISGRI spectrum. This spectral method can be switched ON/OFF by parameter NoisyDetFlag.

Name	Name	Type	Description
(in the main script)	(executable)		
IBIS_II_ChanNum	isgri_e_num	integer	Number of energy intervals for ISGRI.
			possible values: $1 - 10$
			default: 7
IBIS_II_E_band_min	isgri_e_min	string	List of lower energy boundaries (keV).
			default: "15 40 100 200 400 700 1200"
IBIS_II_E_band_max	isgri_e_max	string	List of upper energy boundaries (keV).
			default: "40 100 200 400 700 1200 2500"
IBIS_min_rise	isgri_min_rise	integer	Minimum rise-time.
			default: 7
IBIS_max_rise	isgri_max_rise	integer	Maximum rise-time.
			default: 80
IBIS_NoisyDetMethod	NoisyDetFlag	integer	Defines the way to deal with noisy pixels
			possible values: 0 Time-based noisy detection;
			1 Time-based + Spectral-based method.
			default: 1

Table 11:	ii_shadow_build	parameters	${\rm included}$	into	the	main
script.						

12.4.2 *ip_ev_shadow_build*

 $ip_{ev_{shadow_build}}$ takes as an input PICsIT data received in photon-by-photon mode. For each given energy and time range, intensity and efficiency shadowgrams are produced. Efficiency is defined as:

$$\mathrm{Eff} = (1 - D) * T_{\mathrm{ON}}/T$$

where $T_{\rm ON}/T$ is given by the GTI and the total observation length.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_BIN_cleanTrk	cleanTrk	integer	Cleaning of cosmic-ray induced events.
SCW2_BIN_cleanTrk			possible values:
			0 = No
			1 = Yes
			default: 1
IBIS_IP_ChanNum	picsit_e_bin	integer	Number of Energy bands
			default: 3
IBIS_IP_E_band_min_s	picsit_e_min_s	string	List of lower energy boundaries for PICsIT (single
			events)
			default: "175 600 1000"
IBIS_IP_E_band_max_s	picsit_e_max_s	string	List of higher energy boundaries for PICsIT (single
			events)
			default: "600 1000 10000"
IBIS_IP_E_band_min_s	picsit_e_min_m	string	List of lower energy boundaries for PICsIT (multiple
			events)
			default: "300 600 1000"
IBIS_IP_E_band_max_s	picsit_e_max_m	string	List of higher energy boundaries for PICsIT (multiple
			events)
			default: "600 1000 13500"
SCW1_BIN_P_inDead	inDead	string	DOL of the dead time Data Structure.
			default ""
SCW1_BIN_P_faltStatus	faltStatus	string	DOL of the list of faulty pixels.
			default ""

Table 12: $ip_ev_shadow_build$ parameters included into the main script.

12.4.3 *ip_si_shadow_build*

 $ip_si_shadow_build$ takes as an input PICsIT data received in standard mode. For each given energy and time range, intensity and efficiency shadowgrams are produced. Efficiency is defined as:

$$Eff = (1 - D) * T_{ON}/T$$

Table 13: $ip_si_shadow_build$ parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
IBIS_IPS_ChanNum	picsit_e_bin	integer	Number of Energy bands (0 means to use the default
			ones.)
			default: 0
$IBIS_IPS_E_band_min$	$picsit_e_min$	string	List of lower energy boundary
			default: "170 600 1000"
$IBIS_{IP}_{E}_{band}_{max}$	picsit_e_max	string	List of higher energy boundary
			default: "600 1000 10000"
SCW1_BIN_P_inDead	inDead	string	DOL of the dead time Data Structure.
			default""

12.5 *ii_map_rebin*

ii_map_rebin reads input background and/or Off-axis correction maps and rebins them into new energy bands. Background maps are summed with coefficients equal to the part of each input map that participates in the sum. Off-axis maps are also weighted by a power-law spectrum (with the default slope -2) and ARF (if present). After rebinning, off-axis maps are normalized to 1. Output energy bands are defined from the index of detector shadowgrams (ISGR-DETE-SHD) of a given type.

Name	Name	Type	Description
(in the main script)	(executable)		
corrDol	inpCorrDol	string	DOL of the isgri off-axis corrections
			default ""
rebinned_corrDol_ima	rebCorrDol	string	DOL of the rebinned isgri off-axis corrections for imag-
			ing
			default ""
rebinned_corrDol_spe	rebCorrDol	string	DOL of the rebinned isgri off-axis corrections for spec-
			trum
			default ""
rebinned_corrDol_lcr	rebCorrDol	string	DOL of the rebinned isgri off-axis corrections for lc
			default ""
rebinned_backDol_ima	rebBkgDol	string	DOL of the rebinned isgri background corrections for
			imaging
			default""
rebinned_backDol_spe	rebBkgDol	string	DOL of the rebinned isgri background corrections for
			spectrum
			default ""
rebinned_backDol_lcr	rebBkgDol	string	DOL of the rebinned isgri background corrections for
			lc
			default ""
SCW1_BKG_LisgrBkgDol	inpBkgDol	string	DOL of the isgri background model
			default ""

Table 14: *ii_map_rebin* parameters included into the main script.

12.6 *ibis_background_cor*

This script combines executables which fill the dead zones of the detectors in accordance with the chosen method, creating the extended intensity map I_{ex} . Then with the help of the IC background, B, and uniformity, U, maps (See Section B.4) a corrected intensity map I_{cor} is produced:

$$I_{\rm cor} = \frac{I_{\rm ex} - B}{U}$$

Script *ibis_background_cor* combines the following executables:

- *ii_shadow_ubc*
- *ip_shadow_ubc*

12.6.1 *ii_shadow_ubc*

ii_shadow_ubc reads all raw detector, and corresponding efficiency, shadowgrams, and fills the detector dead zones. *ii_shadow_ubc* is run in three cases; to produce images, spectra and lightcurves. In the imaging case, for each pair of detector-efficiency shadowgrams it performs the background correction for the uniform

(default) and non-uniform spatial distribution. As output it produces 3 shadowgrams of larger dimensions: corrected expanded shadowgram and corresponding variance and efficiency shadowgrams. Method to be applied for the pixel value interpolation in dead zones is defined by the parameter **method_int**.

Name	Name	Type	Description
(in the main script)	(executable)		
ii_shad	low_ubc		
SCW1_BKG_I_method_cor	method_cor	string	Method to be applied for background removal possible values: 0-2 0 – background from IC tree is applied to the whole detector 1 – parameter switches to this value automatically when background map is specified (not an IC one). 2 – background is treated for each module separately default: 0
SCW1_BKG_I_method_int	method_int	string	 default: 0 Method to be applied for the pixel value interpolation possible values: 0 - dead zone pixels will be filled with 0 1 - dead zone pixels will be filled with mean detector value -1 - no dead zones default:1
IBIS_NoisyDetMethod	NoisyPixControl	integer	$0 \Rightarrow$ no Noisy Pixel detection; $1 \Rightarrow$ use SE- LECT_FLAG method. default:1

Table 15: *ii_shadow_ubc* parameters included into the main script.

$12.6.2 \quad ip_shadow_ubc$

ip_shadow_ubc reads raw detector shadowgrams and performs the background correction. Also detector dead zones are filled at this step. Method to be applied for the pixel value interpolation in dead zones is defined by the parameter **method_int**. Expanded intensity and variance shadowgrams are produced as output.

 Table 16:
 ip_shadow_ubc parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW1_BKG_P_picsSUnifDOL	picsSUnifDOL	string	DOL of the PICsIT Detector Uniformity
			model (single).
SCW1_BKG_P_picsMUnifDOL	picsMUnifDOL	string	DOL of the PICsIT Detector Uniformity
			model (multiple).
SCW1_BKG_P_picsSBkgDOL	picsSBkgDOL	string	DOL of the PICsIT Detector Background
			model (single).
SCW1_BKG_P_picsMBkgDOL	picsMBkgDOL	string	DOL of the PICsIT Detector Background
			model (multiple).
SCW1_BKG_P_method	method	integer	Method to use in scaling the background
			maps
			possible values:
			0 – model (in cts) is normalized by the time
			of observation
			1 – model (in cts) is normalized by the mean
			count value
			default: 0

12.7 Catalogs

The catalog extraction selects the sources in the field of view (FOV) from the references catalog.

$12.7.1 \quad cat_extract$

The executable *cat_extract* performs the source selection from a reference catalog. The reference catalog should have the same structure as GNRL-REFR-CAT (see Table 48 in Appendix C.5). Its DOL is passed in the parameter 'refCat'. There is no standard output catalogue for IBIS and the parameter 'outExt' must be filled.

Name	Name	Type	Description
(in the main script)	(executable)		
CAT_refCat	ref_cat	string	DOL of the reference catalogue.
			default value: ''\$ISDC_REF_CAT''
OBS1_CAT_radiusMin	radiusMin	string	Low limit for the position selection.
			default: ''0''
OBS1_CAT_radiusMax	radiusMax	string	High limit for the position selection.
			default: ''20''
OBS1_CAT_fluxDef	fluxDef	string	Column used for flux selection.
			default: ''''
OBS1_CAT_fluxMin	fluxMin	string	Low limit for flux selection (in $ph/cm^2/s$).
			default: ''''
OBS1_CAT_fluxMax	fluxMax	string	High limit for flux selection (in $ph/cm^2/s$).
			default: ''''
OBS1_CAT_class	class	string	Select the sources by class.
			default: '''
OBS1_CAT_date	date	string	IJD for the public data.
			possible values:
			<0 – all public;
			0 – all private;
			>0 – according to 'DAY_ID' date
			default: -1

Table 17: *cat_extract* parameters included into the main script.

12.8 Image analysis

The IBIS telescope is a device based on a coded aperture imaging system. The mask chosen for IBIS is based on a cyclic replication of MURA (Modified Uniformly Redundant Array) of order 53, expanded to 95 pixels. The properties of the MURA patterns are described in the papers [11], [12].

For IBIS, the mask is about 1.8 times larger than the detector. The most important advantage of such configuration is that for a large fraction of the sky a source projects a complete pattern on the detector (the Fully Coded Field Of View, FCFOV). The part of the sky from which only a fraction of the source radiation directed towards the detector plane is modulated is called Partially Coded Field of View (PCFOV). The complete field of view of the telescope (FOV) is therefore composed by the central FCFOV of constant sensitivity and optimum image properties surrounded by the PCFOV of decreasing sensitivity. A source outside the FOV simply contributes to the background level.

