phasefield-accelerator-benchmarks pre-alpha

Generated by Doxygen 1.8.13

Contents

1	Clas	s Index	2				
	1.1	Class List	2				
2	File	e Index 2					
	2.1	File List	2				
3	Clas	ss Documentation 3					
	3.1	Stopwatch Struct Reference	3				
		3.1.1 Detailed Description	3				
		3.1.2 Member Data Documentation	4				
4	File	Documentation	5				
	4.1	analytic_main.c File Reference	5				
		4.1.1 Detailed Description	5				
		4.1.2 Function Documentation	5				
	4.2	boundaries.h File Reference	7				
		4.2.1 Detailed Description	8				
		4.2.2 Function Documentation	8				
	4.3	cuda_boundaries.cu File Reference	10				
		4.3.1 Detailed Description	10				
		4.3.2 Function Documentation	11				
		4.3.3 Variable Documentation	12				
	4.4	cuda_discretization.cu File Reference	12				
		4.4.1 Detailed Description	13				
		4.4.2 Function Documentation	13				
		4.4.3 Variable Documentation	16				
	4.5	cuda_kernels.cuh File Reference	16				
		4.5.1 Detailed Description	17				
		4.5.2 Macro Definition Documentation	17				
		4.5.3 Function Documentation	18				

ii CONTENTS

	4.5.4	Variable Documentation	. 19
4.6	discreti	ization.h File Reference	. 20
	4.6.1	Detailed Description	. 21
	4.6.2	Function Documentation	. 21
4.7	main.c	File Reference	. 24
	4.7.1	Detailed Description	. 24
	4.7.2	Function Documentation	. 24
4.8	mesh.c	File Reference	. 25
	4.8.1	Detailed Description	. 26
	4.8.2	Function Documentation	. 26
4.9	mesh.h	File Reference	. 29
	4.9.1	Detailed Description	. 29
	4.9.2	Function Documentation	. 29
4.10	numeri	cs.c File Reference	. 32
	4.10.1	Detailed Description	. 32
	4.10.2	Function Documentation	. 33
4.11	numeri	cs.h File Reference	. 37
	4.11.1	Detailed Description	. 39
	4.11.2	Macro Definition Documentation	. 39
	4.11.3	Function Documentation	. 39
4.12	openac	cc_boundaries.c File Reference	. 44
	4.12.1	Detailed Description	. 45
	4.12.2	Function Documentation	. 45
4.13	openac	cc_discretization.c File Reference	. 47
	4.13.1	Detailed Description	. 48
	4.13.2	Function Documentation	. 48
4.14	openac	cc_kernels.h File Reference	. 52
	4.14.1	Detailed Description	. 53
	4.14.2	Function Documentation	. 53
4.15	openm	p_boundaries.c File Reference	. 55

	4.15.1 Detailed Description	55
	4.15.2 Function Documentation	55
4.16	openmp_discretization.c File Reference	57
	4.16.1 Detailed Description	57
	4.16.2 Function Documentation	57
4.17	output.c File Reference	60
	4.17.1 Detailed Description	60
	4.17.2 Function Documentation	60
4.18	output.h File Reference	63
	4.18.1 Detailed Description	64
	4.18.2 Function Documentation	64
4.19	serial_boundaries.c File Reference	66
	4.19.1 Detailed Description	67
	4.19.2 Function Documentation	67
4.20	serial_discretization.c File Reference	68
	4.20.1 Detailed Description	68
	4.20.2 Function Documentation	69
4.21	tbb_boundaries.cpp File Reference	71
	4.21.1 Detailed Description	71
	4.21.2 Function Documentation	71
4.22	tbb_discretization.cpp File Reference	72
	4.22.1 Detailed Description	73
	4.22.2 Function Documentation	73
4.23	timer.c File Reference	75
	4.23.1 Detailed Description	76
	4.23.2 Function Documentation	76
	4.23.3 Variable Documentation	77
4.24	timer.h File Reference	77
	4.24.1 Detailed Description	77
	4.24.2 Function Documentation	77
4.25	type.h File Reference	78
	4.25.1 Detailed Description	79
	4.25.2 Typedef Documentation	79

Ind	ex	81
1	Class Index	
1.1	Class List	
Hei	re are the classes, structs, unions and interfaces with brief descriptions:	
	Stopwatch	3
2	File Index	
2.1	File List	
Hei	re is a list of all files with brief descriptions:	
	analytic_main.c Analytical solution to semi-infinite diffusion equation	5
	boundaries.h Declaration of boundary condition function prototypes	7
	cuda_boundaries.cu Implementation of boundary condition functions with OpenMP threading	10
	cuda_discretization.cu Implementation of boundary condition functions with CUDA acceleration	12
	cuda_kernels.cuh Declaration of functions to execute on the GPU (CUDA kernels)	16
	discretization.h Declaration of discretized mathematical function prototypes	20
	main.c Implementation of semi-infinite diffusion equation	24
	mesh.c Implemenatation of mesh handling functions for diffusion benchmarks	25
	mesh.h Declaration of mesh function prototypes for diffusion benchmarks	29
	numerics.c Implementation of Laplacian operator and analytical solution functions	32
	numerics.h Declaration of Laplacian operator and analytical solution functions	37
	openacc_boundaries.c Implementation of boundary condition functions with OpenMP threading	44
	openacc_discretization.c Implementation of boundary condition functions with OpenACC threading	47

3 Class Documentation

openacc_kernels.h Declaration of functions to execute on the GPU (OpenACC kernels)	52
openmp_boundaries.c Implementation of boundary condition functions with OpenMP threading	55
openmp_discretization.c Implementation of boundary condition functions with OpenMP threading	57
output.c Implementation of file output functions for diffusion benchmarks	60
output.h Declaration of output function prototypes for diffusion benchmarks	63
serial_boundaries.c Implementation of boundary condition functions without threading	66
serial_discretization.c Implementation of boundary condition functions without threading	68
tbb_boundaries.cpp Implementation of boundary condition functions with TBB threading	71
tbb_discretization.cpp Implementation of boundary condition functions with TBB threading	72
timer.c High-resolution cross-platform machine time reader	75
timer.h Declaration of timer function prototypes for diffusion benchmarks	77
type.h Definition of scalar data type and Doxygen diffusion group	78

3 Class Documentation

3.1 Stopwatch Struct Reference

#include <type.h>

Public Attributes

- double conv
- double step
- double file
- double soln

3.1.1 Detailed Description

Container for timing data

Definition at line 41 of file type.h.

