

# Alliance

1.0

Generated by Doxygen 1.9.1



<b>1 File Index</b>	<b>1</b>
1.1 File List	1
<b>2 File Documentation</b>	<b>3</b>
2.1 array.c File Reference	3
2.1.1 Detailed Description	4
2.1.2 Function Documentation	4
2.1.2.1 get_flat_c()	4
2.1.2.2 get_flat_r()	4
2.1.2.3 get_flatIndexComplex3D()	5
2.1.2.4 getIndChi()	5
2.1.2.5 getIndChiBufEL_c()	5
2.1.2.6 getIndChiBufEL_r()	7
2.1.2.7 getIndChiBufEM_c()	7
2.1.2.8 getIndChiBufEM_r()	8
2.1.2.9 multiply_ar_c()	8
2.1.2.10 multiply_ar_r()	8
2.2 diagnostics.c File Reference	8
2.2.1 Detailed Description	9
2.2.2 Function Documentation	9
2.2.2.1 diag_compute()	9
2.2.2.2 diag_computeFreeEnergy()	10
2.2.2.3 diag_computeFreeEnergyFields()	10
2.2.2.4 diag_computeKSpectrum()	10
2.2.2.5 diag_computeMSpectrum()	11
2.2.2.6 diag_computeSpectra()	11
2.2.2.7 diag_getShells()	11
2.2.2.8 diag_initSpec()	12
2.3 distrib.c File Reference	12
2.3.1 Detailed Description	12
2.3.2 Function Documentation	12
2.3.2.1 distrib_enforceReality()	12
2.3.2.2 distrib_getG()	13
2.3.2.3 distrib_getH()	13
2.3.2.4 distrib_getXGrad()	13
2.3.2.5 distrib_getYGrad()	14
2.3.2.6 distrib_getZGrad()	14
2.3.2.7 distrib_setZeroNHalf()	14
2.4 equation.c File Reference	15
2.4.1 Detailed Description	15
2.4.2 Function Documentation	16
2.4.2.1 equation_getLinearTerm()	16

2.4.2.2 equation_getNonlinearElectromagnetic()	16
2.4.2.3 equation_getNonlinearElectrostatic()	16
2.4.2.4 equation_getNonlinearProduct()	17
2.4.2.5 equation_getNonlinearTerm()	17
2.5 fields.c File Reference	18
2.5.1 Detailed Description	19
2.5.2 Function Documentation	19
2.5.2.1 fields_getA()	19
2.5.2.2 fields_getAFromH()	20
2.5.2.3 fields_getB()	20
2.5.2.4 fields_getBFromH()	20
2.5.2.5 fields_getChi()	21
2.5.2.6 fields_getChiA()	21
2.5.2.7 fields_getChiB()	21
2.5.2.8 fields_getChiPhi()	21
2.5.2.9 fields_getFields()	21
2.5.2.10 fields_getFieldsFromH()	22
2.5.2.11 fields_getGradX()	22
2.5.2.12 fields_getGradY()	22
2.5.2.13 fields_getPhi()	23
2.5.2.14 fields_getPhiFromH()	23
2.5.2.15 fields_init()	23
2.5.2.16 fields_sendG()	23
2.6 init.c File Reference	24
2.6.1 Detailed Description	25
2.6.2 Macro Definition Documentation	25
2.6.2.1 RANK_IO	25
2.6.3 Function Documentation	25
2.6.3.1 fill_rand()	25
2.6.3.2 fill_randM0()	25
2.6.3.3 fill_randSingleKM()	26
2.6.3.4 init_conditions()	26
2.6.3.5 init_energySpec()	26
2.6.3.6 init_initEnums()	27
2.6.3.7 init_printParameters()	27
2.6.3.8 init_start()	27
2.7 parameters_io.c File Reference	27
2.7.1 Detailed Description	28
2.7.2 Function Documentation	28
2.7.2.1 init_global_size()	28
2.7.2.2 read_parameters()	28
2.7.2.3 read_parametersFromFile()	28

---

2.8 solver.c File Reference . . . . .	29
2.8.1 Detailed Description . . . . .	29
2.8.2 Function Documentation . . . . .	29
2.8.2.1 solver_init() . . . . .	29
2.8.2.2 solver_makeStep() . . . . .	29
<b>Index</b>	<b>31</b>



# Chapter 1

## File Index

### 1.1 File List

Here is a list of all documented files with brief descriptions:

<a href="#">array.c</a>	Array manipulation module . . . . .	3
<a href="#">diagnostics.c</a>	Diagnostics module . . . . .	8
<a href="#">distrib.c</a>	Gyrokinetic distribution function module . . . . .	12
<a href="#">equation.c</a>	Equation module . . . . .	15
<a href="#">fields.c</a>	Field computation and manipulation module . . . . .	18
<a href="#">init.c</a>	Initialization module for alliance . . . . .	24
<a href="#">parameters_io.c</a>	Reads input parameters from parameter file provided by user . . . . .	27
<a href="#">solver.c</a>	Numerical solver . . . . .	29





## Chapter 2

# File Documentation

## 2.1 array.c File Reference

array manipulation module

```
#include "array.h"
#include "utils_fftw.h"
#include "space_config.h"
```

### Macros

- #define **CHI\_EM** 3
- #define **CHI\_EL** 1
- #define **FFT\_OFFSET** 2

### Functions

- size\_t [get\\_flat\\_c](#) (size\_t is, size\_t il, size\_t im, size\_t ix, size\_t iy, size\_t iz)  
*returns flat index of the element of complex 6D array*
- size\_t [getIndChiBufEM\\_c](#) (size\_t ix, size\_t iy, size\_t iz, size\_t is, size\_t ifield)  
*returns flat index of an element of electromagnetic gyrokinetic potential in FOURIER SPACE*
- size\_t [getIndChiBufEM\\_r](#) (size\_t ix, size\_t iy, size\_t iz, size\_t is, size\_t ifield)  
*returns flat index of an element of electromagnetic gyrokinetic potential in POSITION SPACE*
- size\_t [getIndChiBufEL\\_c](#) (size\_t ix, size\_t iy, size\_t iz, size\_t is)  
*returns returns flat index of an element of electrostatic gyrokinetic potential in FOURIER SPACE*
- size\_t [getIndChiBufEL\\_r](#) (size\_t ix, size\_t iy, size\_t iz, size\_t is)  
*returns returns flat index of an element of electrostatic gyrokinetic potential in REAL SPACE*
- size\_t [get\\_flat\\_r](#) (size\_t is, size\_t il, size\_t im, size\_t ix, size\_t iy, size\_t iz)  
*returns flat index of the element of real 6D array*
- size\_t [get\\_flatIndexComplex3D](#) (size\_t ix, size\_t iy, size\_t iz)  
*returns flat array of complex 3D array*
- size\_t [getIndChi](#) (size\_t ix, size\_t iy, size\_t iz, size\_t is)
- void [multiply\\_ar\\_c](#) (COMPLEX \*ar1, COMPLEX \*ar2, COMPLEX \*ret)
- void [multiply\\_ar\\_r](#) (const double \*ar1, const double \*ar2, double \*ret)