Representing the mask with an array M of 1 (open elements) and 0 (opaque ones), the detector array D will be given by the convolution of the sky image S by M plus an unmodulated background array term B,

$$D=S\star M+B$$

MURA patterns have the remarkable property that their cyclic autocorrelation gives a delta function. The decoding array, G, is inverse to M where G = 2M - 1 (i.e., G = +1 for M = 1 and G = -1 for M = 0) and thus is correlation inverse.

With the help of array G we can reconstruct the sky:

$$S' = D \star G = S + B \star G$$

where S' differs from S only by the $B \star G$ term, which for a flat array B is a constant level that can be measured and removed.

To have a sidelobe-free response a source must be able to cast a whole basic pattern on the detector (fully coded source). To make use of all the detector area and to allow more than one source to be fully coded, the mask basic pattern is normally taken as the same size and shape of the detector and the total mask made by a cyclic repetition ($< 2 \times 2$ for rectangular mask) of the basic pattern. For such *optimum systems* a FCFOV source will always project a cyclically shifted version of the basic pattern, and correlating the detector image with the G decoding array will provide a sidelobe-free peak with position-invariant shape at the source position.

A source in the PCFOV will instead cast an incomplete pattern and its contribution cannot be a-priori subtracted and will produce secondary lobes (coding noise). On the other hand the modulated radiation from PC sources can be reconstructed by extending the correlation procedure, with a proper normalization, to the PCFOV.

URA masks also minimize the statistical errors of the reconstructed peaks. Since $V = G^2 \star D = \Sigma D$ the variance associated with each reconstructed sky image pixel is constant in the FCFOV and equal to the total counts recorded by the detector; therefore the source signal to noise is simply

$$\frac{S}{N} = \frac{C_S}{\sqrt{C_S + C_B}}$$

where C_S and C_B are source and background counts. These masks also have nearly equal number of transparent and opaque elements and therefore offer minimum statistical error in condition of high background (typical for the γ -ray domain). However the sensitivity also depends on the detector spatial resolution and an *imaging efficiency* factor must be applied to this maximum S/N to account for this effect.

Discrete cross-correlation to compute sky and variance images can be written

$$S_{ij} = \sum_{kl} G_{i+k,j+l} D_{kl}$$
; $V_{ij} = \sum_{kl} G_{i+k,j+l}^2 D_{kl}$

where Poisson statistics was assumed. This standard deconvolution in FCFOV can be extended in the PCFOV by extending the correlation of the decoding array G with the detector array D in a non-cyclic form, padding G with 0 elements. Since only the detector section modulated by the PC source is used to reconstruct the signal, the statistical error at the source position and significance of the ghost peaks are minimized. However to ensure a flat image in the absence of sources, detector pixels which for a given sky position correspond to opaque mask elements must be balanced before subtraction with the factor $b = n^+/n^-$, where n_+ is the number of pixels corresponding to transparent elements and n_- to opaque ones for that given sky position. This can be written

$$S_{ij} = \sum_{k} G_{i+k,j+l}^{+} W_{kl} D_{kl} - B_{ij} \sum_{kl} G_{i+k,j+l}^{-} W_{kl} D_{kl}$$

where the decoding arrays are obtained from the mask M by $G^+ = M$ and $G^- = 1 - M$, then padded with 0's outside mask region, and where the sum is performed over all detector elements. In the FCFOV we obtain the same result as the standard cross-correlation. To consider effects such as satellite drift corrections (see [7]), dead areas or other specific conditions, a weighting array W is used to weigh properly the detector array before correlating it with the G arrays. The balance array is

$$B_{ij} = \frac{\sum_{kl} G^+_{i+k,j+l} W_{kl}}{\sum_{kl} G^-_{i+k,j+l} W_{kl}}$$

The variance, which is not constant outside the FCFOV, is computed accordingly

$$V_{ij} = \sum D_{kl} \left(G^+_{i+k,j+l} W_{kl} \right)^2$$
$$+ B^2_{ij} \sum D_{kl} \left(G^-_{i+k,j+l} W_{kl} \right)^2$$

since the cross-terms G^+G^- vanish. Note however that when the weights W_{kl} refer to the same pixel in D, the terms G W must be summed before squaring (see [7]). The varying effective area can be calculated by a similar formula and used to renormalize, after background subtraction to FCFOV count rates. All this can be performed for sampling finer than 1 pixel per mask element and using a G array convolved with detector the PSF in order to optimize S/N for point sources, with corresponding normalizations. This procedure can be carried out with a fast algorithm by reducing previous formulae to a set of correlations computed by FFT.

The on-axis System Point Spread Function (SPSF) on the whole FOV for an optimum system and PSF deconvolution is shown in Fig. 32. Note the peak and flat level in the central FCFOV, the secondary lobes (coding noise) in the PCFOV and the 8 main ghosts of the source peak in the PCFOV located at distances, from the source, which are a multiple of the basic pattern.



Figure 32: SPSF for the IBIS/ISGRI telescope.

The average Point Source Location Error (PSLE) for an optimum coded aperture system with a defined SPSF depends on the source signal to noise ratio (S/N) as following

$$PSLE \div \frac{1}{R(S/N)}$$

The IBIS/ISGRI telescope, assuming no error in pointing axis reconstruction or other systematic effects, can locate a 30 σ point-like source at better than $\pm 1'$. Absolute error in attitude reconstruction for INTEGRAL is expected to be < 20''.

In a standard analysis, IBIS events or histograms are binned in detector images, which are then corrected for detector and background non-uniformity [7] and then processed by an iterative algorithm which decodes, cleans and composes sky images. For each detector image a sky image and its variance are obtained using the deconvolution procedure, and then iteratively searched for sources and cleaned of the source side lobes. In this iterative process the source peaks are fitted with the bi-dimensional Gaussian and finely located. Then the source contribution to the image is modeled in detail and subtracted. The images are rotated, projected and summed after being weighted with the variance, and then searched for further contributions.

More details can be found in [8], [9]

12.8.1 *ii_skyimage*

 $ii_skyimage$ deconvolves shadowgrams in the given energy bands for each Science Window, using the balanced cross-correlation method described above. The energy bands should be either the same as in the $ibis_binning$ or the bands created by the merger of those bands. The deconvolution is weighted by efficiency. Thus the weighting array W contains efficiency for normal pixels and 0 elsewhere.

A search for sources is performed in the deconvolved image. The list of sources found is created and the images are cleaned from the PSF ghosts.

In the deconvolved image, the pixel value at the source position is the total source flux in cts/sec units. It is calculated as if the source was in the fully coded field of view (FCFOV). One should NOT do any integration in the source region to estimate the flux from the image. The source flux estimation given in the source list is slightly different as the source flux is given at the fitted source position.

For each output energy band all images of the same type are combined into one mosaic image.

A search for the sources in the mosaic image is then performed and a list of sources found is created.

The current version of software allows you to create the mosaic separately from the main analysis (see Section 7.1 for the details).

The Parameter **PixSpread** sets whether the source flux is spread, or not in the mosaic image.

No-spread mosaic puts the whole input pixel count into one output map pixel. It permits better flux and signal-to-noise ratio approximation as each pixel count and variance is summed without any error. On the other hand it can give some undesired effects as double source peaks because of binning.

In the spread mosaic the input pixel count is spread between some number of output map pixels. This method is better for source position recovery but not so good for source flux estimation, because of the source peak height reduction. Furthermore, neighbouring pixels in the individual Science Window images are correlated and correct spread variance calculations should take this into account. But as exact calculation of the covariance matrix is too heavy for on-line analysis, the approximative variance formula is used.

The *ii_skyimage* parameters are given in the Table 18.

Table 18: *ii_skyimage* parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
OBS1_mask	mask	string	DOL of the MASK pattern fits file
			default: ''',
OBS1_deco	deco	string	DOL of the projected decoding pattern fits file
			default: ''''
OBS1_DataMode	DataMode	integer	Data Simulation mode
			possible values:
			0 - shadowgrams are treated
			1 – shadowgrams are simulated
			default: 0

OBS1_SearchMode	SearchMode	integer	 Source search mode possible values: 0 – search for all significant excesses 1 – search for all catalog sources 2 – search for K sources, where K= ToSearch 3 – search for all catalog sources plus K significant excesses, where K = ToSearch Parameter SearchMode= 1,3 concerns only images at a Science Window level. In the mosaic image <i>ii_skyimage</i> always looks for K = ToSearch sources.
OBS1_ToSearch	ToSearch	integer	Number of sources to be looked for.
OBS1_CleanMode	CleanMode	integer	Ghost cleaning possible values: 1 – source model lobes are subtracted from the decon- volved image -1 – no subtraction default: 1
OBS1_tungAtt	tungAtt	string	DOL of the Tungsten attenuation length fits file default: '' ''
OBS1_aluAtt	aluAtt	string	DOL of the Aluminium attenuation length fits file default: '.'.'
OBS1_leadAtt	leadAtt	string	DOL of the Lead attenuation length fits file default: '.'.'
OBS1_ScwType	ScwType	string	Type of Science Window to be treated possible values: ''POINTING'', 'SLEW'', ''OTHER'', ''ANY''
OBS1_DoPart2	DoPart2	integer	if 1 or absent \rightarrow do mosaic part. Set it to 0 if you don't want to produce a mosaic image.
OBS1_MapAlpha	MapAlpha	real	Mosaic map center [deg] default: 0, 0
OBS1_MapDelta	MapDelta	real	Mosaic map center [deg]
OBS1_MapSize	MapSize	real	Mosaic map radius [deg]
OBS1_PixSpread	PixSpread	integer	0 – no flux spread in mosaic
OBS1_MinCatSouSnr	MinCatSouSnr	real	default: 1 Software detects the catalog source only if its signal- to-noise ratio is higher then this value. This parameter has no meaning for the Science Window level images if OBS1_SearchMode =1,3. default: 6
OBS1_MinNewSouSnr	MinNewSouSnr	real	Software detects a new source only if its signal-to-noise ratio is higher then this value. default: 7
OBS1_SouFit	SouFit	integer	0 for fitting source position and 1 for fixed source po- sition in Scw fit default: 0

OBS1_ExtenType	ExtenType	integer	 exposure/residual maps possible values: 0 or absent - Residual Map at 4*n-th extension of ima idx 1 - Exposure Map only at 4*n-th extension of ima idx 2 - both Residual and Exposure maps at 4,5*n-th extension of ima idx 3 - one Expo map at the end of ima idx if ExtenType=1,2,3 the Mosaic Exposure Map at 4*n-th extension of mosa idx default: 0
OBS1_NegModels	NegModels	integer	0 or absent for no negative models ; 1 - negative models default: 0
OBS1_FastOpen	FastOpen	integer	if 1 then no CommonPreparePars default: 1

12.8.2 *sumhist*

PICsIT needs long integration times to produce a good image. Thus if *INTEGRAL* was stable during several Science Windows it is recommended to sum up PICsIT shadowgrams before the deconvolution. If the *ibis_science_analysis* parameter **staring** is set to **yes**, then *sumhist* checks that *INTEGRAL* was stable during all the Science Windows within the given limits and sums all the PICsIT shadowgrams into one.