3.1.2 Member Data Documentation 3.1.2.1 conv double Stopwatch::conv Cumulative time executing compute_convolution() Definition at line 45 of file type.h. 3.1.2.2 file double Stopwatch::file Cumulative time executing write_csv() and write_png() Definition at line 55 of file type.h. 3.1.2.3 soln double Stopwatch::soln Cumulative time executing check_solution() Definition at line 60 of file type.h. 3.1.2.4 step double Stopwatch::step

The documentation for this struct was generated from the following file:

Cumulative time executing solve_diffusion_equation()

Definition at line 50 of file type.h.

type.h

4 File Documentation 5

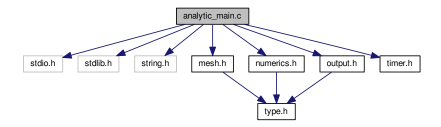
4 File Documentation

4.1 analytic_main.c File Reference

Analytical solution to semi-infinite diffusion equation.

```
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#include "mesh.h"
#include "numerics.h"
#include "output.h"
#include "timer.h"
```

Include dependency graph for analytic_main.c:



Functions

- void solve_diffusion_equation (fp_t **conc, int nx, int ny, int nm, fp_t dx, fp_t dy, fp_t D, fp_t dt, fp_t elapsed)

 Update the scalar composition field using analytical solution.
- int main (int argc, char *argv[])

Find analytical solution at intervals specified in the parameters file.

4.1.1 Detailed Description

Analytical solution to semi-infinite diffusion equation.

4.1.2 Function Documentation

4.1.2.1 main()

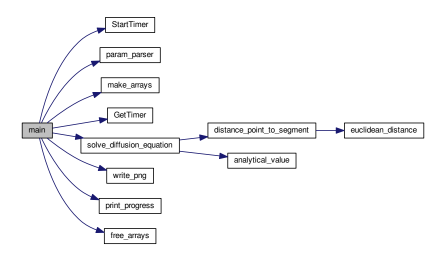
```
int main (
          int argc,
          char * argv[] )
```

Find analytical solution at intervals specified in the parameters file.

Program will write a series of PNG image files to visualize the scalar composition field, useful for qualitative verification of numerical results.

Definition at line 69 of file analytic_main.c.

Here is the call graph for this function:



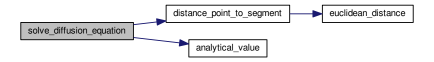
4.1.2.2 solve_diffusion_equation()

```
void solve_diffusion_equation (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t dx,
    fp_t dy,
    fp_t D,
    fp_t dt,
    fp_t elapsed )
```

Update the scalar composition field using analytical solution.

Definition at line 37 of file analytic_main.c.

Here is the call graph for this function:



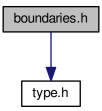
Here is the caller graph for this function:



4.2 boundaries.h File Reference

Declaration of boundary condition function prototypes.

#include "type.h"
Include dependency graph for boundaries.h:



This graph shows which files directly or indirectly include this file:



Functions

```
void set_boundaries (fp_t bc[2][2])
```

Set values to be used along the simulation domain boundaries.

• void apply_initial_conditions (fp_t **conc_old, int nx, int ny, int nm, fp_t bc[2][2])

Initialize flat composition field with fixed boundary conditions.

• void apply_boundary_conditions (fp_t **conc_old, int nx, int ny, int nm, fp_t bc[2][2]) Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

4.2.1 Detailed Description

Declaration of boundary condition function prototypes.

4.2.2 Function Documentation

4.2.2.1 apply_boundary_conditions()

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Definition at line 54 of file serial_boundaries.c.

Here is the call graph for this function:





4.2.2.2 apply_initial_conditions()

Initialize flat composition field with fixed boundary conditions.

The boundary conditions are fixed values of c_{hi} along the lower-left half and upper-right half walls, no flux everywhere else, with an initial values of c_{lo} everywhere. These conditions represent a carburizing process, with partial exposure (rather than the entire left and right walls) to produce an inhomogeneous workload and highlight numerical errors at the boundaries.

Definition at line 37 of file serial boundaries.c.

Here is the caller graph for this function:



4.2.2.3 set_boundaries()

Set values to be used along the simulation domain boundaries.

Indexing is row-major, i.e. A[y][x], so $bc = [[y_{lo}, y_{hi}], [x_{lo}, x_{hi}]].$

Definition at line 28 of file serial_boundaries.c.

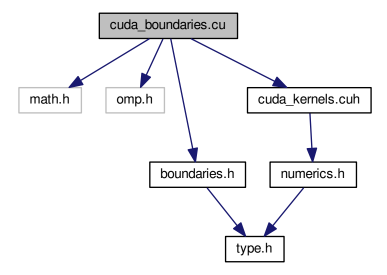


4.3 cuda_boundaries.cu File Reference

Implementation of boundary condition functions with OpenMP threading.

```
#include <math.h>
#include <omp.h>
#include "boundaries.h"
#include "cuda_kernels.cuh"
```

Include dependency graph for cuda_boundaries.cu:



Functions

- void set_boundaries (fp_t bc[2][2])
 - Set values to be used along the simulation domain boundaries.
- void apply_initial_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2])
 - Initialize flat composition field with fixed boundary conditions.
- void boundary_kernel (fp_t *conc, int nx, int ny, int nm)
 - Boundary condition kernel for execution on the GPU.
- $\bullet \ \ void\ apply_boundary_conditions\ (fp_t\ **conc,\ int\ nx,\ int\ ny,\ int\ nm,\ fp_t\ bc[2][2])\\$

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Variables

fp_t d_bc [2][2]

Boundary condition array on the GPU, allocated in protected memory.

