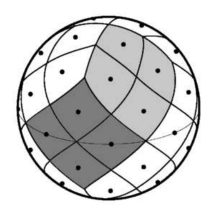
HEALPix Fortran90 Subroutines Overview



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Abstract: This document is an overview of the **HEALPix** For-

tran90 subroutines.

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Conventions

Here we list some conventions which are used in this document.

*	Fortran90 allows generic names which refer to several specific subroutines. Which one of the specific routines is called depends on the type and rank of the arguments supplied in the call. We tag generic names with a \ast in this document.
$ m N_{side}$	$\mathbf{HEALPix} \text{ resolution parameter} \text{ see the } \mathbf{HEALPix} \text{ Primer}.$
map	We use the word "map" referring to a function, defined on the set of all HEALPix pixels.
θ	The polar angle or colatitude on the sphere, ranging from 0 at the North Pole to π at the South Pole.
ϕ	The azimuthal angle on the sphere, $\phi \in [0, 2\pi[$.

Changes between release 2.13 and 2.14

• In alm2map_der routine, a numerical bug affecting the accuracy of the Stokes parameter derivatives $\partial X/\partial\theta$, $\partial^2 X/(\partial\theta\partial\phi\sin\theta)$, $\partial^2 X/\partial\theta^2$, for X=Q,U has been corrected. See "Fortran Facilities" Appendix for details.

Changes between release 2.0 and 2.13

- New functions in version 2.13:
 - get_healpix_data_dir, get_healpix_main_dir, get_healpix_test_dir return full path to HEALPix directories.
- New routines in version 2.10:
 - alm2map_spin: synthesis of maps of arbitrary spin
 - map2alm_iterative: iterative analysis of map
 - map2alm_spin: analysis of maps of arbitrary spin
 - healpix_modules: meta-module
 - write_minimal_header: routine to write minimal FITS header

 parse_check_unused: prints out parameters present in parameter file but not used by the code.

• Improved routines:

- query_strip: the inclusive option now returns *all* (and only) the pixels overlapping, even partially, with the strip
- query_disc: when the disc center is on one of the poles, only the pixels overlapping with the disc are now returned.
- remove_dipole: can now deal with non-uniform pixel weights.
- parse_init: silent mode
- parse_string: can expand environment variables (\${XXX}) and leading ~/

Changes between release 1.2 and 2.0

Some new features have been added

- Most routines dealing with maps and $a_{\ell m}$ (eg, create_alm, map2alm, alm2map, convert_inplace, convert_nest2ring, udgrade_nest, udgrade_ring) or inputting or outputting data (read_*, write_*) now accept both single and double precision arguments.
- The routines map2alm and remove_dipole can now deal with non-symmetric azimuthal cut sky. For backward compatibility, the former calling sequence is still accepted.
- most routines are now parallelized with OpenMP (for shared memory architecture), and some of them are also parallelized with MPI (for distributed memory architecture)

Some new routines have been introduced since version 1.2, as listed below.

- New routines in version 2.0
 - add_dipole
 - alm2cl
 - alm2map_der
 - fits2cl (replaces read_asctab)
 - nside2ntemplates
 - plm_gen
 - rand_gauss, rand_init, rand_uni

- same_shape_pixels_nest, same_shape_pixels_ring
- template_pixel_nest, template_pixel_ring
- write_plm (replaces write_dbintab)
- New modules or modules with new name
 - misc_utils: fatal_error, assert_assert_present, assert_not_present, assert_alloc, file_present, assert_directory_present, string strupcase strlowcase, upcase, low-case, wall_clock_time, brag_openmp
 - rngmod: rand_gauss, rand_init, rand_uni
- The following routines are superseded.
 - read_asctab (replaced by fits2cl)
 - write_dbintab (replaced by write_plm)

Changes between release 1.1 and 1.2

Some new routines have been introduced since version 1.1, as listed below.

- New routines in version 1.2
 - angdist
 - complex_fft
 - concatnl
 - del_card
 - get_card
 - getargument
 - getenvironment
 - $input_tod*$
 - nArguments
 - parse_double, parse_init, parse_lgt, parse_long, parse_real, parse_string (see parse_xxx)
 - query_disc (replaces getdisc_ring)
 - query_polygon
 - query_strip
 - query_triangle

- read_fits_cut4
- real_fft
- scan_directories
- surface_triangle
- vect_prod
- write_bintabh
- write_fits_cut4
- New modules or modules with new name
 - the modules extension (C extensions), healpix_fft (FFT operations), paramfile_io (parameter parsing) have been introduced,
 - the module wrap_fits has been renamed head_fits to reflect its extended capabilities in manipulating FITS headers.
- The following routines are superseded. They have been moved to the obsolete module.
 - ask_inputmap, ask_outputmap, ask_lrange (initially in fitstools module)
 - setpar, getpar, anafast_parser, anafast_setpar, anafast_getpar, hotspots_parser, hotspots_setpar, hotspots_getpar, udgrade_parser, udgrade_setpar, udgrade_getpar, smoothing_parser, smoothing_setpar, smoothing_getpar (initially in utilities module).

alter_alm*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine modifies scalar (and tensor) $a_{\ell m}$ by multiplying them by a beam window function described by a FWHM (in the case of a gaussian beam) or read from an external file (in the more general case of a circular beam) $a_{\ell m} \longrightarrow a_{\ell m} b(\ell)$. It can also be used to multiply the $a_{\ell m}$ by an arbitray function of ℓ .

FORMAT

call alter_alm*(nsmax, nlmax, nmmax, fwhm_arcmin, alm_TGC [, beam_file, window])

name & dimensionality	kind	in/out	description
nsmax	I4B	IN	N_{side} resolution parameter of the map associated with the a_{lm} considered. Currently has no effect on the routine.
nlmax nmmax fwhm_arcmin	I4B I4B SP/	IN IN IN	maximum ℓ value for the $a_{\ell m}$. maximum m value for the $a_{\ell m}$. fwhm size of the gaussian beam in
alm_TGC(1:p,0:nlmax,0:nmmax)	DP SPC/ DPC	INOUT	arcminutes. complex $a_{\ell m}$ values to be altered. The first index here runs from 1:1 for temperature only, and 1:3 for polarisation. In the latter case, 1=T, 2=E, 3=B.

alter_alm*

beam_file(LEN=filenamelen) (OPTIONAL)	CHR	IN	name of the file containing the (non necessarily gaussian) window function B_{ℓ} of a circular beam. If present, it will override the argument fwhm arcmin.
window(0:nlw,1:d) (OPTIONAL)	SP/ DP	IN	arbitrary window by which to multiply the $a_{\ell m}$. If present, it overrides both fwhm_arcmin and beam_file. If nlw < nlmax, the $a_{\ell m}$ with $\ell \in \{\text{nlw+1,nlmax}\}$ are set to 0, and a warning is issued. If $d < p$ the window for temperature is replicated for polarisation.

EXAMPLE:

call alter_alm(64, 128, 128, 1, 5.0, alm_TGC)

Alters scalar and tensor a_{lm} of a map with $N_{\rm side}=64$, $\ell_{\rm max}=m_{\rm max}=128$ by multiplying them by the beam window function of a gaussian beam with FWHM = 5 arcmin.

MODULES & ROUTINES

This section lists the modules and routines used by alter_alm*.

${ m alm_tools}$	module, containing:
$generate_beam$	routine to generate beam window function
$pixel_window$	routine to generate pixel window function

RELATED ROUTINES

This section lists the routines related to alter_alm*.

$create_alm$	Routine to create $a_{\ell m}$ coefficients.
$rotate_alm$	Routine to rotate $a_{\ell m}$ coefficients between 2 different arbitrary coordinate systems.
map2alm	Routines to analyze a HEALPix sky map into its $a_{\ell m}$ coefficients.

alm2map Routines to synthetize a $\mathbf{HEALPix}$ sky map from

its $a_{\ell m}$ coefficients.

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add_card

Location in HEALPix directory tree: src/f90/mod/head_fits.F90

This routine writes a keyword of any kind into a FITS header. It is a wrapper to other routines that write keywords of different kinds.

\mathbf{FORMAT}

call add_card(header, kwd, value [, comment, update])

Arguments appearing in *italic* are optional.

name & dimensionality	kind	in/out	description
header(LEN=80) DIMENSION(:)	CHR	INOUT	The header to write the keyword to.
kwd(LEN=*)	CHR	IN	the FITS keyword to write. Should be shorter or equal to 8 characters.
value	any	IN	the value to give to the keyword.
comment(LEN=*)	CHR	IN	comment to the keyword.
update	LGT	IN	if set to .true., the first occurence of the keyword kwd in header will be updated (and all other occurences removed); otherwise, the keyword will be appended at the end (and any previous occurence removed). If the keyword is either 'HISTORY' or 'COMMENT', update is ignored and the keyword is peacefully appended at the end of the header.

EXAMPLE:

```
character(len=80), dimension(1:120) :: header
header = '' ! very important
call add_card(header,'NSIDE',256,'the nside of the map')
```

Gives the keyword 'NSIDE' the value 256 in the given headerstring. It is important to make sure that the header string array is empty before attempting to write anything in it.

MODULES & ROUTINES

This section lists the modules and routines used by add_card.

write_hl more general routine for adding a keyword to a

header.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to add_card.

write_minimal_header routine to write **HEALPix** compliant baseline

FITS header

get_card general purpose routine to read any keywords

from a header in a FITS file.

del_card routine to discard a keyword from a FITS header

read_par, number_of_alms routines to read specific keywords from a header

in a FITS file.

getsize_fits function returning the size of the data set in a fits

file and reading some other useful FITS keywords

merge_headers routine to merge two FITS headers

add_dipole*

add_dipole*

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

This routine provides a means to add a monopole and dipole to a **HEALPix** map.

FORMAT call add_dipole*(nside, map, ordering, degree, multipoles [, fmissval])

name & dimensionality	kind	in/out	description
nside	I4B	IN	value of N_{side} resolution parameter for input map
map(0:12*nside*nside-1)	SP/ DP	INOUT	HEALPix map to which the monopole and dipole will be added. Those are added to all unflagged pixels.
ordering	I4B	IN	HEALPix scheme 1:RING, 2: NESTED
degree	I4B	IN	multipoles to add. It is either 0 (nothing done), 1 (monopole only) or 2 (monopole and dipole)
$\begin{array}{c} \text{multipoles} (0: \text{degree*degree-} \\ 1) \end{array}$	DP	IN	values of monopole and dipole to add. The monopole is described as a scalar in the same units as the input map, the dipole as a 3D cartesian vector, in the same units.
fmissval (OPTIONAL)	SP/ DP	IN	value used to flag bad pixel on input (default: -1.6375e30). Pixels with that value are left unchanged.

EXAMPLE:

call add_dipole*(128, map, 1, 2, (\ 10.0_dp, 0.0_dp, 1.2_dp, 0.0_dp \))

map is a **HEALPix** map of resolution $N_{\text{side}} = 128$, with the RING ordering scheme. A monopole of amplitude 10 and a dipole of amplitude 1.2 and directed along the y axis will be added to it.

MODULES & ROUTINES

This section lists the modules and routines used by add_dipole*.

pix_tools module, containing:

RELATED ROUTINES

This section lists the routines related to add_dipole*.

remove_dipole routine to remove the best fit monopole and

monopole from a map.

 $alm2cl^*$

alm2cl*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine computes the auto (or cross) power spectra of a one (or two) sets of spherical harmonics coefficients $a_{\ell m}$. $C_{12}(\ell) = \sum_{m=-\ell}^{\ell} a_{1,\ell m} a_{2,\ell m}^* / (2\ell+1)$

FORMAT

call alm2cl*(nlmax, nmmax, alm1, [alm2,] cl)

name & dimensionality	kind	in/out	description
nlmax	I4B	IN	the maximum ℓ value used for the a_{lm} .
nmmax	I4B	IN	the maximum m value used for the
alm1(1:p, 0:nlmax, 0:nmmax)	SPC/ DPC	IN	a_{lm} . First set of a_{lm} values. p is 3 or 1 depending on wether polarisation is included or not. In the former case, the first index runs from 1 to 3 corresponding to (T,E,B) .
alm2(1:p, 0:nlmax, 0:nmmax) (OPTIONAL)	SPC/ DPC	IN	Second set of a_{lm} values.
cl(0:nlmax,1:d)	SP/ DP	OUT	resulting auto or cross power spectra. If both alm1 and alm2 are present, c1 will be their cross power spectrum. If only alm1 is present, c1 will be its power spectrum. If $d=1$, only the temperature spectrum C_l^T will be output. If $d=4$ and $p=3$, the output will be C_l^T , C_l^E , C_l^B , $C_l^{T\times E}$, and if $d\geq 6$ and $p=3$, $C_l^{T\times B}$ $C_l^{E\times B}$ will also be output.

EXAMPLE:

```
lmax = 128 ; mmax = lmax
call alm2cl(lmax, mmax, alm1, cl_auto)
call alm2cl(lmax, mmax, alm1, alm2, cl_cross)
```

cl_auto will contain the (auto) power spectrum of the $a_{\ell m}$ coefficients alm1 up to $\ell=128$, while cl_cross will be the cross power spectra of the two sets of $a_{\ell m}$ coefficients alm1 and alm2.

MODULES & ROUTINES

This section lists the modules and routines used by alm2cl*.

none

RELATED ROUTINES

This section lists the routines related to alm2cl*.

map2alm routine extracting the $a_{\ell m}$ coefficients from a

HEALPix map

create_alm routine to generate randomly distributed $a_{\ell m}$ co-

efficients according to a given power spectrum

 $alm2map^*$

alm2map*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine is a wrapper to 10 other routines: alm2map_sc_X, alm2map_sc_pre_X, alm2map_pol_x, alm2map_pol_pre1_X, alm2map_pol_pre2_X, where X stands for either s or d. These routines synthesize a **HEALPix** RING ordered temperature map (and if specified, polarisation maps) from input a_{lm}^T (and if specified a_{lm}^E and a_{lm}^B) values. The different routines are called dependent on what parameters are passed. Some routines synthesize maps with or without precomputed harmonics and some with or without polarisation. The routines accept both single and double precision arrays for alm_TGC and map_TQU. The precision of these arrays should match.

FORMAT	call	alm2map*(nsmax,	nlmax,	nmmax,
	alm_{-}	ΓGC, map_TQU [, pln	n])	

name & dimensionality	kind	in/out	description
nsmax	I4B	IN	the N_{side} value of the map to synthesize.
nlmax	I4B	IN	the maximum ℓ value used for the a_{lm} .
nmmax	I4B	IN	the maximum m value used for the a_{lm} .
alm_TGC(1:p, 0:nlmax, 0:nmmax)	SPC or DPC	IN	The a_{lm} values to make the map from. p is 3 or 1 depending on wether polarisation is respec- tively included or not. In the former case, the first index runs from 1 to 3 corresponding to (T,E,B).

$map_TQU(0:12*nsmax**2-1)$	SP	OUT	if only a temperature map is to
	or		be synthesized, the map-array
	DP		should be passed with this rank.
$map_TQU(0:12*nsmax**2-1, 1:3)$	SP	OUT	if both temperature an polar-
	or		isation maps are to be syn-
	DP		thesized, the map array should
			have this rank, where the sec-
			ond index is $(1,2,3)$ correspond-
			ing to (T,Q,U) .
$plm(0:n_plm-1),$	DP	IN	If this optional matrix is
OPTIONAL			passed with this rank, pre-
			computed $P_{lm}(\theta)$ are used
			instead of recursion. (n_plm =
			nsmax*(nmmax+1)*(2*nlmax-
			nmmax+2)
$plm(0:n_plm-1,1:3),$	DP	IN	If this optional matrix is passed
OPTIONAL			with this rank, precomputed
			$P_{lm}(\theta)$ AND precomputed
			tensor harmonics are used
			instead of recursion. $(n_plm =$
			nsmax*(nmmax+1)*(2*nlmax-
			nmmax+2)
			,

EXAMPLE:

```
use healpix_types
use pix_tools, only : nside2npix
use alm_tools, only : alm2map
integer(I4B) :: nside, lmax, mmax, npix, n_plm
real(SP), dimension(:,:), allocatable :: map
complex(SPC), dimension(:,:), allocatable :: alm
real(DP), dimension(:,:), allocatable :: plm
...
nside=256; lmax=512; mmax=lmax
npix=nside2npix(nside)
n_plm=nside*(mmax+1)*(2*lmax-mmax+2)
allocate(alm(1:3,0:lmax,0:mmax))
allocate(map(0:npix-1,1:3))
allocate(plm(0:n_plm-1,1:3))
...
call alm2map(nside, lmax, mmax, alm, map, plm)
```

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Make temperature and polarisation maps from the scalar and tensor a_{lm} passed in alm. The maps have N_{side} of 256, and are constructed from a_{lm} values up to 512 in ℓ and m. Since the optional plm array is passed with both precomputed $P_{lm}(\theta)$ AND tensor harmonics, there will be no recursions in the routine and execution will be faster.

MODULES & ROUTINES

This section lists the modules and routines used by alm2map*.

ring_synthesis Performs FFT over m for synthesis of the rings.

compute_lam_mm, get_pixel_layout, gen_lamfac,gen_mfac, gen_normpol, gen_recfac, init_rescale, l_min_ylm

 $\mathcal{L}_{rescale}$, \mathcal{L}_{min_ylm} Ancillary routines used for $Y_{\ell m}$ recursion

misc_utils module, containing:

assert_alloc routine to print error message, when an array can

not be allocated properly

RELATED ROUTINES

This section lists the routines related to alm2map*.

alm2map_der routine generating a map and its derivatives from

its $a_{\ell m}$

alm2map_spin routine generating maps of arbitrary spin from

their $_sa_{\ell m}$

smoothing executable using alm2map* to smooth maps

synfast executable using alm2map* to synthesize maps.

map2alm routine performing the inverse transform of

alm2map*.

create_alm routine to generate randomly distributed $a_{\ell m}$ co-

efficients according to a given power spectrum

pixel_window, generate_beam return the l-space **HEALPix** -pixel and beam

window function respectively

alter_alm modifies a_{lm} to emulate effect of real space filtering

$alm2map_der*$

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine is a wrapper to four other routines that synthesize a **HEALPix** temperature (and polarisation) map(s), its (their) first derivatives, and optionally its (their) second derivatives. The routines accept both single and double precision arrays for alm, map, der1 and der2. The precision of these arrays should match. All maps produced are RING ordered.

See "Fortran Facilities" Appendix for a note on a bug affecting the calculation of polarisation derivatives on past versions of this routine.

FORMAT call alm2map_der*(nsmax, nlmax, nmmax, alm, map, der1 [, der2])

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name & dimensionality	kind	in/out	description
nsmax	I4B	IN	the N_{side} value of the map to synthesize.
nlmax	I4B	IN	the maximum ℓ value used for the
nmmax	I4B	IN	a_{lm} . the maximum m value used for the
alm(1:p, 0:nlmax, 0:nmmax)	SPC/ DPC	IN	a_{lm} . The a_{lm} values to make the map from. p is either 1 (temperature+polarisation).
map(0:12*nsmax**2-1) or $(0:12*nsmax**2-1,1:3)$	SP/ DP	OUT	temperature map $T(p)$ or temperature + polarisation maps $T(p)$, $Q(p)$, $U(p)$ to be synthesized.
der1(0:12*nsmax**2-1, 1:2*p)	SP/ DP	OUT	contains on output the first derivatives of T: $(\partial T/\partial \theta, \partial T/\partial \phi/\sin \theta)$ or the interleaved derivatives of T, Q, and U: $(\partial T/\partial \theta, \partial Q/\partial \theta, \partial U/\partial \theta; \partial T/\partial \phi/\sin \theta,)$
der2(0:12*nsmax**2-1,1:3*p), OPTIONAL	SP/ DP	OUT	If this optional matrix is passed with this rank, it will contain on output the second derivatives $(\partial^2 T/\partial \theta^2, \partial^2 T/\partial \theta \partial \phi/\sin \theta, \partial^2 T/\partial \phi^2/\sin^2 \theta)$ or $(\partial^2 T/\partial \theta^2, \partial^2 Q/\partial \theta^2, \partial^2 Q/\partial \theta^2, \dots)$

EXAMPLE:

```
use healpix_types
use pix_tools, only : nside2npix
use alm_tools, only : alm2map_der
integer(I4B) :: nside, lmax, mmax, npix, n_plm
real(SP), dimension(:), allocatable :: map
real(SP), dimension(:,:), allocatable :: der1, der2
complex(SPC), dimension(:,:,:), allocatable :: alm
...
nside=256; lmax=512; mmax=lmax
npix=nside2npix(nside)
allocate(alm(1:1,0:lmax,0:mmax))
allocate(map(0:npix-1))
```

```
allocate(der1(0:npix-1,1:2), der2(0:npix-1,1:3))
...
call alm2map_der(nside, lmax, mmax, alm, map, der1, der2)
```

Make temperature maps and its derivatives from the a_{lm} passed in alm. The maps have N_{side} of 256, and are constructed from a_{lm} values up to 512 in ℓ and m.

MODULES & ROUTINES

This section lists the modules and routines used by alm2map_der*.

ring_synthesis Performs FFT over m for synthesis of the rings. compute_lam_mm, get_pixel_layout, gen_lamfac_der, gen_mfac, gen_recfac, init_rescale, l_min_ylm Ancillary routines used for ${}_sY_{\ell m}$ recursion

misc_utils module, containing:

assert_alloc routine to print error message, when an array can

not be allocated properly

RELATED ROUTINES

This section lists the routines related to alm2map_der*.

alm2map routine generating maps of temperature and po-

larisation from their $a_{\ell m}$

alm2map_spin routine generating maps of arbitrary spin from

their $sa_{\ell m}$

synfast executable using alm2map_der* to synthesize

maps.

create_alm routine to generate randomly distributed $a_{\ell m}$ co-

efficients according to a given power spectrum

alm2map_spin* 25

alm2map_spin*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine produces the maps of arbitrary spin s and -s given their alm coefficients. A (complex) map S of spin s is a linear combination of the spin weighted harmonics ${}_sY_{lm}$

$${}_{s}S(p) = \sum_{lm} {}_{s}a_{lm} \quad {}_{s}Y_{lm}(p) \tag{1}$$

for $l \ge |m|, l \ge |s|$, and is such that ${}_sS^* = {}_{-s}S$. alm2map_spin* expects the alm coefficients to be provided as

$$a_{ls}^{+}a_{lm}^{+} = -(a_{ls}a_{lm} + (-1)^{s}a_{lm})/2$$
 (2)

$$a_{|s|}a_{lm}^- = -(a_{|s|}a_{lm} - (-1)^s - a_{|s|}a_{lm})/(2i)$$
 (3)

for $m \ge 0$, knowing that, just as for spin 0 maps, the coefficients for m < 0 are given by

$$|s|a_{l-m}^+ = (-1)^m |s|a_{lm}^{+*},$$
 (4)

$$a_{ls}^{-}a_{l-m}^{-} = (-1)^{m}{}_{|s|}a_{lm}^{-*}.$$
 (5)

The two (real) maps produced by alm2map_spin* are defined respectively as

$$|s|S^{+} = (|s|S + -|s|S)/2$$
 (6)

$$|s|S^- = (|s|S - -|s|S)/(2i).$$
 (7)

With these definitions, ${}_2a^+, {}_2a^-, {}_2S^+$ and ${}_2S^-$ match **HEALPix** polarization a^E, a^B, Q and U respectively. However, for $s=0, {}_0a^+_{lm}=-a^T_{lm}, {}_0a^-_{lm}=0, {}_0S^+=T, {}_0S^-=0.$

FORMAT

call alm2map_spin*(nsmax, nlmax, nmmax, spin, alm, map)

name & dimensionality	kind in/out	description
nsmax	I4B IN	the N_{side} value of the map to synthesize.
nlmax	I4B IN	the maximum ℓ value used for the
nmmax	I4B IN	a_{lm} . the maximum m value used for the
spin	I4B IN	a_{lm} . spin s of the maps to be generated (only its absolute value is relevant).
alm(1:2, 0:nlmax, 0:nmmax)	SPC/ IN	The $_{ s }a_{lm}^{+}$ and $_{ s }a_{lm}^{-}$ values to make
map(0:12*nsmax**2-1, 1:2)	DPC SP/ OUT DP	the map from. $ s S^+$ and $ s S^-$ output maps

EXAMPLE:

```
use healpix_types
use pix_tools, only : nside2npix
use alm_tools, only : alm2map_spin
integer(I4B) :: nside, lmax, mmax, npix, spin
real(SP), dimension(:,:), allocatable :: map
complex(SPC), dimension(:,:,:), allocatable :: alm
...
nside=256; lmax=512; mmax=lmax; spin=4
npix=nside2npix(nside)
allocate(alm(1:2,0:lmax,0:mmax))
allocate(map(0:npix-1,1:2))
...
call alm2map_spin(nside, lmax, mmax, spin, alm, map)
```

Make spin-4 maps from the a_{lm} passed in alm. The maps have N_{side} of 256, and are constructed from a_{lm} values up to 512 in ℓ and m.