## Variables

- struct array\_size **array\_local\_size**
- struct array\_size **array\_global\_size**
- struct offset\_size **array\_offset**
- struct offset\_size **array\_offset3D**

### 2.1.1 Detailed Description

array manipulation module

contains functions which are supposed to make array manipulation simpler

### 2.1.2 Function Documentation

#### 2.1.2.1 `get_flat_c()`

```
size_t get_flat_c (
    size_t is,
    size_t il,
    size_t im,
    size_t ix,
    size_t iy,
    size_t iz )
```

returns flat index of the element of complex 6D array

#### Parameters

<i>is</i>	species type
<i>il</i>	Laguerre moment
<i>im</i>	Hermite moment
<i>ix</i>	kx index
<i>iy</i>	ky index
<i>iz</i>	kz index

returns flattened index of a complex array from its 6D index. Flattened index then can be passed to distribution function 6D array to get a required element at position (is,il,im,ix,iy,iz).

#### 2.1.2.2 `get_flat_r()`

```
size_t get_flat_r (
    size_t is,
    size_t il,
    size_t im,
    size_t ix,
```

```

size_t iy,
size_t iz )

```

returns flat index of the element of real 6D array

#### Parameters

<i>is</i>	species type
<i>il</i>	Laguerre moment
<i>im</i>	Hermite moment
<i>ix</i>	x index
<i>iy</i>	y index
<i>iz</i>	z index

returns flattened index of a real array from its 6D index. Flattened index then can be passed to distribution function 6D array to get a required element at position (is,il,im,ix,iy,iz).

#### 2.1.2.3 `get_flatIndexComplex3D()`

```

size_t get_flatIndexComplex3D (
    size_t ix,
    size_t iy,
    size_t iz )

```

returns flat array of complex 3D array

#### Parameters

<i>ix</i>	kx index
<i>iy</i>	ky index
<i>iz</i>	kz index

returns flattened index of a complex array from its 3D position index. Flattened index then can be passed to one of the fields (  $\phi(\mathbf{k})$ ,  $A_{||}(\mathbf{k})$ ,  $B_{||}(\mathbf{k})$  ) 6D array to get a required element at position (ix,iy,iz).

#### 2.1.2.4 `getIndChi()`

```

size_t getIndChi (
    size_t ix,
    size_t iy,
    size_t iz,
    size_t is )

```

[getIndChi\(size\\_t ix,size\\_t iy, size\\_t iz, size\\_t is\)](#)

#### 2.1.2.5 `getIndChiBufEL_c()`

```

size_t getIndChiBufEL_c (
    size_t ix,

```

```
size_t iy,  
size_t iz,  
size_t is )
```

returns returns flat index of an element of electrostatic gyrokinetic potential in FOURIER SPACE

## Parameters

<i>ix</i>	kx index
<i>iy</i>	ky index
<i>iz</i>	kz index
<i>is</i>	particle species index

returns flattened index of a gyrokinetic potential  $\chi^\phi(\mathbf{k})$  from its 4D index in FOURIER SPACE. flattened index is then can be used to access required value of the gyrokinetic potential at position (ix,iy,iz,is).

## 2.1.2.6 getIndChiBufEL\_r()

```
size_t getIndChiBufEL_r (
    size_t ix,
    size_t iy,
    size_t iz,
    size_t is )
```

returns returns flat index of an element of electrostatic gyrokinetic potential in REAL SPACE

## Parameters

<i>ix</i>	x index
<i>iy</i>	y index
<i>iz</i>	z index
<i>is</i>	particle species index

returns flattened index of a gyrokinetic potential  $\chi^\phi(\mathbf{r})$  from its 4D index in REAL SPACE. flattened index is then can be used to access required value of the gyrokinetic potential at position (ix,iy,iz,is).

## 2.1.2.7 getIndChiBufEM\_c()

```
size_t getIndChiBufEM_c (
    size_t ix,
    size_t iy,
    size_t iz,
    size_t is,
    size_t ifield )
```

returns flat index of an element of electromagnetic gyrokinetic potential in FOURIER SPACE

## Parameters

<i>ix</i>	kx index
<i>iy</i>	ky index
<i>iz</i>	kz index
<i>is</i>	particle species index
<i>ifield</i>	field type

returns flattened index of a gyrokinetic potential  $\chi^{\phi,A,B}$  from its 4D index in FOURIER SPACE. flattened index is then can be used to access required value of the gyrokinetic potential at position (ix,iy,iz,is). Type of gyrokinetic potential is specified by ifield parameter. Use 0 is to access  $\chi^{\phi}(\mathbf{k})$ , 1 to access  $\chi^A(\mathbf{k})$  and 2 to access  $\chi^B(\mathbf{k})$ .

### 2.1.2.8 getIndChiBufEM\_r()

```
size_t getIndChiBufEM_r (
    size_t ix,
    size_t iy,
    size_t iz,
    size_t is,
    size_t ifield )
```

returns flat index of an element of electromagnetic gyrokinetic potential in POSITION SPACE

#### Parameters

<i>ix</i>	x index
<i>iy</i>	y index
<i>iz</i>	z index
<i>is</i>	particle species index
<i>ifield</i>	field type

returns flattened index of a gyrokinetic potential  $\chi^{\phi,A,B}(\mathbf{k})$  from its 4D index in POSITION SPACE. flattened index is then can be used to access required value of the gyrokinetic potential at position (ix,iy,iz,is). Type of gyrokinetic potential is specified by ifield parameter. Use 0 is to access  $\chi^{\phi}(\mathbf{r})$ , 1 to access  $\chi^A(\mathbf{r})$  and 2 to access  $\chi^B(\mathbf{r})$ .