4.3.1 Detailed Description

Implementation of boundary condition functions with OpenMP threading.

4.3.2 Function Documentation

4.3.2.1 apply_boundary_conditions()

```
void apply_boundary_conditions (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Definition at line 119 of file cuda boundaries.cu.

4.3.2.2 apply_initial_conditions()

Initialize flat composition field with fixed boundary conditions.

The boundary conditions are fixed values of c_{hi} along the lower-left half and upper-right half walls, no flux everywhere else, with an initial values of c_{lo} everywhere. These conditions represent a carburizing process, with partial exposure (rather than the entire left and right walls) to produce an inhomogeneous workload and highlight numerical errors at the boundaries.

Definition at line 45 of file cuda_boundaries.cu.

4.3.2.3 boundary_kernel()

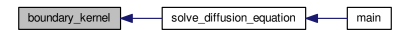
```
void boundary_kernel (
    fp_t * conc,
    int nx,
    int ny,
    int nm )
```

Boundary condition kernel for execution on the GPU.

This function accesses 1D data rather than the 2D array representation of the scalar composition field

Here is the caller graph for this function:

Definition at line 68 of file cuda_boundaries.cu.



4.3.2.4 set_boundaries()

Set values to be used along the simulation domain boundaries.

```
Indexing is row-major, i.e. A[y][x], so bc = [[y_{lo}, y_{hi}], [x_{lo}, x_{hi}]].
```

Definition at line 36 of file cuda boundaries.cu.

4.3.3 Variable Documentation

4.3.3.1 d_bc

```
fp_t d_bc[2][2]
```

Boundary condition array on the GPU, allocated in protected memory.

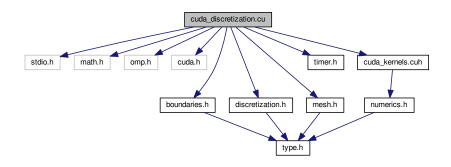
Definition at line 34 of file cuda_boundaries.cu.

4.4 cuda_discretization.cu File Reference

Implementation of boundary condition functions with CUDA acceleration.

```
#include <stdio.h>
#include <math.h>
#include <omp.h>
#include <cuda.h>
#include "boundaries.h"
#include "discretization.h"
#include "mesh.h"
#include "timer.h"
#include "cuda_kernels.cuh"
```

Include dependency graph for cuda_discretization.cu:



Functions

- void convolution_kernel (fp_t *conc_old, fp_t *conc_lap, int nx, int ny, int nm)
 Tiled convolution algorithm for execution on the GPU.
- void compute_convolution (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)

 Perform the convolution of the mask matrix with the composition matrix.
- void diffusion_kernel (fp_t *conc_old, fp_t *conc_new, fp_t *conc_lap, int nx, int ny, int nm, fp_t D, fp_t dt)

 Vector addition algorithm for execution on the GPU.
- void solve_diffusion_equation (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm, fp_t bc[2][2], fp_t D, fp_t dt, fp_t *elapsed, struct Stopwatch *sw, int checks)

Update the scalar composition field using old and Laplacian values.

void check_solution (fp_t **conc_new, fp_t **conc_lap, int nx, int ny, fp_t dx, fp_t dy, int nm, fp_t elapsed, fp_t D, fp_t bc[2][2], fp_t *rss)

Compare numerical and analytical solutions of the diffusion equation.

Variables

fp t d mask [MAX MASK W *MAX MASK H]

Convolution mask array on the GPU, allocated in protected memory.

4.4.1 Detailed Description

Implementation of boundary condition functions with CUDA acceleration.

4.4.2 Function Documentation

4.4.2.1 check_solution()

```
void check_solution (
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    fp_t dx,
    fp_t dy,
    int nm,
    fp_t elapsed,
    fp_t D,
    fp_t * rss )
```

Compare numerical and analytical solutions of the diffusion equation.

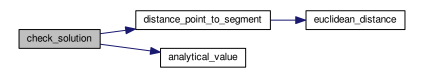
Returns

Residual sum of squares (RSS), normalized to the domain size.

Overwrites *conc_lap*, into which the point-wise RSS is written. Normalized RSS is then computed as the sum of the point-wise values.

Definition at line 215 of file cuda_discretization.cu.

Here is the call graph for this function:



4.4.2.2 compute_convolution()