Table 19:	sumhist parameters	included into	the main script.
-----------	--------------------	---------------	------------------

Name	Name	Type	Description
(in the main script)	(executable)		
tolerance	tol	real	tolerance for staring
			default: 0.0001

12.8.3 *ip_skyimage*

 $ip_skyimage$ performs deconvolution of the shadowgrams with the use of the balanced cross-correlation method described above, for more detailed explanations see Goldwurm et al. 2003, [10]. The weighting array W corresponds to the efficiency map. The values of the decoding G-array are taken from the IC file.

The *ip_skyimage* parameters included into the main script are given in the Table 20.

Table 20: *ip_skyimage* parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
PICSIT_detThr	detThr	real	Detection threshold for the automatic source location.
PICSIT_inCorVar	inCorVar	integer	Corrected variance to be used $[0=NO; 1=YES]$
PICSIT_outVarian	outVarian	integer	Variance maps in output $[0=NO; 1=YES]$
PICSIT_deco	maskDeco	string	DOL of the decoding pattern of PICsIT mask.

12.9 Spectral Analysis

12.9.1 *ii_spectra_extract*

ii_spectra_extract reads the shadowgram for each each input energy band.

For each output energy band, it :

- creates a model (PIF) for each source. The Pixel Illumination Factor (PIF) is a number between 0 and 1, which expresses the theoretical degree of illumination of each pixel in the detector plane of a coded-mask instrument. The PIF is calculated on the basis of the position and is normalized to 1 cts/pix source.
- fits all source intensities together with a background estimate to the data (shadowgrams)
- writes all source estimates to the spectra index

Finally, *ii_spectra_extract* writes to the PIF index one mean model per input catalogue source.

There is a possibility to chose one of several fitting metods. The recommended method is iterative imaginglike Least Squares fit (method 6). In the first step of this method the fit of each source separately in its coded zone is done. Next, in the second step, the calculation of reciprocal source contributions is performed. The first step is aimed to decrease the background modelling error influence. The second one, corresponds to the ghost cleaning. Please note that Maximum Likelihood method is not working in standard OSA configuration (with isdcmath package).

In the fitting process the source positions are considered to be exact. If the input background map is given , then the background map fitting is performed.

Name	Name	Type	Description
(in the main script)	(executable)		
SCW2_ISPE_idx_isgrResp	idx_isgrResp	string	DOL of the index of ISGRI response matrices
			default: '''
SCW2_ISPE_DataMode	DataMode	integer	Data Simulation mode
			possible values:
			0 real data
			1 simulated data
			default: 0
SCW2_ISPE_MethodInt	MethodInt	integer	Method to be applied for the pixel value interpolation
			default: 1
SCW2_ISPE_MethodFit	MethodFit	integer	Method to be applied for background and source in-
			tensity fitting
			possible values:
			MethodFit = 0 = > CHI2
			MethodFit = 1 = > Maximum Likelihood
			$MethodFit = 2 \implies Least Squares$
			MethodFit $= 6 ==>$ iterative imaging-like Least
			Squares fit
			default: 6
SCW2_ISPE_isgrUnifDol	isgrUnifDol	string	DOL of the ISGRI detector uniformity
			default: ''''
SCW2_ISPE_isgrBkgDol	isgrBkgDol	string	DOL of the ISGRI background model
			default: ''''
SCW2_ISPE_tungAtt	tungAtt	string	DOL of the Tungsten attenuation length fits file
			default: ''''
SCW2_ISPE_aluAtt	aluAtt	string	DOL of the Aluminium attenuation length fits file
			default: ''''
SCW2_ISPE_leadAtt	leadAtt	string	DOL of the Lead attenuation length fits file
			default: '' ''
rebinned_corrDol_spe	corrDol	string	DOL of the rebinned ISGRI off-axis corrections for
			spectrum default: '' ''

Table 21: $ii_spectra_extract$ parameters included into the main script.

$12.9.2 \quad ip_spectra_extract$

ip_spectra_extract reads the shadowgram for each each input energy band and the name and coordinates of the source for which a spectrum will be extracted. Since PICsIT operates in an energy range with a few sources, this executable works for one source only.

For each output energy band, it:

- creates a model (PIF) for the source. The Pixel Illumination Factor (PIF) is a number between 0 and 1, which expresses the theoretical degree of illumination of each pixel in the detector plane of a coded-mask instrument. The PIF is calculated on the basis of the position and is normalized to 1 cts/pix source. The present PIF is purely geometrical.
- by taking into account that there is one single source with ideal PIF, than the Equation at pag. L228 of Goldwurm et al. (2003) can be simplified. In this case, all the pixel counts, depending on the PIF values, are due to the selected single source. That is, deviations from the average counts in the observed shadowgrams are due entirely to the source.
- writes the source count rates in the selected energy bands and the PIF.

Table 22: $ip_spectra_extract$ parameters included into the main script.

Name	Name	Type	Description
(in the main script)	(executable)		
PICSIT_source_name	inName	string	Name of the source for extracting PICsIT spectrum
			default: ''''
PICSIT_source_RA	inRA	real	RA of the source for extracting PICsIT spectrum
			default: '' ''
PICSIT_source_DEC	inDEC	real	DEC of the source for extracting PICsIT spectrum
			default: '' ''

12.10 Timing Analysis

$12.10.1 \quad ii_lc_extract$

ii_lc_extract performs similar tasks to *ii_spectra_extract*, dealing with time bins instead of spectra ones.

Table 23:	Parameters	for the	$ii_lc_extract.$
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Name	Name	Type	Description
(in the main script)	(executable)		
ISGRI_mask	mask	string	DOL of the MASK pattern fits file
			default: ''''
SCW2_cat_for_extract	inCat	string	DOL of the subset of the source catalogue
		_	default: ''''
SCW2_ISPE_DataMode	DataMode	integer	Data Simulation mode
		_	$DataMode = 0 \implies OG$ shadowgrams are treated
			1 = > shadowgrams are simulated

SCW2_ISPE_MethodInt	MethodInt	integer	Method to be applied for the pixel value interpolation
			This parameter should have the same value as in the
			ii_shadow_build
			In the source model procedure
			MethodInt = 0 = > dead zone pixels will be filled with
			0
			1 = > dead zone pixels will be filled with mean de-
			tector value
			-1 => no dead zones
SCW2_ISPE_MethodFit	MethodFit	integer	Method to be applied for background and source in-
			tensity fitting
			default: 6
SCW2_ISPE_isgrUnifDol	isgrUnifDol	string	DOL of the ISGRI detector uniformity
			default: '' ''
SCW2_BKG_L_isgrBkgDol	isgrBkgDol	string	DOL of the ISGRI background model
			default: '' ''
SCW2_ISPE_tungAtt	tungAtt	string	DOL of the Tungsten attenuation length fits file
			default: '''
SCW2_ISPE_aluAtt	aluAtt	string	DOL of the Aluminium attenuation length fits file
			default: '''
SCW2_ISPE_leadAtt	leadAtt	string	DOL of the Lead attenuation length fits file
			default: '' ''
rebinned_corrDol_lc	corrDo	string	DOL of the rebinned ISGRI off-axis corrections for lc
			default: '' ''

$12.10.2 \quad ip_st_lc_extract$

This program builds PICsIT detector light curves and related errors starting from spectral timing data of PICsIT.

Table 24:	Parameters	for the	$ip_st_lc_extract.$
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Name	Type	Description
inGRP	string	DOL of the input group.
inCorSPT	string	DOL of the input Corrected SPT histogram.
inDead	string	DOL of the dead times.
outGRP	string	DOL of the output group.
outLCR	string	DOL of the light curves.

12.11 Summing up the results

$12.11.1 \quad ip_skymosaic$

 $ip_skymosaic$ creates a mosaic image of all the science windows within an Observation Group, and performs a source location

Name	Name	Type	Description
In the main script			
OBS2_detThr	detThr	real	Detection Threshold in Sigmas
			default: 3
OBS2_projSel	projSel	string	Selection of projection (default: TAN)

Table 25: *ip_skymosaic* parameters included into the main script.

12.12 Tools not included in the pipeline

$12.12.1 \quad mosaic_spec$

mosaic_spec is a tool that extracts a count spectrum at a given sky position from a set of mosaic sky images. Intensity could be measured in a specific or in the most significant pixel or derived from a gaussian fit with free or frozen position and width.

Note that if the significance of the point is less than five, then *mosaic_spec* will assign a non-zero value to its quality flag.

Name	Type	Description
hline DOL_inp	string	DOL of the group containing the input images.
-	Ŭ	default: ""
DOL		
DOL_out	string	DOL of the group containing the output images.
		default: "/
EXTNAME	string	EXTNAME of the input images
	5011118	default: "ISGR-MOSA-IMA"
DOL_idx	string	DOL of the index of input images.
		default: ""
DOI	<i>,</i> .	
DOL_spec	string	DOL of the index of output images.
		default:
ximg	real	h,0,0,10000,"Enter source x in pixels.
-		possible values: 0 10000
		default: 0
	,	
yımg	real	Enter source y in pixels.
		default: 0
ra	real	Enter source RA
		possible values: $-180 - 360$
		default: 0
dec	real	Enter source DEC
uce	1000	possible values: -90 90
		default: 0
posmode	integer	Position fitting mode
		0: position is always left free:
		1: freeze position as found in first energy band
		default: 1
widthmode	integer	Gaussian width fitting mode
		posible values: -1: radial width frozen as input
		1. x and y width left free
		2: x and y width fixed as found for first band
		default: 2

Table 26: *mosaic_spec* parameters.

outmode	integer	Choose data for output posible values: 1: intensity from the most significant pixel; 2: fitted intensity default: 2
psf	real	HWHM of PSF in arcmin possible values: 0.1 20 default: 6
size	integer	h,20,1,1000,"Enter source box half size in pixels on which the fit is performed. possible values: 1 1000 default: 20
back	boolean	Should a constant intensity background be considered in the fit default: no
allEnergies	boolean	Enter if the program has to select automatically energies default: yes
emin	string	Enter vector of energy bands minimum. default: "25 30 40"
chatty	integer	Enter reporting-level (10: includes MINUIT log). possible values: 0 10 default: 4

12.12.2 *ii_light*

For every PIF found in the Science Window, ii_light extracts simultaneously a lightcurve for each source and one light curve for background in all specified energy bands. Dead pixel, data gaps, off-axis correction, energy low threshold and illumination factors are taken into account.