### 2.1.2.9 multiply\_ar\_c()

```
void multiply_ar_c (
    COMPLEX * ar1,
    COMPLEX * ar2,
    COMPLEX * ret )
```

[multiply\\_ar\\_c\(COMPLEX \\*ar1, COMPLEX \\*ar2, COMPLEX \\*ret\)](#)

### 2.1.2.10 multiply\_ar\_r()

```
void multiply_ar_r (
    const double * ar1,
    const double * ar2,
    double * ret )
```

[multiply\\_ar\\_r\(const double \\*ar1, const double \\*ar2, double \\*ret\)](#)

## 2.2 diagnostics.c File Reference

diagnostics module

```
#include "diagnostics.h"
#include "parameters_io.h"
```

## Macros

- #define **TO\_ROOT** 0
- #define **BUFFER\_SIZE** 1

## Functions

- void [diag\\_computeSpectra](#) (const COMPLEX \*g, const COMPLEX \*h, int timestep)  
*general function to compute k or m spectra*
- void [diag\\_initSpec](#) ()  
*initialize spectra computation*
- void [diag\\_computeFreeEnergy](#) (COMPLEX \*g, COMPLEX \*h)  
*compute free energy*
- void [diag\\_computeKSpectrum](#) (const COMPLEX \*g, const COMPLEX \*h, double \*spec)
- void [diag\\_computeMSpectrum](#) (const COMPLEX \*g, const COMPLEX \*h, double \*spec)  
*computes free energy spectra in m space*
- void [diag\\_getShells](#) ()  
*computes shells from parameters*
- double [diag\\_computeFreeEnergyFields](#) (COMPLEX \*g, COMPLEX \*fields)  
*to be done later*
- void [diag\\_compute](#) (COMPLEX \*g, COMPLEX \*h, int timestep)  
*computes all diagnostics*

## Variables

- double \* [diag\\_kSpec](#) = 0  
*used to store free energy k spectra*
- double \* [diag\\_mSpec](#) = 0  
*used to store free energy m spectra*
- double \* [diag\\_shells](#) = 0  
*used to store positions of k shells required to compute k spectra*
- double [diag\\_freeEnergy](#)  
*free energy*

### 2.2.1 Detailed Description

diagnostics module

different diagnostic tools are gathered in this module

### 2.2.2 Function Documentation

#### 2.2.2.1 [diag\\_compute\(\)](#)

```
void diag_compute (  
    COMPLEX * g,  
    COMPLEX * h,  
    int timestep )
```

computes all diagnostics

## Parameters

<i>g</i>	modified distribution function
<i>h</i>	distribution function
<i>iter</i>	current time step

**2.2.2.2 diag\_computeFreeEnergy()**

```
void diag_computeFreeEnergy (
    COMPLEX * g,
    COMPLEX * h )
```

compute free energy

## Parameters

<i>g</i>	modified gyrokinetic distribution function
<i>h</i>	gyrokinetic distribution function

computes free energy as  $W = 2.\Re(\sum_{k_x, k_y, k_z > 0, m, l, s} g * \bar{h})$ , taking into account reality condition.

**2.2.2.3 diag\_computeFreeEnergyFields()**

```
diag_computeFreeEnergyFields (
    COMPLEX * g,
    COMPLEX * fields )
```

to be done later

## Parameters

<i>g</i>	
<i>fields</i>	computes free energy from the fields and distribution function.

**2.2.2.4 diag\_computeKSpectrum()**

```
void diag_computeKSpectrum (
    const COMPLEX * g,
    const COMPLEX * h,
    double * spec )
```

## Parameters

<i>g</i>	modified gyrokinetic distribution function
<i>h</i>	gyrokinetic distribution function
<i>spec</i>	spectra array



computes free energy  $k_{\perp}$  spectra  $W(k_i^{shell}) = \frac{1}{N} \sum_{k_{i-1}^{shell} < |k_{\perp}| < k_i^{shell}} \sum_{k_z, l, m, s} g \bar{h}$  where  $N$  is a number of wave vectors between shells  $k_{i-1}^{shell}$  and  $k_i^{shell}$

### 2.2.2.5 diag\_computeMSpectrum()

```
void diag_computeMSpectrum (
    const COMPLEX * g,
    const COMPLEX * h,
    double * spec )
```

computes free energy spectra in m space

#### Parameters

<i>g</i>	modified gyrokinetic distribution function
<i>h</i>	gyrokinetic distribution function
<i>spec</i>	spectra array

computes free energy m spectra as  $W(m) = \sum_{k_x, k_y, k_z, l, s} g \bar{h}$

### 2.2.2.6 diag\_computeSpectra()

```
void diag_computeSpectra (
    const COMPLEX * g,
    const COMPLEX * h,
    int timestep )
```

general function to compute k or m spectra

#### Parameters

<i>g</i>	gyrokinetic distribution function
<i>h</i>	distribution function
<i>timestep</i>	current time step

function computes spectra at timestep as given in parameter file.  $k_{\perp}$  spectra is computed using [diag\\_computeKSpectrum](#), and m spectra is computed using [diag\\_computeMSpectrum](#)

### 2.2.2.7 diag\_getShells()

```
diag_getShells ( )
```

computes shells from parameters

computes positions of k\_shells in between last\_shell and first\_shell as provided by user in parameter file. Position of  $i^{th}$  shell is computed as  $k_i^{shell} = (last\_shell - first\_shell)/(k\_shells) \cdot i$

### 2.2.2.8 diag\_initSpec()

```
void diag_initSpec ( )
```

initialize spectra computation

Prepares free energy spectra computation. For spectra in k: allocates diag\_kSpec array used to store k spectra. Allocates diag\_shells array and fills it with shell positions  $k^{shells}$ , used for binning of wave vectors when computing  $k_{\perp}$  spectra. For spectra in m: allocates diag\_mSpec array used to store m spectra. Called in [init\\_start](#) function

## 2.3 distrib.c File Reference

gyrokinetic distribution function module

```
#include "distrib.h"
```

### Functions

- void [distrib\\_getH](#) (COMPLEX \*h, const COMPLEX \*g)  
*computes h from g*
- void [distrib\\_getG](#) (COMPLEX \*g, const COMPLEX \*h)  
*computes g from h*
- void [distrib\\_getXGrad](#) (const COMPLEX \*in, COMPLEX \*out)  
*Computes gradient in kx direction.*
- void [distrib\\_getYGrad](#) (const COMPLEX \*in, COMPLEX \*out)  
*Computes gradient in ky direction.*
- void [distrib\\_getZGrad](#) (const COMPLEX \*in, COMPLEX \*out)  
*Computes gradient in kz direction.*
- void [distrib\\_enforceReality](#) (COMPLEX \*f)  
*enforces reality condition on distribution function array*
- void [distrib\\_setZeroNHalf](#) (COMPLEX \*f)  
*sets all Nk/2 modes to zero*