```
void compute_convolution (
    fp_t ** conc_old,
    fp_t ** conc_lap,
    fp_t ** mask_lap,
    int nx,
    int ny,
    int nm )
```

Perform the convolution of the mask matrix with the composition matrix.

If the convolution mask is the Laplacian stencil, the convolution evaluates the discrete Laplacian of the composition field. Other masks are possible, for example the Sobel filters for edge detection. This function is general purpose: as long as the dimensions nx, ny, and nm are properly specified, the convolution will be correctly computed.

Definition at line 99 of file cuda_discretization.cu.

4.4.2.3 convolution_kernel()

Tiled convolution algorithm for execution on the GPU.

This function accesses 1D data rather than the 2D array representation of the scalar composition field, mapping into 2D tiles on the GPU with halo cells before computing the convolution.

Note:

- The source matrix (conc_old) and destination matrix (conc_lap) must be identical in size
- · One CUDA core operates on one array index: there is no nested loop over matrix elements
- The halo (nm/2 perimeter cells) in conc_lap are unallocated garbage
- The same cells in conc_old are boundary values, and contribute to the convolution
- · conc tile is the shared tile of input data, accessible by all threads in this block

Definition at line 41 of file cuda_discretization.cu.

4.4.2.4 diffusion_kernel()

Vector addition algorithm for execution on the GPU.

This function accesses 1D data rather than the 2D array representation of the scalar composition field

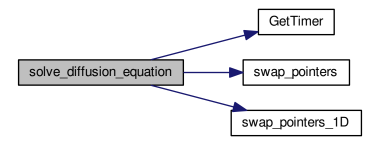
Definition at line 134 of file cuda_discretization.cu.

4.4.2.5 solve_diffusion_equation()

Update the scalar composition field using old and Laplacian values.

Definition at line 156 of file cuda_discretization.cu.

Here is the call graph for this function:



4.4.3 Variable Documentation

4.4.3.1 d_mask

fp_t d_mask[MAX_MASK_W *MAX_MASK_H]

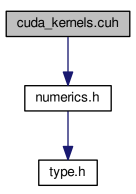
Convolution mask array on the GPU, allocated in protected memory.

Definition at line 39 of file cuda_discretization.cu.

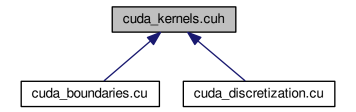
4.5 cuda_kernels.cuh File Reference

Declaration of functions to execute on the GPU (CUDA kernels)

#include "numerics.h"
Include dependency graph for cuda_kernels.cuh:



This graph shows which files directly or indirectly include this file:



Macros

• #define MAX TILE W 32

Maximum width of an input tile, including halo cells, for GPU memory allocation.

• #define MAX_TILE_H 32

Maximum height of an input tile, including halo cells, for GPU memory allocation.

Functions

- void boundary_kernel (fp_t *conc, int nx, int ny, int nm)
 - Boundary condition kernel for execution on the GPU.
- void convolution_kernel (fp_t *conc_old, fp_t *conc_lap, int nx, int ny, int nm)

Tiled convolution algorithm for execution on the GPU.

• void diffusion_kernel (fp_t *conc_old, fp_t *conc_new, fp_t *conc_lap, int nx, int ny, int nm, fp_t D, fp_t dt)

Vector addition algorithm for execution on the GPU.

Variables

fp_t d_mask [MAX_MASK_W *MAX_MASK_H]

Convolution mask array on the GPU, allocated in protected memory.

fp_t d_bc [2][2]

Boundary condition array on the GPU, allocated in protected memory.

4.5.1 Detailed Description

Declaration of functions to execute on the GPU (CUDA kernels)

4.5.2 Macro Definition Documentation

4.5.2.1 MAX_TILE_H

```
#define MAX_TILE_H 32
```

Maximum height of an input tile, including halo cells, for GPU memory allocation.

Definition at line 42 of file cuda_kernels.cuh.

4.5.2.2 MAX_TILE_W

```
#define MAX_TILE_W 32
```

Maximum width of an input tile, including halo cells, for GPU memory allocation.

Definition at line 37 of file cuda_kernels.cuh.

4.5.3 Function Documentation

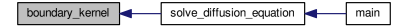
4.5.3.1 boundary_kernel()

```
void boundary_kernel (
    fp_t * conc,
    int nx,
    int ny,
    int nm )
```

Boundary condition kernel for execution on the GPU.

This function accesses 1D data rather than the 2D array representation of the scalar composition field

Definition at line 68 of file cuda_boundaries.cu.



4.5.3.2 convolution_kernel()

Tiled convolution algorithm for execution on the GPU.

This function accesses 1D data rather than the 2D array representation of the scalar composition field, mapping into 2D tiles on the GPU with halo cells before computing the convolution.

Note:

- The source matrix (conc_old) and destination matrix (conc_lap) must be identical in size
- One CUDA core operates on one array index: there is no nested loop over matrix elements
- The halo (nm/2 perimeter cells) in conc_lap are unallocated garbage
- The same cells in conc_old are boundary values, and contribute to the convolution
- · conc_tile is the shared tile of input data, accessible by all threads in this block

Definition at line 41 of file cuda_discretization.cu.

4.5.3.3 diffusion_kernel()

```
void diffusion_kernel (
    fp_t * conc_old,
    fp_t * conc_new,
    fp_t * conc_lap,
    int nx,
    int ny,
    int nm,
    fp_t D,
    fp_t dt )
```

Vector addition algorithm for execution on the GPU.

This function accesses 1D data rather than the 2D array representation of the scalar composition field

Definition at line 134 of file cuda_discretization.cu.

4.5.4 Variable Documentation

4.5.4.1 d_bc

```
fp_t d_bc[2][2]
```

Boundary condition array on the GPU, allocated in protected memory.

Definition at line 34 of file cuda_boundaries.cu.

4.5.4.2 d_mask

```
fp_t d_mask[MAX_MASK_W *MAX_MASK_H]
```

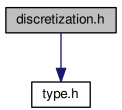
Convolution mask array on the GPU, allocated in protected memory.

Definition at line 39 of file cuda_discretization.cu.

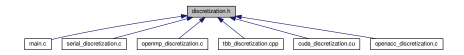
4.6 discretization.h File Reference

Declaration of discretized mathematical function prototypes.

```
#include "type.h"
Include dependency graph for discretization.h:
```



This graph shows which files directly or indirectly include this file:



Functions

- void compute_convolution (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)

 Perform the convolution of the mask matrix with the composition matrix.
- void solve_diffusion_equation (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm, fp_t bc[2][2], fp_t D, fp_t dt, fp_t *elapsed, struct Stopwatch *sw, int checks)

Update the scalar composition field using old and Laplacian values.

void check_solution (fp_t **conc_new, fp_t **conc_lap, int nx, int ny, fp_t dx, fp_t dy, int nm, fp_t elapsed, fp_t D, fp_t bc[2][2], fp_t *rss)

Compare numerical and analytical solutions of the diffusion equation.

4.6.1 Detailed Description

Declaration of discretized mathematical function prototypes.

4.6.2 Function Documentation

4.6.2.1 check_solution()