The method used is a fit of hyperplane through the cloud of points formed by the number of counts versus PIF fraction for every source. In the case of one source this is just a linear regression. The intercept gives the background, while the slope gives the flux of the source in one fully illuminated pixel. Normalization is a number of counts in a perfect detector if source is on axis (half of all detector pixels illuminated). Care has been taken so that structure is compatible with HEASARC tools.

There are no limits on the size of time bin (up to 0.1s), and number of energy bands.

ii_light is not the official lightcurve extraction tool and should be used mainly to check relative variability of bright sources within a given Science Window, rather than for a long term absolute flux estimate.

Name	Type	Description
inSwg	string	DOL of the input Observation Group.
		default: ""
outSwg	string	DOL of the output Observation Group default: ""
outLC	string	Dol of the light curves file default: ""

Table 27: *ii_light* parameters.

context	string	DOL of the context where we can find low threshold default: ""
GTIname	string	Name of the GTI to be applied default: ""
select	string	Event selection condition default: ""
pifDOL	string	Dol of the PIF (or of the index of PIF) default: ""
deadDOL	string	Dol of the ISGR-DEAD-SCP" default: ""
corrDol	string	DOL of the isgri off-axis corrections default: ""
backDol	string	DOL of the background maps default: ""
delta_t	real	Time bin in seconds. possible values: 0.1 – 10000 default: 100
num_e	integer	Number of energy channels possible values: 1 – 10 default: 2
e_min	string	List of low energy boundaries default: "15 40"
e_max	string	List of high energy boundaries default: "40 300"

A Low Level Processing Data Products

A.1 Raw Data

Table 28 summarizes all Data Structures with the raw events measured by IBIS. The description of these Data Structures can be found below.

		5	D 1	<u> </u>
Observing	Type of event	Raw	Prepared	Corrected
mode		Data Structure	Data Structure	Data Structure
Photon-	ISGRI [I]	ISGR-EVTS-RAW	ISGR-EVTS-PRP	ISGR-EVTS-COR
	PICsIT single [PS]	PICS-SGLE-RAW	PICS-SGLE-PRP	PICS-SGLE-COR
by-	PICsIT multiple [PM]	PICS-MULE-RAW	PICS-MULE-PRP	PICS-MULE-COR
	Compton single [CS]	COMP-SGLE-RAW	COMP-SGLE-PRP	COMP-SGLE-COR
photon	Compton multiple [CM]	COMP-MULE-RAW	COMP-MULE-PRP	COMP-MULE-COR
Spectral				
Imaging	PICsIT single	PICS-SISH-RAW	PICS-SISH-PRP	PICS-SISH-COR
Histogram	PICsIT multiple	PICS-SIMH-RAW	PICS-SIMH-PRP	
Spectral				
Timing		PICS-SPTI-RAW	PICS-SPTI-PRP	
Histogram				

A.1.1 Photon-by-photon mode

In this mode Data Structures with similar contents are created for all types of events. Data Structures with names finished by **RAW** contain the information about the event itself. See Table 29 for details.

Data Structures with names finished by **PRW** and **SRW** contain some technical information about the structure of the telemetry packet and the Local On-Board Time (LOBT) of the events.

Column Name	Description	Event Type
DELTA_TIME	Delta time to previous event	I, PS, PM
RISE_TIME	Event rise-time (describes the shape of the registered	I, CS, CM
	pulse)	
ISGRI_PHA	Pulse height in the ISGRI layer	I, CS, CM
PICSIT_PHA	Pulse height in the PICsIT layer	PS, PM, CS, CM
ISGRI_Y	Y location in the ISGRI layer	I, CS, CM
ISGRI_Z	Z location in the ISGRI layer	I, CS, CM
PICSIT_Y	Y location in the PICsIT layer	PS, PM, CS, CM
PICSIT_Z	Z location in the PICsIT layer	PS, PM, CS, CM
CAL_FLAG	Calibration flag (1 - when neither of the events are	CS, CM
	calibration ones, only in this case the event is used in	
	the following analysis.)	
TIME_TAG	Compton delta time to previous event	CS, CM
DUMMY_COUNTER	Dummy counter - some technical information used to	CS, CM
	reconstruct the On-Board Time.	

Table 29: Contest of Photon-by-Photon Mode Raw Data.

A.1.2 PICsIT Standard Mode

In this mode the information is accumulated on-board and transmitted to the Earth in a kind of histograms. Spectral-image histograms are written to the Data Structures **PICS-SISH-RAW** and **PICS-SIMH-RAW** for single and multiple events correspondingly. These Data Structures represent a three dimensional table with total counts during integration time. The axes of the grid are directed along the following axes: AXIS 1 - Channel number, AXIS 2 - Y position, AXIS 3 - Z position.

Spectral-timing histograms are written to the Data Structure **PICS-SPTI-RAW** and contain the information on the number of the events accumulated in the up to 8 energy bins during a given time amount, see Table 30

Column Na	me Description
CELL_1	1st cell of the spectrum
CELL_2	2nd cell of the spectrum
CELL_3	3rd cell of the spectrum
CELL_4	4th cell of the spectrum
CELL_5	5th cell of the spectrum
CELL_6	6th cell of the spectrum
CELL_7	7th cell of the spectrum
CELL_8	8th cell of the spectrum

Table 30: Content of PICS-SPTI-RAW Data Structure.

The local on-board time and the channels definitions can be found in the **PICS-SPTI-PRW** Data Structure.

A.2 Prepared Data

The main task of the Science Window Pipeline is to prepare raw data for the following Scientific Analysis. It converts the housekeeping parameters into the physical units and makes some corrections and transformations of the raw data that are not included in Pre-Processing. The Summary of all the prepared Data Structures with scientific information can be found in Table 28. All these Data Structures has the only column **OB_TIME** with the on-board time.

B Instrument Characteristics used in Data Analysis.

B.1 Noisy Pixels

It is possible that with time some of the pixels of the detector start to produce an output not triggered by an income photon, i.e. to become "noisy". If the particular pixel countrate is too high relatively to the module countrate, then the on-board electronics switch it off. In ISGRI the noisy pixels can recover after being switched off for some time and disabled pixels are periodically reset to check their status. Data Structure **ISGR-SWIT-STA** with the list of noisy pixel switches in the ISGRI detector layer is produced during the pre-processing basing on the rise-time information and pixels light-curves transmitted, see Table 31.

Column Name	Description
ISGRI_Y	Y location in the ISGRI layer
ISGRI_Z	Z location in the ISGRI layer
OBT_DETECT	First time when the pixel can be declared as noisy
OBT_SWITCH	OBT when the pixel is switched off
NOIS_FLAG	Noisy flag

In PICsIT pixels cannot be recovered that easy. A PICsIT pixel will remain off once killed. Only if half of the detector (or so) will be off pixels will be attempted to turn on. The history of the disable pixels can be found in Data Structure **PICS-FALT-STA**, see Table 32.

Table 32:	Content of PI	CS-FALT-STA	. Data Structure
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Column Name	Description
PICSIT_Y	Y location in the PICsIT layer
PICSIT_Z	Z location in the PICsIT layer
OBT_DETECT	First Time when the pixel can be declared as noisy
OBT_FAULT	OBT when the pixel is switched off
FALT_FLAG	FLAG reflecting the changes of the pixel status.
	Possible values are ON-OFF and OFF-ON

B.2 Calibration Corrections

B.2.1 ISGRI

To perform a rise-time correction for each raw ISGRI event, the Rise-Time correction table is used (**ISGR-RISE-MOD** Data Structure, see details in Table 33). This table is composed of NUM_ENER lines, giving for each incident energy the correction factor **corr** for a given value of the rise-time.

Table 33: Content of ISGR-RISE-MOD Data Structure

Column Name	Description
ENERGY	Energy at which the gain-offset relationship is measured
CHANNEL	Channel at which the gain-offset relationship is measured
CORR	Rise-time correction for a given rise-time value $(0-127)$

The rise-time itself and the gain offset are kept at the **ISGR-OFFS-MOD** Data Structure.

Column Name	Description
AGAIN	Amplitude Gain
AOFFSET	Amplitude Offset
RTGAIN	Risetime Gain
RTOFFSET	Risetime Offset
PIXTYPE	Pixel type

B.2.2 PICsIT

Data Structure **PICS-ENER-MOD** is created by the PICsIT Automatic Calibration Analysis and contains parameters for gain and offset correction of PICsIT raw events. Content of this Data Structure is given in the Table 35. The values given in the Table 35 are normalized to the average values, given by the keywords **AVGAIN** for average gain value (in units keV/channel) and **AVOFFSET** for average offset value (in keV).

Table 35: Content of PICS-ENER-MOD Data Structure

Column Name	Description
PICSIT_Y	Y location in the PICsIT layer
PICSIT_Z	Z location in the PICsIT layer
GAIN	Gain
OFFSET	Offset

B.3 Limit Tables

Instrument GTIs depending on HK and other parameters are defined by a limit in a limit table **IBIS-GOOD-LIM**, see details in Table 36.

Column Name	Description	
PAR_NAME	Parameter name	
OBT_START	Start of validity of the limit values	
OBT_END	End of validity of the limit values	
MIN_VAL	Minimum value allowed	
MAX_VAL	Maximum value allowed	
GTI_NAME	Name of the group to which the parameter belongs	
SUB_ASSEMBLY	Identifier of the instrument sub-assembly	
CHECK_MODE	Modes in which the parameters must be checked	

Table 36: Content of IBIS-GOOD-LIM limit table.

B.4 Instrument Background

Table 37 lists Data Structures with the instrument background models.

Table 37: Instrument Background Model Data Structures.