### 2.3.1 Detailed Description

gyrokinetic distribution function module

everything required to perform different manipulations to distribution functions

### 2.3.2 Function Documentation

#### 2.3.2.1 distrib\_enforceReality()

```
void distrib_enforceReality (
    COMPLEX * f )
```

enforces reality condition on distribution function array

## Parameters

<i>f</i>	complex array for which reality condition will be forced.
----------	---

Enforces reality condition  $f(k) = \text{conj}(f(-k))$  in plane  $k_z = 0$ . For a given  $k_x$ , it first checks where modes  $-k_x$  are located using the `#mpi_whereIsX` function:

```
where_neg = mpi_whereIsX[kxNeg * 2];
```

If  $-k_x$  is stored on a different processor, `MPI_VECTOR` with a 4D data slice  $f(k_x, k_z = 0)$  is sent to this processor, to the buffer array:

```
mpi_sendVector(&f[ind6D], buffer, where_pos, where_neg);
```

if the data stored on the same processor, no vector is being sent. Reality condition is fulfilled in a loop over all other coordinates:

```
f[ind6D_neg] = conj(buffer[ind6D_pos]);
```

**2.3.2.2 distrib\_getG()**

```
distrib_getG (
    COMPLEX * g,
    const COMPLEX * h )
```

computes  $g$  from  $h$

## Parameters

<i>g</i>	complex array to store $g$
<i>h</i>	complex array with $h$

computes modified gyrokinetic distribution function  $g$  from gyrokinetic distribution function  $h$ . Please note that before calling this function gyrokinetic potentials must be computed

**2.3.2.3 distrib\_getH()**

```
void distrib_getH (
    COMPLEX * h,
    const COMPLEX * g )
```

computes  $h$  from  $g$

## Parameters

<i>h</i>	complex array to store $h$
<i>g</i>	complex array with $g$

computes gyrokinetic distribution function  $h$  from modified gyrokinetic distribution function  $g$ . Please note that before calling this function gyrokinetic potentials must be computed

**2.3.2.4 distrib\_getXGrad()**

```
void distrib_getXGrad (
```

```
const COMPLEX * in,
COMPLEX * out )
```

Computes gradient in kx direction.

#### Parameters

<i>in</i>	complex array. Distribution function of which gradient will be taken
<i>out</i>	complex array, where gradient is stored

Computes gradient in kx direction as following:

$$\text{grad}(f) = i * kx * f$$

#### 2.3.2.5 distrib\_getYGrad()

```
void distrib_getYGrad (
    const COMPLEX * in,
    COMPLEX * out )
```

Computes gradient in ky direction.

#### Parameters

<i>in</i>	complex array. Distribution function of which gradient will be taken
<i>out</i>	complex array, where gradient is stored

Computes gradient in ky direction as following:

$$\text{grad}(f) = i * ky * f$$

#### 2.3.2.6 distrib\_getZGrad()

```
void distrib_getZGrad (
    const COMPLEX * in,
    COMPLEX * out )
```

Computes gradient in kz direction.

#### Parameters

<i>in</i>	complex array. Distribution function of which gradient will be taken
<i>out</i>	complex array, where gradient is stored

Computes gradient in kz direction as following:

$$\text{grad}(f) = i * kz * f$$

#### 2.3.2.7 distrib\_setZeroNHalf()

```
void distrib_setZeroNHalf (
    COMPLEX * f )
```

sets all  $N_k/2$  modes to zero

#### Parameters

$f$	complex array
-----	---------------

sets  $N_{kx}/2$ ,  $N_{ky}/2$  and  $N_z/2$  modes of distribution function to zero. Due to reality condition, for  $k_z$  the last mode should be set to zero.

## 2.4 equation.c File Reference

equation module

```
#include "equation.h"
#include "array.h"
#include "fields.h"
#include "distrib.h"
```

### Macros

- `#define CHI_EM 3`
- `#define CHI_EL 1`
- `#define CHI_PHI 0`
- `#define CHI_A 1`
- `#define CHI_B 2`

### Functions

- void [equation\\_getLinearTerm](#) (const COMPLEX \*in, const COMPLEX \*plus\_boundary, const COMPLEX \*minus\_boundary, COMPLEX \*out)  
*computes linear term*
- void [equation\\_getNonlinearElectromagnetic](#) (double \*in, double \*chiAr, double \*out, double sign)  
*returns nonlinear electromagnetic term*
- void [equation\\_getNonlinearElectrostatic](#) (double \*in, double \*chiAr, double \*out, double sign)  
*returns nonlinear electrostatic term*
- void [equation\\_getNonlinearProduct](#) (double \*in, double \*chiAr, double \*out, double sign)  
*chooses between computing electrostatic or electromagnetic term*
- void [equation\\_getNonlinearTerm](#) (const COMPLEX \*h, COMPLEX \*out)  
*computes nonlinear term*
- void [equation\\_getRHS](#) (const COMPLEX \*in\_g, COMPLEX \*in\_h, COMPLEX \*out)

### 2.4.1 Detailed Description

equation module

module required to compute RHS of the equation.

## 2.4.2 Function Documentation

### 2.4.2.1 `equation_getLinearTerm()`

```
void equation_getLinearTerm (
    const COMPLEX * in,
    const COMPLEX * plus_boundary,
    const COMPLEX * minus_boundary,
    COMPLEX * out )
```

computes linear term

#### Parameters

<i>in</i>	complex array
<i>out</i>	complex array
<i>plus_boundary</i>	complex array
<i>minus_boundary</i>	complex array

computes linear term *out* from distribution function *in* .

### 2.4.2.2 `equation_getNonlinearElectromagnetic()`

```
void equation_getNonlinearElectromagnetic (
    double * in,
    double * chiAr,
    double * out,
    double sign )
```

returns nonlinear electromagnetic term

#### Parameters

<i>in</i>	input double array
<i>chiAr</i>	input double array
<i>out</i>	output double array
<i>sign</i>	should be 1 or -1

performs multiplication between input 6D complex array *in* and gyrokinetic potential array *chiAr*, in such way that the structure of the product is the same as nonlinear term of drift-kinetic equations. Used by [equation\\_getNonlinearProduct](#). *sign* is used to determine the sign of the resulting product. See [equation\\_getNonlinearTerm](#) for explanation.