```
void check_solution (
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    fp_t dx,
    fp_t dy,
    int nm,
    fp_t elapsed,
    fp_t D,
    fp_t * rss )
```

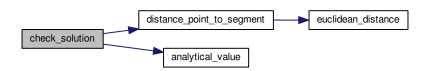
Compare numerical and analytical solutions of the diffusion equation.

Returns

Residual sum of squares (RSS), normalized to the domain size.

Overwrites *conc_lap*, into which the point-wise RSS is written. Normalized RSS is then computed as the sum of the point-wise values.

Definition at line 78 of file serial_discretization.c.



Here is the caller graph for this function:



4.6.2.2 compute_convolution()

Perform the convolution of the mask matrix with the composition matrix.

If the convolution mask is the Laplacian stencil, the convolution evaluates the discrete Laplacian of the composition field. Other masks are possible, for example the Sobel filters for edge detection. This function is general purpose: as long as the dimensions nx, ny, and nm are properly specified, the convolution will be correctly computed.

Definition at line 32 of file serial_discretization.c.

Here is the call graph for this function:



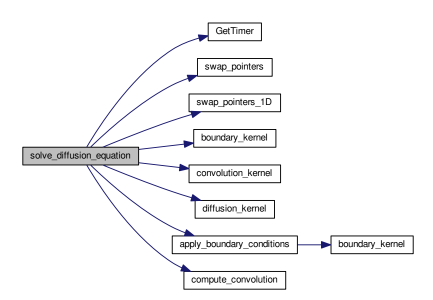


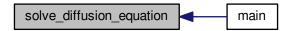
4.6.2.3 solve_diffusion_equation()

Update the scalar composition field using old and Laplacian values.

Definition at line 51 of file serial_discretization.c.

Here is the call graph for this function:



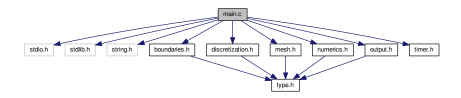


4.7 main.c File Reference

Implementation of semi-infinite diffusion equation.

```
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#include "boundaries.h"
#include "discretization.h"
#include "mesh.h"
#include "numerics.h"
#include "output.h"
#include "timer.h"
```

Include dependency graph for main.c:



Functions

• int main (int argc, char *argv[])

Run simulation using input parameters specified on the command line.

4.7.1 Detailed Description

Implementation of semi-infinite diffusion equation.

4.7.2 Function Documentation

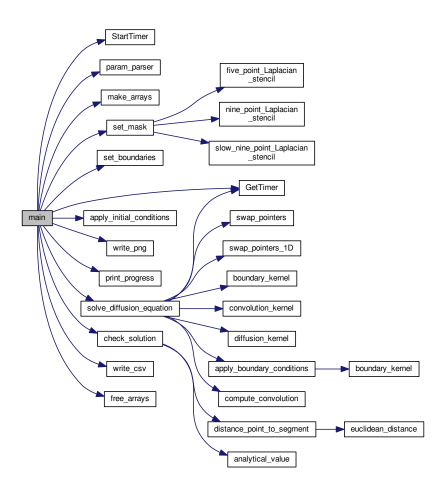
4.7.2.1 main()

Run simulation using input parameters specified on the command line.

Program will write a series of PNG image files to visualize scalar composition field, plus a final CSV raw data file and CSV runtime log tabulating the iteration counter (*iter*), elapsed simulation time (*sim_time*), system free energy (*energy*), error relative to analytical solution (*wrss*), time spent performing convolution (*conv_time*), time spent updating fields (*step_time*), time spent writing to disk (*IO_time*), time spent generating analytical values (*soln_time*), and total elapsed (*run_time*).

Definition at line 47 of file main.c.

Here is the call graph for this function:

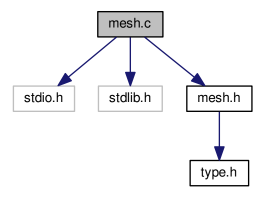


4.8 mesh.c File Reference

Implemenatation of mesh handling functions for diffusion benchmarks.

```
#include <stdio.h>
#include <stdlib.h>
#include "mesh.h"
```

Include dependency graph for mesh.c:



Functions

void make_arrays (fp_t ***conc_old, fp_t ***conc_new, fp_t ***conc_lap, fp_t ***mask_lap, int nx, int ny, int nm)

Allocate 2D arrays to store scalar composition values.

- void free_arrays (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap)

 Free dynamically allocated memory.
- void swap_pointers (fp_t ***conc_old, fp_t ***conc_new)

Swap pointers to 2D arrays.

 $\bullet \ \ void \ swap_pointers_1D \ (fp_t \ **conc_old, fp_t \ **conc_new)\\$

Swap pointers to data underlying 1D arrays.

4.8.1 Detailed Description

Implemenatation of mesh handling functions for diffusion benchmarks.

4.8.2 Function Documentation

4.8.2.1 free_arrays()

Free dynamically allocated memory.

4.8 mesh.c File Reference 27

Definition at line 58 of file mesh.c.

Here is the caller graph for this function:



4.8.2.2 make_arrays()