MODULES & ROUTINES

This section lists the modules and routines used by alm2map_spin*.

ring_synthesis Performs FFT over m for synthesis of the rings. compute_lam_mm, get_pixel_layout,

alm2map_spin* 27

gen_lamfac_der, gen_mfac_spin, do_lam_lm_spin,

gen_recfac_spin, init_rescale, l_min_ylm Ancillary routines used for

 $Y_{\ell m}$ recursion

misc_utils module, containing:

assert_alloc routine to print error message, when an array can

not be allocated properly

RELATED ROUTINES

This section lists the routines related to alm2map_spin*.

alm2map routine generating maps of temperature and po-

larisation from their $a_{\ell m}$

alm2map_der routine generating maps of temperature and po-

larisation, and their spatial derivatives, from their

 $a_{\ell m}$

map2alm_spin routine performing the inverse transform of

alm2map.

create_alm routine to generate randomly distributed $a_{\ell m}$ co-

efficients according to a given power spectrum

alms2fits*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine stores a_{lm} values in a binary FITS file. Each FITS file extension created will contain one integer column with $index = \ell^2 + \ell + m + 1$, and 2 or 4 single (or double) precision columns with real/imaginary a_{lm} values and real/imaginary standard deviation. One can store temperature a_{lm} or temperature and polarisation, a_{lm}^T , a_{lm}^E and a_{lm}^B . If temperature is specified, a FITS file with one extension is created. If polarisation is specified, a FITS file with 3 extensions one for each set of a_{lm} , a_{lm}^T , a_{lm}^E and a_{lm}^B is created.

FORMAT call alms2fits*(filename, nalms, alms, ncl, header, nlheader, next)

name & dimensionality	kind	in/o	utdescription
filename(LEN = filenamelen)	CHR	IN	filename for the FITS file to store the a_{lm} in.
nalms	I4B	IN	number of a_{lm} to store.
ncl	I4B	IN	number of columns in the FITS file.
			If an standard deviation is given, this number is 5, otherwise it is 3.
next	I4B	IN	the number of extensions. 1 for temperature only, 3 for temperature and polarisation.

alms2fits*

name & dimensiona	lity	kind	in/o	utdescription
alms(1:nalms,1:ncl+1,1:next)		SP/ DP	IN	the a_{lm} to write to the file. alms(i,1,j) and alms(i,2,j) contain the ℓ and m values for the ith a_{lm} (j=1,2,3 for (T,E,B)). alms(i,3,j) and alms(i,4,j) contain the real and imaginary value of the ith a_{lm} . Finally, the standard deviation for the ith a_{lm} is contained in alms(i,5,j) (real) and alms(i,6,j) (imaginary).
nlheader		I4B	IN	number of header lines to write to
header(LEN=80) 1:next)	(1:nlheader,	CHR	IN	the file. the header to the FITS file.

EXAMPLE:

call alms2fits ('alms.fits', 65*66/2, alms, 3, header, 80, 3)

Creates a FITS file with the a_{lm}^T , a_{lm}^E and a_{lm}^B values given in alms(1:65*66/2,1:4,1:3). The last index specifies (T,E,B). The second index gives l, m, real(a_{lm}), imaginary(a_{lm}) for each of the a_{lm} . The number 65*66/2 is the number of a_{lm} values up to an ℓ value of 64. 80 lines from header(1:80,1:3) is written to each extension.

MODULES & ROUTINES

This section lists the modules and routines used by alms2fits*.

$write_alms$	routine called by alms2fits* for each extension.
fitstools	module, containing:
printerror	routine for printing FITS error messages.
cfitsio	library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to alms2fits*.

fits2alms, read_conbintab routines to read a_{lm} from a FITS file

dump_alms has the same function as alms2fits* but with pa-

rameters passed differently.

ang2vec 31

ang2vec

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to convert the position angles (θ, ϕ) of a point on the sphere into its 3D position vector (x, y, z) with $x = \sin \theta \cos \phi$, $y = \sin \theta \sin \phi$, $z = \cos \theta$.

FORMAT

call ang2vec(theta, phi, vector)

ARGUMENTS

name & dimensionality	kind	in/out	description
theta	DP	IN	colatitude in radians measured southward from north pole (in
phi	DP	IN	$[0, \pi]$). longitude in radians measured
vector(3)	DP	OUT	eastward (in $[0, 2\pi]$). three dimensional cartesian position vector (x, y, z) nor- malised to unity. The north pole is $(0, 0, 1)$

RELATED ROUTINES

This section lists the routines related to ang2vec.

vec2ang

converts the 3D position vector of point into its position angles on the sphere.

angdist

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Returns the angular distance in radians between two vectors. The input vectors do not have to be normalised. For almost colinear or anti-colinear vectors, renders numerically more accurate results than the \cos^{-1} of the scalar product.

FORMAT

call angdist(v1, v2, dist)

ARGUMENTS

name & dimension-	kind	in/out	description
ality			
v1(3) v2(3) dist	DP DP DP	IN IN OUT	cartesian vector. cartesian vector. angular distance in radians between the 2 vectors.

EXAMPLE:

```
use healpix_types
use pix_tools, only : angdist
real(DP) :: dist, one = 1.0_dp
call angdist((/1,2,3/)*one, (/1,2,4/)*one, dist)
print*, dist
```

Returns the angular distance between 2 vectors.

RELATED ROUTINES

This section lists the routines related to **angdist**.

vect_prod computes the vector product between two 3D vectors

angdist 33

assert_assert_alloc, assert_directory_present, . . .

Location in HEALPix directory tree: src/f90/mod/misc_utils.F90

The Fortran90 module misc_utils contains a few routines to test an assertion and return an error message if it is false.

FUNCTIONS:

call assert(test [, msg, errcode])

if test is true, proceeds with normal code execution. If test is false, issues a standard error message (unless msg is provided) and stops the code execution with the status errcode (or 1 by default).

call assert_alloc(status, code, array)

if status is 0, proceeds with normal code execution. If not, issues an error message indicating a problem during memory allocation of array in program code, and stops the code execution.

call assert_directory_present(directory)

issues an error message and stops the code execution if the directory named directory can not be found

call assert_present(filename)

issues an error message and stops the code execution if the file named filename can not be found.

call assert_not_present(filename)

issues an error message and stops the code execution if a file with name filename already exists.

ARGUMENTS

name & di	mensionality	kind	in/out	description
test msg errorcode	OPTIONAL OPTIONAL	LGT CHR I4B	IN IN IN	result of a logical test character string describing nature of error error status given to code interruption
status		I4B	IN	value of the stat flag returned by the F90
code		CHR	IN	allocate command name of program or code in which alloca- tion is made
array		CHR	IN	name of array allocated
directory		CHR	IN	directory name (contains a '/')
$_{ m filename}$		CHR	IN	file name

EXAMPLE:

```
program my_code
use misc_utils
real, allocatable, dimension(:) :: vector
integer :: status
real :: a = -1.

allocate(vector(12345), stat=status)
call assert_alloc(status, 'my_code', 'vector')

call assert_directory_present('/home')

call assert(a > 0., 'a is NEGATIVE !!!')
end program my_code
```

Will issue a error message and stops the code if vector can not be allocated, will stop the code if '/home' is not found, and will stop the code and complain loudly about it because a is actually negative.

brag_openmp

Location in HEALPix directory tree: src/f90/mod/misc_utils.F90

If compiled with shared memory libraries (OpenMP), this routine prints out the number of CPUs used (controlled by the environment variable OMP_NUM_THREADS) and the number of CPUs available.

FORMAT

call brag_openmp()

EXAMPLE:

use misc_utils
call brag_openmp()

Will print out:

Number of OpenMP threads in use: 2 Number of CPUs available: 2

on bi-pro (or dual core) computer

complex_fft 37

complex_fft

Location in HEALPix directory tree: src/f90/mod/healpix_fft.F90

This routine performs a forward or backward Fast Fourier Transformation on its argument data.

FORMAT

call complex_fft(data, backward)

ARGUMENTS

name&dimensionality	kind in/out	description
data(:)	XXX INOUT	array containing the input and output data. It can be of type real(sp), real(dp), complex(spc) or complex(dpc). If it is of
backward	LGT IN	type real, it is interpreted as an array of size(data)/2 complex variables. Optional argument. If present and true, perform backward transformation, else forward

EXAMPLE:

use healpix_fft
call complex_fft (data, backward=.true.)

Performs a backward FFT on data.

RELATED ROUTINES

This section lists the routines related to **complex_fft**.

real_fft routine for FFT of real data

compute_statistics*

 $Location \ in \ HEALPix \ directory \ tree: \ src/f90/mod/statistics.f90$

This routine computes the min, max, absolute deviation and first four order moment of a data set

FORMAT

call compute_statistics*(data ,stats [, badval])

ARGUMENTS

name & dimensionality	kind	in/ou	ıtdescription
data(:)	SP/	IN	data set
	DP		
stats	tstats	OUT	structure containing the statistics of the data.
			The respective fields (stats%field) are:
ntot	I4B	_	total number of data points
nvalid	I4B	_	number n of valid data points
\min d, \max d	DP	_	minimum and maximum valid data
average	DP	_	average of valid points $m = \sum x/n$
absdev	DP	_	absolute deviation $a = \sum x - m /n$
var	DP	_	variance $\sigma^2 = \sum (x-m)^2/(n-1)$
${ m rms}$	DP	_	standard deviation σ
skew	DP	_	skewness factor $s = \sum (x - m)^3 / (n\sigma^3)$
kurt	DP	_	kurtosis factor $k = \sum (x - m)^4 / (n\sigma^4) - 3$
badval	SP/	IN	sentinel value given to bad data points. Data
(OPTIONAL)	DP		points with this value will be ignored during
			calculation of the statistics. If not set, all
			points will be considered. Do not set to 0! .

EXAMPLE:

use statistics, only: compute_statistics, print_statistics, tstats type(tstats) :: stats

compute_statistics(map, stats)
print*,stats%average, stats%rms
print_statistics(stats)

Computes the statistics of map, prints its average and rms and prints the whole list of statistical measures.

RELATED ROUTINES

This section lists the routines related to **compute_statistics***.

median routine to compute median of a data set

concatnl

Location in HEALPix directory tree: src/f90/mod/paramfile_io.F90

Function to concatenate up to 10 subtrings interspaced with LineFeed character. Upon printing each subtring will be on a different line.

FORMAT var=concatnl(string1[, string2, string3, ...])

ARGUMENTS

name & dimensionality	kind	in/out	description
string1 string2	CHR CHR		the first substring to be concatenated. the second substring (if any) to be con-
string3		optional IN	catenated up to 10 substrings can be concate-
var	CHR	optional OUT	nated. concatenation of the substrings interspaced with LineFeed character.

EXAMPLE:

RELATED ROUTINES

This section lists the routines related to **concatnl**.

parse_xxx parse an ASCII file for parameters definition

concatnl 41

convert_inplace*

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to convert a **HEALPix** map from NESTED to RING scheme or vice versa. The conversion is done inplace, meaning that it doesn't require memory for a temporary map, like the *convert_nest2ring* or *convert_ring2nest* routines. But for that reason, this routine is slower and not parallelized. The routine is a wrapper for 6 different routines and can threfore process integer, single precision and double precision maps as well as mono or bi dimensional arrays.

FORMAT

call convert_inplace*(subcall, map)

ARGUMENTS

name & dimensionality	kind	in/out	description
subcall	_	IN	routine to be called by convert_inplace_real. Set this to ring2nest or nest2ring dependent on wether the conversion is RING to NESTED or vice versa.
map(0:npix-1)	I4B/ SP/ DP	INOUT	mono-dimensional full sky map to be converted, the routine finds the size itself.
map(0:npix-1,1:nd)	I4B/ SP/ DP	INOUT	bi-dimensional (nd>0) full sky map to be converted, the routine finds both dimensions itself. Processing a bidimensional map with nd> 1 should be faster than each of the nd 1D-maps consecutively.

convert_inplace*

EXAMPLE:

call convert_inplace(ring2nest,map)

Converts an map from RING to NESTED scheme.

MODULES & ROUTINES

This section lists the modules and routines used by **convert_inplace***.

nest2ring routine to convert a NESTED pixel index to

RING pixel number.

ring2nest routine to convert a RING pixel index to

NESTED pixel number.

RELATED ROUTINES

This section lists the routines related to **convert_inplace***.

convert_nest2ring convert from NESTED to RING scheme using a

temporary array. Requires more space then con-

vert_inplace, but is faster.

convert_ring2nest convert from RING to NESTED scheme using a

temporary array. Requires more space then con-

vert_inplace, but is faster.

convert_nest2ring*

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to convert a **HEALPix** map from NESTED to RING scheme.

The routine is a wrapper for 6 different routines and can threfore process integer, single precision and double precision maps as well as mono or bi dimensional arrays.

This routine is fast, and is parallelized for shared memory architecture, but requires extra memory to store a temporary map in.

FORMAT

call convert_nest2ring*(nside, map)

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	the N_{side} parameter of the map to be converted.
map(0:12*nside**2-1)	I4B/ SP/ DP	INOUT	mono-dimensional full sky map to be converted to RING scheme.
map(0:12*nside**2-1,1:nd)	I4B/ SP/ DP	INOUT	bi-dimensional full sky map to be converted to RING scheme. The routine finds the second di- mension (nd) by itself. Process- ing a bidimensional map with nd> 1 should be faster than each of the nd 1D-maps consec- utively.

EXAMPLE:

call convert_nest2ring(256,map)

convert_nest2ring* 45

Converts an $N_{side} = 256$ map given in array map from NESTED to RING scheme.

MODULES & ROUTINES

This section lists the modules and routines used by **convert_nest2ring***.

nest2ring

routine to convert a NESTED pixel index to RING pixel number.

RELATED ROUTINES

This section lists the routines related to **convert_nest2ring***.

convert_ring2nest convert_inplace

convert between RING and NESTED schemes. convert between NESTED and RING schemes inplace. This routine is slower than convert_nest2ring*, but doesn't require as much memory.

convert_ring2nest*

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to convert a **HEALPix** map from RING to NESTED scheme.

The routine is a wrapper for 6 different routines and can threfore process integer, single precision and double precision maps as well as mono or bi dimensional arrays.

This routine is fast, and is parallelized for shared memory architecture, but requires extra memory to store a temporary map in.

FORMAT

call convert_ring2nest*(nside, map)

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	the N_{side} parameter of the map to be converted.
map(0:12*nside**2-1)	I4B/ SP/ DP	INOUT	mono-dimensional full sky map to be converted to RING scheme.
map(0:12*nside**2-1,1:nd)	I4B/ SP/ DP	INOUT	bi-dimensional full sky map to be converted to RING scheme. The routine finds the second di- mension (nd) by itself. Process- ing a bidimensional map with nd> 1 should be faster than each of the nd 1D-maps consec- utively.

EXAMPLE:

call convert_ring2nest(256,map)

convert_ring2nest* 47

Converts an $N_{side} = 256$ map given in array map from RING to NESTED scheme.

MODULES & ROUTINES

This section lists the modules and routines used by **convert_ring2nest***.

ring2nest

routine to convert a RING pixel index to NESTED pixel number.

RELATED ROUTINES

This section lists the routines related to **convert_ring2nest***.

convert_nest2ring convert_inplace convert between NESTED and RING schemes. convert between RING and NESTED schemes inplace. This routine is slower than convert_ring2nest*, but doesn't require as much memory.

coordsys2euler_zyz

$Location \ in \ HEALPix \ directory \ tree: \ src/f90/mod/coord_v_convert.f90$

This routine returns the three Euler angles ψ , θ , φ , corresponding to a rotation between standard astronomical coordinate systems. This angles can then be used in rotate_alm

FORMAT call coordsys2euler_zyz(iepoch, oepoch, isys, osys, psi, theta, phi)

ARGUMENTS

name & dimension-	kind	in/out	description
ality			
iepoch	DP	IN	epoch of the input astronomical coordinate system.
oepoch	DP	IN	epoch of the output astronomical coordinate
			system.
isys(len=*)	CHR	IN	input coordinate system, should be
			one of 'E'=Ecliptic, 'G'=Galactic,
			'C'/'Q'=Celestial/eQuatorial.
osys(len=*)	CHR	IN	output coordinate system, same choice as
			above.
psi	DP	OUT	first Euler angle: rotation ψ about the z-axis.
theta	DP	OUT	second Euler angle: rotation θ about the orig-
			inal (unrotated) y-axis;
phi	DP	OUT	third Euler angle: rotation φ about the orig-
			inal (unrotated) z-axis;

EXAMPLE:

```
use coord_v_convert, only: coordsys2euler_zyz
```

use alm_tools, only: rotate_alm

. . .

call coordsys2euler_zyz(2000.0_dp, 2000.0_dp, 'E', 'G', psi, theta, phi)

coordsys2euler_zyz 49

call rotate_alm(64, alm_TGC, psi, theta, phi)

Rotate the a_{lm} from Ecliptic to Galactic coordinates.

RELATED ROUTINES

This section lists the routines related to **coordsys2euler_zyz**.

rotate_alm apply arbitrary sky rotation to a set of a_{lm} coeffi-

cients.

xcc_v_convert rotates a 3D coordinate vector from one astronom-

ical coordinate system to another.

$create_alm*$

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine generates scalar (and tensor) a_{lm} for a temperature (and polarisation) power spectrum read from an input FITS file. The a_{lm} are gaussian distributed with a zero mean, and their amplitude is multiplied with the ℓ -space window function of a gaussian beam characterized by its FWHM or an arbitrary circular beam and a pixel window read from an external file.

FORMAT

call create_alm*(nsmax, nlmax, nmmax, polar, filename, iseed, fwhm_arcmin, alm_TGC, header [, windowfile, units, beam_file, rng_handle])

ARGUMENTS

create_alm* 51

name & dimensionality	kind	in/out	description
nsmax	I4B	IN	N_{side} of the map to be synthetized from the $a_{\ell m}$ created by this routine.
nlmax	I4B	IN	maximum ℓ value to be considered $(MAX = 3 \times N_{side})$.
nmmax	I4B	IN	maximum m value for the $a_{\ell m}$.
polar	I4B	IN	equals 1 if polarisation is used, 0 otherwise.
filename(LEN = filenamelen)	CHR	IN	name of FITS file containing power spectrum.
rng_handle	planc	k_rng INOUT	structure containing information necessary to continue a random sequence initiated <i>previously</i> with the subroutine rand_init. Consecutive calls to create_alm*can be made after a single invocation to rand_init.
$fwhm_arcmin$	SP/ DP	IN	FWHM size of the gaussian beam in arcminutes.

$alm_TGC(1:p,0:nlmax,0:nmmax)$	SPC/ OUT	complex $a_{\ell m}$ values generated
	DPC	from the powerspectrum in the
		FITS-file. The first index here
		runs form 1:1 for temperature
		only, and 1:3 for polarisation.
		• • • • • • • • • • • • • • • • • • • •
		In the latter case, 1=T, 2=E, 3=B.
header(LEN=80),dimension(60)	CHR OUT	part of header which will be in-
,,,		cluded in the FITS-file contain-
		ing the map synthesised from
		the $a_{\ell m}$ which create_alm gen-
		erates.
windowfile(LEN=filenamelen)	CHR IN	full filename specification of the
,	CIII(IIV	-
(OPTIONAL)		FITS file with the pixel window
(1777 00) 11 (1)	CIID OIIT	function.
units(LEN=80), dimension(1:)	CHR OUT	physical units of the created $a_{\ell m}$
(OPTIONAL)		(square-root of the input power
		spectrum units).
$beam_file(LEN=filenamelen)$	CHR IN	name of the file containing
(OPTIONAL)		the (non necessarily gaussian)
()		window function B_{ℓ} of a cir-
		cular beam. If present, it
		- ,
		will override the argument
		${ t fwhm_arcmin.}$

EXAMPLE:

```
use alm_tools, only: create_alm
use rngmod, only: rand_init, rng_handle
type(planck_rng) :: rng_handle

call rand_init(rng_handle, -1)
call create_alm(64, 128, 128, 1, 'cl.fits', rng_handle, 5.0, alm_TGC, header, 'data/pixel_window_n0064.fits')
```

create_alm* 53

Creates scalar and tensor a_{lm} from the power spectrum given in the file 'cl.fits'. The map to be created from these a_{lm} is assumed to have $N_{side}=64$. C_{l} s from the power spectrum are used up to an ℓ value of 128. Corresponding a_{lm} values up to l=128 and m=128 are created as gaussian distributed complex numbers. Their are drawn from a sequence of pseudorandom numbers initiated with a seed of -1. The produced a_{lm} are convolved with a gaussian beam of FWHM 5 arcminutes and a pixel window read from 'data/pixel_window_n0064.fits'. It is assumed that after the return from this routine, a map is generated from the created a_{lm} . For this purpose, header is updated with FITS format information describing the origin and history of these a_{lm} .

MODULES & ROUTINES

This section lists the modules and routines used by **create_alm***.

${ m alm_tools}$	<u>module</u> , containing:
$pow2alm_units$	routine to convert from power spectrum units to
	$a_{\ell m}$ units
$generate_beam$	routine to generate beam window function
$pixel_window$	routine to read in pixel window function
utilities	module, containing:
die_alloc	routine that prints an error message if there is not enough space for allocation of variables.
fitstools	module, containing:
fits2cl	routine to read a FITS file containing a power spectrum.
read_dbintab	routine to read a FITS-binary file containing the pixel window functions.
$\mathbf{head_fits}$	module, containing:
add_card	routine to add a keyword to a FITS header.
$\operatorname{get_card}$	routine to read a keyword value from FITS header.
$merge_headers$	routine to merge two FITS headers.
rngmod	module, containing:
${\rm rand_gauss}$	function which returns a gaussian distributed ran-

dom number.

RELATED ROUTINES

This section lists the routines related to **create_alm***.

rand_init subroutine to initiate a random number sequence.

synfast executable using create_alm* to synthesize CMB

maps from a given power spectrum.

alm2map Routine to transform a set of a_{lm} created by cre-

ate_alm* to a **HEALPix** map.

alms2fits, dump_alms Routines to save a set of a_{lm} in a FITS file.

del_card 55

del_card

Location in HEALPix directory tree: src/f90/mod/head_fits.F90

This routine removes one or several keywords from a FITS header.

FORMAT

call del_card(header, kwds)

ARGUMENTS

name & dimensionality	kind in/out	description
header(LEN=80)(1:nlheader)	CHR INOUT	The header to remove the keyword(s) from. The routines
kwds(LEN=20)(1:nkws)	CHR IN	finds out the header size. list of FITS keywords to re- move. The routine accepts ei-
kwds(LEN=20)	CHR IN	ther a vector a keywords or a single one in a scalar variable the one FITS keyword to remove.