### 2.4.2.3 `equation_getNonlinearElectrostatic()`

```
void equation_getNonlinearElectrostatic (
    double * in,
```

```
double * chiAr,
double * out,
double sign )
```

returns nonlinear electrostatic term

#### Parameters

<i>in</i>	input double array
<i>chiAr</i>	input double array
<i>out</i>	output double array
<i>sign</i>	should be 1 or -1

see [equation\\_getNonlinearElectromagnetic](#) for explanation

#### 2.4.2.4 equation\_getNonlinearProduct()

```
equation_getNonlinearProduct (
    double * in,
    double * chiAr,
    double * out,
    double sign )
```

chooses between computing electrostatic or electromagnetic term

#### Parameters

<i>in</i>	input double array
<i>chiAr</i>	input double array
<i>out</i>	output double array
<i>sign</i>	should be 1 or -1

depending on flag `systemType` provided by user in parameter file, chooses between [equation\\_getNonlinearElectrostatic](#) and [equation\\_getNonlinearElectromagnetic](#)

#### 2.4.2.5 equation\_getNonlinearTerm()

```
void equation_getNonlinearTerm (
    const COMPLEX * h,
    COMPLEX * out )
```

computes nonlinear term

#### Parameters

<i>h</i>	input complex array
<i>out</i>	output complex array

function returns nonlinear term. First it takes y gradient of distribution function *h*, and x gradient of gyrokinetic

potentials chi, and transforms them to real space:

```
distrib_getYGrad(h, fftw_hBuf);
fields_getGradX(fftw_chiBuf);
fftw_c2r();
fftw_c2r_chi();
```

after that, it computes  $\frac{\partial h}{\partial y} \frac{\partial \chi}{\partial x}$  part of the poisson brackets:

```
equation_getNonlinearProduct((double *)fftw_hBuf, (double *)fftw_chiBuf,
buffer, 1.);
```

with the result stored in `buffer` after that, it computes x gradient of h and y gradient of gyrokinetic potential chi, and transforms results to real space:

```
distrib_getXGrad(h, fftw_hBuf);
fields_getGradY(fftw_chiBuf);
fftw_c2r();
fftw_c2r_chi();
```

and computes second part of the poisson brackets  $-\frac{\partial h}{\partial x} \frac{\partial \chi}{\partial y}$  and adds the result to `buffer`. `buffer` is then transformed back to Fourier space, and dealiasing is performed.

## 2.5 fields.c File Reference

field computation and manipulation module

```
#include "fields.h"
```

### Macros

- `#define CHI_PHI 0`
- `#define CHI_A 1`
- `#define CHI_B 2`

### Functions

- void [fields\\_init](#) ()  
*intializes fields*
- void [fields\\_getA](#) (const COMPLEX \*g)  
*compute A field*
- void [fields\\_getB](#) (const COMPLEX \*g0, const COMPLEX \*g1)  
*computes B potential*
- void [fields\\_getPhi](#) (const COMPLEX \*g0, const COMPLEX \*g1)  
*computes phi potential*
- void [fields\\_getFields](#) (COMPLEX \*g00, COMPLEX \*g10, COMPLEX \*g01)  
*wrapper to get all the fields simultaneously*
- void [fields\\_getChi](#) ()  
*computes gyrokinetic potentials chi*
- void [fields\\_getChiPhi](#) ()  
*computes chiPhi gyrokinetic potential from phi potential*
- void [fields\\_getChiB](#) ()  
*computes chiB gyrokinetic potential from B potential*
- void [fields\\_getChiA](#) ()  
*computes chiA gyrokinetic potential from A potential*



- void `fields_sendG` (COMPLEX \*g)  
*sends moments of gyrokinetic distribution function which are required to compute fields*
- void `fields_getFieldsFromH` (COMPLEX \*h00, COMPLEX \*h10, COMPLEX \*h01)  
*wrapper to get all the fields simultaneously, computed from gyrokinetic distribution function*
- void `fields_getAFromH` (const COMPLEX \*h)  
*compute A field*
- void `fields_getBFromH` (const COMPLEX \*h0, const COMPLEX \*h1)  
*computes B potential*
- void `fields_getPhiFromH` (const COMPLEX \*h)  
*computes phi potential*
- void `fields_getGradX` (COMPLEX \*out)  
*computes chi gradient in x direction*
- void `fields_getGradY` (COMPLEX \*out)  
*computes chi gradient in y direction*

## Variables

- struct fields\_fields **fields\_fields**
- struct fields\_chi **fields\_chi**
- double \* **A\_denom**
- double \* **qnvTsJ**
- double \* **I\_B**
- double \* **I\_phi**
- double \* **a\_pot**
- double \* **b\_pot**
- double \* **c\_pot**
- double \* **phiB\_denom**
- int \* **global\_nm\_index**
- COMPLEX \* **g00**
- COMPLEX \* **g10**
- COMPLEX \* **g01**

### 2.5.1 Detailed Description

field computation and manipulation module

Required to compute  $A_{||}(\mathbf{k})$ ,  $B_{||}(\mathbf{k})$ ,  $\phi(\mathbf{k})$  potentials, as well as gyrokinetic potentials  $\chi_s^A(\mathbf{k})$ ,  $\chi_s^B(\mathbf{k})$ ,  $\chi_s^\phi(\mathbf{k})$

### 2.5.2 Function Documentation

#### 2.5.2.1 fields\_getA()

```
void fields_getA (
    const COMPLEX * g )
```

compute A field

**Parameters**

<i>g</i>	4D complex array (kx,ky,kz,s). Must be first Hermite and zeroth Laguerre moment of modified gyrokinetic distribution function g.
----------	--

computes  $A_{||}(\mathbf{k})$  potential from  $g_{s0}^1$  (g parameter)

**2.5.2.2 fields\_getAFromH()**

```
void fields_getAFromH (
    const COMPLEX * h )
```

compute A field

**Parameters**

<i>h</i>	4D complex array (kx,ky,kz,s). Must be first Hermite and zeroth Laguerre moment of gyrokinetic distribution function h.
----------	---

computes  $A_{||}(\mathbf{k})$  potential from  $h_{s0}^1$  (h parameter)

**2.5.2.3 fields\_getB()**

```
void fields_getB (
    const COMPLEX * g0,
    const COMPLEX * g1 )
```

computes B potential

**Parameters**

<i>g0</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and Laguerre moment of modified gyrokinetic distribution function.
<i>g1</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and first Laguerre moment of the modified distribution function.