Data Structure	Description
ISGR-BACK-BKG	ISGRI instrument background array.
ISGR-UNIF-BKG	ISGRI instrument background uniformity array.
PICS-SBAC-BKG	PICsIT instrument background array for single events.
PICS-SUNI-BKG	PICsIT instrument background uniformity array for single events.
PICS-MBAC-BKG	PICsIT instrument background array for multiple events.
PICS-MUNI-BKG	PICsIT instrument background uniformity array for multiple events.

COMP-SBAC-BKG	COMPTON instrument background array for single events.
COMP-SUNI-BKG	COMPTON instrument background uniformity array for single events.
COMP-MBAC-BKG	COMPTON instrument background array for multiple events.
COMP-MUNI-BKG	COMPTON instrument background uniformity array for multiple events.

For each type of Data Structures in Table 37 there is an Index. Its content is given in Table 38. Two last columns (RISE_MIN and RISE_MAX) are present only in **ISGR-BACK-BKG-IDX** and **ISGR-UNIF-BKG-IDX** Indexes.

Table 38:	Content	of Indexes	for Table	37 Data	Structures.
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Column Name	Description
VERSION	Version of the instrument characteristic file
VSTART	Start of validity time in IJD
VSTOP	End of validity time in IJD
E_MIN	Lower bound of the energy range
E_MAX	Upper bound of the energy range
EXPOSURE	Effective exposure time
VETO_THR	Veto threshold
METH_BKG	Method used to produce this background image
RISE_MIN	Minimum event rise time channel 0-127
RISE_MAX	Maximum event rise time channel 0-127

C Science Data Products

C.1 ibis_correction

This script converts photons energy into keV with the help of the calibration data. Table 39 summarises the output Data Structures.

Observing mode	Type of event	Corrected Data Structure
Photon –	ISGRI [I]	ISGR-EVTS-COR
	PICsIT single [PS]	PICS-SGLE-COR
by –	PICsIT multiple [PM]	PICS-MULE-COR
	Compton single [CS]	COMP-SGLE-COR
photon	Compton multiple [CM]	COMP-MULE-COR
Spectral-Imaging	PICsIT single	PICS-SISH-COR

Table 39: List of Data Structures produced at COR level

The content of the level COR Data Structures for the photon-by-photon mode is given in Table 40. The abbreviations for the events types are taken from Table 39.

Table 40: Content the level COR Data Structures for the photonby-photon mode.

Column Name	Description	Event Type
ISGRI_PI	corrected rise-time for ISGRI	I,CS,CM
ISGRI_ENERGY	Deposited energy in the ISGRI layer	I,CS,CM
PICSIT_ENERGY	Deposited energy in the PICsIT layer	PS,PM,CS,CM
SELECT_FLAG	Selection flag - shows whether the event was	I,CS,CM
	noisy (1) or not (0) .	

The **PICS-SISH-COR** Data Structure for spectral-imaging mode contains the total number of counts measured by each detection unit in each energy channels. The time information is in the index **PICS-SISH-COR-IDX**, see Table 41 for details.

Table 41: Content the PICS-SISH-COR-IDX Data Structure.

Column Name	Description
OBTSTART	OBT start of integration
OBTEND	OBT end of integration
HIST_NUM	Histogram number
HIST_ID	Histogram identifier
MEM_AREA	Histogram memory area

C.2 *ibis_gti*

This script builds Good Time Intervals from housekeeping data, information about satellite stability, and data gaps. The resulted GTIs are written to the **IBIS-GNRL-GTI**, see details in Table 42. Index of all **IBIS-GNRL-GTI** Data Structures for all categories is written to the **IBIS-GNRL-GTI-IDX**.

Table 42: Content of **IBIS-GNRL-GTI** Data Structures.

Column Name	Description
OBT_START	On-board time of start of the GTI
OBT_END	On-board time of end of the GTI

C.3 ibis_dead

This script calculates the dead times for ISGRI, PICsIT and Compton cases.

The results of the executables are written to Data Structures **ISGR-DEAD-SCP**, **PICS-DEAD-SCP** and **COMP-DEAD-SCP**. This Data Structures contains the information about the on-board time and the dead time for each module, see Tables 43 - 45.

Column Name	Description
OB_TIME	On-board time
II_DEADTIME_0	Dead time for module 0
II_DEADTIME_1	Dead time for module 1
II_DEADTIME_2	Dead time for module 2
II_DEADTIME_3	Dead time for module 3
II_DEADTIME_4	Dead time for module 4
II_DEADTIME_5	Dead time for module 5
II_DEADTIME_6	Dead time for module 6
II_DEADTIME_7	Dead time for module 7

Table 43: Content of ISGR-DEAD-SCP Data Structures.

Table 44: Content of **PICS-DEAD-SCP** Data Structures.

Column Name	Description
OB_TIME	On-board time
IP_DEADTIME_0	Dead time for Semi-module 0
IP_DEADTIME_1	Dead time for Semi-module 1
IP_DEADTIME_2	Dead time for Semi-module 2
IP_DEADTIME_3	Dead time for Semi-module 3
IP_DEADTIME_4	Dead time for Semi-module 4
IP_DEADTIME_5	Dead time for Semi-module 5
IP_DEADTIME_6	Dead time for Semi-module 6
IP_DEADTIME_7	Dead time for Semi-module 7
IP_DEADTIME_0	Dead time for Semi-module 8
IP_DEADTIME_1	Dead time for Semi-module 9
IP_DEADTIME_2	Dead time for Semi-module 10
IP_DEADTIME_3	Dead time for Semi-module 11
IP_DEADTIME_4	Dead time for Semi-module 12
IP_DEADTIME_5	Dead time for Semi-module 13
IP_DEADTIME_6	Dead time for Semi-module 14
IP_DEADTIME_7	Dead time for Semi-module 15

Table 45: Content of COMP-DEAD-SCP Data Structures.

Column Name	Description
OB_TIME	On-board time
IC_DEADTIME_0	Dead time for module 0
IC_DEADTIME_1	Dead time for module 1
IC_DEADTIME_2	Dead time for module 2
IC_DEADTIME_3	Dead time for module 3
IC_DEADTIME_4	Dead time for module 4
IC_DEADTIME_5	Dead time for module 5
IC_DEADTIME_6	Dead time for module 6
IC_DEADTIME_7	Dead time for module 7

C.4 *ibis_binning*

This script prepares IBIS data for scientific analysis. Its main function is to split the data into energy bins and time bins. For each time and energy ranges detector shadowgram and a corresponding efficiency shadowgram are created. Output Data Structures are listed in the Table 46. These Data Structures keep the position of each pixel and the total counts in given energy band during integration time or efficiency correspondingly. The boundaries of the energy and time bins can be found in the corresponding index Data Structures, see Table 47 for details, the abbreviations used in this table were introduced in Table 46.

${\rm C.4.1} \quad ii_shadow_build$

ii_shadow_build prepares ISGRI data for scientific analysis.

During the run of this executable Data Structures **ISGR-DETE-SHD** for a detector shadowgrams and **ISGR-EFFI-SHD** for ISGRI detector efficiency shadowgrams are filled. These Data Structures keep the position of each pixel and the total counts in given energy band during integration time or efficiency correspondingly.

During PICsIT analysis the same information about the PICsIT detector is written to **PICS-DETE-SHD** and **PICS-EFFI-SHD** Data Structures.

Instrument	Type of shadowgram	Output Data Structure
ISGRI	detector [ID]	ISGR-DETE-SHD
	efficiency [IE]	ISGR-EFFI-SHD
PICsIT	detector [PD]	PICS-DETE-SHD
	efficiency [PE]	PICS-EFFI-SHD

Table 46: List of Data Structures produced at BIN level

Cable 47: Content	of ****-*	****-SHD-IDX	Data	Structure
able 47: Content	OI	-SHD-IDA	Data	Structure

Column Name	Description	Shadowgram Type
ISDCLEVL	ISDC level of data processing	ID, IE, PD, PE
TFIRST	Time of the first data element	ID, IE, PD, PE
TLAST	Time of the last data element	ID, IE, PD, PE
TELAPSE	Total elapsed time of the data	ID, IE, PD, PE
CHANMIN	Lowest channel of the energy range	ID, IE
CHANMAX	Highest channel of the energy range	ID, IE
E_MIN	Lower bound of the energy range	ID, IE, PD, PE
$E_{-}MAX$	Upper bound of the energy range	ID, IE, PD, PE
BANDTYPE	Type of energy band	ID, IE
HIS_TYPE	Type of histogram data	PD
SHD_TYPE	Shadowgram type	PE

C.5 $cat_{-}extract$

The catalogue extraction selects the sources in the FOV from the reference catalogue. The output Data Structures **ISGR-SRCL-CAT** and **PICS-SRCL-CAT** have the same structure as the reference catalogue **GNRL-REFR-CAT**, see Table 48.

Table 48: Content of GNRL-REFR-CAT Data Structures.

Column Name	Description
SOURCE_ID	ISDC unique source identifier

DAY_ID	Modified Julian Date of source's first identification
NAME	One commonly used name for the source
CLASS	source classification code
RA_OBJ	Source right ascension in degrees
DEC_OBJ	Source declination in degrees
ERR_RAD	Error radius
SPA_MODL	Model for source spatial extension (point, disk, ellipse, square,
	gaussian, Bspline, etc)
SPA_NPAR	Number of parameters for source spatial extension
SPA_PARS	Parameters for source spatial extension
SPE_MODL	Model for source spectrum (XSPEC syntax)
SPE_NPAR	Number of parameters for source spectrum
SPE_PARS	Parameters for source spectrum
VAR_MODL	Model for source intensity variability (constant, sin, burst)
VAR_NPAR	Number of parameters for source intensity variability
VAR_PARS	Parameters for source intensity variability
COMMENTS	Comments
SPI_FLUX_1	SPI flux in the soft SPI energy band
SPI_FLUX_2	SPI flux in the hard SPI energy band
ISGR_FLUX_1	ISGRI flux in the soft ISGRI energy band
ISGR_FLUX_2	ISGRI flux in the hard ISGRI energy band
PICS_FLUX_1	PICsIT flux in the soft PICsIT energy band
PICS_FLUX_2	PICsIT flux in the hard PICsIT energy band
JEMX_FLUX_1	JEMX flux in the soft JEMX energy band
JEMX_FLUX_2	JEMX flux in the hard JEMX energy band
E_MIN	Lower energy boundaries
E_MAX	Upper energy boundaries
FLUX	Flux values
FLUX_ERR	Flux errors
SEL_FLAG	Source selection flag
FLAG	Generic flag

C.6 ibis_background_cor

This script combines executables performing the background correction in accordance with the chosen method.