EXAMPLES: #1

call del_card(header,(/ 'NSIDE ','COORD ','ORDERING' /))

Removes the keywords 'NSIDE', 'COORD' and 'ORDERING' from Header

EXAMPLES: #2

call del_card(header, 'ORDERING')

Removes the keyword 'ORDERING' from Header

MODULES & ROUTINES

This section lists the modules and routines used by del_card.

write_hl more general routine for adding a keyword to a

header.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **del_card**.

add_card general purpose routine to write any keywords

into a FITS file header

get_card general purpose routine to read any keywords

from a header in a FITS file.

read_par, number_of_alms routines to read specific keywords from a header

in a FITS file.

getsize_fits function returning the size of the data set in a fits

file and reading some other useful FITS keywords

merge_headers routine to merge two FITS headers

 $dump_alms^*$ 57

dump_alms*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine stores a_{lm} values in a binary FITS file. The FITS file created will contain one integer column with $index = \ell^2 + \ell + m + 1$ and 2 single precision columns with real/imaginary a_{lm} values. One can store temperature a_{lm} or polarisation, a_{lm}^E or a_{lm}^B . If temperature is specified, a FITS file is created. If polarisation is specified, an old FITS file is opened and extra extensions is created.

FORMAT call dump_alms*(filename, alms, nlmax, header, nlheader, extno)

ARGUMENTS

name & dimensionality	kind in/ou	t description
$\mathrm{filename}(\mathrm{LEN} {=} \mathtt{filenamelen})$	CHR IN	filename for the FITS-file to store the a_{lm} in.
$\begin{array}{l} nlmax \\ alms(0:nlmax,0:nlmax) \end{array}$	I4B IN SPC/ IN	maximum ℓ value to store. array with a_{lm} . alms(l,m) cor-
extno	DPC I4B IN	responds to a_{lm} extension number. If 0 is spec-
		ified, a FITS file is created and a_{lm} is stored in the first FITS extension as temperature a_{lm} .
		If 1 or 2 is specified, an already existing file is opened and
		a 2nd or 3rd extension is created, treating a_{lm} as a_{lm}^E or a_{lm}^B .
nlheader	I4B IN	number of header lines to write to the file.
header(LEN=80) (1:nlheader)	CHR IN	the header to the FITS-file.

EXAMPLE:

call dump_alms ('alms.fits', alms, 64, header, 80, 1)

Opens an already existing FITS file which contains temperature a_{lm} . An extra extension is added to the file where the a_{lm} array are written in a three-column format as described above. 80 header lines are written to the file from the array header(1:80).

MODULES & ROUTINES

This section lists the modules and routines used by dump_alms*.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to dump_alms*.

fits2alms, read_conbintab routines to read a_{lm} from a FITS-file

alms2fits has the same function as dump_alms* but is more

general.

fits2alms*

fits2alms*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a_{lm} values from a binary FITS file. Each FITS file extension is supposed to contain one integer column with $index = \ell^2 + \ell + m + 1$ and 2 or 4 single (or double) precision columns with real/imaginary a_{lm} values and real/imaginary standard deviation. One can read temperature a_{lm} or temperature and polarisation, a_{lm}^T , a_{lm}^E and a_{lm}^B .

FORMAT	call fits2alms*(filename,	nalms,	alms,	ncl,
	header, nlheader, next)			

ARGUMENTS

name & dimensionality	kind	in/outdescription
filename(LEN=filenamelen)	CHR	IN filename of the FITS-file to read the a_{lm} from.
nalms ncl	I4B I4B	IN number of a_{lm} to read. IN number of columns to read in the FITS file. If an standard deviation is to be read, this number is 5, oth-
next	I4B	erwise it is 3. IN the number of extensions to read. 1 for temperature only, 3 for temperature and polarisation.

alms(1:nalms,1:(ncl+1),1:next)	SP/ ODP	OUT the a_{lm} to read from the file. alms(i,1,j) and $alms(i,2,j)$ contain the ℓ and m values for the ith a_{lm} (j=1,2,3 for (T,E,B)). $alms(i,3,j)$ and $alms(i,4,j)$ contain the real and imaginary value of the ith a_{lm} . Fi- nally, the standard deviation for the ith a_{lm} is contained in $alms(i,5,j)$
nlheader	I4B I	(real) and alms(i,6,j) (imaginary). N number of header lines to read from the file.
header(LEN=80) (1:nlhe 1:next)	ader, CHR (

EXAMPLE:

call fits2alms ('alms.fits', 65*66/2, alms, 3, header, 80, 3)

Reads a FITS file with the a_{lm}^T , a_{lm}^E and a_{lm}^B values read into alms(1:65*66/2,1:4,1:3). The last index specifies (T,E,B). The second index gives l, m, real(a_{lm}), imaginary(a_{lm}) for each of the a_{lm} . The number 65*66/2 is the number of a_{lm} values up to an ℓ value of 64. 80 lines is read from the header in each extension and returned in header(1:80,1:3).

MODULES & ROUTINES

This section lists the modules and routines used by fits2alms*.

read_alms routine called by fits2alms* for each extension.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

fits2alms*

RELATED ROUTINES

This section lists the routines related to fits2alms*.

alms2fits, dump_alms routines to store a_{lm} in a FITS-file

read_conbintab has the same function as fits2alms* but with pa-

rameters passed differently.

number_of_alms,[getsize_fits can be used to find out the number of a_{lm} avail-

able in the file.

fits2cl*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a power spectrum from a FITS ASCII or binary table. The routine can read temperature coeffecients C_l^T or both temperature and polarisation coeffecients C_l^T , C_l^E , C_l^B , $C_l^{T \times E}$. If the keyword PDMTYPE is found in the header, fits2cl assumes the table to be in the special format used by Planck and will ignore the first data column.

\mathbf{FORMAT}

call fits2cl*(filename, clin, lmax, ncl, header, [units])

ARGUMENTS

name & dimensionality	kind	in/out	description
filename(LEN=filenamelen)	CHR	IN	the FITS file containing the
lmax ncl	I4B I4B	IN IN	power spectrum. Maximum ℓ value to be read. 1 for temperature coeffecients
clin(0:lmax,1:ncl)	SP/ DP	OUT	only, 4 for polarisation. the power spectrum read from the file.
header(LEN=80) (1:)	CHR	OUT	the header read from the FITS-file.
units(LEN= 80) (1:)	CHR	OUT	the column units read from the FITS-file.

EXAMPLE:

call fits2cl ('cl.fits',cl,64,4,header,units)

Reads a power spectrum from the FITS file 'cl.fits' and stores the result in cl(0:64,1:4) which are the C_l coeffecients up to l = 64 for $(T, E, B, T \times E)$. The FITS header is returned in header, the column units in units.

 $fits2cl^*$

MODULES & ROUTINES

This section lists the modules and routines used by fits2cl*.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to fits2cl*.

create_alm Routine to create $a_{\ell m}$ values from an input power

spectrum.

write_asctab Routine to create an ascii FITS file containing a

power spectrum.

gaussbeam

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine generates the beam window function in multipole space of a gaussian beam parametrized by its FWHM. The polarization beam is also provided assuming a perfectly co-polarized beam (eg, Challinor et al 2000, astro-ph/0008228)

FORMAT

call gaussbeam(fwhm_arcmin, lmax, beam)

ARGUMENTS

name & dimensionality	kind	in/oı	in/outdescription		
fwhm_arcmin	DP	IN	FWHM of the gaussian beam in arcmin-		
lmax	I4B	IN	utes. maximum ℓ value of the window function.		
beam(0:lmax,1:p)	DP	OUT			

EXAMPLE:

call gaussbeam(5.0_dp, 1024, beam)

Generates the window function of a gaussian beam of FWHM = 5 arcmin, for $\ell \leq 1024$.

RELATED ROUTINES

This section lists the routines related to **gaussbeam**.

generate_beam

Routine returning a beam window function.

gaussbeam 65

 $pixel_window$

Routine returning a pixel window function.

$generate_beam$

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine generates the beam window function in multipole space. It is either a gaussian parametrized by its FWHM in arcmin in real space, or it is read from an external file.

FORMAT call generate_beam(fwhm_arcmin, lmax, beam [, beam_file])

ARGUMENTS

name & dimensionality	kind	in/out	description
fwhm_arcmin	DP	IN	fwhm size of the gaussian beam in arcminutes.
lmax	I4B	IN	maximum ℓ value of the window function.
beam(0:lmax,1:p)	DP	OUT	beam window function generated. The second index runs form 1:1 for temperature only, and 1:3 for polarisation. In the latter case, 1=T, 2=E, 3=B.
beam_file(LEN=filenamelen) (OPTIONAL)	CHR	IN	name of the file containing the (non necessarily gaussian) window function B_{ℓ} of a circular beam. If present, it will override the argument fwhm_arcmin.

EXAMPLE:

call generate_beam(5.0_dp, 1024, beam)

Generates the window function of a gaussian beam of FWHM = 5 arcmin, for $\ell \leq 1024$.

generate_beam 67

MODULES & ROUTINES

This section lists the modules and routines used by **generate_beam**.

alm_tools module, containing:

gaussbeam routine to generate a gaussian beam

RELATED ROUTINES

This section lists the routines related to **generate_beam**.

create_alm Routine to create $a_{\ell m}$ coefficients using gener-

 $ate_beam.$

alter_alm Routine to alter $a_{\ell m}$ coefficients using gener-

 $ate_beam.$

pixel_window Routine returning a pixel window function.

get_card

Location in HEALPix directory tree: src/f90/mod/head_fits.F90

This routine reads a keyword of any kind from a FITS header. It is a wrapper to other routines that read keywords of different kinds.

FORMAT

call get_card(header, kwd, value, comment)

ARGUMENTS

name & dimensionality	kind	in/out	description
header(LEN=80) DIMENSION(:)	CHR	IN	The header to read the keyword from.
kwd(LEN=8)	CHR	IN	the FITS keyword to read (NOT case sensitive).
value	any	OUT	the value read for the keyword. The type of the fortran variable 'value' (double, real, integer, logical or character) should match the type under which the value is written in the FITS file, except if 'value' is a character string, in which case it can read any keyword value, or if 'value' if real or double, in which case it can read any numerical value
comment(LEN=*)	CHR	OUT	comment read for the keyword.

EXAMPLE:

call get_card(header,'NsIdE',nside,comment)

if nside is defined as an integer, it will contain on output the value of NSIDE (say 256) found in header

 get_card 69

EXAMPLE:

call get_card(header, 'ORDERING', ordering, comment)

if ordering is defined as an character string, it will contain on output the value of ORDERING (say 'RING') found in header

MODULES & ROUTINES

This section lists the modules and routines used by **get_card**.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to $\mathbf{get_card}$.

add_card	general purpose routine to write any keywords into a FITS file header
$\operatorname{del_card}$	routine to discard a keyword from a FITS header
read_par, number_of_alms	routines to read specific keywords from a header in a FITS file.
getsize_fits	function returning the size of the data set in a fits file and reading some other useful FITS keywords
$merge_headers$	routine to merge two FITS headers

get_healpix_main_dir, ...

Location in HEALPix directory tree: src/f90/mod/paramfile_io.F90

A few functions are available to return the full path to **HEALPix** main directory and its data and test subdirectories. This allow those paths to be controlled by preprocessing macros or environment variables in case of non-standard installation of the **HEALPix** directory structure.

FUNCTIONS:

hmd = get_healpix_main_dir()

returns the full path to the main **HEALPix** directory. It will be determined, in this order, from the value of the preprocessing macros <code>HEALPIX</code> and <code>HEALPIXDIR</code> if they are defined or the environment variable <code>\$HEALPIX</code> otherwise

hdd = get_healpix_data_dir()

returns the full path to **HEALPix** data subdirectory. It will be determined from the preprocessing macro <code>HEALPIXDATA</code> or the environment variable <code>\$HEALPIXDATA</code>. If both fail, it will return the list of directories {.../data ./data .. \$HEALPIX <code>\$HEALPIX/data</code> \$HEALPIX/.../data \$HEALPIX\data} separated by LineFeed.

htd = get_healpix_test_dir()

returns the full path to **HEALPix** test subdirectory. It will be determined, in this order, from the preprocessing macro HEALPIXTEST, the environment variable \$HEALPIXTEST or \$HEALPIX/test.

getArgument 71

getArgument

Location in HEALPix directory tree: src/f90/mod/extension.F90

This subroutine emulates the C routine getarg, which returns the value of a given command line argument.

FORMAT call getArgument(index, value)

ARGUMENTS

name & dimensionality	kind	in/out	description
index	I4B	IN	index of the command line argument
value	CHR	OUT	(where the first argument has index 1) value of the argument

RELATED ROUTINES

This section lists the routines related to **getArgument**.

getEnvironment returns value of environment variable

nArguments returns number of command line arguments

getEnvironment

Location in HEALPix directory tree: src/f90/mod/extension.F90

This subroutine emulates the C routine getenv, which returns the value of an environment variable.

FORMAT

call getEnvironment(name, value)

ARGUMENTS

name & dimensionality	kind in	out description	
name value	CHR IN CHR OU	name of the environment value of the environment v	

EXAMPLE:

use extension

character(len=128) :: healpixdir

call getEnvironment('HEALPIX', healpixdir)

print*,healpixdir

Will return the value of the \$HEALPIX system variable (if it is

defined)

RELATED ROUTINES

This section lists the routines related to **getEnvironment**.

getArgument returns list of command line arguments

nArguments returns number of command line arguments

getdisc_ring 73

$\underline{\mathbf{getdisc_ring}}$

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

This routine is obsolete, use query_disc instead

$getnumext_fits$

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine returns the number of extensions present in a given FITS file.

FORMAT var=getnumext_fits(filename)

ARGUMENTS

name & dimensionality	kind	in/outdescription		
var	I4B	OUT	number of extensions in the FITS file (excluding the primary unit). According to the current format, HEALPix files have at least one	
filename(LEN=filenamelen))CHR	IN	extension. filename of the FITS file.	

getnumext_fits 75

EXAMPLE:

next = getnumext_fits('map.fits')

Returns in **next** the number of extensions present in the FITS file 'map.fits'.

MODULES & ROUTINES

This section lists the modules and routines used by **getnumext_fits**.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **getnumext_fits**.

getsize_fits routine returning the number of data points in a

FITS file, as well as much more information on

the file.

input_map routine to read a **HEALPix** FITS file

getsize_fits

 $Location \ in \ HEALPix \ directory \ tree: \ src/f90/mod/fitstools.f90$

This routine reads the number of maps and/or the pixel ordering of a FITS file containing a **HEALPix** map.

FORMAT

var=getsize_fits(filename [, nmaps, ordering, obs_npix, nside, mlpol, type, polarisation, fwhm_arcmin, beam_leg, coordsys, polcconv, extno])

ARGUMENTS

getsize_fits 77

name & dimensionality	kind	in/ou	itdescription
var	I8B	OUT	number of pixels or time samples in the chosen extension of the FITS file
filename(LEN=filenamele)	n)CHR	IN	filename of the FITS-file containing HEALPix map(s).
nmaps (OPTIONAL)	I4B	OUT	number of maps in the extension.
ordering (OPTIONAL)	I4B	OUT	pixel ordering, 0=unknown, 1=RING, 2=NESTED
obs_npix (OPTIONAL)	I4B	OUT	number of non blanck pixels. It is set to -1 if it can not be determined from header information alone
nside (OPTIONAL)	I4B	OUT	Healpix resolution parameter Nside. Returns a negative value if not found.
mlpol (OPTIONAL)	I4B	OUT	maximum multipole used to generate the map (for simulated map). Returns a negative value if not found.
type (OPTIONAL) polarisation (OPTIONAL)	I4B	OUT	
polarisation (Of HONAL)	14D	001	 <0: can not find out 0: no polarisation 1: contains polarisation (Q,U or G,C)
fwhm_arcmin (OP-TIONAL)	DP	OUT	returns the beam FWHM read from FITS header, translated from Deg (hopefully) to arcmin. Returns a negative value if not found.
$\begin{array}{c} \text{beam_leg(LEN=filenamelo}\\ \text{(OPTIONAL)} \end{array}$	en¢HR	OUT	filename of beam or filtering window function applied to data (FITS keyword BEAM_LEG). Returns a empty string if not found.
coordsys(LEN=20) (OP-TIONAL)	CHR	OUT	- • •
polcconv (OPTIONAL)	I4B	OUT	polarisation coordinate convention (see Healpix primer for details) 0=unknown, 1=COSMO, 2=IAU
$\underset{\scriptscriptstyle{\text{HEALPix 2.14}}}{\operatorname{extno}}(\underset{\scriptscriptstyle{\text{OPTIONAL}}}{\operatorname{OPTIONAL}})$	I4B	IN	extension number (0 based) for which information is provided. Default = 0 (first extension).

EXAMPLE:

npix= getsize_fits('map.fits', nmaps=nmaps, ordering=ordering,
obs_npix=obs_npix, nside=nside, mlpol=mlpol, type=type,
polarisation=polarisation)

Returns 1 or 3 in nmaps, dependent on wether 'map.fits' contain only temperature or both temperature and polarisation maps. The pixel ordering number is found by reading the keyword ORDERING in the FITS file. If this keyword does not exist, 0 is returned.

MODULES & ROUTINES

This section lists the modules and routines used by getsize_fits.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **getsize_fits**.

getnumext_fits routine returning the number of extension in a

FITS file

input_map routine to read a **HEALPix** FITS file

healpix_modules

Location in HEALPix directory tree: src/f90/mod/healpix_modules.f90

This module is a meta module containing most of the **HEALPix** modules. It currently includes

- alm_tools,
- bit_manipulation,
- coord_v_convert,
- extension,
- fitstools,
- head_fits,
- healpix_fft,
- healpix_types,
- misc_utils,
- num_rec,
- obsolete,
- paramfile_io,
- pix_tools,
- ran_tools,
- rngmod,
- statistics,
- udgrade_nr,
- utilities.

Note that mpi_alm_tools is not included since it requires the MPI library for compilation.

EXAMPLE:

```
use healpix_modules
print*,' pi = ',PI
print*,' number of pixels in a Nside=64 map:',nside2npix(64)
```

Invoking healpix_modules gives access to all **HEALPix** routines and parameters.

healpix_types

Location in HEALPix directory tree: src/f90/mod/healpix_types.F90

This module defines a set of parameters used by most other **HEALPix** modules.

The parameters defined in healpix_types include

• 'kind' parameters, used when defining the type of a variable,

name	type	$value^a$	definition
I1B	integer	1	number of bytes in the hardware-supported signed inte-
			gers covering the range -99 to 99 with the least margin
I2B	integer	2	same as above for the range -9999 to 9999 (ie, 4 digits)
I4B	integer	4	same as above for 9 digits
I8B	integer	8	same as above for 16 digits b
SP	integer	4	number of bytes in the hardware-supported floating-point
			numbers covering the range 10^{-30} to 10^{30} with the least
			margin (hereafter single precision)
DP	integer	8	same as above for the range 10^{-200} to 10^{200} (double pre-
			cision)
SPC	integer	4	number of bytes in real (or imaginary) part of single pre-
			cision complex numbers
DPC	integer	8	same as above for double precision complex numbers
LGT	integer	4	number of bytes in logical variables

 $^{^{}a}$ actual value may depend on hardware or compiler

• largest accessible numbers,

name	type or kind	value^a	definition
MAX_I1B	integer	127	largest number accessible to inte-
			gers of kind I1B
$MAX_{12}B$	integer	32767	same as above for I2B integers
MAX_I4B	integer	$2^{31} - 1 \simeq 2.1 \ 10^9$	same as above for I4B integers
MAX_I8B	I8B	$2^{63} - 1 \simeq 9.2 \ 10^{18}$	same as above for I4B integers
MAX_SP	SP	$\simeq 3.40 \ 10^{38}$	same as above for SP floating-point
MAX_DP	DP	$\simeq 1.80 \ 10^{308}$	same as above for DP floating-point

^aactual value may depend on hardware or compiler

 $[^]b$ may not be supported by some hardware or compiler; on those systems, the user should set the preprocessing variable N064BITS to 1 during compilation to demote automatically I8B to I4B

• mathematical definitions,

name	kind	value	definition
QUARTPI	DP	$\pi/4$	
HALFPI	DP	$\pi/2$	
PΙ	DP	π	
TWOPI	DP	2π	
FOURPI	DP	4π	
SQRT2	DP	$\sqrt{2}$	
EULER	DP	$\gamma \simeq 0.577\dots$	Euler constant
SQ4PI_INV	DP	$1/\sqrt{4\pi}$	
TWOTHIRD	DP	2/3	
DEG2RAD	DP	$\pi/180$	Degrees to Radians conversion factor
RAD2DEG	DP	$180/\pi$	Radians to Degrees conversion factor

• and **HEALPix** specific definitions,

name	type or kind	value	definition
HPX_SBADVAL	SP	$-1.6375 \ 10^{30}$	default sentinel value given to
			missing pixels in single precision
			data sets
HPX_DBADVAL	DP	$-1.6375 \ 10^{30}$	same as above for double preci-
			sion data sets
FILENAMELEN	integer	1024	default length in character of file
			names.

EXAMPLE:

use healpix_types
real(kind=DP) :: dx
print*,' pi = ',PI

The value of PI, as well as all other healpix_types parameters are made known to the code

in_ring

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to find the pixel index of all pixels on a slice of a given ring. The output indices can be either in the RING or NESTED scheme, depending on the nest keyword.

FORMAT

call in_ring(nside, iz, phi0, dphi, listir, nir, nest)

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	the N_{side} parameter of the map.
iz	I4B	IN	ring number, counted south-
			wards from the north pole.
phi0	DP	IN	central ϕ position in the slice.
dphi	DP	IN	defines the size of the slice. The
•			slice has length $2 \times dphi$ along
			the ring with center at $phi0$.
listir(0:4*nside-1)	I4B	OUT	The pixel indexes in the slice.
nir	I4B	OUT	the number of pixels in the slice.
nest (OPTIONAL)	I4B	IN	The pixel indexes are in the
,			NESTED numbering scheme if
			nest=1, and in RING scheme
			otherwise.

EXAMPLE:

call in_ring(256, 10, 0, 0.1, listir, nir, nest=1)

Returns the NESTED pixel index of all pixels within 0.1 radians on each side of $\phi = 0$ on the 10th ring.

in_ring 83

MODULES & ROUTINES

This section lists the modules and routines used by in_ring.

ring2nest conversion from RING scheme pixel index to

NESTED scheme pixel index

next_in_line_nest returns NESTED index of pixel lying to the East

of the current pixel and on the same ring

RELATED ROUTINES

This section lists the routines related to in_ring.

pix2ang, ang2pix convert between angle and pixel number.

pix2vec, vec2pix convert between a cartesian vector and pixel num-

ber.

getdisc_ring find all pixels within a certain radius.

input_map*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a **HEALPix** map from a FITS file. This can deal with full sky as well as cut sky maps

FORMAT

call input_map*(filename, map, npixtot, nmaps [, fmissval, header, units, extno])

ARGUMENTS

name & dimensionality	kind	in/ou	utdescription
filename(len=filenamelen)	CHR	IN	FITS file to be read from, containing a full sky or cut sky map
map(0:npixtot-1,1:nmaps)	SP/ DP	OUT	
npixtot	I4B	IN	number of pixels in the full sky map
nmaps	I4B	IN	number of maps in the file
fmissval (OPTIONAL) header(LEN=80)(1:) (OPTIONAL)	SP/ DP CHR	IN OUT	value to be given to missing pixels, its default value is 0 FITS extension header
units(LEN=20)(1:nmaps) (OPTIONAL)	CHR	OUT	maps units
extno (OPTIONAL)	I4B	IN	extension number to read the data from (0 based).(default: 0) (the first extension is read)

EXAMPLE:

use pix_tools, only: nside2npix

use fitstools, only: getsize_fits, input_map

. . .

input_map*

```
npixtot = getsize_fits('map.fits',nmaps=nmaps, nside=nside)
npix = nside2npix(nside)
allocate(map(0:npix-1,1:nmaps))
call input_map('map.fits', map, npix, nmaps)
```

Reads into map the content of the FITS file 'map.fits'

MODULES & ROUTINES

This section lists the modules and routines used by **input_map***.

${ m fitstools}$	module, containing:
printerror	routine for printing FITS error messages.
${\rm read_bintab}$	routine to read a binary table from a FITS file
$read_fits_cut4$	routine to read cut sky map from a FITS file
cfitsio	library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **input_map***.

anafast	executable that reads a $\mathbf{HEALPix}$ map and anal-
	yses it.
synfast	executable that generate full sky $\mathbf{HEALPix}$ maps
$getsize_fits$	subroutine to know the size of a FITS file.
$output_map$	subroutine to write a FITS file from a ${\bf HEALPix}$
	map
$write_bintabh$	subroutine to write a large array into a FITS file piece by piece
$\mathrm{input_tod}^*$	subroutine to read an arbitrary subsection of a large binary table

$input_tod*$

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a large binary table (for instance a Time Ordered Data set) from a FITS file. The user can choose to read only a section of the table, starting from an arbitrary position. The data can be read into a single or double precision array.