Computes  $B_{\perp}(\mathbf{k})$  from  $g_{s0}^0$  (g0 parameter) and  $g_{s0}^1$  (g1 parameter).

**2.5.2.4 fields\_getBFromH()**

```
void fields_getBFromH (
    const COMPLEX * h0,
    const COMPLEX * h1 )
```

computes B potential

**Parameters**

<i>h0</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and Laguerre moment of gyrokinetic distribution function.
<i>h1</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and first Laguerre moment of distribution function.

Computes  $B_{\perp}(\mathbf{k})$  from  $h_{s0}^0$  (h0 parameter) and  $h_{s0}^1$  (h1 parameter).

### 2.5.2.5 fields\_getChi()

```
void fields_getChi ( )
```

computes gyrokinetic potentials chi

Wrapper for functions [fields\\_getChiPhi](#), [fields\\_getChiA](#), [fields\\_getChiB](#)

### 2.5.2.6 fields\_getChiA()

```
void fields_getChiA ( )
```

computes chiA gyrokinetic potential from A potential

computes  $chi_s^A(\mathbf{k})$

### 2.5.2.7 fields\_getChiB()

```
void fields_getChiB ( )
```

computes chiB gyrokinetic potential from B potential

computes  $\chi_s^B(\mathbf{k})$

### 2.5.2.8 fields\_getChiPhi()

```
fields_getChiPhi ( )
```

computes chiPhi gyrokinetic potential from phi potential

computes  $chi_s^{\phi}(\mathbf{k})$

### 2.5.2.9 fields\_getFields()

```
void fields_getFields (
    COMPLEX * g00,
    COMPLEX * g10,
    COMPLEX * g01 )
```

wrapper to get all the fields simultaneously

#### Parameters

<i>g00</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and Laguerre moment of modified gyrokinetic distribution function.
<i>g10</i>	4D complex array (kx,ky,kz,s). Must be first Hermite and zeroth Laguerre moment of modified gyrokinetic distribution function g.
<i>g01</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and first Laguerre moment of the modified distribution function.

Wrapper for functions [fields\\_getPhi](#), [fields\\_getB](#), [fields\\_getA](#).

### 2.5.2.10 fields\_getFieldsFromH()

```
void fields_getFieldsFromH (
    COMPLEX * h00,
    COMPLEX * h10,
    COMPLEX * h01 )
```

wrapper to get all the fields simultaneously, computed from gyrokinetic distribution function

#### Parameters

<i>h00</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and Laguerre moment of gyrokinetic distribution function.
<i>h10</i>	4D complex array (kx,ky,kz,s). Must be first Hermite and zeroth Laguerre moment of gyrokinetic distribution function h.
<i>h01</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and first Laguerre moment of the distribution function.

Wrapper for functions [fields\\_getPhiFromH](#), [#fields\\_get\\_BFromH](#), [fields\\_getAFromH](#).

### 2.5.2.11 fields\_getGradX()

```
void fields_getGradX (
    COMPLEX * out )
```

computes chi gradient in x direction

#### Parameters

<i>out</i>	output complex array of size (kx,ky,kz,s,Nfields).
------------	--

computes gradient in x direction for chi potentials. Nfields<\tt> can be 1 or 3, and chosen automatically at start of the simulation depending on the simulation type (electrostatic or electromagnetic)

### 2.5.2.12 fields\_getGradY()

```
void fields_getGradY (
    COMPLEX * out )
```

computes chi gradient in y direction

#### Parameters

<i>out</i>	output complex array of size (kx,ky,kz,s,Nfields).
------------	--

computes gradient in y direction for chi potentials. Nfields<\tt> can be 1 or 3, and chosen automatically at start of the simulation depending on the simulation type

(electrostatic or electromagnetic)

### 2.5.2.13 fields\_getPhi()

```
void fields_getPhi (
    const COMPLEX * g0,
    const COMPLEX * g1 )
```

computes phi potential

#### Parameters

<i>g0</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and Laguerre moment of modified gyrokinetic distribution function.
<i>g1</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and first Laguerre moment of the modified distribution function.

Computes  $\phi(\mathbf{k})$  from  $g_{s0}^0$  ( $g0$  parameter) and  $g_{s0}^1$  ( $g1$  parameter).

### 2.5.2.14 fields\_getPhiFromH()

```
void fields_getPhiFromH (
    const COMPLEX * h )
```

computes phi potential

#### Parameters

<i>h</i>	4D complex array (kx,ky,kz,s). Zeroth Hermite and Laguerre moment of gyrokinetic distribution function.
----------	---

Computes  $\phi(\mathbf{k})$  from  $h_{s0}^0$  ( $h$  parameter)

### 2.5.2.15 fields\_init()

```
void fields_init ( )
```

initializes fields

pre-computes some constants required to compute fields. Called in [init\\_start](#) function

### 2.5.2.16 fields\_sendG()

```
fields_sendG (
    COMPLEX * g )
```

sends moments of gyrokinetic distribution function which are required to compute fields

## Parameters

$g$	complex array. Modified or non-modified gyrokinetic distribution function
-----	---

sends  $g_{s0}^1(\mathbf{k})f$ ,  $g_{s1}^0(\mathbf{k})f$ ,  $g_{s0}^0(\mathbf{k})f$

to all processes to compute potentials locally.

## 2.6 init.c File Reference

initialization module for alliance.

```
#include "init.h"
#include "distrib.h"
```

### Macros

- `#define RANK_IO 0`

### Functions

- void `init_start` (char \*filename)  
*initialization of ALLIANCE*
- void `init_printParameters` ()  
*parameter output*
- void `init_initEnums` ()  
*enumerator initialization*
- void `fill_rand` (COMPLEX \*ar1)  
*fills the initial conditions randomly*
- void `fill_randM0` (COMPLEX \*ar1)  
*fill zeroth Hermite moment with random values*
- void `fill_randSingleKM` (COMPLEX \*ar1)  
*fill single chosen wavevector and Hermite moment*
- void `init_conditions` (COMPLEX \*data)  
*distribution function initialization*
- double `init_energySpec` (double k, double m, double amp, double disp)  
*returns energy spectrum*

### Variables

- enum adiabatic **kinetic**
- enum electromagnetic **systemType**
- enum initial **initialConditions**

## 2.6.1 Detailed Description

initialization module for alliance.

all the initialization routines are here.