```
void make_arrays (
    fp_t *** conc_old,
    fp_t *** conc_new,
    fp_t *** conc_lap,
    fp_t *** mask_lap,
    int nx,
    int ny,
    int nm )
```

Allocate 2D arrays to store scalar composition values.

Arrays are allocated as 1D arrays, then 2D pointer arrays are mapped over the top. This facilitates use of either 1D or 2D data access, depending on whether the task is spatially dependent or not.

Definition at line 29 of file mesh.c.



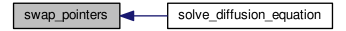
4.8.2.3 swap_pointers()

Swap pointers to 2D arrays.

Rather than copy data from $fp_t** conc_old$ into $fp_t** conc_new$, an expensive operation, simply trade the top-most pointers. New becomes old, old becomes new, with no data lost and in almost no time.

Definition at line 73 of file mesh.c.

Here is the caller graph for this function:

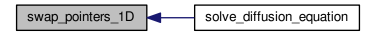


4.8.2.4 swap_pointers_1D()

Swap pointers to data underlying 1D arrays.

Rather than copy data from $fp_t*conc_old[0]$ into $fp_t*conc_new[0]$, an expensive operation, simply trade the top-most pointers. New becomes old, old becomes new, with no data lost and in almost no time.

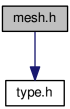
Definition at line 82 of file mesh.c.



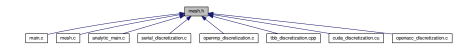
4.9 mesh.h File Reference

Declaration of mesh function prototypes for diffusion benchmarks.

#include "type.h"
Include dependency graph for mesh.h:



This graph shows which files directly or indirectly include this file:



Functions

void make_arrays (fp_t ***conc_old, fp_t ***conc_new, fp_t ***conc_lap, fp_t ***mask_lap, int nx, int ny, int nm)

Allocate 2D arrays to store scalar composition values.

 $\bullet \ \ \text{void free_arrays (fp_t } ** \text{conc_old, fp_t} ** \text{conc_new, fp_t} ** \text{conc_lap, fp_t} ** \text{mask_lap)}\\$

Free dynamically allocated memory.

void swap_pointers (fp_t ***conc_old, fp_t ***conc_new)

Swap pointers to 2D arrays.

void swap_pointers_1D (fp_t **conc_old, fp_t **conc_new)
 Swap pointers to data underlying 1D arrays.

4.9.1 Detailed Description

Declaration of mesh function prototypes for diffusion benchmarks.

4.9.2 Function Documentation

4.9.2.1 free_arrays()

Free dynamically allocated memory.

Definition at line 58 of file mesh.c.

Here is the caller graph for this function:



4.9.2.2 make_arrays()

Allocate 2D arrays to store scalar composition values.

Arrays are allocated as 1D arrays, then 2D pointer arrays are mapped over the top. This facilitates use of either 1D or 2D data access, depending on whether the task is spatially dependent or not.

Definition at line 29 of file mesh.c.



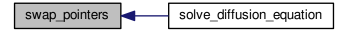
4.9.2.3 swap_pointers()

Swap pointers to 2D arrays.

Rather than copy data from $fp_t** conc_old$ into $fp_t** conc_new$, an expensive operation, simply trade the top-most pointers. New becomes old, old becomes new, with no data lost and in almost no time.

Definition at line 73 of file mesh.c.

Here is the caller graph for this function:



4.9.2.4 swap_pointers_1D()

Swap pointers to data underlying 1D arrays.

Rather than copy data from $fp_t*conc_old[0]$ into $fp_t*conc_new[0]$, an expensive operation, simply trade the top-most pointers. New becomes old, old becomes new, with no data lost and in almost no time.

Definition at line 82 of file mesh.c.

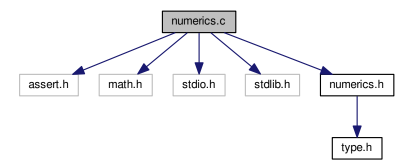


4.10 numerics.c File Reference

Implementation of Laplacian operator and analytical solution functions.

```
#include <assert.h>
#include <math.h>
#include <stdio.h>
#include <stdlib.h>
#include "numerics.h"
```

Include dependency graph for numerics.c:



Functions

- void set_mask (fp_t dx, fp_t dy, int code, fp_t **mask_lap, int nm)
 - Specify which stencil (mask) to use for the Laplacian (convolution)
- void five_point_Laplacian_stencil (fp_t dx, fp_t dy, fp_t **mask_lap, int nm)

Write 5-point Laplacian stencil into convolution mask.

• void nine_point_Laplacian_stencil (fp_t dx, fp_t dy, fp_t **mask_lap, int nm)

Write 9-point Laplacian stencil into convolution mask.

• void slow_nine_point_Laplacian_stencil (fp_t dx, fp_t dy, fp_t **mask_lap, int nm)

Write 9-point Laplacian stencil into convolution mask.

- fp_t euclidean_distance (fp_t ax, fp_t ay, fp_t bx, fp_t by)
 - Compute Euclidean distance between two points, a and b.
- fp_t manhattan_distance (fp_t ax, fp_t ay, fp_t bx, fp_t by)

Compute Manhattan distance between two points, a and b.

fp_t distance_point_to_segment (fp_t ax, fp_t ay, fp_t bx, fp_t by, fp_t px, fp_t py)

Compute minimum distance from point p to a line segment bounded by points a and b.

• void analytical_value (fp_t x, fp_t t, fp_t D, fp_t bc[2][2], fp_t *c)

Analytical solution of the diffusion equation for a carburizing process.

4.10.1 Detailed Description

Implementation of Laplacian operator and analytical solution functions.

4.10.2 Function Documentation

4.10.2.1 analytical_value()

Analytical solution of the diffusion equation for a carburizing process.

For 1D diffusion through a semi-infinite domain with initial and far-field composition c_{∞} and boundary value $c(x=0,t)=c_0$ with constant diffusivity D, the solution to Fick's second law is

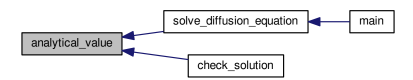
$$c(x,t) = c_0 - (c_0 - c_\infty) \operatorname{erf}\left(\frac{x}{\sqrt{4Dt}}\right)$$

which reduces, when $c_{\infty}=0$, to

$$c(x,t) = c_0 \left[1 - \operatorname{erf}\left(\frac{x}{\sqrt{4Dt}}\right) \right].$$

Definition at line 121 of file numerics.c.

Here is the caller graph for this function:



4.10.2.2 distance_point_to_segment()

Compute minimum distance from point p to a line segment bounded by points a and b.

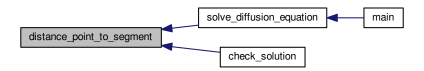
This function computes the projection of p onto ab, limiting the projected range to [0, 1] to handle projections that fall outside of ab. Implemented after Grumdrig on Stackoverflow, https://stackoverflow.com/a/1501725.

Definition at line 108 of file numerics.c.

Here is the call graph for this function:



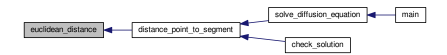
Here is the caller graph for this function:



4.10.2.3 euclidean_distance()

Compute Euclidean distance between two points, a and b.

Definition at line 98 of file numerics.c.



4.10.2.4 five_point_Laplacian_stencil()

```
void five_point_Laplacian_stencil (
    fp_t dx,
    fp_t dy,
    fp_t ** mask_lap,
    int nm )
```

Write 5-point Laplacian stencil into convolution mask.