FORMAT call input_tod*(filename, tod, npix, ntods [, header, firstpix, fmissval])

ARGUMENTS

name & dimensionality	kind	in/outdescription		
filename (LEN = filenamelen)	CHR	IN	FITS file to be read from	
tod(0:npix-1,1:ntods)	SP/ DP	OUT	array constructed from the data present in the file (from the sample firstpix to firstpix + npix - 1. Missing pixels or time samples are filled with fmissval.	
npix	I8B	IN	number of pixels or samples to be read. See Note below.	
ntods	I4B	IN	number of columns to read	
header(LEN=80)(1:) (OPTIONAL)	CHR	OUT	FITS extension header	
firstpix (OPTIONAL)	I8B	IN	first pixel (or time sample) to read from (0 based). (default: 0). See Note below.	
fmissval (OPTIONAL)	SP/ DP	IN	value to be given to missing pixels, its default value is 0. Should be of the same type as tod.	

Note: Indices and number of data elements larger than 2^{31} are only accessible in FITS files on computers with 64 bit enabled compilers and with some specific compilation options of cfitsio (see cfitsio documentation).

MODULES & ROUTINES

 $input_tod*$ 87

This section lists the modules and routines used by **input_tod***.

fitstools module, containing:

routine for printing FITS error messages. printerror

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **input_tod***.

anafast executable that reads a **HEALPix** map and anal-

vses it.

executable that generate full sky $\mathbf{HEALPix}$ maps synfast

 $getsize_fits$ subroutine to know the size of a FITS file. subroutine to write large arrays into FITS files write_bintabh output_map

subroutine to write a FITS file from a **HEALPix**

map

subroutine to read a **HEALPix** map (either full input_map

sky of cut sky) from a FITS file

map2alm*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine is a wrapper to 5 internal routines:map2alm_sc, map2alm_sc_pre, map2alm_pol, map2alm_pol_pre1, map2alm_pol_pre2. These routines analyse a **HEALPix** RING ordered map and return a_{lm}^T (and if specified a_{lm}^E and a_{lm}^B) values up to the desired order in ℓ (maximum $3*N_{side}$). The different routines are called depending on what parameters are passed. Some routines analyse with or without precomputed harmonics and some with or without polarisation.

\mathbf{FORMAT}

call map2alm*(nsmax, nlmax, nmmax, map_TQU, alm_TGC, zbounds, w8ring_TQU [, plm])

ARGUMENTS

name & dimensionality	kind	in/outde	escription
nsmax	I4B	IN th	ne N_{side} value of the map to anal-
nlmax	I4B	v	ne maximum ℓ value for the analy-
nmmax	I4B		ne maximum m value for the analsis.
map_TQU(0:12*nsmax**2-1)	SP/ DP	be	only the temperature map is to e analyse, the map-array should be assed with this rank.
map_TQU(0:12*nsmax**2-1, 1:3)	SP/ DP	m ra se	both temperature an polarisation aps are to be analysed, the map arely should have this rank, where the econd index is $(1,2,3)$ corresponding to (T,Q,U) .

map2alm*

alm_TGC(1:p, 0:nlmax, 0:nmmax)	SPC/ DPC	OUT	The a_{lm} values output from the analysis. p is 1 or 3 dependent on wether polarisation is included or not. In the former case, the first index is $(1,2,3)$ corresponding to (T,E,B) .
zbounds(1:2)	DP	IN	section of the map on which to perform the a_{lm} analysis, expressed in terms of $z = \sin(\text{latitude}) = \cos(\theta)$. If zbounds(1) <zbounds(2), analysis="" is="" on="" performed="" strip="" the="" zbounds(1)<<math="">z<zbounds(2); if="" is="" it="" not,="" of="" outside="" performed="" td="" the<=""></zbounds(2);></zbounds(2),>
w8ring_TQU(1:2*nsmax, 1:p)	DP	IN	strip zbounds(2) $< z <$ zbounds(1). ring weights for quadrature corrections. If ring weights are not used, this array should be 1 everywhere. p is 1 for a temperature analysis and 3 for (T,Q,U) .
plm(0:(nlmax+1)(nlmax+2)nsmax-1), OPTIONAL	DP	IN	If this optional matrix is passed with this rank, precomputed $P_{lm}(\theta)$ are used instead of recursion.
plm(0:(nlmax+1)(nlmax+2)nsmax-1,1:3), TIONAL	DP	IN	If this optional matrix is passed with Ω Ris rank, precomputed $P_{lm}(\theta)$ AND precomputed tensor harmonics are used instead of recursion.

EXAMPLE:

```
use healpix_types
use alm_tools
use pix_tools
integer(i4b) :: nside, lmax
real(dp), allocatable, dimension(:,:) :: dw8
real(dp), dimension(2) :: z
real(sp), allocatable, dimension(:,:) :: map
complex(spc), allocatable, dimension(:,:,:) :: alm

nside = 256
lmax = 512
allocate(dw8(1:2*nside, 1:3))
allocate(map(0:nside2npix(nside)-1,1:3))
allocate(alm(1:3, 0:lmax, 0:lmax))
```

Analyses temperature and polarisation maps passed in map. The map has an N_{side} of 256, and the analysis is performed up to 512 in ℓ and m. The resulting a_{lm} coefficients for temperature and polarisation are returned in alm. A 10° cut on each side of the equator is applied. Uniform weights are used. Since the optional plm array is provided with rank one, precomputed scalar $P_{lm}(\theta)$ are used while tensor harmonics are computed with a recursion.

MODULES & ROUTINES

This section lists the modules and routines used by **map2alm***.

ring_analysis Performs FFT for the ring analysis.

misc_util module, containing:

assert_alloc routine to print error message when an array is

not properly allocated

RELATED ROUTINES

This section lists the routines related to map2alm*.

anafast executable using map2alm*to analyse maps.

alm2map routine performing the inverse transform of

map2alm*.

map2alm_iterative similar to map2alm* with iterative scheme.

map2alm_iterative*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine covers and extends the functionalities of map2alm: it analyzes a (polarised) **HEALPix** RING ordered map and returns its a_{lm} coefficients for temperature (and polarisation) up to a specified multipole, and use precomputed harmonics if those are provided, but it also can also perform an iterative (Jacobi) determination of the a_{lm} , and apply a pixel mask if one is provided.

FORMAT

call map2alm_iterative*(nsmax, nlmax, nm-max, iter_order, map_TQU, alm_TGC [, zbounds, w8ring_TQU , plm, mask])

ARGUMENTS

name & dimensionality	kind	in/outdescription
nsmax	I4B	IN the N_{side} value of the map to anal-
		yse.
nlmax	I4B	IN the maximum ℓ value for the analy-
		sis.
nmmax	I4B	IN the maximum m value for the anal-
		ysis.
iter_order	I4B	IN the order of Jacobi iteration. In-
		creasing that order improves the ac-
		curacy of the final a_{lm} but increases
		the computation time $T_{\rm CPU} \propto 1 +$
		$2 \times \text{iter_order}$. iter_order = 0 is a
		straight analysis, while iter_order =
TIOI (0.10*	CD /	3 is usually a good compromise.
$map_TQU(0:12*nsmax**2-1, 1:p)$	SP/	INOUTinput map. p is 1 or 3 depending
	DP	if temperature (T) only or tempera-
		ture and polarisation (T, Q, U) are
		to be analysed. It will be altered on
		output if a mask is provided.

1 TCCC/1 0 1 0	CDC /	OTTO	
alm_TGC(1:p, 0:nlmax, 0:nmmax)	DPC	OUT	The a_{lm} values output from the analysis. p is 1 or 3 depending on whether polarisation is included or not. In the former case, the first index is $(1,2,3)$ corresponding to (T,E,B) .
zbounds(1:2), OPTIONAL	DP	IN	section of the map on which to perform the a_{lm} analysis, expressed in terms of $z = \sin(\text{latitude}) = \cos(\theta)$. If zbounds(1) <zbounds(2), analysis="" is="" on="" performed="" strip="" the="" zbounds(1)<<math="">z<zbounds(2); if="" is="" it="" not,="" of="" outside="" performed="" strip="" the="" zbounds(2)<<math="">z<zbounds(1). absent,="" analyzed<="" if="" is="" map="" td="" the="" whole=""></zbounds(1).></zbounds(2);></zbounds(2),>
w8ring_TQU(1:2*nsmax,1:p), OPTIONAL	DP	IN	ring weights for quadrature corrections. p is 1 for a temperature analysis and 3 for (T,Q,U). If absent, the ring weights are all set to 1.
plm(0:,1:p), OPTIONAL	DP	IN	If this optional matrix is passed, precomputed scalar (and tensor) $P_{lm}(\theta)$ are used instead of recursion.
mask(0:12*nsmax**2-1,1:q), OPTIONAL	SP/ DP	IN	pixel mask, assumed to have the same resolution (and RING ordering) as the map. The map map_TQU is multiplied by that mask before being analyzed, and will therefore be altered on output. q should be in $\{1,2,3\}$. If $p=q=3$, then each of the 3 masks is applied to the respective map. If $p=3$ and $q=2$, the first mask is applied to the first map, and the second mask to the second (Q) and third (U) map. If $p=3$ and $q=1$, the same mask is applied to the 3 maps. Note: the output a_{lm} are computed directly on the masked map, and are not corrected for the loss of power, correlation or leakage created by the mask.

EXAMPLE:

```
use healpix_types
use alm_tools
use pix_tools
integer(i4b) :: nside, lmax, npix, iter
real(sp), allocatable, dimension(:,:) ::
real(sp), allocatable, dimension(:) :: mask
complex(spc), allocatable, dimension(:,:,:)
                                                 alm
nside = 256
lmax = 512
iter = 2
npix = nside2npix(nside)
allocate(map(0:npix-1,1:3))
allocate(mask(0:npix-1))
mask(0:) = 0.! set unvalid pixels to 0
mask(0:10000-1) = 1. ! valid pixels
allocate(alm(1:3, 0:lmax, 0:lmax)
call map2alm_iterative(nside, lmax, lmax, iter, map, alm, mask=mask)
```

Analyses temperature and polarisation signals in the first 10000 pixels of map (as determined by mask). The map has an N_{side} of 256, and the analysis is supposed to be performed up to 512 in ℓ and m. The resulting a_{lm} coefficients for temperature and polarisation are returned in alm. Uniform weights are assumed. In order to improve the allm accuracy, 2 Jacobi iterations are performed.

MODULES & ROUTINES

This section lists the modules and routines used by map2alm_iterative*.

ring_analysis Performs FFT for the ring analysis.

map2alm Perform the alm analysis

misc_util module, containing:

assert_alloc routine to print error message when an array is

not properly allocated

RELATED ROUTINES

This section lists the routines related to map2alm_iterative*.

anafast executable using map2alm_iterative* to analyse

maps.

alm2map routine performing the inverse transform of

map2alm_iterative*.

 $alm 2 map_spin \hspace{1cm} synthesize \hspace{0.1cm} spin \hspace{0.1cm} weighted \hspace{0.1cm} maps.$

map2alm_spin analyze spin weighted maps.

map2alm_spin*

map2alm_spin*

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine extracts the alm coefficients out of maps of spin s and -s. A (complex) map S of spin s is a linear combination of the spin weighted harmonics ${}_sY_{lm}$

$${}_{s}S(p) = \sum_{lm} {}_{s}a_{lm} \quad {}_{s}Y_{lm}(p) \tag{8}$$

for $l \ge |m|, l \ge |s|$, and is such that ${}_sS^* = {}_{-s}S$.

The two (real) input maps for map2alm_spin* are defined respectively as

$$|s|S^{+} = (|s|S + -|s|S)/2$$
 (9)

$$|s|S^{-} = (|s|S - -|s|S)/(2i).$$
 (10)

map2alm_spin* outputs the alm coefficients defined as

$$_{|s|}a_{lm}^{+} = -(_{|s|}a_{lm} + (-1)^{s}_{-|s|}a_{lm})/2$$
 (11)

$$_{|s|}a_{lm}^{-} = -(_{|s|}a_{lm} - (-1)^{s}_{-|s|}a_{lm})/(2i)$$
 (12)

for $m \ge 0$, knowing that, just as for spin 0 maps, the coefficients for m < 0 are given by

$$_{|s|}a_{l-m}^{+} = (-1)^{m}{}_{|s|}a_{lm}^{+*},$$
 (13)

$$a_{l-m}^- = (-1)^m{}_{|s|}a_{lm}^{-*}.$$
 (14)

With these definitions, ${}_2a^+, {}_2a^-, {}_2S^+$ and ${}_2S^-$ match **HEALPix** polarization a^E, a^B, Q and U respectively. However, for $s=0, {}_0a^+_{lm}=-a^T_{lm}, {}_0a^-_{lm}=0, {}_0S^+=T, {}_0S^-=0.$

FORMAT

call map2alm_spin*(nsmax, nlmax, nmmax, spin, map, alm [, zbounds, w8ring_TQU])

ARGUMENTS

name & dimensionality	kind	in/oı	utdescription
nsmax	I4B	IN	the N_{side} value of the map to analyse.
nlmax	I4B	IN	the maximum ℓ value for the analysis.
nmmax	I4B	IN	the maximum m value for the analysis.
spin	I4B	IN	the spin s of the maps to be analysed (only its absolute value is relevant).
map(0:12*nsmax**2-1, 1:2)	SP/ DP	IN	$_{ s }S^{+}$ and $_{ s }S^{-}$ input maps
alm(1:2, 0:nlmax, 0:nmmax)	SPC/ DPC	OUT	The $_{ s }a_{lm}^+$ and $_{ s }a_{lm}^-$ output values.
zbounds(1:2), OPTIONAL	DP	IN	section of the map on which to perform the a_{lm} analysis, expressed in terms of $z = \sin(\text{latitude}) = \cos(\theta)$. If zbounds(1) <zbounds(2), analysis="" is="" on="" performed="" strip="" the="" zbounds(1)<<math="">z<zbounds(2); if="" is="" it="" not,="" of="" outside="" performed="" strip="" the="" zbounds(2)<<math="">z<zbounds(1).< td=""></zbounds(1).<></zbounds(2);></zbounds(2),>
w8ring(1:2*nsmax,1:2), OPTIONAL	DP	IN	ring weights for quadrature corrections. If ring weights are not used, this array should be 1 everywhere.

EXAMPLE:

```
use healpix_types
use alm_tools
use pix_tools
integer(i4b) :: nside, lmax, spin
real(sp), allocatable, dimension(:,:) :: map
complex(spc), allocatable, dimension(:,:,:) :: alm

nside = 256
lmax = 512
spin = 5
allocate(map(0:nside2npix(nside)-1,1:2))
allocate(alm(1:2, 0:lmax, 0:lmax)
...
call map2alm_spin(nside, lmax, lmax, spin, map, alm)
```

map2alm_spin*

Analyses spin 5 and -5 maps. The maps have an N_{side} of 256, and the analysis is performed up to 512 in ℓ and m. The resulting a_{lm} coefficients for are returned in alm.

MODULES & ROUTINES

This section lists the modules and routines used by map2alm_spin*.

ring_analysis Performs FFT for the ring analysis.

compute_lam_mm, get_pixel_layout,

gen_lamfac_der, gen_mfac,

gen_recfac, init_rescale, l_min_ylm Ancillary routines used for ${}_sY_{\ell m}$ recursion

misc_util module, containing:

assert_alloc routine to print error message when an array is

not properly allocated

RELATED ROUTINES

This section lists the routines related to map2alm_spin*.

alm2map_spin routine performing the inverse transform of

map2alm_spin*.

map2alm routine analyzing temperature and polarization

maps

medfiltmap*

 $Location \ in \ HEALPix \ directory \ tree: \ src/f90/mod/pix_tools.f90$

This routine performs the median filtering of a **HEALPix** full sky map for a given neighborhood radius

FORMAT call medfiltmap*(in_map, radius, med_map [, nest, fmissval, fill_holes])

ARGUMENTS

name &	dimensionality	kind	in/outdescription		
$in_map(0)$:npix-1)	SP/	IN	Full sky HEALPix map to filter. npix	
		DP		should be valid HEALPix pixel number.	
radius		DP	IN	Radius in RADIANS of the disk on which the	
				median is computed.	
med_map	0 = 0 = 0	SP/	OUT	Median filtered map: each pixel is the median	
		$\overline{\mathrm{DP}}$		of the input map valid neighboring pixels con-	
				tained within a distance radius	
nest	OPTIONAL	I4B	IN	set to 1 if the map ordering is NESTED, set	
				to 0 if it is RING.	
fmissval	OPTIONAL	SP/	IN	sentinel value given to missing or non-	
		$\overline{\mathrm{DP}}$		valid pixels. Default: HPX_SBADVAL or	
				${\tt HPX_DBADVAL} = -1.6375 \ 10^{30}$	
fill_holes	OPTIONAL	LGT	IN	if set to .true. will replace non-valid pixels	
				by median of neighbors; if set to .false. will	
				leave non-valid pixels unchanged. Default:	
				.false.	

EXAMPLE:

```
use healpix_types
use pix_tools
...
call medfiltmap(map, 0.5*DEG2RAD, med)
```

medfiltmap*

Output in \mathtt{med} the median filter of $\mathtt{map},$ using a filter radius of $0.5~\mathrm{Deg}$

MODULES & ROUTINES

This section lists the modules and routines used by **medfiltmap***.

statistics module, containing:

median routine to compute the median of a data set

pix_tools module, containing:

pix2vec_ring, pix2vec_nest routines to find the location of a pixel on the sky

query_disc routine to find pixels lying within a radius of a

given point

median*

Location in HEALPix directory tree: src/f90/mod/statistics.f90

This function computes the median of a data set

FORMAT var=median*(data [, badval, even])

ARGUMENTS

name & dimensionality	kind	in/ou	ıtdescription
var	SP/ DP	OUT	median of the data set, defined as the middle number (or the average of the 2 middle num- bers) once the valid data points are sorted in monotonous order
data(:)	SP/ DP	IN	data set
badval (OPTIONAL)	SP/ DP	IN	sentinel value given to bad data points. Data points with this value will be ignored during calculation of the median. If not set, all points will be considered. Do not set to 0! .
even (OPTIONAL)	LGT	IN	if set to .true. and the number of valid data points is even, will output the average of the 2 middle points (which doubles the calculation time). If the number of points is odd, the single middle point is output and this keyword is ignored.

EXAMPLE:

```
use statistics, only: median
...
med = median(map, even=.true.)
```

Outputs in med the median of map

median*

MODULES & ROUTINES

This section lists the modules and routines used by **median***.

m_indmed module of the Orderpack 2.0 package, writ-

ten by: Michel Olagnon, http://www.fortran-

2000.com/rank/

indmed routine to output rank of median

RELATED ROUTINES

This section lists the routines related to **median***.

compute_statistics routine min, max, absolute deviation, and first

four order moments of a data set

merge_headers

Location in HEALPix directory tree: src/f90/mod/head_fits.F90 This routine merges two FITS headers.

FORMAT

call merge_headers(header1, header2)

ARGUMENTS

name&dimensionality	kind in/out	$\operatorname{description}$
header1(LEN=80) DIMENSION(:) header2(LEN=80) DIMENSION(:)	CHR IN CHR INOUT	First header. Second header. On output, will contain the concatenation of (in that order) header2 and header1. If header2 is too short to allow the merging the output will be truncated

EXAMPLE:

call merge_headers(header1, header2)

On output header2 will contain the original header2, followed by the content of header1

MODULES & ROUTINES

This section lists the modules and routines used by merge_headers.

$\mathrm{write_hl}$	more general routine for adding a keyword to a
	header.
cfitsio	library for FITS file handling.

merge_headers 103

RELATED ROUTINES

This section lists the routines related to merge_headers.

add _card	general purpose routine to write any keywords
	into a FITS file header
$\operatorname{get_card}$	general purpose routine to read any keywords from a header in a FITS file.
del _card	routine to discard a keyword from a FITS header
read_par, number_of_alms	routines to read specific keywords from a header

in a FITS file.

getsize_fits $\,$ function returning the size of the data set in a fits

file and reading some other useful FITS keywords

mpi_alm_tools*

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This module implements MPI parallelization of the alm2map and map2alm routines. It is not compiled by default during installation, but rather intended for users who need massive parallelization in their own programming. Typical applications are Monte Carlo simulations and Markov chain type analyses.

The routines can be called in two modes, either simple or advanced. The former mimics the interface of the standard routines, but with an additional MPI handle as a first argument, and is intended for applications which requires only one or a few transforms. The latter interface provides both more flexibility (in particular the option of pre-computation of the Legendre polynomials) and a simpler interface when multiple transforms are required. This interface is particularly well suited for Monte Carlo simulations and Markov chain type analyses.

EXAMPLE:

- Simple one-line interfaces:
 - mpi_map2alm_simple
 - mpi_alm2map_simple
- Three-step advanced interfaces:
 - 1. Initialization:
 mpi_initialize_alm_tools
 - 2. Execution of spherical harmonics transforms
 - mpi_map2alm (root processor)
 - mpi_alm2map (root processor)
 - mpi_map2alm_slave (slave processor)
 - mpi_alm2map_slave (slave processor)
 - 3. Finalizing:

 ${\tt mpi_cleanup_alm_tools}$

mpi_alm_tools*

mpi_alm2map*

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine implements MPI parallelization of the serial alm2map routine. It supports both temperature and polarization inputs in both single and double precision. It must only be run by the root node of the MPI communicator.

FORMAT

call mpi_alm2map*(alms, map)

ARGUMENTS

name & dimensionality	kind	in/outdescription
alms(1:nmaps,0:lmax,0:nmax)	SPC or DPC	IN Input alms. If nmaps=1, only temperature information is included; it nmaps=3, polarization information
map(0:npix,1:nmaps)	SP or DP	is included OUT Output map. nmaps must match that of the input alms array.

EXAMPLE:

mpi_alm2map*

This example 1) initializes the mpi_alm_tools module (i.e., allocates internal arrays and defines required parameters), 2) executes a parallel alm2map operation, and 3) frees the previously allocated memory.

MODULES & ROUTINES

This section lists the modules and routines used by mpi_alm2map*.

alm_tools module

RELATED ROUTINES

This section lists the routines related to mpi_alm2map*.

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current routine.
$mpi_initialize_alm_tools$	Allocates memory and defines variables for the mpi_alm_tools module.
$mpi_alm2map_slave$	Routine for executing a parallel inverse spherical harmonics transform (slave processor interface)
${ m mpi_map2alm}$	Routine for executing a parallel spherical harmonics transform (root processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmonics transform (slave processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical harmonics transform
mpi_map2alm_simple	One-line interface to the parallel spherical harmonics transform

mpi_alm2map_simple*

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine provides a simplified (one-line) interface to the MPI version of alm2map. It supports both temperature and polarization inputs in both single and double precision. It must only be run by all nodes in the MPI communicator.

FORMAT

call mpi_alm2map_simple*(comm, alms, map)

ARGUMENTS

name & dimensionality	kind	in/oı	utdescription
$\begin{array}{c} \text{comm} \\ \text{alms} (1:\text{nmaps}, 0:\text{lmax}, 0:\text{nmax}) \end{array}$	I4B SPC or	IN IN	MPI communicator. Input alms. If nmaps=1, only temperature information is included; if
	DPC		nmaps=3, polarization information is included
map(0:npix,1:nmaps)	SP or DP	OUT	Output map. nmaps must match that of the input alms array.

EXAMPLE:

call mpi_alm2map_simple(comm, map, alms)

This example executes a parallel map2alm operation through the one-line interface. Although all processors must supply allocated arrays to the routine, only the root processor's information will be used as input, and only the root processor's alms will be complete after execution.

MODULES & ROUTINES

This section lists the modules and routines used by mpi_alm2map_simple*.

alm_tools module

RELATED ROUTINES

This section lists the routines related to mpi_alm2map_simple*.