## 2.6.2 Macro Definition Documentation

### 2.6.2.1 RANK\_IO

```
#define RANK_IO 0
```

defines rank of the processor used to output information to console

## 2.6.3 Function Documentation

### 2.6.3.1 fill\_rand()

```
void fill_rand (
    COMPLEX * data )
```

fills the initial conditions randomly

#### Parameters

<i>data</i>	complex 6d array to fill initializes distribution with spectrum defined in <a href="#">init_energySpec</a> This function is supposed to be used in-module only and should not be used elsewhere outside <a href="#">init.c</a> file.
-------------	--

### 2.6.3.2 fill\_randM0()

```
void fill_randM0 (
    COMPLEX * data )
```

fill zeroth Hermite moment with random values

#### Parameters

<i>data</i>	complex 6D array to fill
-------------	--------------------------

fills 0-th Hermite moment of a distribution function `ar1` with random values. This function is supposed to be used in-module only and should not be used elsewhere outside `init.c` file.

### 2.6.3.3 `fill_randSingleKM()`

```
void fill_randSingleKM (
    COMPLEX * ar1 )
```

fill single chosen wavevector and Hermite moment

#### Parameters

<i>data</i>	complex 6D array
-------------	------------------

initializes single wavevector and Hermite moment of a distribution function with random variable. This function is only for in-module use and should not be used elsewhere outside `init.c` file.

### 2.6.3.4 `init_conditions()`

```
void init_conditions (
    COMPLEX * data )
```

distribution function initialization

#### Parameters

<i>data</i>	complex 6D array
-------------	------------------

initializes distribution function with chosen method (see `fill_rand`, `fill_randM0`, `fill_randSingleKM`)

### 2.6.3.5 `init_energySpec()`

```
double init_energySpec (
    double k,
    double m,
    double amp,
    double disp )
```

returns energy spectrum

#### Parameters

<i>k</i>	a wavenumber at which spectrum is computed
<i>m</i>	Hermite moment at which amplitude is computed
<i>amp</i>	amplitude of the spectrum
<i>disp</i>	dispersion of the spectrum

computes spectrum of form  $A \cdot k^2 \exp(-2k^2/\sigma^2)$ , where  $\sigma = \text{disp}$ , and  $A = \text{amp}$ . This function is supposed to be



used in-module only and should not be used elsewhere outside [init.c](#) file.

### 2.6.3.6 init\_initEnums()

```
void init_initEnums ( )
```

enumerator initialization

initializes enumerators, which are then used to define if system is adiabatic or kinetic, electromagnetic or electrostatic, and type of initial conditions

### 2.6.3.7 init\_printParameters()

```
void init_printParameters ( )
```

parameter output

prints parameters of the simulation

### 2.6.3.8 init\_start()

```
void init_start (
    char * filename )
```

initialization of ALLIANCE

#### Parameters

<i>filename</i>	specifies parameter filename
-----------------	------------------------------

initializes all the modules required for ALLIANCE to work.

## 2.7 parameters\_io.c File Reference

reads input parameters from parameter file provided by user

```
#include "parameters_io.h"
#include "utils_fftw.h"
```

### Macros

- `#define VERBOSE 0`

## Functions

- void `init_global_size` ()  
*initializes global size of the 6D array*
- void `read_parameters` (char \*filename)  
*reads parameters from user parameter file.*
- void `read_parametersFromFile` (char \*filename)

## Variables

- struct system\_param **parameters**

### 2.7.1 Detailed Description

reads input parameters from parameter file provided by user

### 2.7.2 Function Documentation

#### 2.7.2.1 `init_global_size()`

```
void init_global_size ( )
```

initializes global size of the 6D array

initializes `array_local_size` structure with global simulation size.

#### 2.7.2.2 `read_parameters()`

```
read_parameters (
    char * filename )
```

reads parameters from user parameter file.

Reads parameters from user parameter file. All the parameters are stored in the `parameters` structure

#### 2.7.2.3 `read_parametersFromFile()`

```
void read_parametersFromFile (
    char * filename )
```

`read_parametersFromFile(char *filename):`

## 2.8 solver.c File Reference

numerical solver

```
#include "solver.h"
```

### Macros

- #define **SOLVERTYPE** RK4
- #define **IORANK** 0

### Functions

- void [solver\\_init](#) ()  
*initializes solver*
- void [solver\\_makeStep](#) (COMPLEX \*\*g, COMPLEX \*h)  
*iterate solver forward*

### Variables

- enum solverType **solverType**
- struct solver **solver**
- struct rk4 **rk4**

### 2.8.1 Detailed Description

numerical solver

### 2.8.2 Function Documentation

#### 2.8.2.1 solver\_init()

```
void solver_init ( )
```

initializes solver

initializes solver with the `solverType`.

#### 2.8.2.2 solver\_makeStep()

```
void solver_makeStep (  
    COMPLEX ** g,  
    COMPLEX * h )
```

iterate solver forward

**Parameters**

<i>g</i>	address of the 6D complex array. Modified gyrokinetic distribution function
<i>h</i>	6D complex array. Gyrokinetic distribution function