As the output for each energy range for ISGRI shadowgrams of larger dimensions (corrected expanded shadowgram and corresponding variance and efficiency expanded shadowgrams) are written to **ISGR-CEXP-SHD**. The energy range is given in the keywords **E_MIN**, **E_MAX**, **CHAN_MIN**, **CHAN_MAX**, and a shadowgram type is given by **SHD_TYPE** keyword.

For PICsIT Data Structures **PICS-CEXP-SHD** and **PICS-VEXP-SHD** are filled for the corrected expanded detector shadowgram and the corrected expanded variance detector shadowgram. The energy range is again given in the keywords.

C.7 Image Analysis

$C.7.1 \quad ii_skyimage$

ii_skyimage deconvolves shadowgrams in the given energy bands. The index of deconvolved and cleaned images is written to **ISGR-SKY.-IMA-IDX** Data Structure. For each output energy band defined four images are created and attached to this index. The content of this Data Structure is given in the Table 49.

Column Name	Description
IMATYPE	Type of image. Possible values are:
	"IMAGE" - deconvolved and cleaned sky image
	"VARIANCE" - variance image
	"SIGNIFICANCE" - significance map
	"RESIDUAL" - difference between raw (deconvolved only) and
	cleaned image. "EXPOSURE" - true exposure map can be cre-
	ated.
CHANMIN	Lowest channel of the energy range
CHANMAX	Highest channel of the energy range
E_MIN	Lower bound of the energy range
E_MEAN	Mean energy of the energy range
E_MAX	Upper bound of the energy range
TFIRST	Time of the first data element
TLAST	Time of the last data element
TELAPSE	Total elapsed time of the data
EXPOSURE	Effective exposure time
CRVAL1	LONG at the reference value
CRVAL2	LAT at the reference value

Table 49: Content of **ISGR-SKY.-IMA-IDX** Data Structure.

For each deconvolved image the list of found sources is created (Data Structure **ISGR-SKY.-RES**) and attached to the **ISGR-SKY.-RES-IDX** Data Structure. See content of these Data Structures in Tables 51, 50.

Column Name	Description
NEW_SOURCE	New source flag (0 if old, 1 if new)
SOURCE_ID	ISDC unique source identifier
RA_OBJ	Source right ascension in degrees
DEC_OBJ	Source declination in degrees
Y_FIN	Y axis fine position of the source in pixels
Z_FIN	Z axis fine position of the source in pixels
FIN_YZ_ERR	Error of the fine position along the Y and Z axis in pixels
RA_FIN	Right Ascension of the fine position of the source
DEC_FIN	Declination of the fine position of the source
FIN_RD_ERR	Error of the fine position of the source in RA and DEC
FLUX	Flux values for a given energy band
FLUX_ERR	Flux errors for a given energy band
DETSIG	Source detection significance in ISGRI

Table 50: Content of **ISGR-SKY.-RES** Data Structure.

Table 51: Content of IS	SGR-SKYRES-IDX	Data Structure.
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Column Name	Description
TFIRST	Time of the first data element
TLAST	Time of the last data element
TELAPSE	Total elapsed time of the data
E_MIN	Lower bound of the energy range
E_MAX	Upper bound of the energy range

Search for the sources in the FOV is then performed and a list of sources found is created (Data Structure **ISGR-MOSA-RES** has the same structure as the **ISGR-SKY.-RES**, Table50). The index of lists of sources found in each map can be found in Data Structure **ISGR-MOSA-RES-IDX** (structure analogous

to the one described in the Table 51).

The output catalog containing the sources description is written to the data structure **ISGR-SRCL-RES**. This Data Structure has structure similar to input catalog **ISGR-SRCL-CAT**. Each row of this Data Structure contains the description of a source in the OG FOV. If it was present in the input catalog all the input info is copied. For each source found during the analysis the new info is added, see Table 52

The DETSIG in ISGR-SRCL-RES is calculated from the results of the mosaic image and the results of the analysis at Science Window level

$$detsig = \sqrt{\Sigma detsig_i^2},$$

where i is for all energy bands and all Science Windows and mosaic image

For the FLUX and err_flux of ISGR-SRCL-RES you have:

$$flux = \frac{\Sigma flux_i}{N}$$

$$fluxerr = \frac{\Sigma fluxerr_i}{N}$$

where i - for all Science Windows and a mosaic image for a given energy band

Table 52: New information added to the **ISGR-SRCL-RES** Data Structure.

Column Name	Description
RA_FIN	Right Ascension of the fine position of the source
DEC_FIN	Declination of the fine position of the source
FIN_RD_ERR	Error of the fine position of the source in RA and DEC
SCW_NUM_C	number of times that the source was in ScW FOV
	If $SCW_NUM_C = 0$ then it is a new source
SCW_NUM_F	number of times that the source was found in ScW image
OG_NUM	number of times that the source was found in OG image

With the help of stand alone program *src_collect* it si possible to collect results from different science windows. Results are written to the **ISGR-OBS.-RES** Data Structure, see Table 53.

Table 53: Content of the ISGR-OBS.-RES Data Structure.

Column Name	Description	
SWID	Science Window identifier	
SOURCE_ID	ISDC unique source identifier	
NAME	One commonly used name for the source	
RA_FIN	Source right ascension in degrees	
DEC_FIN	Source declination in degrees	
FIN_RD_ERR	Error of the fine position of the source in RA and DEC	
Y_FIN	Y axis fine position of the source in pixels	
Z_FIN	Z axis fine position of the source in pixels	
FIN_YZ_ERR	Error of the fine position along Y and Z axis in pixels	
DETSIG	Source detection significance in ISGRI	
E_MIN	Lower energy boundaries	
E_MAX	Upper energy boundaries	
FLUX	Flux values	
FLUX_ERR	Flux errors	
DEADC	Mean deadtime (and greyfilter) correction factor	
EXPOSURE	Mean exposure time over the detector plane	
TSTART	Start time of the observation (IJD)	
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TSTOP	End time of the observation (IJD)	

C.7.2 $ip_{-skyimage}$

 $ip_skyimage$ performs deconvolution of the PICsIT shadowgrams with the use of the balanced crosscorrelation method described in Goldwurm et al. 2003 [10], see also Section 12.8. The values of the decoding *G*-array are taken from **PICS-DECO-MOD** Data Structure.

The output Data Structure **PICS-SKY.-IMA** store either a cartesian (CAR) or a tangential (TAN) projection of the celestial sphere. The index of deconvolved images is written to the **PICS-SKY.-IMA-IDX** Data Structure, it has the same content as **ISGR-SKY.-IMA-IDX** (Table 49). IMATYPE can be either "IMAGE", either "VARIANCE", or "SIGNIFICANCE".

The list of sources that were found during the analysis are written to the **PICS-SKY.-RES**

C.8 Spectral Analysis

C.8.1 $ii_spectra_extract$

ii_spectra_extract reads the shadowgram for each input energy band and creates a spectrum for background and each source from isgri_srcl_res.fits file. The result is written to the ISGR-EVTS-SPE and ISGR-EVTS-SPE-IDX Data Structures.

Column Name	Description	
CHANNEL	Channel number	
RATE	Countrate in the given channel	
SYS_ERR	Systematic error	
STAT_ERR	Statistical error	
QUALITY	Quality flag	
GROUPING	Grouping flag	

Table 54: Content of the ISGR-EVTS-SPE Data Structure.

Table 55: Content of the **ISGR-EVTS-SPE-IDX** Data Structure.

Column Name	Description	
SOURCEID	ISDC unique source identifier	
RA_OBJ	Source right ascension in degrees	
DEC_OBJ	Source declination in degrees	
TFIRST	Time of the first data element	
TLAST	Time of the last data element	
TELAPSE	Total elapsed time of the data	

C.9 Timing Analysis

${\rm C.9.1} \quad ip_st_lc_extract$

This program build light curves and related errors starting from spectral timing data of PICsIT. The content of the filled Data Structures is given in the Tables 56, 57.

Table 56:	$\operatorname{Content}$	of the	PICS-EVTS-LCR-IDX	Data	Struc-
ture.					

Column Name	Description
SOURCEID	ISDC unique source identifier
CHANMIN	Lowest channel of the energy range
CHANMAX	Highest channel of the energy range
E_MIN	Lower bound of the energy range
E_MAX	Upper bound of the energy range
PICSMODE	Mode of the PICsIT detector layer

Table 57: Content of the **PICS-EVTS-LCR** Data Structure.

Column Name	Description	
TIME	Time of measurement for the bin	
TOT_COUNTS	Total counts of the source region	
BACKV	Background counts scaled to the source region	
BACKE	Background count errors	
ERROR	Net count error in the source region	
RATE	Countrate in the given energy band	
FRACEXP	Fraction of integration bin time for exposure correction	

C.9.2 $ii_{-}lc_{-}extract$

For all sources from the isgri_srcl_res.fits file, *ii_lc_extract* extracts the ISGRI lightcurves. The results are written to the ISGR-SRC.-LCR and ISGR-SRC.-LCR-IDX Data Structures. ISGR-SRC.-LCR has the same structure as PICS-EVTS-LCR, see Table 57

Table 58: C	Content of the ISGF	R-SRCLCR-II)X Data Structure.
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Column Name Description	
SOURCEID ISDC unique source identifier	
RA_OBJ Source right ascension in degrees	
DEC_OBJ Source declination in degrees	
CHANMIN Lowest channel of the energy range	
CHANMAX	Highest channel of the energy range
E_MIN Lower bound of the energy range	
E_MAX	Upper bound of the energy range

C.9.3 Timing Analysis without the deconvolution

During the run of stand alone program $evts_extract$ two data structures **GNRL-EVTS-GTI** and **GNRL-EVTS-LST** are filled.

GNRL-EVTS-LST (Table 59) combines all available information for photon-by-photon events from different instruments during a given time interval.

Column Name Description	
DETY Y location in the detector layer (offset from center)	
DETZ	Z location in the detector layer (offset from center)
ENERGY	Energy deposited by the event
EVNT_TYPE	Type and origin of event (bit coded)
TIME	Time of event in INTEGRAL Julian Date units
TIMEDEL	Uncertainty of time stamp
DEADC	Dead time correction factor
BARYTIME_N	Barycenter time for source number N
PIF_N	Pixel Illumination Factor for source number N
AREASCAL_N	Nominal effective area for source number N

Table 59: Content of the **GNRL-EVTS-LST** Data Structure.