```
3 \times 3 mask, 5 values, truncation error \mathcal{O}(\Delta x^2)
```

Definition at line 51 of file numerics.c.

Here is the caller graph for this function:



4.10.2.5 manhattan_distance()

Compute Manhattan distance between two points, a and b.

Definition at line 103 of file numerics.c.

4.10.2.6 nine_point_Laplacian_stencil()

```
void nine_point_Laplacian_stencil (
    fp_t dx,
    fp_t dy,
    fp_t ** mask_lap,
    int nm )
```

Write 9-point Laplacian stencil into convolution mask.

 3×3 mask, 9 values, truncation error $\mathcal{O}(\Delta x^4)$

Definition at line 62 of file numerics.c.

Here is the caller graph for this function:



4.10.2.7 set_mask()

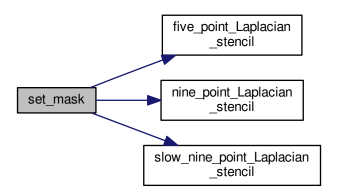
Specify which stencil (mask) to use for the Laplacian (convolution)

The mask corresponding to the numerical code will be applied. The suggested encoding is mask width as the ones digit and value count as the tens digit, *e.g.* 53 specifies five_point_Laplacian_stencil(), while 93 specifies nine_point_Laplacian_stencil().

To add your own mask (stencil), add a case to this function with your chosen numerical encoding, then specify that code in the input parameters file (params.txt by default). Note that, for a Laplacian stencil, the sum of the coefficients must equal zero and *nm* must be an odd integer.

If your stencil is larger than 5×5 , you must increase the values defined by MAX_MASK_W and MAX_MASK_H.

Definition at line 31 of file numerics.c.



Here is the caller graph for this function:



4.10.2.8 slow_nine_point_Laplacian_stencil()

Write 9-point Laplacian stencil into convolution mask.

```
5 \times 5 mask, 9 values, truncation error \mathcal{O}(\Delta x^4)
```

Provided for testing and demonstration of scalability, only: as the name indicates, this 9-point stencil is computationally more expensive than the 3×3 version. If your code requires $\mathcal{O}(\Delta x^4)$ accuracy, please use nine_point_ \leftarrow Laplacian_stencil().

Definition at line 79 of file numerics.c.

Here is the caller graph for this function:

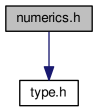


4.11 numerics.h File Reference

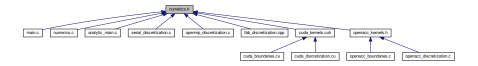
Declaration of Laplacian operator and analytical solution functions.

#include "type.h"

Include dependency graph for numerics.h:



This graph shows which files directly or indirectly include this file:



Macros

• #define MAX_MASK_W 5

Maximum width of the convolution mask (Laplacian stencil) array.

• #define MAX_MASK_H 5

Maximum height of the convolution mask (Laplacian stencil) array.

Functions

void set_mask (fp_t dx, fp_t dy, int code, fp_t **mask_lap, int nm)

Specify which stencil (mask) to use for the Laplacian (convolution)

void five_point_Laplacian_stencil (fp_t dx, fp_t dy, fp_t **mask_lap, int nm)

Write 5-point Laplacian stencil into convolution mask.

• void nine_point_Laplacian_stencil (fp_t dx, fp_t dy, fp_t **mask_lap, int nm)

Write 9-point Laplacian stencil into convolution mask.

• void slow_nine_point_Laplacian_stencil (fp_t dx, fp_t dy, fp_t **mask_lap, int nm)

Write 9-point Laplacian stencil into convolution mask.

fp_t euclidean_distance (fp_t ax, fp_t ay, fp_t bx, fp_t by)

Compute Euclidean distance between two points, a and b.

• fp_t manhattan_distance (fp_t ax, fp_t ay, fp_t bx, fp_t by)

Compute Manhattan distance between two points, a and b.

• fp_t distance_point_to_segment (fp_t ax, fp_t ay, fp_t bx, fp_t by, fp_t px, fp_t py)

Compute minimum distance from point p to a line segment bounded by points a and b.

void analytical_value (fp_t x, fp_t t, fp_t D, fp_t bc[2][2], fp_t *c)

Analytical solution of the diffusion equation for a carburizing process.

4.11.1 Detailed Description

Declaration of Laplacian operator and analytical solution functions.

4.11.2 Macro Definition Documentation

4.11.2.1 MAX MASK H

```
#define MAX_MASK_H 5
```

Maximum height of the convolution mask (Laplacian stencil) array.

Definition at line 40 of file numerics.h.

4.11.2.2 MAX MASK W