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current routine.
$mpi_initialize_alm_tools$	Allocates memory and defines variables for the mpi_alm_tools module.
mpi_alm2map	Routine for executing a parallel inverse spherical harmonics transform (root processor interface)
$mpi_alm2map_slave$	Routine for executing a parallel inverse spherical harmonics transform (slave processor interface)
$\rm mpi_map2alm$	Routine for executing a parallel spherical harmonics transform (root processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmonics transform (slave processor interface)
$mpi_map2alm_simple$	One-line interface to the parallel spherical harmonics transform

mpi_alm2map_slave

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine complements the master routine mpi_alm2map, and should be run by all slaves in the current MPI communicator. It is run without arguments, but after an appropriate call to initialize_mpi_alm_tools.

FORMAT

call mpi_alm2map_slave()

ARGUMENTS

None.

EXAMPLE:

This example 1) initializes the mpi_alm_tools module (i.e., allocates internal arrays and defines required parameters), 2) executes a parallel alm2map operation, and 3) frees the previously allocated memory.

MODULES & ROUTINES

This section lists the modules and routines used by mpi_alm2map_slave.

alm_tools module

RELATED ROUTINES

This section lists the routines related to ${\bf mpi_alm2map_slave}.$

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current routine.
$mpi_initialize_alm_tools$	Allocates memory and defines variables for the mpi_alm_tools module.
${ m mpi_alm2map}$	Routine for executing a parallel inverse spherical harmonics transform (root processor interface)
${\rm mpi_map2alm}$	Routine for executing a parallel spherical harmonics transform (root processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmonics transform (slave processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical harmonics transform
mpi_map2alm_simple	One-line interface to the parallel spherical harmonics transform

$\underline{mpi_cleanup_alm_tools}$

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine deallocates any private arrays previously allocated in the mpi_alm_tools module. It should be run (without arguments) by all processors in the current communicator after the last call to any of the working routines.

FORMAT

call mpi_cleanup_alm_tools()

ARGUMENTS

None.

EXAMPLE:

This example 1) initializes the mpi_alm_tools module (i.e., allocates internal arrays and defines required parameters), 2) executes a parallel map2alm operation, and 3) frees the previously allocated memory.

RELATED ROUTINES

This section lists the routines related to mpi_cleanup_alm_tools.

$mpi_initialize_alm_tools$	Allocates memory and defines variables for the mpi_alm_tools module.
$\mathrm{mpi_alm2map}$	Routine for executing a parallel inverse spherical harmonics transform (root processor interface)
$mpi_alm2map_slave$	Routine for executing a parallel inverse spherical harmonics transform (slave processor interface)
$mpi_map2alm$	Routine for executing a parallel spherical harmonics transform (root processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmonics transform (slave processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical harmonics transform
$mpi_map2alm_simple$	One-line interface to the parallel spherical harmonics transform

mpi_initialize_alm_tools

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine initializes the mpi_alm_tools module, and must be run prior to any of the advanced interface working routines by all processors in the MPI communicator. The root processor must supply all arguments, while it is optional for the slaves. However, the information is disregarded if they do.

A major advantage of MPI parallelization is large quantities of memory, allowing for pre-computation of the Legendre polynomials even with high $N_{\rm side}$ and $\ell_{\rm max}$, since each processor only needs a fraction $(1/N_{\rm procs})$ of the complete table. This feature is controlled by the "precompute_plms" parameter. In general, the CPU time can be expected to decrease by roughly 50% using pre-computed Legendre polynomials for temperature calculations, and by about 30% for polarization calculations.

FORMAT

call mpi_initialize_alm_tools(comm, [nsmax], [nlmax], [nmmax], [zbounds], [polarization], [precompute_plms], [w8ring])

ARGUMENTS

name & dimensionality	kind	in/o	utdescription
comm	I4B	IN	MPI communicator.
nsmax	I4B	IN	the N_{side} value of the HEALPix
			map. (OPTIONAL)
nlmax	I4B	IN	the maximum ℓ value used for the
			a_{lm} . (OPTIONAL)
nmmax	I4B	IN	the maximum m value used for the
			a_{lm} . (OPTIONAL)

zbounds $(1:2)$	DP	IN	section of the map on which to
			perform the a_{lm} analysis, expressed
			in terms of $z = \sin(\text{latitude}) =$
			$\cos(\theta)$. If zbounds(1) <zbounds(2),< td=""></zbounds(2),<>
			the analysis is performed on the
			strip zbounds(1) < z < zbounds(2);
			if not, it is performed <i>outside</i> of the
			strip zbounds(2) $< z >$ zbounds(1).
			(OPTIONAL)
polarization	LGT	IN	if polarization is required, this
			should be set to true, else it should
			be set to false. (OPTIONAL)
$precompute_plms$	I4B	IN	$0 = \text{do not pre-compute any } P_{\ell m}$'s;
			$1 = \text{pre-compute } P_{\ell m}^{\text{T}}; 2 = \text{pre-}$
			compute $P_{\ell_m}^{\mathrm{T}}$ and $P_{\ell_m}^{\mathrm{P}}$. (OP-
			TIONAL)
w8ring_TQU(1:2*nsmax, 1:p)	DP	IN	ring weights for quadrature correc-
,			tions. If ring weights are not used,
			this array should be 1 everywhere. p
			is 1 for a temperature analysis and
			3 for (T,Q,U). (OPTIONAL)
			, , , , ,

EXAMPLE:

This example 1) initializes the mpi_alm_tools module (i.e., allocates internal arrays and defines required parameters), 2) executes a parallel map2alm operation, and 3) frees the previously allocated memory.

RELATED ROUTINES

This section lists the routines related to ${\bf mpi_initialize_alm_tools}.$

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current rou-
	tine.
${ m mpi_alm2map}$	Routine for executing a parallel inverse spherical
	harmonics transform (root processor interface)
mpi_alm2map_slave	Routine for executing a parallel inverse spherical harmonics transform (slave processor interface)
	` -
mpi_map2alm	Routine for executing a parallel spherical harmonics transform (root processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmonics transform (slave processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical harmonics transform
$mpi_map2alm_simple$	One-line interface to the parallel spherical harmonics transform

mpi_map2alm*

mpi_map2alm*

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine implements MPI parallelization of the serial map2alm routine. It supports both temperature and polarization inputs in both single and double precision. It must only be run by the root node of the MPI communicator.

FORMAT call mpi_map2alm*(map, alms)

ARGUMENTS

name & dimensionality	kind	in/outdescription	cription
map(0:npix,1:nmaps)	SP or DP	IN map to analyse. If nmaps=1 only temperature information is in cluded; if nmaps=3, polarization in	temperature information
alms(1:nmaps,0:lmax,0:nmax)	SPC	formation is included OUT output alms. nmaps must equal the	nation is included
,	or DPC	of the input map	ne input map

EXAMPLE:

This example 1) initializes the mpi_alm_tools module (i.e., allocates internal arrays and defines required parameters), 2) executes a parallel map2alm operation, and 3) frees the previously allocated memory.

MODULES & ROUTINES

This section lists the modules and routines used by mpi_map2alm*.

alm_tools module

RELATED ROUTINES

This section lists the routines related to mpi_map2alm*.

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current rou-
	tine.
$mpi_initialize_alm_tools$	Allocates memory and defines variables for the mpi_alm_tools module.
$mpi_alm2map$	Routine for executing a parallel inverse spherical harmonics transform (root processor interface)
$mpi_alm2map_slave$	Routine for executing a parallel inverse spherical harmonics transform (slave processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmonics transform (slave processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical harmonics transform
$mpi_map2alm_simple$	One-line interface to the parallel spherical harmonics transform

mpi_map2alm_simple*

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine provides a simplified (one-line) interface to the MPI version of map2alm. It supports both temperature and polarization inputs in both single and double precision. It must only be run by all processors in the MPI communicator.

FORMAT

call mpi_map2alm_simple*(comm, map, alms, [zbounds], [w8ring])

ARGUMENTS

name & dimensionality	kind	in/outdescription
comm map(0:npix-1,1:nmaps)	I4B SP or DP	IN MPI communicator. IN input map. If nmaps=1, only temperature information is included; if nmaps=3, polarization information is included
alms(1:nmaps,0:lmax,0:nmax)	SPC or DPC	IN output alms. nmaps must equal that of the input map
zbounds(1:2)	DP	IN section of the map on which to perform the a_{lm} analysis, expressed in terms of $z = \sin(\text{latitude}) = \cos(\theta)$. If zbounds(1) <zbounds(2), analysis="" is="" on="" performed="" strip="" the="" zbounds(1)<<math="">z<zbounds(2); if="" is="" it="" not,="" of="" outside="" performed="" strip="" the="" zbounds(2)<<math="">z<zbounds(1). (optional)<="" td=""></zbounds(1).></zbounds(2);></zbounds(2),>
w8ring_TQU(1:2*nsmax, 1:p)	DP	IN ring weights for quadrature corrections. If ring weights are not used, this array should be 1 everywhere. p is 1 for a temperature analysis and 3 for (T,Q,U). (OPTIONAL)

EXAMPLE:

call mpi_map2alm_simple(comm, map, alms)

This example executes a parallel map2alm operation through the one-line interface. Although all processors must supply allocated arrays to the routine, only the root processor's information will be used as input, and only the root processor's alms will be complete after execution.

MODULES & ROUTINES

This section lists the modules and routines used by mpi_map2alm_simple*.

alm_tools module

RELATED ROUTINES

This section lists the routines related to mpi_map2alm_simple*.

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current rou-
	tine.
$mpi_initialize_alm_tools$	Allocates memory and defines variables for the
	mpi_alm_tools module.
${ m mpi_alm2map}$	Routine for executing a parallel inverse spherical
	harmonics transform (root processor interface)
$mpi_alm2map_slave$	Routine for executing a parallel inverse spherical
	harmonics transform (slave processor interface)
${ m mpi_map2alm}$	Routine for executing a parallel spherical harmon-
	ics transform (root processor interface)
$mpi_map2alm_slave$	Routine for executing a parallel spherical harmon-
	ics transform (slave processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical
	harmonics transform

mpi_map2alm_slave

Location in HEALPix directory tree: src/f90/mod/mpi_alm_tools.f90

This subroutine complements the master routine mpi_map2alm, and should be run by all slaves in the current MPI communicator. It is run without arguments, but after an appropriate call to initialize_mpi_alm_tools.

FORMAT

call mpi_map2alm_slave()

ARGUMENTS

None.

EXAMPLE:

This example 1) initializes the mpi_alm_tools module (i.e., allocates internal arrays and defines required parameters), 2) executes a parallel map2alm operation, and 3) frees the previously allocated memory.

MODULES & ROUTINES

This section lists the modules and routines used by mpi_map2alm_slave.

alm_tools module

RELATED ROUTINES

This section lists the routines related to ${\bf mpi_map2alm_slave}.$

$mpi_cleanup_alm_tools$	Frees memory that is allocated by the current routine.
$mpi_initialize_alm_tools$	Allocates memory and defines variables for the mpi_alm_tools module.
$mpi_alm2map$	Routine for executing a parallel inverse spherical harmonics transform (root processor interface)
$mpi_alm2map_slave$	Routine for executing a parallel inverse spherical harmonics transform (slave processor interface)
${ m mpi_map2alm}$	Routine for executing a parallel spherical harmonics transform (root processor interface)
$mpi_alm2map_simple$	One-line interface to the parallel inverse spherical harmonics transform
$mpi_map2alm_simple$	One-line interface to the parallel spherical harmonics transform

nArguments 123

nArguments

Location in HEALPix directory tree: src/f90/mod/extension.F90

This function emulates the C routine iargc, which returns the number of command line arguments provided.

FORMAT var=nArguments()

ARGUMENTS

name&dimensionality	kind	in/out	description
var	I4B	OUT	number of command line arguments

RELATED ROUTINES

This section lists the routines related to nArguments.

getEnvironment returns value of environment variable getArgument returns list of command line arguments

$neighbours_nest$

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

This subroutine returns the number and locations (in terms of pixel numbers) of the topological neighbours of a central pixel. The pixels are ordered in a clockwise sense about the central pixel with the southernmost pixel in first element. For the 4 pixels in the southern corners of the equatorial faces which have two equally southern neighbours the routine returns the southwestern pixel first and proceeds clockwise.

FORMAT

call neighbours_nest(nside, ipix, list, nneigh)

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	The N_{side} parameter of the map.
ipix	I4B	IN	The pixel number of the central pixel.
list(8)	I4B	OUT	On exit, the vector of neighbouring pixels. This contains nneigh relevant elements.
nneigh	I4B	OUT	The number of neighbours (mostly 8, except at 8 sites, where there are only 7 neighbours).

EXAMPLE:

use pix_tools

integer :: n, list(1:8)

call neighbours_nest(4, 1, list, nneigh)

print*,nneigh

print*,list(1:nneigh)

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This returns nneigh= 8 and a vector list, which contains the pixel numbers (90, 0, 2, 3, 6, 4, 94, 91).

MODULES & ROUTINES

This section lists the modules and routines used by **neighbours_nest**.

mk_xy2pix, mk_pix2xy precomputing arrays for the conversion of

NESTED pixel numbers to Cartesian coords in

each face.

pix2xy_nest, xy2pix_nest Conversion between NESTED pixel numbers to

Cartesian coords in each face.

bit_manipulation module, containing:

invMSB, invLSB,swapLSBMSB,invswapLSBMSB functions which manip-

ulate the bit vector which represents the NESTED pixel numbers. They correspond to NorthWest;-¿SouthEast, SouthWest;-¿NorthEast, East;-¿West and North-South flips of the diamond

faces, respectively.

RELATED ROUTINES

This section lists the routines related to **neighbours_nest**.

pix2ang, ang2pix convert between angle and pixel number.

pix2vec, vec2pix convert between a cartesian vector and pixel num-

ber.

npix2nside

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Function to provide the resolution parameter N_{side} corresponding to N_{pix} pixels over the full sky.

FORMAT var=npix2nside(npix)

ARGUMENTS

name & dimensionality	kind	in/out	description
npix	I4B	IN	the number N_{pix} of pixels over the whole sky.
var	I4B	OUT	the parameter $N_{\rm side}$. If $N_{\rm pix}$ is valid (12 times a power of 2 in $\{1, \dots, 8192\}$), $N_{\rm side} = \sqrt{N_{\rm pix}/12}$ is returned; if not, an error message is issued and -1 is returned.

EXAMPLE:

nside= npix2nside(786432)

Returns the resolution parameter N_{side} (256) corresponding to 786432 pixels on the sky.

RELATED ROUTINES

This section lists the routines related to **npix2nside**.

nside2npix returns the number of pixels $N_{\rm pix}$ corresponding to resolution parameter $N_{\rm side}$

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nside2npix

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Function to provide the number of pixels $N_{\rm pix}$ over the full sky corresponding to resolution parameter $N_{\rm side}$.

FORMAT var=nside2npix(nside)

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	the $N_{\rm side}$ parameter of the map. the number of pixels $N_{\rm pix}$ of the map. If $N_{\rm side}$ is valid (a power of 2 in $\{1,\ldots,8192\}$), $N_{\rm pix}=12N_{\rm side}^2$ is returned; if not, an error message is issued and -1 is returned.
var	I4B	OUT	

EXAMPLE:

npix= nside2npix(256)

Returns the number of **HEALPix** pixels (786432) for the resolution parameter 256.

RELATED ROUTINES

This section lists the routines related to **nside2npix**.

npix2nside returns resolution parameter corresponding to the number of pixels.

nside2ntemplates

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Function to provide the number of template pixels

$$N_{\text{templates}} = \frac{1 + N_{\text{side}}(N_{\text{side}} + 6)}{4}$$

corresponding to resolution parameter N_{side} . Each template pixel has a different shape that $can \ not$ be matched (by rotation or reflexion) to that of any of the other templates.

FORMAT

var=nside2ntemplates(nside)

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	the N_{side} parameter.
var	I4B	OUT	the number of template pixels $N_{\text{templates}}$.

EXAMPLE:

ntpl= nside2ntemplates(256)

Returns in ntpl the number of **HEALPix** template pixels (16768) for the resolution parameter 256.

RELATED ROUTINES

This section lists the routines related to **nside2ntemplates**.

 $template_pixel_ring$

template_pixel_nest return the template pixel associated with any

HEALPix pixel

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 $same_shape_pixels_ring \\ same_shape_pixels_nest$

return the ordered list of pixels having the same shape as a given pixel template

number_of_alms

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This function returns the number of a_{lm} values stored in each FITS extension in a FITS file containing a_{lm}

FORMAT var=number_of_alms(filename[, extnum])

ARGUMENTS

name & dimensionality	kind	in/out	description
filename(LEN = filenamelen)	CHR	IN	filename of the FITS-file con-
extnum	I4B	OUT	taining $a_{\ell m}$. number of extensions in the file

EXAMPLE:

print*,number_of_alms('alms.fits')

Prints the number of a_{lm} stored in each extension of the file 'alms.fits'

MODULES & ROUTINES

This section lists the modules and routines used by **number_of_alms**.

fitstools	module, containing:
printerror	routine for printing FITS error messages.
${f cfitsio}$	library for FITS file handling.

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RELATED ROUTINES

This section lists the routines related to **number_of_alms**.

 $fits 2 alms, \ read_conbintab$

routines that read a_{lm} values from FITS files.

output_map*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine writes a full sky **HEALPix** map into a FITS file. The map can be either single or double precision real. It *has* to be 2-dimensional.

ARGUMENTS

name & dimensionality	kind	in/out	description
map(0:,1:)	SP/ DP	IN	full sky map(s) to be output
header(LEN=80)(1:)	CHR	IN	string array containing the FITS header to be included in the file
filename(LEN=*)	CHR	IN	filename of the FITS-file to contain HEALPix map(s).
extno	I4B	IN	extension number in which to write the data (0 based). (default: 0)

EXAMPLE:

output_map* 133

MODULES & ROUTINES

This section lists the modules and routines used by **output_map***.

fitstools module, containing:

routine for printing FITS error messages. printerror

 $write_bintab$ routine to write a binary table into a FITS file.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **output_map***.

executable that reads a **HEALPix** map from a anafast

FITS file and analyses it.

synfast executable that generate full sky **HEALPix** maps input_map

subroutine to read a **HEALPix** map from a a

FITS file

write_bintabh subroutine to write a large array into a FITS file

piece by piece

input_tod* subroutine to read an arbitrary subsection of a

large binary table

write_minimal_header routine to write minimal FITS header

$parse_xxx$

Location in HEALPix directory tree: src/f90/mod/paramfile_io.F90

The Fortran 90 module paramfile \perp io contains functions to obtain parameters from parameter files or interactively

ARGUMENTS

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name&dimensionality	kind	in/out	description
fname	CHR	IN	file containing the simulation parameters. If empty, parameters are obtained interactively.
handle	PMF	INOUT	Object of type (paramfile_handle) used to store parameter information
keyname	CHR	IN	name of the required parameter
default	XXX	IN	optional argument containing the default value for a given simulation parameter; must be of appropriate type.
vmin	XXX	IN	optional argument containing the minimum value for a given simulation parameter; must be of appropriate type.
vmax	XXX	IN	optional argument containing the maximum value for a given simulation parameter; must be of appropriate type.
descr	CHR	IN	optional argument containing a description of the required simulation parameter
filestatus	CHR	IN	optional argument. If present, the parameter must be a valid filename. If filestatus=='new', the file must not exist; if filestatus=='old', the file must exist.
code	CHR	IN	optional argument. Contains the name of the executable.
silent	LGT	IN	optional argument. If set to .true. the parsing routines will run silently in non-interactive mode (except for warning or error messages, which will always appear). This is mainly intended for MPI usage where many processors parse the same parameter file: silent can be set to .true. on all CPUs except one.

ROUTINES:

```
handle = parse_init (fname [,silent])
  initializes the parser to work on the file fname, or interactively, if fname is empty
intval = parse_int (handle, keyname [, default, vmin, vmax, descr])
longval = parse_long (handle, keyname [, default, vmin, vmax, descr])
realval = parse_real (handle, keyname [, default, vmin, vmax, descr])
doubleval = parse_double (handle, keyname [, default, vmin, vmax, descr])
stringval = parse_string (handle, keyname [, default, descr, filestatus])
logicval = parse_lgt (handle, keyname [, default, descr])
```

These routines obtain integer(i4b), integer(i8b), real(sp), real(dp), character(len=*) and logical values, respectively.

Note: parse_string will expand all environment variables of the form \${XXX} (eg: \${HOME}). It will also replace a leading ~/ by the user's home directory.

```
call parse_summarize (handle [, code])
```

if the parameters were set interactively, this routine will print out a parameter file performing the same settings.

```
call parse_check_unused (handle [, code])
```

if a parameter file was read, this routine will print out all the parameters found in the file but not used by the code. Intended at detecting typos in parameter names.

```
call parse_finish (handle)
```

frees the memory

EXAMPLE:

```
program who_r_u
use healpix_types
use paramfile_io
use extension

implicit none
type(paramfile_handle) :: handle
character(len=256) :: parafile, name
real(DP) :: age

parafile = ''
if (nArguments() == 1) call getArgument(1, parafile)
handle = parse_init(parafile)
name = parse_string(handle, 'name',descr='Enter your last name: ')
age = parse_double(handle, 'age', descr='Enter your age in years: ', &
```