 \mathbf{GNRL} - \mathbf{EVTS} - \mathbf{GTI} (Table 60) contains good time intervals for selecting events.

Table 60: Content of the GNRL-EVTS-GTI I	Data Structure.
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Column Name	Description		
START IJD of start of the GTI			
STOP	IJD of end of the GTI		
OBT_START On-board time of start of the GTI			
OBT_END On-board time of end of the GTI			
UTC_START UTC of start of the GTI			
UTC_END	UTC of end of the GTI		

D List of $ibis_science_analysis$ parameters

Name	Туре	Description
	Ge	neral parameters
ogDOL	string	DOL of the Observational Group to be analyzed. default: ''og_ibis.fits[1]''
startLevel	string	Analysis level at which the analysis begins. The names of the possible analysis levels are given in section 5. Possible values: 'COR'' - 'COMP''. Also 'CLEAN'' value is possible. default: 'COR''
endLevel	string	Analysis level at which analysis finishes. The names of the possible analysis levels are given in the section 5. possible values: COR - CLEAN. We recommend you to proceed by steps, as described in the cookbook. default: "IMA2"
staring	boolean	Enter yes if it was a staring observation. It has an influence on PICsIT analysis - for staring observation all shadowgrams are summed before the deconvolution. default: no
tolerance	real	tolerance for staring. In case of staring=yes the check is performed that during the Science Windows within the Observation Group the spacecraft was stable within the given tolerance. default: 0.0001
sum_spectra	boolean	Do the summing of spectra? default: "no"
CAT_refCat	string	DOL of Reference Catalog default: '`\$ISDC_REF_CAT[ISGRI_FLAG==1]''
CAT_usrCat	string	DOL of User Catalog (CURRENTLY IG- NORED!!!)
chatter	integer	Verbosity level possible values: $0 - 5$, $0 = \text{errors only}$, $1 = \text{warnings}$, $2 = \text{normal}$, default: 2
IC_Group	string	DOL of Instrument Characteristics master group default: '`\$REP_BASE_PROD/idx/ic/ic_master_file.fits[1]''
IC_Alias	string	Selection alias for Instrument Characteristics default: ''OSA''

Table 61: $ibis_science_analysis$ parameters description. Query parameters are marked with bold font.

corrDol	string	DOL of the isgri off-axis corrections default: ""
rebinned_corrDol_ima	string	DOL of the rebinned is gri off-axis corrections for imaging default: ""
rebinned_corrDol_spe	string	DOL of the rebinned is gri off-axis corrections for spectrum default: ""
rebinned_corrDol_lc	string	DOL of the rebinned is gri off-axis corrections for lc default: ""
rebinned_backDol_ima	string	DOL of the rebinned is gri back corrections for imaging default: ""
rebinned_backDol_spe	string	DOL of the rebinned is gri back corrections for spectrum default: ""
rebinned_backDol_lc	string	DOL of the rebinned is gri back corrections for lc default: ""
GENERAL_clobber	boolean	Clobber existing output data? default: YES
GENERAL_levelList	boolean	List of all levels default: cor,gti,dead,bin_i,bkg_i,cat_i,ima,ima2,bin_S,spe,lcr,comp,clean
Parameters t	o define w	hich part of data should be analyzed
SWITCH_disableIsgri	boolean	Disable ISGRI analysis? possible values: YES, NO default: NO
SWITCH_disablePICsIT	boolean	Disable PICsIT analysis? possible values: YES, NO default: YES
SWITCH_disableCompton	boolean	Disable Compton analysis? possible values: YES, NO default: YES
SWITCH_osimData	boolean	Data are from simulator? possible values: YES, NO default: NO
Parameters	for ibis bii	nning and ibis image reconstruction
IBIS_II_ChanNum	integer	Number of output energy bands for ISGRI possible values: $-1 - 10$ value -1 is for more than 10 energy bands, in this case IBIS_II_inEnergyValues should be set. default: 4

IBIS_II_E_band_min	string	List of lower limits of output energy bands (keV) for ISGRI default: ''20 40 60 100''
IBIS_II_E_band_max	string	List of upper limits of output energy bands (keV) for ISGRI default: ''40 60 100 200''
IBIS_II_inEnergyValues	string	DOL of the energy values when required. default: '' ''
default:0		
IBIS_IPS_corrPDH	integer	Packets threshold for partially downloaded his- tograms. default:0
IBIS_IPS_ChanNum	integer	Number of Energy bands for PICsIT in standard mode. possible values: 0 – 300 You should set it to 0 to build automatically the only binning for which background maps are present. It corresponds to the following energy ranges (keV): 203 -252, 252 - 329, 329 - 455, 455 - 655, 655 - 1057, 1057 - 1841, 1841 - 3570, 3570 - 6510. default: 0
IBIS_IPS_E_band_min_s	string	List of lower energy boundary for PICsIT in standard mode for SINGLE. Do not touch this parameter! default: "170 600 1000"
IBIS_IPS_E_band_max_s	string	List of higher energy boundary for PICsIT in standard mode for SINGLE. Do not touch this parameter! default: "600 1000 10000"
IBIS_IPS_E_band_min_m	string	List of lower energy boundary for PICsIT in standard mode for MULTIPLE. Do not touch this parameter! default: "170 600 1000"
IBIS_IPS_E_band_max_m	string	List of higher energy boundary for PICsIT in standard mode for MULTIPLE. Do not touch this parameter! default: "600 1000 10000"
IBIS_IP_ChanNum	integer	Number of Energy bands for PICsIT in photon-by- photon mode. possible values: 1 – 300 Do not touch this parameter! default: 3
IBIS_IP_E_band_min_s	string	List of lower energy boundary (single) Do not touch this parameter! default (keV): ''175 600 1000''

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IBIS_IP_E_band_max_s	string	List of higher energy boundary (single) Do not touch this parameter! default (keV): ''600 1000 10000''
IBIS_IP_E_band_min_m	string	List of lower energy boundary (multiple) Do not touch this parameter! default (keV): ''350 600 1000''
IBIS_IP_E_band_max_m	string	List of higher energy boundary (multiple) Do not touch this parameter! default (keV): ''600 1000 13500''
IBIS_NoisyDetMethod	integer	Noisy Pixel detection method possible values: 0 (only normal noisy detection), (add also spectral noisy detection) default: 0
These parameters are re	ecommende	ed by the IBIS team, better not to change them.
IBIS_min_rise	integer	Minimum rise time. default: 7
IBIS_max_rise	integer	Maximum rise time. default:90
IBIS_P_convFact	real	Conversion factor channel-to-energy. default: 7.0
Р	arameters	specific to SCW1 pipeline.
SCW1_ISGRI_event_select	string	CFITSIO event selection string default: ''''
$SCW1_GTI_LimitTable$	string	DOL of the limit table for GTIs default: '' ''
$SCW1_GTI_attTolerance$	real	Accepted attitude variability [arc min] possible values: 0 – 180.0 default: 0.5
$SCW1_GTI_gtiUserP$	string	DOL of the user GTI table for PICsIT default: '' ''
$SCW1_GTI_gtiUserI$	string	DOL of the user GTI table for ISGRI default: '' ''
SCW1_GTL_TimeFormat	string	Time format to be used possible values: ''IJD'', ''UTC'', ''OBT'' default: ''IJD''
SCW1_GTI_Accuracy	string	Used accuracy for OBT to IJD conversion and vic versa possible values: '`any'', '`inaccurate'' '`accurate'' default: '`any''

SCW1_GTI_SCP	string	Names of spacecraft GTIs to be merged for PICsIT
		default: '' ''
SCW1_GTL_SCI	string	Names of spacecraft GTIs to be merged for ISGRI default: '' ''
SCW1_GTI_PICsIT	string	GTIs to be merged for PICsIT default: ' 'VETO ATTITUDE P_SGLE_DATA_GAPS
		P_MULE_DATA_GAPS'
SCW1_GTI_ISGRI	string	GTIs to be merged for ISGRI default: ''VETO ATTITUDE ISGRI_DATA_GAPS''
SCW1_GTI_BTI_Dol	string	DOL of a bad time interval table (GNRL-INTL-BTI) default: '' ''
$SCW1_GTI_BTI_Names$	string	Input BTI names to be considered default: '' ''
$SCW1_ICOR_idxSwitch$	string	DOL of the index of pixels switches list. default: '' ''
SCW1_ICOR_GODOL	string	DOL of the gain-offset table default: '' ''
SCW1_ICOR_riseDOL	string	DOL of the rise time correction table default: '' ''
SCW1_ICOR_icDOL	string	DOL of the IC file for calibration parameters default: '' ''
$SCW1_ICOR_probShot$	real	Probability of shot time decay. default: 0.01
SCW1_PCOR_enerDOL	string	DOL of the energy correction table (PICsIT) default: '' ''
SCW1_BIN_cleanTrk	integer	Cleaning of cosmic-ray induced events. possible values: 0 = No 1 = Yes default: 1
SCW1_veto_mod	string	DOL of the IC file for VETO model and width of Compton window default: '' ''
SCW1_BIN_L_idxNoisy	string	DOL of the index of noisy maps. default: '' ''
SCW1_BIN_LidxLowThre	string	index of Low Threshold default: '' ''
SCW1_BIN_P_inDead	string	DOL of the dead time data structure. default: '' ''

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SCW1_BIN_P_inGTI	string	DOL of the good time interval data structure. default: '' ''
SCW1_BIN_P_HepiLut	string	DOL of the hepi lut default: '' ''
SCW1_BKG_divide	boolean	Divide by efficiency? Do not touch it! default: no
SCW1_BKG_badpix	boolean	Remove bad pixels? Do not touch it! default: yes
$SCW1_BKG_flatmodule$	boolean	flatten modules? Do not touch it! default: no
SCW1_BKG_L_isgrUnifDol	string	DOL of the ISGRI detector uniformity possible values: DOL for a specific model ''-'' - no uniformity correction '' '' - DOL is taken from the IC tree default: ''-''
SCW1_BKG_L_isgrBkgDol	string	DOL of the ISGRI background model possible values: DOL for a specific model ''-'' - no background subtraction '' '' - DOL is taken from the IC tree default: '' ''
$SCW1_BKG_I_method_cor$	string	Method to be applied for background removal default: ''0''
$SCW1_BKG_I_method_int$	string	Method to be applied for the pixel value interpolation default: ''1''
SCW1_BKG_P_method	integer	Method applied in filling gaps possible values: 0 - model (in cts) is normalized by the time of obser- vation 1 - model (in cts) is normalized by the mean count value default: 0
$SCW1_BKG_picsSUnifDOL$	string	DOL of the PICsIT Detector Uniformity model (sin- gle). default: ''-''
$SCW1_BKG_picsSBkgDOL$	string	DOL of the PICsIT Detector Background model (single). default: '' ''
SCW1_BKG_picsMUnifDOL	string	DOL of the PICsIT Detector Uniformity model (mul- tiple). default: ''-''