solves one simulation time step

# Index

- array.c, [3](#)
  - [get\\_flat\\_c](#), [4](#)
  - [get\\_flat\\_r](#), [4](#)
  - [get\\_flatIndexComplex3D](#), [5](#)
  - [getIndChi](#), [5](#)
  - [getIndChiBufEL\\_c](#), [5](#)
  - [getIndChiBufEL\\_r](#), [7](#)
  - [getIndChiBufEM\\_c](#), [7](#)
  - [getIndChiBufEM\\_r](#), [8](#)
  - [multiply\\_ar\\_c](#), [8](#)
  - [multiply\\_ar\\_r](#), [8](#)
- diag\_compute
  - [diagnostics.c](#), [9](#)
- diag\_computeFreeEnergy
  - [diagnostics.c](#), [10](#)
- diag\_computeFreeEnergyFields
  - [diagnostics.c](#), [10](#)
- diag\_computeKSpectrum
  - [diagnostics.c](#), [10](#)
- diag\_computeMSpectrum
  - [diagnostics.c](#), [11](#)
- diag\_computeSpectra
  - [diagnostics.c](#), [11](#)
- diag\_getShells
  - [diagnostics.c](#), [11](#)
- diag\_initSpec
  - [diagnostics.c](#), [11](#)
- diagnostics.c, [8](#)
  - [diag\\_compute](#), [9](#)
  - [diag\\_computeFreeEnergy](#), [10](#)
  - [diag\\_computeFreeEnergyFields](#), [10](#)
  - [diag\\_computeKSpectrum](#), [10](#)
  - [diag\\_computeMSpectrum](#), [11](#)
  - [diag\\_computeSpectra](#), [11](#)
  - [diag\\_getShells](#), [11](#)
  - [diag\\_initSpec](#), [11](#)
- distrib.c, [12](#)
  - [distrib\\_enforceReality](#), [12](#)
  - [distrib\\_getG](#), [13](#)
  - [distrib\\_getH](#), [13](#)
  - [distrib\\_getXGrad](#), [13](#)
  - [distrib\\_getYGrad](#), [14](#)
  - [distrib\\_getZGrad](#), [14](#)
  - [distrib\\_setZeroNHalf](#), [14](#)
- distrib\_enforceReality
  - [distrib.c](#), [12](#)
- distrib\_getG
  - [distrib.c](#), [13](#)
- distrib\_getH
  - [distrib.c](#), [13](#)
- distrib.c, [13](#)
  - [distrib\\_getXGrad](#)
    - [distrib.c](#), [13](#)
  - [distrib\\_getYGrad](#)
    - [distrib.c](#), [14](#)
  - [distrib\\_getZGrad](#)
    - [distrib.c](#), [14](#)
  - [distrib\\_setZeroNHalf](#)
    - [distrib.c](#), [14](#)
- equation.c, [15](#)
  - [equation\\_getLinearTerm](#), [16](#)
  - [equation\\_getNonlinearElectromagnetic](#), [16](#)
  - [equation\\_getNonlinearElectrostatic](#), [16](#)
  - [equation\\_getNonlinearProduct](#), [17](#)
  - [equation\\_getNonlinearTerm](#), [17](#)
- equation\_getLinearTerm
  - [equation.c](#), [16](#)
- equation\_getNonlinearElectromagnetic
  - [equation.c](#), [16](#)
- equation\_getNonlinearElectrostatic
  - [equation.c](#), [16](#)
- equation\_getNonlinearProduct
  - [equation.c](#), [17](#)
- equation\_getNonlinearTerm
  - [equation.c](#), [17](#)
- fields.c, [18](#)
  - [fields\\_getA](#), [19](#)
  - [fields\\_getAFromH](#), [20](#)
  - [fields\\_getB](#), [20](#)
  - [fields\\_getBFromH](#), [20](#)
  - [fields\\_getChi](#), [21](#)
  - [fields\\_getChiA](#), [21](#)
  - [fields\\_getChiB](#), [21](#)
  - [fields\\_getChiPhi](#), [21](#)
  - [fields\\_getFields](#), [21](#)
  - [fields\\_getFieldsFromH](#), [22](#)
  - [fields\\_getGradX](#), [22](#)
  - [fields\\_getGradY](#), [22](#)
  - [fields\\_getPhi](#), [23](#)
  - [fields\\_getPhiFromH](#), [23](#)
  - [fields\\_init](#), [23](#)
  - [fields\\_sendG](#), [23](#)
- fields\_getA
  - [fields.c](#), [19](#)
- fields\_getAFromH
  - [fields.c](#), [20](#)
- fields\_getB
  - [fields.c](#), [20](#)

- fields\_getBFromH
  - fields.c, [20](#)
- fields\_getChi
  - fields.c, [21](#)
- fields\_getChiA
  - fields.c, [21](#)
- fields\_getChiB
  - fields.c, [21](#)
- fields\_getChiPhi
  - fields.c, [21](#)
- fields\_getFields
  - fields.c, [21](#)
- fields\_getFieldsFromH
  - fields.c, [22](#)
- fields\_getGradX
  - fields.c, [22](#)
- fields\_getGradY
  - fields.c, [22](#)
- fields\_getPhi
  - fields.c, [23](#)
- fields\_getPhiFromH
  - fields.c, [23](#)
- fields\_init
  - fields.c, [23](#)
- fields\_sendG
  - fields.c, [23](#)
- fill\_rand
  - init.c, [25](#)
- fill\_randM0
  - init.c, [25](#)
- fill\_randSingleKM
  - init.c, [26](#)
- get\_flat\_c
  - array.c, [4](#)
- get\_flat\_r
  - array.c, [4](#)
- get\_flatIndexComplex3D
  - array.c, [5](#)
- getIndChi
  - array.c, [5](#)
- getIndChiBufEL\_c
  - array.c, [5](#)
- getIndChiBufEL\_r
  - array.c, [7](#)
- getIndChiBufEM\_c
  - array.c, [7](#)
- getIndChiBufEM\_r
  - array.c, [8](#)
- init.c, [24](#)
  - fill\_rand, [25](#)
  - fill\_randM0, [25](#)
  - fill\_randSingleKM, [26](#)
  - init\_conditions, [26](#)
  - init\_energySpec, [26](#)
  - init\_initEnums, [27](#)
  - init\_printParameters, [27](#)
  - init\_start, [27](#)
  - RANK\_IO, [25](#)
  - init\_conditions
    - init.c, [26](#)
  - init\_energySpec
    - init.c, [26](#)
  - init\_global\_size
    - parameters\_io.c, [28](#)
  - init\_initEnums
    - init.c, [27](#)
  - init\_printParameters
    - init.c, [27](#)
  - init\_start
    - init.c, [27](#)
  - multiply\_ar\_c
    - array.c, [8](#)
  - multiply\_ar\_r
    - array.c, [8](#)
  - parameters\_io.c, [27](#)
    - init\_global\_size, [28](#)
    - read\_parameters, [28](#)
    - read\_parametersFromFile, [28](#)
  - RANK\_IO
    - init.c, [25](#)
  - read\_parameters
    - parameters\_io.c, [28](#)
  - read\_parametersFromFile
    - parameters\_io.c, [28](#)
  - solver.c, [29](#)
    - solver\_init, [29](#)
    - solver\_makeStep, [29](#)
  - solver\_init
    - solver.c, [29](#)
  - solver\_makeStep
    - solver.c, [29](#)