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```
& default=18.d0,vmin=0.d0)
call parse_summarize(handle, 'who_r_u')
end program who_r_u
```

If a file is provided as command line argument when running the executable who_r_u, that file will be parsed in search of the lines starting with 'name =' and 'age =', otherwise the same questions will be asked interactively.

RELATED ROUTINES

This section lists the routines related to parse_xxx.

concatnl generates from a set of strings the multi-line description

nArguments returns the number of command line arguments getArgument returns the list of command line arguments

pixel_window

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine returns the averaged ℓ -space window function $w_{\text{pix}}(\ell)$ (for temperature and polarisation) associated to **HEALPix** pixels of resolution parameter N_{side} . Because of the integration of the signal over the pixel area, the $a_{lm}^{(\text{pix})}$ coefficients of a pixelized map are related to the unpixelized underlying a_{lm} by $a_{lm}^{(\text{pix})} = a_{lm}w_{\text{pix}}(\ell)$.

Unless specified otherwise, the $w_{\rm pix}(\ell)$ are read from the files $HEALPIX/data/pixel_window_n????.fits.$

FORMAT

call pixel_window(pixlw, nside [, windowfile])

ARGUMENTS

name & dimensionality	kind	in/ou	utdescription
pixlw(0:lmax,1:p)	DP	OUT	pixel window function(s) $w_{\rm pix}(\ell)$ generated. The first index must be $\ell_{\rm max} \leq 4N_{\rm side}$. The second index runs from 1:1 for temperature only, and 1:3 for polarisation. In the latter case, 1=T, 2=E, 3=B.
nside	I4B	IN	HEALPix N_{side} resolution parameter. Unless windowfile is set, the file associated with N_{side} and shipped with the package is read by default. If $\text{nside} = 0$, the pixel is assumed infinitely small and pixlw is returned with value 1.
windowfile (OPTIONAL)	CHR	IN	FITS file containing the pixel window to be read instead of the default.

EXAMPLE:

call pixel_window(pixlw, 64)

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returns in pixlw the pixel window function for $N_{\text{side}} = 64$.

MODULES & ROUTINES

This section lists the modules and routines used by **pixel_window**.

misc_utils module, containing:

assert, fatal_error interrupt code in case of error

extension module, containing:

getEnvironment read environment variable

fitstools module, containing:

read_dbintab reads double precision binary table

RELATED ROUTINES

This section lists the routines related to **pixel_window**.

gaussbeam routine to generate a gaussian beam window func-

tion

generate_beam returns a beam window function

alter_alm, rotate_alm modifies a_{lm} to emulate effect of real space filtering

and coordinate rotation respectively

alm2map synthetize a **HEALPix** map from its a_{lm} (or

 $a_{lm}^{(\text{pix})}$).

alm2map_der synthetize a map and its derivatives from its a_{lm}

(or $a_{lm}^{(pix)}$).

$\begin{array}{l} pix2xxx, ang2xxx, vec2xxx, \\ nest2ring, ring2nest \end{array}$

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

The Fortran90 module pix_tools contains some subroutines to convert between pixel number in the **HEALPix** map and (θ, ϕ) or (x, y, z) coordinates on the sphere. Some of these routines are listed here.

ARGUMENTS

name & dimensional-	kind	in/out	description
ity			
nside	I4B	IN	$N_{\rm side}$ parameter for the HEALPix map.
ipnest	I4B		pixel identification number in NESTED
			scheme over the range $\{0, N_{\text{pix}} - 1\}$.
ipring	I4B		pixel identification number in RING scheme
-			over the range $\{0, N_{\text{pix}} - 1\}$.
theta	DP		colatitude in radians measured southward
			from north pole in $\{0,\pi\}$.
phi	DP		longitude in radians, measured eastward in
-			$[0, 2\pi].$
vector(3)	DP		three dimensional cartesian position vector
、 /			(x, y, z). The north pole is $(0, 0, 1)$. An out-
			put vector is normalised to unity.
vertex(3,4)	DP	OUT	three dimensional cartesian position vectors
OPTIONAL			(x, y, z) (normalised to unity) pointing to the
			4 vertices of a given pixel. The four vertices
			are listed in the order North, West, South,
			East.

ROUTINES:

call pix2ang_ring(nside, ipring, theta, phi)

renders theta and phi coordinates of the nominal pixel center given the pixel number ipring and a map resolution parameter nside.

call pix2vec_ring(nside, ipring, vector [,vertex])

renders cartesian vector coordinates of the nominal pixel center given the pixel number *ipring* and a map resolution parameter *nside*. Optionally renders cartesian vector coordinates of the considered pixel four vertices.

call ang2pix_ring(nside, theta, phi, ipring)

renders the pixel number *ipring* for a pixel which, given the map resolution parameter *nside*, contains the point on the sphere at angular coordinates *theta* and *phi*.

call vec2pix_ring(nside, vector, ipring)

renders the pixel number *ipring* for a pixel which, given the map resolution parameter *nside*, contains the point on the sphere at cartesian coordinates *vector*.

call pix2ang_nest(nside, ipnest, theta, phi)

renders theta and phi coordinates of the nominal pixel center given the pixel number ipnest and a map resolution parameter nside.

call pix2vec_nest(nside, ipnest, vector [,vertex])

renders cartesian vector coordinates of the nominal pixel center given the pixel number *ipnest* and a map resolution parameter *nside*. Optionally renders cartesian vector coordinates of the considered pixel four vertices.

call ang2pix_nest(nside, theta, phi, ipnest)

renders the pixel number *ipnest* for a pixel which, given the map resolution parameter *nside*, contains the point on the sphere at angular coordinates *theta* and *phi*.

call vec2pix_nest(nside, vector, ipnest)

renders the pixel number ipnest for a pixel which, given the map resolution parameter nside, contains the point on the sphere at cartesian coordinates vector.

call nest2ring(nside, ipnest, ipring)

performs conversion from NESTED to RING pixel number.

call ring2nest(nside, ipring, ipnest)

performs conversion from RING to NESTED pixel number.

MODULES & ROUTINES

This section lists the modules and routines used by pix2xxx,ang2xxx,vec2xxx, nest2ring,ring2nest.

mk_pix2xy, mk_xy2pix routine

routines used in the conversion between pixel values and "cartesian" coordinates on the Healpix face.

RELATED ROUTINES

This section lists the routines related to pix2xxx,ang2xxx,vec2xxx, nest2ring,ring2nest.

neighbours_nest find neighbouring pixels.

ang2vec convert (θ, ϕ) spherical coordinates into (x, y, z)

cartesian coordinates.

vec2ang convert (x, y, z) cartesian coordinates into (θ, ϕ)

spherical coordinates.

convert_inplace in-place conversion between RING and NESTED

for integer/real/double maps.

convert_nest2ring convert from NESTED to RING scheme using a

temporary array.

planck_rng

Location in HEALPix directory tree: src/f90/mod/rngmod.f90

The derived type planck_rng is used by the Random Number Generation routines rand_init, rand_uni, rand_gauss to describe fully the current RNG sequence.

Most users do not need to know about the planck_rng definition. It may be useful for those wanting to take a snapshot of the RNG sequence they are using (by eg, dumping the latest values of planck_rng structure on disk) so that the same sequence can be resumed later on from that same point.

The type planck_rng is a structure defined as

name	type	definition
x, y, z, w	I4B	internal variables of uniform RNG
gset	DP	internal variable for Gaussian RNG
empty	LGT	flag used by Gaussian RNG

RELATED ROUTINES

This section lists the routines related to **planck_rng**.

${ m rand_gauss}$	function which returns a random normal deviate.
${ m rand_uni}$	function which returns a random uniform deviate.
$\operatorname{rand_init}$	subroutine to initiate a random number sequence.

plm_gen

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This routine computes the latitude dependent part $\lambda_{\ell m}$ of the spherical harmonics $(Y_{\ell m}(\theta,\phi)=\lambda_{\ell m}(\theta)e^{im\phi})$ of spin 0 and 2 (see **HEALPix** primer) used to synthetize or analyze **HEALPix** maps of temperature and polarisation.

FORMAT

call plm_gen(nsmax, nlmax, nmmax, plm)

ARGUMENTS

name & dimensionality	kind	in/out	description
nsmax	I4B	IN	The N_{side} value for which to
nlmax	I4B	IN	compute the $\lambda_{\ell m}$. The maximum multipole order ℓ of the generated $\lambda_{\ell m}$.
nmmax	I4B	IN	The maximum degree m of the generated λ_{lm} .
plm(0:n_plm-1, 1:p)	DP	OUT	The λ_{lm} values, either for temperature only $(p=1)$ or temperature and polarisation $(p=3)$. The number of $\lambda_{\ell m}$ is n-plm = nsmax*(nmmax+1)*(2*nlmax-nmmax+2). They are stored in the order of increasing order ℓ , increasing degree m , for all the HEALPix ring colatitudes θ from North Pole to Equator, ie $\lambda_{00}(\theta_1)$, $\lambda_{10}(\theta_1)$, $\lambda_{20}(\theta_1)$,, $\lambda_{11}(\theta_1)$, $\lambda_{21}(\theta_1)$;, $\lambda_{00}(\theta_2)$

EXAMPLE:

plm_gen 145

```
use healpix_types
use alm_tools, only : plm_gen
integer(I4B) :: nside, lmax, mmax, n_plm
real(DP), dimension(:,:), allocatable :: plm
...
nside=256 ; lmax=512 ; mmax=lmax
npix=nside2npix(nside)
n_plm=nside*(mmax+1)*(2*lmax-mmax+2)
allocate(plm(0:n_plm-1,1:3))
...
call plm_gen(nside, lmax, mmax, plm)
```

Computes the spherical harmonics for temperature and polarisation for $N_{side} = 256$, up to 512 in ℓ and m.

MODULES & ROUTINES

This section lists the modules and routines used by **plm_gen**.

compute_lam_mm, get_pixel_layout, gen_lamfac,gen_mfac, gen_normpol, gen_recfac, init_rescale, l_min_ylm

Ancillary routines used for $\lambda_{\ell m}$ recursion

 $misc_utils$ module, containing:

assert_alloc routine to print error message, when an array can

not be allocated properly

RELATED ROUTINES

This section lists the routines related to plm_gen.

alm2map routine generating maps of temperature and po-

larisation from their $a_{\ell m}$ that can use precom-

puted $\lambda_{\ell m}$ generated by plm_gen

map2alm routine analysing maps of temperature and polar-

isation that can use precomputed $\lambda_{\ell m}$ generated

by plm_gen

plmgen executable using plm-gen to compute the $\lambda_{\ell m}$ and

writting them on disc

query_disc 147

$query_disc$

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to find the index of all pixels within an angular distance radius from a defined center. The output indices can be either in the RING or NESTED scheme

FORMAT

call query_disc(nside, vector0, radius, listpix, nlist [, nest, inclusive])

ARGUMENTS

name & dimensional-	kind	in/out	description
ity			
nside	I4B	IN	the N_{side} parameter of the map.
vector0(3)	DP	IN	cartesian vector pointing at the disc center.
radius	DP	IN	disc radius in radians.
listpix(0:*)	I4B	OUT	the index for all pixels within radius.
			Make sure that the size of the array is big
			enough to contain all pixels.
nlist	I4B	OUT	The number of pixels listed in <i>listpix</i> .
nest (OPTIONAL)	I4B	IN	The pixel indices are in the NESTED num-
,			bering scheme if nest=1, and in RING
			scheme otherwise.
inclusive (OPTIONAL)	I4B	IN	If set to 1, all the pixels overlapping (even
,			partially) with the disc are listed, other-
			wise only those whose center lies within the
			disc are listed.

EXAMPLE:

call query_disc(256,(/0,0,1/),pi/2,listpix,nlist,nest=1)

Returns the NESTED pixel index of all pixels north of the equatorial line in a $N_{side}=256$ map.

MODULES & ROUTINES

This section lists the modules and routines used by query_disc.

in_ring routine to find the pixels in a certain slice of a

given ring.

ring_num function to return the ring number corresponding

to the coordinate z

RELATED ROUTINES

This section lists the routines related to query_disc.

pix2ang, ang2pix convert between angle and pixel number.

pix2vec, vec2pix convert between a cartesian vector and pixel num-

ber.

 $query_disc,\ query_polygon,$

query_strip, query_triangle render the list of pixels enclosed respectively in a

given disc, polygon, latitude strip and triangle

query_polygon 149

query_polygon

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to find the index of all pixels enclosed in a polygon. The polygon should be convex, or have only one concave vertex. The edges should not intersect each other. The output indices can be either in the RING or NESTED scheme

FORMAT call query_polygon(nside, vlist, nv, listpix, nlist [, nest, inclusive])

ARGUMENTS

name & dimension-	kind	in/out	description	
ality				
nside	I4B	IN	the N_{side} parameter of the map.	
vlist(1:3,0:*)	DP	IN	cartesian vector pointing at polygon respec- tive vertices.	
nv	I4B	IN	number of vertices, should be equal to 3 or larger.	
listpix(0:*)	I4B	OUT	the index for all pixels enclosed in the triangle. Make sure that the size of the array is big enough to contain all pixels.	
nlist	I4B	OUT	The number of pixels listed in <i>listpix</i> .	
nest (OPTIONAL)	I4B	IN	The pixel indices are in the NESTED numbering scheme if nest=1, and in RING scheme otherwise.	
inclusive (OP-TIONAL)	I4B	IN	If set to 1, all the pixels overlapping (even partially) with the polygon are listed, otherwise only those whose center lies within the polygon are listed.	

EXAMPLE:

real(dp), dimension(1:3,0:9) :: vertices vertices(:,0) = (/0.,0.,1./) ! +z

```
vertices(:,1) = (/1.,0.,0./) ! +x
vertices(:,2) = (/1.,1.,-1./) ! x+y-z
vertices(:,3) = (/0.,1.,0./) ! +y
call query_polygon(256,vertices,4,listpix,nlist,nest=0)
```

Returns the RING pixel index of all pixels in the polygon with vertices of cartesian coordinates (0,0,1), (1,0,0), (1,1,-1) and (0,1,0) in a $N_{side}=256$ map.

MODULES & ROUTINES

This section lists the modules and routines used by query_polygon.

isort	routine to sort integer number
${\it query_triangle}$	render the list of pixels enclosed in a given triangle
$surface_triangle$	computes the surface of a spherical triangle defined by 3 vertices
$\operatorname{vect_prod}$	routine to compute the vectorial product of two 3D vectors

RELATED ROUTINES

This section lists the routines related to query_polygon.

pix2ang, ang2pix	convert between angle and pixel number.
pix2vec, vec2pix	convert between a cartesian vector and pixel num-
	ber.
query_disc, query_polygon,	
query_strip, query_triangle	render the list of pixels enclosed respectively in a given disc, polygon, latitude strip and triangle

query_strip 151

query_strip

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to find the index of all pixels enclosed in a latitude strip. The output indices can be either in the RING or NESTED scheme

FORMAT call query_strip(nside, theta1, theta2, listpix, nlist [, nest, inclusive])

ARGUMENTS

name&dimensionalitykind		in/out	description	
nside	I4B	IN	the N_{side} parameter of the map.	
theta1	DP	IN	colatitude lower bound in radians measured	
			from North Pole (between 0 and π).	
theta2	DP	IN	colatitude upper bound in radians mea-	
			sured from North Pole (between 0 and	
			π). If theta1< theta2, the pixels lying in	
			[theta1,theta2] are output, otherwise, the	
			pixel lying in [0, theta2] and those lying in	
			[theta1, π] are output.	
listpix(0:*)	I4B	OUT	the index for all pixels enclosed in the strip(s).	
			Make sure that the size of the array is big	
	T.(D)	0.1177	enough to contain all pixels.	
nlist	I4B	OUT	The number of pixels listed in <i>listpix</i> .	
nest (OPTIONAL)	I4B	IN	The pixel indices are in the NESTED num-	
			bering scheme if nest=1, and in RING scheme	
	TAD	T. T.	otherwise.	
inclusive (OP-	I4B	IN	If set to 1, all the pixels overlapping (even	
TIONAL)			partially) with the strip are listed; otherwise	
			only those whose center lies within the strip	
			are listed.	

EXAMPLE:

call query_strip(256,0.75*PI,0.2*PI,listpix,nlist,nest=1)

Returns the NESTED pixel index of all pixels with colatitude in $[0,\pi/5]$ and those with colatitude in $[3\pi/4,\pi]$

MODULES & ROUTINES

This section lists the modules and routines used by query_strip.

$\operatorname{in_ring}$	routine to find the pixels in a certain slice of a given ring.
$intrs_intrv$	routine to compute the intersection of 2 intervals on a circle
ring_num	function to return the ring number corresponding to the coordinate \boldsymbol{z}
$\operatorname{vect_prod}$	routine to compute the vectorial product of two 3D vectors

RELATED ROUTINES

This section lists the routines related to query_strip.

pix2ang, ang2pix	convert between angle and pixel number.
pix2vec, vec2pix	convert between a cartesian vector and pixel number.
query_disc, query_polygon,	
query_strip, query_triangle	render the list of pixels enclosed respectively in a given disc, polygon, latitude strip and triangle
$surface_triangle$	computes the surface of a spherical triangle defined by 3 vertices

query_triangle 153

query_triangle

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to find the index of all pixels enclosed in a spherical triangle described by its three vertices. The output indices can be either in the RING or NESTED scheme

FORMAT call query_triangle(nside, v1, v2, v3, listpix, nlist [, nest, inclusive])

ARGUMENTS

name&dimensionality	kind	in/out	out description	
nside	I4B	IN	the N_{side} parameter of the map.	
v1(3)	DP	IN	cartesian vector pointing at the triangle	
			first vertex.	
v2(3)	DP	IN	cartesian vector pointing at the triangle	
			second vertex.	
v3(3)	DP	IN	cartesian vector pointing at the triangle	
			third vertex.	
listpix(0:*)	I4B	OUT	the index for all pixels enclosed in the tri-	
			angle. Make sure that the size of the array	
			is big enough to contain all pixels.	
nlist	I4B	OUT	The number of pixels listed in $listpix$.	
nest (OPTIONAL)	I4B	IN	The pixel indices are in the NESTED num-	
			bering scheme if nest=1, and in RING	
			scheme otherwise.	
inclusive (OPTIONAL)	I4B	IN	If set to 1, all the pixels overlapping (even	
			partially) with the triangle are listed, oth-	
			erwise only those whose center lies within	
			the triangle are listed.	

EXAMPLE:

call query_triangle(256, (/1,0,0/), (/0,1,0/), (/0,0,1/), listpix, nlist)

Returns the RING pixel index of the (98560) pixels in the octant (x, y, z > 0) in a $N_{side} = 256$ map.

MODULES & ROUTINES

This section lists the modules and routines used by query_triangle.

in_ring routine to find the pixels in a certain slice of a given ring.
intrs_intrv routine to compute the intersection of 2 intervals on a circle
ring_num function to return the ring number corresponding to the coordinate z

vect_prod routine to compute the vectorial product of two

3D vectors

RELATED ROUTINES

This section lists the routines related to query_triangle.

pix2ang, ang2pix convert between angle and pixel number.

pix2vec, vec2pix convert between a cartesian vector and pixel number.

query_disc, query_polygon,

query_strip, query_triangle render the list of pixels enclosed respectively in a given disc, polygon, latitude strip and triangle

surface_triangle computes the surface of a spherical triangle de-

fined by 3 vertices

rand_gauss 155

rand_gauss

Location in HEALPix directory tree: src/f90/mod/rngmod.f90

This routine returns a number out of a pseudo-random normal deviate (ie, its probability distribution is a Gaussian of mean 0 and variance 1).

FORMAT

var=rand_gauss(rng_handle)

ARGUMENTS

name & dimensionality	kind	in/out	description
rng_handle	planck_rng	INOUT	structure of type planck_rng containing on all information
var	DP	OUT	necessary to continue same ran- dom sequence. number belonging to a pseudo- random normal deviate.

EXAMPLE:

use healpix_types

use rngmod

type(planck_rng) :: rng_handle

real(dp) :: gauss

call rand_init(rng_handle, 12345, 6789012)

gauss = rand_gauss(rng_handle)

initiates a random sequence with the pair of seeds (12345, 6789012), and generates one number out of the normal deviate.

RELATED ROUTINES

This section lists the routines related to ${\bf rand_gauss}$.

planck_rng	derived type describing RNG state
${ m rand}_{ m uni}$	function which returns a random uniform deviate.
$\operatorname{rand_init}$	subroutine to initiate a random number sequence.

rand_init 157

rand_init

Location in HEALPix directory tree: src/f90/mod/rngmod.f90

This routine initializes, with up to 4 seeds, a randomn number sequence. The generator being primed is an F90 port of an xorshift generator described in Marsaglia, Journal of Statistical Software 2003, vol 8. It has a theoretical period of $2^{128} - 1 \approx 3.410^{38}$. Please refer to the "Comment on Random Number Generator" in the Fortran90 facilities guidelines.

\mathbf{FORMAT}

call rand_init(rng_handle, [seed1, seed2, seed3, seed4])

ARGUMENTS

name & dimensionality	kind	in/out	description
rng_handle	planck_rng	OUT	structure of type planck_rng containing on output all infor- mation necessary to continue
seed1 (OPTIONAL)	I4B	IN	same random sequence. first seed of the random sequence. Can be of arbitray sign. If set to zero or not provided will be replaced internally by a
seed2 (OPTIONAL)	I4B	IN	non-zero hard coded value. second seed. Same properties as above
seed3 (OPTIONAL) seed4 (OPTIONAL)	I4B I4B	IN IN	third seed. Same as above. fourth seed. Same as above.

EXAMPLE:

use rngmod

type(planck_rng) :: rng_handle

call rand_init(rng_handle, 12345, 6789012)

initiates a random sequence with the pair of seeds (12345, 6789012).

RELATED ROUTINES

This section lists the routines related to rand_init.

planck_rng derived type describing RNG state

rand_gauss function which returns a random normal deviate.
rand_uni function which returns a random uniform deviate.

rand_uni 159

rand_uni

Location in HEALPix directory tree: src/f90/mod/rngmod.f90

This routine returns a number out of a pseudo-random uniform deviate (ie, its probability distribution is uniform in the range]0,1[).

FORMAT

var=rand_uni(rng_handle)

ARGUMENTS

name & dimensionality	kind	in/out	description
rng_handle	planck_rng	INOUT	structure of type planck_rng containing on all information
var	DP	OUT	necessary to continue same ran- dom sequence. number belonging to a pseudo- random uniform deviate.

EXAMPLE:

use healpix_types

use rngmod

type(planck_rng) :: rng_handle

real(dp) :: uni

call rand_init(rng_handle, 12345, 6789012)
uni = rand_uni(rng_handle)

initiates a random sequence with the pair of seeds (12345, 6789012), and generates one number out of the uniform deviate.

RELATED ROUTINES

This section lists the routines related to rand_uni.

planck_rng rand_gauss rand_init derived type describing RNG state function which returns a random normal deviate. subroutine to initiate a random number sequence. read_asctab*

$\underline{read}_asctab^*$

read_bintab*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a **HEALPix** map from a binary FITS-file. The routine can read a temperature map or both temperature and polarisation maps (T,Q,U)

FORMAT

call read_bintab*(filename, map, npixtot, nmaps, nullval, anynull [,header, units, extno])

ARGUMENTS

name &d imensionality	kind	in/oı	utdescription
filename(LEN=filenamelen)	CHR	IN	filename of FITS-file containing the map(s).
npixtot	I4B	IN	Number of pixels to be read from map.
nmap	I4B	IN	number of maps to be read, 1 for temperature only, and 3 for (T,Q,U) .
map(0:npixtot-1,1:nmap)	SP/ DP	OUT	the map read from the FITS-file.
nullval	SP/ DP	OUT	value of missing pixels in the map.
anynull	LGT	OUT	TRUE, if there are missing pixels, and FALSE otherwise.
header(LEN=80)(1:) (OPTIONAL)	CHR	OUT	character string array containing the FITS header read from the file. Its dimension has to be defined prior to calling the routine
units(LEN=*)(1:nmaps) (OPTIONAL)	CHR	OUT	
extno (OPTIONAL)	I4B	IN	extension number to read the data from (0 based).(default: 0) (the first extension is read)

read_bintab*

EXAMPLE:

call read_bintab ('map.fits',map,12*32**2,1,nullval,anynull)

Reads a **HEALPix** temperature map from the file 'map.fits' to the array map(0:12*32**2-1,1:1). The pixel number 12*32**2 is the number of pixels in a $N_{\rm side}=32$ **HEALPix** map. If there are missing pixels in the file, anynull is TRUE and these pixels get the value returned in nullval.

MODULES & ROUTINES

This section lists the modules and routines used by read_bintab*.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to read_bintab*.

input_map Routine which reads a map using read_bintab*and

fills missing pixels with a given value.

map2alm Routine which analyse a map and returns the a_{lm}

coefficients.

read_fits_cut4 Routine to read cut sky **HEALPix** FITS maps

write_plm, write_bintab Routines to write **HEALPix** FITS maps

read_conbintab*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a FITS file containing a_{lm} values for constained realisation. The FITS file is supposed to contain one integer column with $index = \ell^2 + \ell + m + 1$ and 2 or 4 single (or double) precision columns with real/imaginary a_{lm} values and real/imaginary standard deviation on these a_{lm} . It is supposed to contain either 1 or 3 extension(s) containing respectively the a_{lm} for T and if applicable E and B.

\mathbf{FORMAT}

call read_conbintab*(filename, alms, nalms [, units, extno])

ARGUMENTS

name&dimensionality	kind	in/oı	utdescription
filename(LEN=filenamelen) nalms	CHR I4B	IN IN	filename of FITS file containing a_{lm} . Number of a_{lm} values to read from the file.
alms(0:nalms-1,1:6)	SP/ DP	OUT	the a_{lm} read from the file. alms(i,1) and alms(i,2) contain the ℓ and m values for the ith a_{lm} . alms(i,3) and alms(i,4) contain the real and imaginary value of the ith a_{lm} . Finally, the standard deviation for the ith a_{lm} is contained in alms(i,5) (real) and alms(i,6) (imaginary).
units(len=20)(1:) (OPTIONAL)	CHR	OUT	character string containing the units of the $a_{\ell m}$
extno TIONAL)	I 4⊕ P-	- IN	extension (0 based) of the FITS file to be read

read_conbintab*

EXAMPLE:

call read_conbintab ('alms.fits',alms,65*66/2)

Read 65*66/2 (the number of a_{lm} needed to fill the whole range from l=0 to l=64) a_{lm} values from the file 'alms.fits' into the array alms(0:65*66/2-1,1:6).

MODULES & ROUTINES

This section lists the modules and routines used by **read_conbintab***.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to **read_conbintab***.

alms2fits, dump_alms routines to write a_{lm} to a FITS-file

fits2alms has the same function as read_conbintab but is

more general.

number_of_alms,[getsize_fits] can be used to find out the number of a_{lm} avail-

able in the file.

$read_dbintab$

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a double precision binary array from a FITS-file. It is used by **HEALPix** to read precomputed $P_{lm}(\theta)$ values and pixel window functions.

FORMAT call read_dbintab(filename, map, npixtot, nmaps, nullval, anynull, units)

ARGUMENTS

name & dimensionality	kind	$_{ m in/out}$	$\operatorname{description}$
filename(LEN=filenamelen)	CHR	IN	filename of FITS-file containing the double precision array.
npixtot	I4B	IN	Number of values to be read from the file.
nmaps	I4B	IN	number of 1-dim. arrays, 1 for scalar P_{lm} s and pixel windows,
map(0:npixtot-1,1:nmaps)	DP	OUT	3 for scalar and tensor P_{lm} s. the array read from the FITS-file.
nullval	DP	OUT	value of missing pixels in the array.
anynull	LGT	OUT	TRUE, if there are missing pixels, and FALSE otherwise.
units(len=20)(1:nmaps)	CHR	OUT	respective physical units of the maps in the FITS file.

read_dbintab

EXAMPLE:

call read_dbintab ('plm_32.fits',plm,65*66*32,1,nullval,anynull)

Reads precomputed scalar $P_{lm}(\theta)$ from the file 'plm_32.fits'. The values are returned in the array plm(0:65*66*32,1:1). The number of values 65*66*32 is the number of precomputed $P_{lm}(\theta)$ for a $N_{side}=32$, lmax=64 map. If there are missing values in the file, anynull is TRUE and nullval contains the values of these pixels.

MODULES & ROUTINES

This section lists the modules and routines used by read_dbintab.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to read_dbintab.

plmgen Executable to create files with precomputed

 $P_{lm}(\theta)$.

write_dbintab Routine to create a file to be read by read_dbintab.

read_fits_cut4

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads a cut sky **HEALPix** map from a FITS file. The format used for the FITS file follows the one used for Boomerang98 and is adapted from COBE/DMR

FORMAT call read_fits_cut4(filename, np, pixel, [signal, n_obs, serror, header, units, extno])

ARGUMENTS

name&dimensionality	kind	in/outdescription		
filename(LEN=filenamelen)CHR		IN	FITS file to be read from, containing a cut sky map	
np	I4B	IN	number of pixels to be read from the file	
pixel(0:np-1)	I4B	OUT	index of observed (or valid) pixels	
signal(0:np-1)	SP	OUT	value of signal in each observed pixel	
(OPTIONAL)				
$n_{-}obs(0:np-1)$	I4B	OUT	number of observation per pixel	
(OPTIONAL)				
serror(0:np-1)	SP	OUT	rms of signal in pixel. (For white noise, this	
(OPTIONAL)			would be $\propto 1/\sqrt{\text{n_obs}}$)	
header(LEN=80)(1:)	CHR	OUT	FITS extension header	
(OPTIONAL)				
units(LEN=20)	CHR	R OUT maps units (applies only to Signal and		
(OPTIONAL)		which are assumed to have the same un		
extno (OPTIONAL)	I4B	IN	extension number (0 based) for which map is	
			read. Default $= 0$ (first extension).	

read_fits_cut4

MODULES & ROUTINES

This section lists the modules and routines used by read_fits_cut4.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to read_fits_cut4.

anafast executable that reads a **HEALPix** map and anal-

yses it.

synfast executable that generate full sky **HEALPix** maps

getsize_fits routine to know the size of a FITS file and its type

(eg, full sky vs cut sky)

input_map all purpose routine to input a map of any kind

from a FITS file

output_map subroutine to write a FITS file from a **HEALPix**

map

write_fits_cut4 subroutine to write a cut sky map into a FITS file

read_par

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine reads the 'NSIDE', 'TFIELDS', 'MAX-LPOL', and optionally 'MAX-MPOL' keywords from a FITS-file. These keywords desribe N_{side} , number of datasets (maps) and maximum multipole ℓ (order) and m (degree) value for the file. If a keyword is not found in the FITS file, a value of -1 is returned instead. The file could eg. be a **HEALPix** map, or a set of a_{lm} or precomputed $P_{lm}(\theta)$

FORMAT

call read_par(filename, nside, lmax, tfields [, mmax])

ARGUMENTS

name & dimensionality	kind	in/oı	in/outdescription			
filename(LEN=filenamelen)	CHR	IN	filename of the FITS file.			
nside	I4B	OUT	'NSIDE' keyword value from the FITS			
			header.			
lmax	I4B	OUT	'MAX-LPOL' keyword value from the			
			FITS header.			
tfields	I4B	OUT	'TFIELDS' keyword value from the FITS			
			header.			
mmax (OPTIONAL)	I4B	OUT	'MAX-MPOL' keyword value from the			
,			FITS header.			

EXAMPLE:

call read_par('plm_128p.fits', nside, lmax, nhar)

Checks the N_{side} and maximum ℓ value used for the precomputed $P_{\ell m}(\theta)$ that are stored in the file 'plm_128p.fits'. If the file also contains tensor harmonics, nhar is returned with the value 3, otherwise it is 1.

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MODULES & ROUTINES

This section lists the modules and routines used by **read_par**.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to ${\bf read_par}.$

synfast, plmgen executables that produce FITS-files with key-

words relevant to this routine.

$real_{fft}$

Location in HEALPix directory tree: src/f90/mod/healpix_fft.F90

This routine performs a forward or backward Fast Fourier Transformation on its argument data.

FORMAT call real_fft(data, backward)

ARGUMENTS

name & dimensionality	kind	in/out	description
data(:)	XXX	INOUT	array containing the input and output data. Can be of type real(sp) or real(dp)
backward	LGT	IN	Optional argument. If present and true, perform backward transformation, else forward

EXAMPLE:

use healpix_fft
call real_fft (data, backward=.true.)

Performs a backward FFT on data.

RELATED ROUTINES

This section lists the routines related to real_fft.

complex_fft routine for FFT of complex data

remove_dipole*

remove_dipole*

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

This routine provides a means to fit and remove the dipole and monopole from a **HEALPix** map. The fit is obtained by solving the linear system

$$\sum_{j=0}^{d^2-1} A_{ij} f_j = b_i \tag{15}$$

with, d = 1 or 2, and

$$b_i = \sum_{p \in \mathcal{P}} s_i(p)w(p)m(p), \tag{16}$$

$$A_{ij} = \sum_{p \in \mathcal{P}} s_i(p)w(p)s_j(p), \tag{17}$$

where \mathcal{P} is the set of valid, unmasked pixels, m is the input map, w is pixel weighting, while $s_0(p) = 1$ and $s_1(p) = x$, $s_2(p) = y$, $s_3(p) = z$ are respectively the monopole and dipole templates. The output map is then

$$m'(p) = m(p) - \sum_{i=0}^{d^2 - 1} f_i s_i(p).$$
 (18)

FORMAT

call remove_dipole*(nside, map, ordering, degree, multipoles, zbounds [, fmissval, mask, weights])

ARGUMENTS

name & dimensionality	kind	in/out	description
nside	I4B	IN	value of $N_{\rm side}$ resolution parameter for in-
map(0:12*nside*nside-1)	SP/ DP	INOUT	put map HEALPix map from which the monopole and dipole will be removed. Those are removed from all unflagged pixels, even those excluded by the cut
ordering	I4B	IN	zounds or the mask. HEALPix scheme 1:RING, 2: NESTED
degree	I4B	IN	multipoles to fit and remove. It is either 0 (nothing done), 1 (monopole only) or 2 (monopole and dipole).
multipoles(0:degree*degree-1)	DP	OUT	values of best fit monopole and dipole. The monopole is described as a scalar in the same units as the input map, the dipole as a 3D cartesian vector, in the same units.
zbounds(1:2)	DP	IN	section of the map on which to perform the fit, expressed in terms of $z = \sin(\operatorname{latitude}) = \cos(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ in $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$. If $z = \sin(\theta)$ is $z = \sin(\theta)$.
fmissval (OPTIONAL)	SP/ DP	IN	value used to flag bad pixel on input (default: -1.6375e30). Pixels with that value are ignored during the fit, and left unchanged on output.
mask(0:12*nside*nside-1) (OPTIONAL)	SP/ DP	IN	mask of valid pixels. Pixels with $ \text{mask} < 10^{-10}$ are not used for fit. Note: the map is <i>not</i> multiplied by the mask.
weights(0:12*nside*nside-1) (OPTIONAL)	SP/ DP	IN	weight to be given to each map pixel before doing the fit. By default pixels are given a uniform weight of 1. Note: the output map is <i>not</i> multiplied by the weights.

remove_dipole*

EXAMPLE:

```
s = sin(15.0_dp * PI / 180.0_dp) call remove_dipole*(128, map, 1, 2, multipoles, (\ s, -s \) )
```

Will compute and remove the best fit monopole and dipole from a map with $N_{\rm side} = 128$ in RING ordering scheme. The fit is performed on pixels with $|b| > 15^{\circ}$.

MODULES & ROUTINES

This section lists the modules and routines used by **remove_dipole***.

pix_tools module, containing:

RELATED ROUTINES

This section lists the routines related to **remove_dipole***.

 add_dipole

routine to add a dipole and monopole to a map.

ring_analysis

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

This subroutine computes the Fast Fourier Transform of a single ring of pixels and extends the computed coefficients up to the maximum m of the transform.

FORMAT	call	ring_analysis(nsmax,	nlmax,	nmmax,
	data	in, nph, dataout, kphi0)		

ARGUMENTS

name & dimensionality	kind	in/out	description
nsmax	I4B	IN	$N_{\rm side}$ of the map.
nlmax	I4B	IN	Maximum ℓ of the analysis.
nmmax	I4B	IN	Maximum m of the analysis.
nph	I4B	IN	The number of points on the
			ring.
datain(0:nph-1)	DP	IN	Function values on the ring.
dataout(0:nmmax)	DPC	OUT	Fourier components, replicated
,			to Nmmax.
kphi0	I4B	IN	0 if the first pixel on the ring is
			at $\phi = 0$; 1 otherwise.

EXAMPLE:

call ring_analysis(64,128,128,datain,8,dataout,0)

Returns the periodically extended complex Fourier Transform of data in in dataout. They are returned in the following order: 0 1 2 3 4 5 6 7 6 5 4 3 2 1 0.... The value kphi0=0 specifies that no phase factor needed to be applied, because the ring starts at $\phi=0$. ring_analysis 177

MODULES & ROUTINES

This section lists the modules and routines used by **ring_analysis**.

healpix_fft module.

RELATED ROUTINES

This section lists the routines related to ring_analysis.

ring_synthesis Inverse transform (complex-to-real), used in

alm2map, alm2map_der and synfast

ring_num

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

This function returns the ring number for a z coordinate.

FORMAT var=ring_num(nside, z)

ARGUMENTS

name&dimensionality	kind	in/out	description
nside z	I4B DP	IN IN	the N_{side} parameter of the map. the z coordinate to find the ring number for.

EXAMPLE:

print*,ring_num(256, 0.5)

Prints the ring number of the ring at position z = 0.5.

MODULES & ROUTINES

This section lists the modules and routines used by ring_num.

None

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RELATED ROUTINES

This section lists the routines related to ${\bf ring_num}.$

in_ring Returns the pixels in a slice on a given ring.

$\underline{\mathbf{ring_syn}}$ thesis

Location in HEALPix directory tree: src/f90/mod/alm_tools.f90

FORMAT	call ring_synthesis(nsmax,	nlmax,	nmmax,
	datain, nph, dataout, kphi0)		

ARGUMENTS

name & dimensionality	kind	in/out	description
nsmax	I4B	IN	$N_{\rm side}$ of the map.
nlmax	I4B	IN	Maximum ℓ of the analysis.
nmmax	I4B	IN	Maximum m of the analysis.
nph	I4B	IN	The number of points on the
			ring.
datain(0:nmmax)	DPC	IN	Fourier components as com-
			puted from the a_{lm} .
dataout(0:nph-1)	DP	OUT	Synthesized function values on
			the ring.
kphi0	I4B	IN	0 if the first pixel on the ring is
			at $\phi = 0$; 1 otherwise.

EXAMPLE:

call ring_synthesis(64,128,128,datain,8,dataout,1)

This computes the inverse (complex-to-real) Fast Fourier Transform for the second ring from the pole, containing 8 pixels, for a map resolution of $N_{\rm side}=64$. 128 complex Fourier components contribute to these 8 pixels. The value kphi0=1 specifies that a phase factor needed to be applied to correctly rotate the ring into position on the **HEALPix** grid.

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MODULES & ROUTINES

This section lists the modules and routines used by **ring_synthesis**.

healpix_fft module.

RELATED ROUTINES

This section lists the routines related to ring_synthesis.

ring_analysis Forward transform, used in map2alm and anafast

$rotate_alm*$

$Location \ in \ HEALPix \ directory \ tree: \ src/f90/mod/alm_tools.f90$

This routine transform the scalar (and tensor) $a_{\ell m}$ coefficients to emulate the effect of an arbitrary rotation of the underlying map. The rotation is done directly on the $a_{\ell m}$ using the Wigner rotation matrices, computed by recursion. To rotate the $a_{\ell m}$ for $\ell \leq \ell_{\rm max}$ the number of operations scales like $\ell_{\rm max}^3$.

FORMAT call rotate_alm*(lmax, alm_TGC, psi, theta, phi)

name & dimensionality	kind	in/out	description
nlmax alm_TGC(1:p,0:nlmax,0:nlmax)	I4B SPC/ DPC	IN INOUT	maximum ℓ value for the $a_{\ell m}$. complex $a_{\ell m}$ values before and after rotation of the coordinate system. The first index here runs from 1:1 for temperature only, and 1:3 for po- larisation. In the latter case, 1=T, 2=E, 3=B.
psi	DP	IN	first rotation: angle ψ about the z-axis. All angles are in radians and should lie in $[-2\pi,2\pi]$, the rotations are active and the referential system is assumed to be right handed, the routine coordsys2euler_zyz can be used to generate the Euler angles ψ, θ, φ for rotation between standard astronomical coordinate systems;
theta	DP	IN	second rotation: angle θ about the original (unrotated) y-axis;
phi	DP	IN	third rotation: angle φ about the original (unrotated) z-axis;

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EXAMPLE:

```
use alm_tools, only: rotate_alm
...
call rotate_alm(64, alm_TGC, PI/3., 0.5_dp, 0.0_dp)
```

Transforms scalar and tensor a_{lm} for $\ell_{max} = m_{max} = 64$ to emulate a rotation of the underlying map by $(\psi = \pi/3, \theta = 0.5, \varphi = 0)$.

EXAMPLE:

```
use coord_v_convert, only: coordsys2euler_zyz
use alm_tools, only: rotate_alm
...
call coordsys2euler_zyz(2000.0_dp, 2000.0_dp, 'E', 'G', psi, theta, phi)
call rotate_alm(64, alm_TGC, psi, theta, phi)
```

Rotate the a_{lm} from Ecliptic to Galactic coordinates.

RELATED ROUTINES

This section lists the routines related to **rotate_alm***.

coordsys2euler_zyz	can be used to generate the Euler angles ψ, θ, φ for rotation between standard astronomical coordinate systems
$create_alm$	Routine to create $a_{\ell m}$ coefficients.
$alter_alm$	Routine to modify $a_{\ell m}$ coefficients to apply or remove the effect of an instrumental beam.
map2alm	Routines to analyze a HEALPix sky map into its $a_{\ell m}$ coefficients.
alm2map	Routines to synthetize a HEALPix sky map from its $a_{\ell m}$ coefficients.
$alms2 fits, \ dump_alms$	Routines to save a set of a_{lm} in a FITS file.
xcc_v_convert	rotates a 3D coordinate vector from one astronom-

ical coordinate system to another.

same_shape_pixels_nest, same_shape_pixels_ring

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

These routines provide the ordered list of all **HEALPix** pixels having the same shape as a given template, for a resolution parameter N_{side} . Depending on the template considered the number of such pixels is either 8, 16, $4N_{\text{side}}$ or $8N_{\text{side}}$.

The template pixels are all located in the Northern Hemisphere, or on the Equator. They are chosen to have their center located at

$$\begin{split} z &= \cos(\theta) \geq 2/3, \qquad 0 < \phi \leq \pi/2, \\ 2/3 > z \geq 0, \qquad \phi = 0, \quad \text{or} \quad \phi = \frac{\pi}{4N_{\text{side}}}. \end{split}$$

They are numbered continuously from 0, starting at the North Pole, with the index increasing in ϕ , and then increasing for decreasing z.

FORMAT	call same_shape_pixels_nest(nside,	template
	[, list, reflexion, nrep])		

FORMAT call same_shape_pixels_ring(nside, template [, list, reflexion, nrep])

name & dimensionality	kind	in/out	description
	LID	T. 7	- I TIPAT DI W
nside template	I4B I4B	IN OUT	the HEALPix N_{side} parameter. identification number(s) of the template matching in shape the pixel(s) provided (the numbering scheme of the pixel templates is the same for both routines).
list(0:nrep-1) OPTIONAL	I4B	OUT	pointer containing the ordered list of NESTED/RING scheme identification numbers (in $\{0.12N_{\rm side}^2 - 1\}$) of all pixels having the same shape as the template provided. The routines will allocate the list array if it is not allocated upon calling.
reflexion(0:nrep-1) OPTIONAL	I4B	OUT	pointer containing the transformation(s) (in {0, 3}) to apply to each of the returned pixels to match exactly in shape and position its respective template. 0: rotation around the polar axis only, 1: rotation + East-West swap (ie, reflexion around meridian), 2: rotation + North-South swap (ie, reflexion around Equator), 3: rotation + East-West and North-South swaps. The routines will allocate the list array if it is not allocated upon calling.
nrep OPTIONAL	I4B	OUT	number of pixels having the same template (either 8, 16, $4N_{\text{side}}$ or $8N_{\text{side}}$).

EXAMPLE:

call same_shape_pixels_ring(256, 1234, list, reflexion, np)

Returns in list the RING-scheme index of the all the pixels having the same shape as the template #1234 for $N_{\rm side}=256$. Upon return reflexion will contain the rotation/reflexions to apply to each pixel returned to match the template, and np will contain the number of pixels having that same shape (16 in that case).

RELATED ROUTINES

This section lists the routines related to **same_shape_pixels_ring**.

nside2templates returns the number of template pixel shapes avail-

able for a given N_{side} .

template_pixel_ring

template_pixel_nest return the template shape matching the pixel pro-

vided

scan_directories

Location in HEALPix directory tree: src/f90/mod/paramfile_io.F90 Function to scan a set of directories for a given file

FORMAT var=scan_directories(directories, filename, full-path)

ARGUMENTS

name&dimensionality	kind in/ou	t description
directories	CHR IN	contains the set of directories (up to 20), separated by an ASCII character of value < 32 (see concatn1). During the search, it is assumed that the given directories and filename can be separated by nothing as a file of the character of the cha
filename fullpath	CHR IN CHR OUT	ing, a / (slash) or a \ (backslash) the file to be found. returns the full path to the first occur- rence of the file among the directories
var	LGT OUT	provided. Empty if the file is not found. The search is not recursive. set to true if the file is found, to false otherwise.

EXAMPLE:

```
use paramfile_io
character(len=filenamelen) :: dirs, full
logical(lgt) :: found
dirs = concatnl('dir1','/dir2','/dir2/subdir1/') ! build directories
list.
found = scan_directories(dirs, 'myfile', full) ! do the search
if (found) print*,trim(full)
```

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Search for 'myfile' in the directories 'dir1', '/dir2', '/dir2/subdir1/'

RELATED ROUTINES

This section lists the routines related to scan_directories.

parse_xxx parse an ASCII file for parameters definition concatnl concatenates a set of substrings into one string,

interspaced with LineFeed character

string, strlowcase, strupcase

Location in HEALPix directory tree: src/f90/mod/misc_utils.F90

The Fortran90 module misc_utils contains three functions to create or manipulate character strings.

ARGUMENTS

name & dimensionality	kind in/out	description
Tty		
number	LGT/ IN	number or boolean flag to be turned into
	I4B/	a character string.
	SP/	
	DP	
instring	CHR IN	arbitrary character string.
outstring	CHR —	output character string.
format	CHR IN	character string describing Fortran format
OPTIONAL		of output.
		-

FUNCTIONS:

outstring = string(number [,format])

returns in outstring its argument number converted to a character string. If format is provided it is used to format the output, if not, the fortran default format matching number's type is used.

outstring = strlowcase(instring)

returns in outstring its argument instring converted to lowercase. ASCII characters in the [A-Z] range are mapped to [a-z], while all others remain unchanged.

outstring = strupcase(instring)

returns in outstring its argument instring converted to uppercase. ASCII characters in the [a-z] range are mapped to [A-Z], while all others remain unchanged.

EXAMPLE:

```
use misc_utils
character(len=24) :: s1
s1 = string(123,'(i5.5)')
print*, trim(s1)
print*,trim(strupcase('*aBcD-123'))
print*,trim(strlowcase('*aBcD-123'))
```

Will printout 00123, *ABCD-123 and *abcd-123.

surface_triangle

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Returns the surface in steradians of the spherical triangle described by its three vertices

FORMAT

call surface_triangle(v1, v2, v3, surface)

ARGUMENTS

name&dimensionalitykin	d in/out	description
v1(3) DP	IN	cartesian vector pointing at the triangle first vertex.
v2(3) DP	IN	cartesian vector pointing at the triangle second vertex.
v3(3) DP	IN	cartesian vector pointing at the triangle third vertex.
surface DP	OUT	surface of the triangle in steradians.

EXAMPLE:

use healpix_types

use pix_tools, only : surface_triangle

real(DP) :: surface, one = 1.0_dp

call $surface_triangle((/1,0,0/)*one, (/0,1,0/)*one, (/0,0,1/)*one,$

surface)

print*, surface

Returns the surface in steradians of the triangle defined by the octant (x, y, z > 0): 1.5707963267948966

RELATED ROUTINES

This section lists the routines related to **surface_triangle**.

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pix2ang, ang2pix pix2vec, vec2pix convert between angle and pixel number. convert between a cartesian vector and pixel number.

query_disc, query_polygon, query_strip, query_triangle

render the list of pixels enclosed respectively in a given disc, polygon, latitude strip and triangle

template_pixel_nest, template_pixel_ring

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routines to provide the index of the template pixel associated with a given **HEALPix** pixel, for a resolution parameter N_{side} .

Any pixel can be *matched in shape* to a single of these templates by a combination of a rotation around the polar axis with reflexion(s) around a meridian and/or the equator.

The template pixels are all located in the Northern Hemisphere, or on the Equator. They are chosen to have their center located at

$$\begin{split} z &= \cos(\theta) \geq 2/3, \qquad 0 < \phi \leq \pi/2, \\ 2/3 > z \geq 0, \qquad \phi = 0, \quad \text{or} \quad \phi = \frac{\pi}{4N_{\text{side}}}. \end{split}$$

They are numbered continuously from 0, starting at the North Pole, with the index increasing in ϕ , and then increasing for decreasing z.

FORMAT	call template_pixel_nest(nside, pixel_nest, template, reflexion)
FORMAT	call template_pixel_ring(nside, pixel_ring, template, reflexion)

0 1' ' 1'	1 • 1	• / 1	1 * /*
name & dimensionality	kına	in/out	$\operatorname{description}$
nside	I4B	IN	the HEALPix $N_{\rm side}$ parameter.
$pixel_nest$	I4B	IN	NESTED scheme pixel identification
			number over the range $\{0.12N_{\text{side}}^2 - 1\}$.
pixel_ring	I4B	IN	RING scheme pixel identification num-
			ber over the range $\{0.12N_{\text{side}}^2 - 1\}$.
template	I4B	OUT	identification number(s) of the template
			matching in shape the pixel(s) provided
			(the numbering scheme of the pixel tem-
			plates is the same for both routines).
reflexion	I4B	OUT	in {0, 3} encodes the transformation(s)
			to apply to each pixel provided to match
			exactly in shape and position its respec-
			tive template. 0: rotation around the
			polar axis only, 1: rotation $+$ East-West
			swap (ie, reflexion around meridian), 2:
			rotation + North-South swap (ie, reflex-
			ion around Equator), 3: rotation $+$ East-
			West and North-South swaps

EXAMPLE:

call template_pixel_ring(256, 500000, template, reflexion)

Returns in template the index of the template pixel (16663) whose shape matches that of the pixel #500000 for $N_{\rm side}=256$. Upon return reflexion will contain 2, meaning that the template must be reflected around a meridian and around the equator (and then rotated around the polar axis) in order to match the pixel.

RELATED ROUTINES

This section lists the routines related to template_pixel_ring.

${\it nside 2 temp lates}$	returns the number of template pixel shapes available for a given N_{side} .
$same_shape_pixels_ring$	
$same_shape_pixels_nest$	return the ordered list of pixels having the same shape as a given pixel template

udgrade_nest*

$udgrade_nest*$

Location in HEALPix directory tree: src/f90/mod/udgrade_nr.f90

Routine to degrade or prograde the pixel size of a **HEALPix** map indexed with the NESTED scheme. The degradation/progradation is done assuming an intensive quantity (like temperature) that does NOT scale with surface area.

In case of degradation, a big pixel that contains one or several bad pixels will take the average of the valid small pixels, unless a 'pessimistic' behavior is assumed in which case the big pixel will take the bad pixel sentinel value. In case of progradation, a bad pixel only spawns bad pixels.

The routine accepts both mono and bi-dimensional maps.

FORMAT

call udgrade_nest*(map_in, nside_in, map_out, nside_out [, fmissval, pessimistic])

name & dimensionality	kind	in/out	description
		/	1
map_in(0:12*nside_in**2- 1)	SP/ DP	IN	mono-dimensional full sky map to be prograded or degraded.
map_in(0:12*nside_in**2- 1,1:nd)	SP/ DP	IN	bi-dimensional full sky map to be prograded or degraded. The routine finds the second dimension (nd) by itself.
nside_in	I4B	IN	the N_{side} resolution parameter of the input map. Must be a power of 2.
map_out(0:12*nside_out**2- 1)	SP/ DP	OUT	mono-dimensional full sky map after degradation or progradation.
map_out(0:12*nside_out**2-1,1:nd)	SP/ DP	OUT	bi-dimensional full sky map after degradation or progradation. The second dimension (nd) should match that of the input map.
nside_out	I4B	IN	the N_{side} resolution parameter of the output map. Must be a power of 2. If nside_out > nside_in, the map is prograded (ie, more and smaller pixels) with each pixel having the same value as its parent; otherwise, the map in degraded (ie, fewer larger pixels), with each pixel being the average of its (nside_in/nside_out) ² components.
fmissval (OPTIONAL)	SP/ DP	IN	sentinel value given to bad pixels in input and output maps. (default: $-1.6375 \ 10^{30}$)
pessimistic (OPTIONAL)	LGT	IN	if set to .true., during a degradation, a big pixel containing at least a small bad pixel will be returned as bad as well, instead of taking the average of the remaing valid pixels. (default: .false.)

EXAMPLE:

call udgrade_nest(map_hi, 256, map_low, 64)

Degrades a NESTED ordered map with $N_{side}=256$ into a NESTED map with $N_{side}=64$

RELATED ROUTINES

udgrade_nest*

This section lists the routines related to $\mathbf{udgrade_nest*}$.

udgrade_ring

prograde or degrade a RING ordered map.

udgrade_ring*

Location in HEALPix directory tree: src/f90/mod/udgrade_nr.f90

Routine to degrade or prograde the pixel size of a **HEALPix** map indexed with the RING scheme. The degradation/progradation is done assuming an intensive quantity (like temperature) that does NOT scale with surface area.

In case of degradation, a big pixel that contains one or several bad pixels will take the average of the valid small pixels, unless a 'pessimistic' behavior is assumed in which case the big pixel will take the bad pixel sentinel value. In case of progradation, a bad pixel only spawns bad pixels.

The routine accepts both mono and bi-dimensional maps.

FORMAT

call udgrade_ring*(map_in, nside_in, map_out, nside_out [, fmissval, pessimistic])

udgrade_ring* 201

name & dimensionality	kind	in/out	description
map_in(0:12*nside_in**2-1)	SP/ DP	INOUT	mono-dimensional full sky map to be prograded or degraded. The routine finds the second dimension (nd) by itself. Note that the map is modified on output
map_in(0:12*nside_in**2-1,1:nd)	SP/ DP	INOUT	(reordered into NESTED scheme). bi-dimensional full sky map to be prograded or degraded. Note that the map is modified on output (reordered into NESTED scheme).
$nside_in$	I4B	IN	the N_{side} resolution parameter of the input map. Must be a power of 2.
map_out(0:12*nside_out**2-1)	SP/ DP	OUT	mono-dimensional full sky map after degradation or progradation.
map_out(0:12*nside_out**2-1,1:nd)		OUT	bi-dimensional full sky map after degradation or progradation. The second dimension (nd) should match that of the input map.
$nside_out$	I4B	IN	the N_{side} resolution parameter of the output map. Must be a power of 2. If nside_out > nside_in, the map is prograded (ie, more and smaller pixels) with each pixel having the same value as its parent; otherwise, the map in degraded (ie, fewer larger pixels), with each pixel being the average of its (nside_in/nside_out) ² components.
fmissval (OPTIONAL)	SP/ DP	IN	sentinel value given to bad pixels in input and output maps. (default: $-1.6375 \ 10^{30}$)
pessimistic (OPTIONAL)	LGT	IN	if set to .true., during a degradation, a big pixel containing at least a small bad pixel will be returned as bad as well, instead of taking the average of the remaing valid pixels. (default: .false.)

EXAMPLE:

call udgrade_ring(map_hi, 256, map_low, 64)

Degrades a RING ordered map with $N_{side}=256$ into a RING map with $N_{side}=64$

RELATED ROUTINES

This section lists the routines related to $\mathbf{udgrade_ring*}$.

 $udgrade_nest$

prograde or degrade a NESTED ordered map.

vec2ang 203

vec2ang

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90

Routine to convert the 3D position vector (x, y, z) of point into its position angles (θ, ϕ) on the sphere with $x = \sin \theta \cos \phi$, $y = \sin \theta \sin \phi$, $z = \cos \theta$.

FORMAT

call vec2ang(vector, theta, phi)

ARGUMENTS

name&dimensionality	kind	in/out	description
vector(3)	DP	IN	three dimensional cartesian position vector (x, y, z) . The
theta	DP	OUT	north pole is $(0,0,1)$ colatitude in radians measured southward from north pole (in
phi	DP	OUT	$[0,\pi]$). longitude in radians measured eastward (in $[0, 2\pi]$).

RELATED ROUTINES

This section lists the routines related to **vec2ang**.

ang2vec

converts the position angles of a point on the sphere into its 3D position vector.

$\mathbf{vect}_{-}\mathbf{prod}$

Location in HEALPix directory tree: src/f90/mod/pix_tools.f90 Returns the vectorial product of two vectors.

FORMAT

call vect_prod(v1, v2, v3)

ARGUMENTS

name&dimensionalitykind	in/out	description
v1(3) DP v2(3) DP v3(3) DP	IN IN OUT	cartesian vector \mathbf{v}_1 . cartesian vector \mathbf{v}_2 . cartesian vector $\mathbf{v}_3 = \mathbf{v}_1 \times \mathbf{v}_2$

EXAMPLE:

use healpix_types

use pix_tools, only : vect_prod
real(DP), dimension(3) :: vec

 $real(DP) :: one = 1.0_dp$

call $vect_prod((/2,0,0/)*one, (/0,1,0/)*one, vec)$

print*, vec

will return: 0.00E+000 0.00E+000 2.00

RELATED ROUTINES

This section lists the routines related to **vect_prod**.

angdist

computes the angular distance between 2 vectors

write_asctab*

write_asctab*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine stores a power spectrum in an ascii FITS-file. The routine can store temperature coeffecients C_l^T or both temperature and polarisation coeffecients C_l^T , C_l^B , C_l^B , $C_l^{T \times E}$.

FORMAT

call write_asctab*(clout, lmax, ncl, header, nl-header, filename)

ARGUMENTS

name & dimensionality	kind in/out	description
filename(LEN=filenamelen)	CHR IN	the FITS file to which the power spectrum is written.
lmax ncl	I4B IN I4B IN	Maximum ℓ value to be written. 1 for temperature coeffecients only, 4 for polarisation.
clout(0:lmax,1:ncl)	SP/ IN DP	the powerspectrum to be saved in the file.
nlheader	I4B IN	number of header lines to write to the file.
header(LEN=80) (1:nlheader)	CHR IN	the header to the FITS-file.

EXAMPLE:

call write_asctab (cl,64,1,header,80,'cl.fits')

Writes a powerspectrum in the array cl(0:64,1:1) to a FITS-file called 'cl.fits'. The cl array contains the temperature power-spectrum C_l^T up to an ℓ value of 64. 80 header lines are written to the file from the array header(1:80).

MODULES & ROUTINES

This section lists the modules and routines used by write_asctab*.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to write_asctab*.

alm2cl Routine computing the power spectrum from

spherical harmonics coefficients $a_{\ell m}$

fits2cl Routine to read a FITS file created by

 $write_asctab.$

write_minimal_header routine to write minimal FITS header

write_bintab*

write_bintab*

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine creates a binary FITS-file from a **HEALPix** map. The routine can save a temperature map or both temperature and polarisation maps (T,Q,U) to the file.

FORMAT call write_bintab*(map, npix, nmap, header, nl-header, filename [, extno])

Arguments appearing in *italic* are optional.

ARGUMENTS

name & dimensionality	kind	in/out	description
map(0:npix-1,1:nmap)	SP/ DP	IN	the map to write to the FITS-file.
npix	I4B	IN	Number of pixels in the map.
nmap	I4B	IN	number of maps to be written, 1 for temperature only, and 3 for (T,Q,U) .
header(LEN=80) (1:nl-header)	CHR	IN	The header for the FITS-file.
nlheader	I4B	IN	number of header lines to write to the file.
filename(LEN=8)	CHR	IN	the map(s) is (are) written to a FITS-file with this filename.
extno	I4B	IN	extension number in which to write the data (0 based). (default: 0)

EXAMPLE:

call write_bintab (map,12*32**2,3,header,120,'map.fits')

Makes a binary FITS-file called 'map.fits' from the **HEALPix** maps (T,Q,U) in the array map(0:12*32**2-1,1:3). The number of pixels 12*32**2 corresponds to the number of pixels in a $N_{side} = 32$ **HEALPix** map. The header for the FITS-file is given in the string array header and the number of lines in the header is 120.

MODULES & ROUTINES

This section lists the modules and routines used by write_bintab*.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to write_bintab*.

input_map, read_bintab routines which read a file created by

write_bintab*.

map2alm subroutine which analyse a map and returns the

 a_{lm} coeffecients.

output_map subroutine which calls write_bintab*

write_bintabh subroutine to write a large array into a FITS file

piece by piece

input_tod* subroutine to read an arbitrary subsection of a

large binary table

write_minimal_header routine to write minimal FITS header

write_bintabh 209

write_bintabh

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine is designed to write large (or huge) arrays into a binary table extension of a FITS file. The user can choose to write the array piece by piece. This is designed to deal with Time Ordered Data set (tod).

FORMAT

call write_bintabh(tod, npix, ntod, header, nl-header, filename, [extno, firstpix, repeat])

Arguments appearing in *italic* are optional.

name & dimensionality	kind	in/out	description
tod(0:npix-1,1:ntod)	SP	IN	the map or tod to write to the FITS-
			file. It will be written in the file at the location corresponding to pixels (or time
			samples) firstpix to firtpix + npix -1.
npix	I8B	IN	Number of pixels or time samples in the map or TOD. See Note below.
ntod	I4B	IN	number of maps or tods to be written.
header(LEN=80) (1:nlheader)	CHR	IN	The header for the FITS-file.
nlheader	I4B	IN	number of header lines to write to the file.
filename(LEN = filenamelen)	CHR	IN	the array is written into a FITS-file with this filename.
extno	I4B	IN	extension number in which to write the
			data (0 based). (default: 0)
firstpix	I8B	IN	0 location in the FITS file of the first
			pixel (or time sample) to be written (0
			based). (default: 0). See Note below.
repeat	I4B	IN	length of the element vector used in the
			binary table. (default: 1024) if npix
			$\propto 1024$; 12000 if npix > 12000 and 1
			otherwise. Choosing a large repeat for
			multi-column tables (ntod > 1) generally
			speeds up the I/O. It also helps bringing
			the number of rows of the table under
			2 ³¹ , which is a hard limit of cfitsio.

Note: Indices and number of data elements larger than 2^{31} are only accessible in FITS files on computers with 64 bit enabled compilers and with some specific compilation options of cfitsio (see cfitsio documentation).

EXAMPLE:

```
use healpix_types
use fitstools, only : write_bintabh
character(len=80), dimension(1:128) :: hdr
real(SP), dimension(0:49,1) :: tod
hdr(:) = ' '
tod(:,1) = 1.
call write_bintabh(tod, 50_i8b, 1, header, 128, 'tod.fits', firstpix=0_i8b, repeat=10)
```

write_bintabh 211

```
tod = tod * 3.
call write_bintabh(tod, 20_i8b, 1, header, 128, 'tod.fits',
firstpix=40_i8b)
```

Writes into the FITS file 'tod.fits' a 1 column binary table, where the first 40 data samples have the value 1. and the next 20 have the value 3. (Note that in this example the second call to write_bintabh overwrites some of the pixels written by the first call). The samples will be written in element vectors of length 10. The header for the FITS-file is given in the string array hdr and the number of lines in the header is 128.

MODULES & ROUTINES

This section lists the modules and routines used by write_bintabh.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to write_bintabh.

input_tod* routine that reads a file created by write_bintabh.

input_map, read_bintab routines to read **HEALPix** sky map, write_minimal_header routine to write minimal FITS header

$write_dbintab$

Location in HEALPix directory tree: src/f90/mod/fitstools.f90 This routine is obsolete. Use write_plm instead.

write_fits_cut4 213

write_fits_cut4

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine writes a cut sky **HEALPix** map into a FITS file. The format used for the FITS file follows the one used for Boomerang98 and is adapted from COBE/DMR. This routine can be used to store polarized maps, where the information relative to the Stokes parameters I, Q and U are placed in extension 0, 1 and 2 respectively by successive invocation of the routine.

FORMAT

call write_fits_cut4(filename, np, pixel, signal, n_obs, serror[, header, coord, nside, order, units, extno, polarisation])

name&dimensionality	kind	in/oı	utdescription
filename(LEN=filenamele:	n)CHR	IN	FITS file to be read from, containing a cut sky map
np	I4B	IN	number of pixels to be written in the file
pixel(0:np-1)	I4B	IN	index of observed (or valid) pixels
signal(0:np-1)	SP	IN	value of signal in each observed pixel
$n_{\text{obs}}(0:\text{np-1})$	I4B	IN	number of observation per pixel
serror(0:np-1)	SP	IN	rms of signal in pixel, for white noise, this is $\propto 1/\sqrt{\text{n}_{-}\text{obs}}$.
header(LEN=80)(1:) (OPTIONAL)	CHR	IN	FITS extension header
coord(LEN=1) (OPTIONAL)	CHR	IN	astrophysical coordinates ('C' or 'Q' Celestial/eQuatorial, 'G' for Galactic, 'E' for Ecliptic)
nside (OPTIONAL)	I4B	IN	HEALPix resolution parameter of data set
order (OPTIONAL)	I4B	IN	HEALPix ordering scheme, 1: RING, 2: NESTED
header(LEN=80) (OPTIONAL)	CHR	IN	FITS header to be included in the FITS file
units(LEN=20)	CHR	IN	maps units (applies only to Signal and Serror)
(OPTIONAL)			1 (11))
extension (OPTIONAL)	I4B	IN	(0 based) extension number in which to write data. (default: 0). If set to 0 (or not set) a new file is written from scratch. If set to a value larger than 1, the corresponding extension is added or updated, as long as all previous extensions already exist. All extensions of the same file should use the same Nside, Order and Coord
polarisaton (OPTIONAL)	I4B	IN	if set to a non zero value, specifies that file will contain the I, Q and U polarisation Stokes parameter in extensions 0, 1 and 2 respectively, and sets the FITS header keywords accordingly. If not set, the keywords found in header will prevail. Note: the information relative to Nside, Order and Coord has to be given, either thru these keyword or via the FITS Header.

write_fits_cut4 215

MODULES & ROUTINES

This section lists the modules and routines used by write_fits_cut4.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to write_fits_cut4.

anafast executable that reads a **HEALPix** map and anal-

yses it.

synfast executable that generate full sky **HEALPix** maps

getsize_fits routine to know the size of a FITS file and its type

(eg, full sky vs cut sky)

input_map all purpose routine to input a map of any kind

from a FITS file

output_map subroutine to write a FITS file from a **HEALPix**

map

read_fits_cut4 subroutine to read a **HEALPix** cut sky map from

a FITS file

write_minimal_header routine to write minimal FITS header

write_minimal_header

Location in HEALPix directory tree: src/f90/mod/head_fits.F90

This routine writes the baseline FITS header for the most common **HEALPix** data sets: (cut sky or full sky) map, C(l) power spectra and a_{lm} coefficients.

FORMAT

call write_minimal_header(header, dtype, [append, nside, order, ordering, coordsys, creator, version, randseed, beam_leg, fwhm_degree, units, nlmax, polar, nmmax, bcross, deriv])

Arguments appearing in *italic* are optional.

write_minimal_header 217

name & dimensionality	kind	in/out	description
header(LEN=80) DIMENSION(:)	CHR	INOUT	The FITS header to fill in.
dtype(LEN=*)	CHR	IN	data to be put in the FITS file, must be one of 'ALM', 'CL', 'MAP', 'CUTMAP' (case un-sensitive).
append	LGT	IN	if set to TRUE, the keywords will be appended to the content of header instead of written from scrath
nside	I4B	IN	map resolution parameter; required for dtype='MAP' and dtype='CUTMAP'
order	I4B	IN	map ordering, either 1 (=ring) or 2 (=nested); see ordering
ordering(LEN=*)	CHR	IN	map ordering, either 'RING' or 'NESTED' (case un-sensitive); either order or ordering is required for dtype='MAP' and dtype='CUTMAP'
coordsys(LEN=*)	CHR	IN	map coordinate system; Valid choices are 'G' = Galactic, 'E' = Ecliptic, 'C'/'Q' = Celestial = eQuatorial
creator(LEN=*)	CHR	IN	name of software generating the data set
version(LEN=*)	CHR	IN	version of creator software
randseed	I4B	IN	random number generator seed used to generate the data
$beam_leg(LEN=*)$	CHR	IN	File containing Legendre transform of symmetric beam
$fwhm_degree$	DP	IN	FWHM in degrees of gaussian symmetric beam (FITS keyword: FWHM)
units(LEN=*)	CHR	IN	physical units of the data set (FITS keyword: TUNIT*)
nlmax	I4B	IN	maximum multipole order l of the data set (FITS keyword: MAX-LPOL)
polar	LGT	IN	if set to .TRUE., the file to be written contains polarized data
nmmax	I4B	IN	maximum degree m of data set (FITS keyword: MAX-MPOL)
bcross	LGT	IN	if set to .TRUE., the magnetic cross terms power spectra (TB and EB) are included; only applies to dtype='CL'
deriv	I4B	IN	order of derivatives to included in FITS file (0, 1 or 2); only applies to dtype='MAP'

EXAMPLE:

use head_fits
character(len=80), dimension(1:60) :: header
call write_minimal_header(header, 'MAP', nside=256, ordering='Nested')
call add_card(header, 'HISTORY', 'Dummy map')

Writes in header a **HEALPix** compliant FITS header for a $N_{\text{side}} = 256$ map with NESTED ordering. Further HISTORY information is added with add_card

MODULES & ROUTINES

This section lists the modules and routines used by write_minimal_header.

write_hl more general routine for adding a keyword to a

header.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to write_minimal_header.

add_card general purpose routine to write/edit an arbitrary

keyword into a FITS file header.

get_card general purpose routine to read any keywords

from a header in a FITS file.

del_card routine to discard a keyword from a FITS header

read_par, number_of_alms routines to read specific keywords from a header

in a FITS file.

getsize_fits function returning the size of the data set in a fits

file and reading some other useful FITS keywords

merge_headers routine to merge two FITS headers

write_plm 219

$write_plm$

Location in HEALPix directory tree: src/f90/mod/fitstools.f90

This routine creates a double precision binary FITS-file from a given array. The routine is used by the **HEALPix** facility plungen to store precomputed $P_{lm}(\theta)$.

FORMAT call write_plm(plm, nplm, nhar, header, nl-header, filename, nsmax, nlmax)

name&dimensionality	kind	in/o	utdescription
1 (0 1 1 1 1)	DD	TNI	
plm(0:nplm-1,1:nhar)	DP	IN	the array with the precomputed $P_{lm}(\theta)$ values.
nplm	I4B	IN	Number of P_{lm} values to store.
прии	14D	111	Number of I_{lm} values to store.
nhar	I4B	IN	1 for scalar P_{lm} only and 3 for tensor
			harmonics.
header(LEN=80) (1:nlheader)	CHR	IN	The header for the FITS-file.
nlheader	I4B	IN	number of header lines to write to
			the file.
filename(LEN=filenamelen)	CHR	IN	the precomputed $P_{lm}(\theta)$ values are
,			written to this file.
nsmax	I4B	IN	N_{side} for the precomputed P_{lm} s.
nlmax	I4B	IN	maximum ℓ value for the precom-
			puted P_{lm} s.
			_

EXAMPLE:

call write_plm (plm, 65*66*32, 1, header, 120, 'plm_32.fits', 32, 64)

Makes a double precision binary FITS-file called 'plm_32.fits' from the precomputed $P_{lm}(\theta)$ in the array plm(0:65*66*32-1,1:1). The number 65*66*32 corresponds to the number of precomputed P_{lm} s needed for a $N_{side} = 32$ **HEALPix** map synthesis/analysis. The header for the FITS-file is given in the string array header and the number of lines in the header is 120.

MODULES & ROUTINES

This section lists the modules and routines used by write_plm.

fitstools module, containing:

printerror routine for printing FITS error messages.

cfitsio library for FITS file handling.

RELATED ROUTINES

This section lists the routines related to write_plm.

read_dbintab, read_bintab routines which reads a file created by write_plm.

map2alm, alm2map routines using precomputed $P_{lm}(\theta)$.

xcc_v_convert 221

xcc_v_convert

 $Location \ in \ HEALPix \ directory \ tree: \ src/f90/mod/coord_v_convert.f90$

This routine rotates a 3D coordinate vector from one astronomical coordinate system to another.

FORMAT call xcc_v_convert(ivector, iepoch, oepoch, isys, osys, ovector)

ARGUMENTS

name & dimension-	kind	in/out	$\operatorname{description}$
ality			
ivector(1:3)	DP	IN	3D coordinate vector of one astronomical ob-
			ject, in the input coordinate system.
iepoch	DP	IN	epoch of the input astronomical coordinate
			system.
oepoch	DP	IN	epoch of the output astronomical coordinate
			system.
isys(len=*)	CHR	IN	input coordinate system, should be
			one of 'E'=Ecliptic, 'G'=Galactic,
(5 ds)	0		'C'/'Q'=Celestial/eQuatorial.
osys(len=*)	CHR	IN	output coordinate system, same choice as
(1.0)	D D	***	above.
ovector(1:3)	DP	IN	3D coordinate vector of the same object, in
			the output coordinate system.

EXAMPLE:

```
use healpix_types
use coord_v_convert, only: xcc_v_convert
real(dp) :: vecin(1:3), vecout(1:3)
vecin = (/ 0_dp, 0_dp, 1_dp /)
call xcc_v_convert(vecin, 2000.0_dp, 2000.0_dp, 'g', 'c', vecout)
```

Will produce in vecout the location in Celestial coordinates (2000 epoch) of the North Galactic Pole (defined in vecin)

RELATED ROUTINES

This section lists the routines related to xcc_v_convert.

coordsys2euler_zyz produces the Euler angles ψ, θ, φ in (Z,Y,Z) con-

vention for rotation between standard astronomi-

cal coordinate systems.

ang2vec, vec2ang Routine to convert spherical coordinates (co-

latitude and longitude) into 3D vector coordinates

and vice-versa.