SCW1_BKG_picsMBkgDOL	string	DOL of the PICsIT Detector Background model (mul- tiple). default: '' ''
P	arameters :	specific to OBS1 pipeline.
	ii_sk	yimage parameters
OBS1_DataMode	integer	Data Simulation mode possible values: 0 - shadowgrams are treated 1 - shadowgrams are simulated default: 0
rebin_slope	real	Spectral slope to be used default: -2
rebin_arfDol	string	DOL of the ISGRI arf for rebinning. default: '' ''
OBS1_SearchMode	integer	Source search mode possible values for the Science Window analysis: 1 - search for all catalog sources 2 - search for K sources, where K= ToSearch 3 - search for all catalog sources plus for K significant excesses, where K = ToSearch For <i>all</i> the above, when the mosaic is created, the software looks for K = ToSearch sources. default: 2
OBS1_ToSearch	integer	Number of sources to be looked for default: 15
OBS1_CleanMode	integer	Ghost cleaning possible values: 1 - source model lobes are subtracted from the decon- volved image -1 - no subtraction default: 1
OBS1_ExtenType	integer	Save exposure/residual maps possible values: absent, 0 - 3, 0 or absent - no action 1 - true exposure maps are saved in 4*n extension of isgri_sky_ima.fits instead of residual images 2 - true exposure maps are saved in the 5*n extension of isgri_sky_ima.fits after the residual images 3 - one true exposure map is created for the last energy band and saved in the last extension of is- gri_sky_ima.fits default: 0
OBS1_NegModels	integer	0 or absent for no negative models ; 1 - negative models default: 0
OBS1_FastOpen	integer	if 1 then no CommonPreparePars default: 1

ISGRI_mask	string	DOL of the MASK pattern fits file default: '' ''
OBS1_deco	string	DOL of the projected decoding pattern fits file default: '' ''
OBS1_tungAtt	string	DOL of the Tungsten attenuation length fits file default: '' ''
OBS1_aluAtt	string	DOL of the Aluminium attenuation length fits file default: ''''
OBS1_leadAtt	string	DOL of the Lead attenuation length fits file default: ''''
OBS1_covrMod	string	DOL of the covariance fits file default:
OBS1_ScwType	string	Type of Science Window to be treated possible values: ''POINTING'', ''SLEW'', ''OTHER'', ''ANY'' default: ''POINTING''
OBS1_DoPart2	integer	if 1 or absent \rightarrow do mosaic part. Set it to 0 if don't want to produce a mosaic image. Set it to 2 if you want to produce a mosaic image from different existing runs. default: 1
OBS1_MapAlpha	real	Mosaic map center [deg] default: 0.0
OBS1_MapDelta	real	Mosaic map center [deg] default: 0.0
OBS1_MapSize	real	Mosaic map radius [deg] default: 40.0
OBS1_PixSpread	integer	0 – no flux spread in mosaic default: 1
OBS1_MinCatSouSnr	real	The software detects a catalog source only if its signal- to-noise ratio is higher than this value. default: 6
OBS1_MinNewSouSnr	real	The software detects a catalog source only if its signal- to-noise ratio is higher than this value. default: 7
OBS1_SouFit	integer	defines whether to fit or not the source position 0 - for fitting source position in Scw fit 1 - for fixed source position in Scw fit default: 0
PICSIT_detThr	real	Detection Threshold

PICSIT_inCorVar	integer	Corrected variance to be used [0=NO; 1=YES] default: 1		
PICSIT_outVarian	integer	Variance maps in output [0=NO; 1=YES] default: 0		
cat_extract	parameter	s, see Section 12.7 for more details.		
OBS1_CAT_radiusMin	string	Low limit for the position selection. default: ''0''		
OBS1_CAT_radiusMax	string	High limit for the position selection. default: ''20''		
OBS1_CAT_fluxDef	string	Column used for flux selection. default: '' ''		
OBS1_CAT_fluxMin	string	Low limit for flux selection. default: '' ''		
OBS1_CAT_fluxMax	string	High limit for flux selection. default: '' ''		
OBS1_CAT_class	string	Select the object by class. default: '' ''		
OBS1_CAT_date	real	IJD for the public data. default: -1		
Parameters specific to SCW2 pipeline.				
PICSIT_deco	string	DOL of the MASK decoding fits file default: '' ''		
SCW2_cat_for_extract	string	Catalog for spectral and lightcurve extraction (if empty then it is taken from the imaging result of the Science Window) default: '' ''		
SCW2_racolumn	string	Name of the column where to get Ra default: ''RA_FIN''		
SCW2_deccolumn	string	Name of the column where to get Dec default: ''DEC_FIN''		
SCW2_catalog	string	Catalog for PICsIT imaging default: '' ''		
SCW2_PIF_filter	string	filter to apply on the source default: '' ''		
$SCW2_ISGRI_event_select$	string	CFITSIO event selection string default: '' ''		
parameter for building light curves from PICsIT spectral timing data				
Parameters for spectra extraction.				

IBIS_SI_ChanNum	integer	Number of Channels possible values: $-1 - 10$, -1 = take from ISGR-EBDS-MOD structure default: -1
IBIS_SI_E_band_min	string	List of lower limits of output energy bands default: '' ''
IBIS_SI_E_band_max	string	List of upper limits of output energy bands default: '' ''
IBIS_SI_inEnergyValues	string	DOL of the energy values when required. default: '' ''
PICSIT_source_name	string	Name of the source for extracting PICsIT spectrum default: '' ''
PICSIT_source_RA	real	RA of the source for extracting PICsIT spectrum default: '' ''
PICSIT_source_DEC	real	DEC of the source for extracting PICsIT spectrum default: '' ''
IBIS_SS_inEnergyValues	string	DOL of the energy values for single events default: '' ''
$IBIS_SM_inEnergyValues$	string	DOL of the energy values for multiple events default: '' ''
IBIS_SP_ChanNum	integer	Number of Channels possible values: 0 – 300 default: 51
IBIS_SP_E_band_min_s	string	List of lower limits of output energy bands (single) default: '' ''
IBIS_SP_E_band_max_s	string	List of upper limits of output energy bands (single) default: '' ''
IBIS_SP_E_band_min_m	string	List of lower limits of output energy bands (multiple) default: '' ''
IBIS_SP_E_band_max_m	string	List of upper limits of output energy bands (multiple) default: '' ''
IBIS_SPS_ChanNum	integer	Number of Channels possible values: 0 – 300 default: 51
IBIS_SPS_E_band_min_s	string	List of lower limits of output energy bands for SINGLE default: '' ''
IBIS_SPS_E_band_max_s	string	List of upper limits of output energy bands for SIN- GLE default: '' ''

IBIS_SPS_E_band_min_m	string	List of lower limits of output energy bands for multiple default: '' ''
IBIS_SPS_E_band_max_m	string	List of upper limits of output energy bands for multiple default: '' ''
$SCW2_ISPE_idx_isgrResp$	string	DOL of the index of ISGRI response matrices default: ''''
SCW2_ISPE_isgrarfDOL	string	DOL of the ARFs for XSPEC default: ''''
SCW2_ISPE_DataMode	integer	Data Simulation mode default: 0
$SCW2_ISPE_MethodInt$	integer	Method to be applied for the pixel value interpolation default: 1
$SCW2_ISPE_MethodFit$	integer	Method to be applied for background and source in- tensity fitting default: 6
$SCW2_ISPE_isgrUnifDol$	string	DOL of the ISGRI detector uniformity default: ''''
$SCW2_ISPE_tungAtt$	string	DOL of the Tungsten attenuation length fits file default: ''''
$SCW2_ISPE_aluAtt$	string	DOL of the Aluminium attenuation length fits file default: ''''
$SCW2_ISPE_leadAtt$	string	DOL of the Lead attenuation length fits file default: ''''
SCW2_BIN_cleanTrk	integer	Cleaning of cosmic-ray induced events. possible values: 0 = No 1 = Yes default: 0
SCW2_BIN_L_idxNoisy	string	DOL of the index of noisy maps. default: ''''
SCW2_BIN_L_idxLowThre	string	index of Low Threshold default: ''''
SCW2_BIN_P_inDead	string	DOL of the dead time data structure. default: ''''
SCW2_BIN_P_inGTI	string	DOL of the good time interval data structure. default: '''
SCW2_BIN_P_HepiLut	string	DOL of the hepi lut default: ''''

string	DOL of the isgri bckg model or - if you want none or empty if you want DOL automatic from the IC tree default: ''''
boolean	Divide by efficiency default: no
boolean	Remove bad pixels default: yes
boolean	flatten modules default: no
string	Method to be applied for background removal default: ''0''
string	Method to be applied for the pixel value interpolation default: ''1''
integer	Method applied in filling gaps possible values: 0 – 2 default: 1
string	DOL of the PICsIT Detector Uniformity model (sin- gle). default: '''
string	DOL of the PICsIT Detector Background model (sin- gle). default: ''-''
string	DOL of the PICsIT Detector Uniformity model (mul- tiple). default: ''''
string	DOL of the PICsIT Detector Background model (mul- tiple). default: ''-''
 Parameters	for lightcurve extraction.
string	Event selection condition default: '''
real	Time bin in seconds. possible values: 0.1 – 10000 default: 100
integer	Number of energy channels possible values: 1 – 10 default: 4
string	List of low energy boundaries default: "20 40 60 100"
string	List of high energy boundaries default: "40 60 100 200"
	string boolean boolean boolean string string integer string string string string string string string integer string string

OBS2_detThr	real	Detection Threshold (in sigmas). default: 3.0
OBS2_projSel	string	Selection of projection (default: TAN) default: TAN
OBS2_imgSel	string	Selection criteria of images. default: "EVT_TYPE=='SINGLE' && E_MIN==252 && E_MAX==336"

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