```
#define MAX_MASK_W 5
```

Maximum width of the convolution mask (Laplacian stencil) array.

Definition at line 35 of file numerics.h.

4.11.3 Function Documentation

4.11.3.1 analytical_value()

Analytical solution of the diffusion equation for a carburizing process.

For 1D diffusion through a semi-infinite domain with initial and far-field composition c_{∞} and boundary value $c(x=0,t)=c_0$ with constant diffusivity D, the solution to Fick's second law is

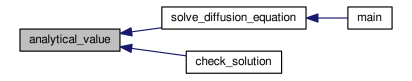
$$c(x,t) = c_0 - (c_0 - c_\infty) \operatorname{erf}\left(\frac{x}{\sqrt{4Dt}}\right)$$

which reduces, when $c_{\infty}=0$, to

$$c(x,t) = c_0 \left[1 - \operatorname{erf}\left(\frac{x}{\sqrt{4Dt}}\right) \right].$$

Definition at line 121 of file numerics.c.

Here is the caller graph for this function:



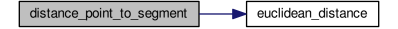
4.11.3.2 distance_point_to_segment()

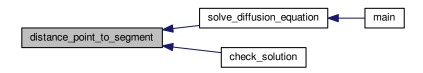
Compute minimum distance from point p to a line segment bounded by points a and b.

This function computes the projection of p onto ab, limiting the projected range to [0, 1] to handle projections that fall outside of ab. Implemented after Grumdrig on Stackoverflow, https://stackoverflow.com/a/1501725.

Definition at line 108 of file numerics.c.

Here is the call graph for this function:



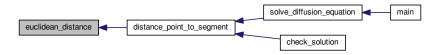


4.11.3.3 euclidean_distance()

Compute Euclidean distance between two points, a and b.

Definition at line 98 of file numerics.c.

Here is the caller graph for this function:



4.11.3.4 five_point_Laplacian_stencil()

Write 5-point Laplacian stencil into convolution mask.

 3×3 mask, 5 values, truncation error $\mathcal{O}(\Delta x^2)$

Definition at line 51 of file numerics.c.



4.11.3.5 manhattan_distance()

Compute Manhattan distance between two points, a and b.

Definition at line 103 of file numerics.c.

4.11.3.6 nine_point_Laplacian_stencil()

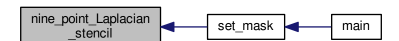
```
void nine_point_Laplacian_stencil (
    fp_t dx,
    fp_t dy,
    fp_t ** mask_lap,
    int nm )
```

Write 9-point Laplacian stencil into convolution mask.

```
3 \times 3 mask, 9 values, truncation error \mathcal{O}(\Delta x^4)
```

Definition at line 62 of file numerics.c.

Here is the caller graph for this function:



4.11.3.7 set_mask()

Specify which stencil (mask) to use for the Laplacian (convolution)

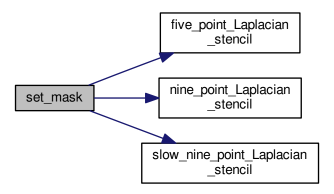
The mask corresponding to the numerical code will be applied. The suggested encoding is mask width as the ones digit and value count as the tens digit, *e.g.* 53 specifies five_point_Laplacian_stencil(), while 93 specifies nine_point_Laplacian_stencil().

To add your own mask (stencil), add a case to this function with your chosen numerical encoding, then specify that code in the input parameters file (params.txt by default). Note that, for a Laplacian stencil, the sum of the coefficients must equal zero and *nm* must be an odd integer.

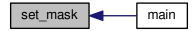
If your stencil is larger than 5×5 , you must increase the values defined by MAX_MASK_W and MAX_MASK_H.

Definition at line 31 of file numerics.c.

Here is the call graph for this function:



Here is the caller graph for this function:



4.11.3.8 slow_nine_point_Laplacian_stencil()

Write 9-point Laplacian stencil into convolution mask.

```
5 \times 5 mask, 9 values, truncation error \mathcal{O}(\Delta x^4)
```

Provided for testing and demonstration of scalability, only: as the name indicates, this 9-point stencil is computationally more expensive than the 3×3 version. If your code requires $\mathcal{O}(\Delta x^4)$ accuracy, please use nine_point_ \leftarrow Laplacian stencil().

Definition at line 79 of file numerics.c.

Here is the caller graph for this function:

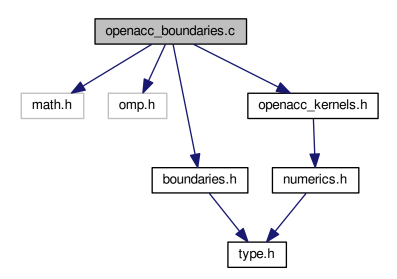


4.12 openacc_boundaries.c File Reference

Implementation of boundary condition functions with OpenMP threading.

```
#include <math.h>
#include <omp.h>
#include "boundaries.h"
#include "openacc_kernels.h"
```

Include dependency graph for openacc_boundaries.c:



Functions

- void set_boundaries (fp_t bc[2][2])
 - Set values to be used along the simulation domain boundaries.
- void apply_initial_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2])

Initialize flat composition field with fixed boundary conditions.

- void boundary_kernel (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2])

 Boundary condition kernel for execution on the GPU.
- $\bullet \ \ void\ apply_boundary_conditions\ (fp_t\ **conc,\ int\ nx,\ int\ ny,\ int\ nm,\ fp_t\ bc[2][2])$

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

4.12.1 Detailed Description

Implementation of boundary condition functions with OpenMP threading.

4.12.2 Function Documentation

4.12.2.1 apply_boundary_conditions()

```
void apply_boundary_conditions (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Definition at line 108 of file openacc_boundaries.c.

Here is the call graph for this function:





4.12.2.2 apply_initial_conditions()

```
void apply_initial_conditions (
    fp_t ** conc_old,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Initialize flat composition field with fixed boundary conditions.

The boundary conditions are fixed values of c_{hi} along the lower-left half and upper-right half walls, no flux everywhere else, with an initial values of c_{lo} everywhere. These conditions represent a carburizing process, with partial exposure (rather than the entire left and right walls) to produce an inhomogeneous workload and highlight numerical errors at the boundaries.

Definition at line 39 of file openacc_boundaries.c.

Here is the caller graph for this function:



4.12.2.3 boundary_kernel()

```
void boundary_kernel (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Boundary condition kernel for execution on the GPU.

Definition at line 62 of file openacc_boundaries.c.



4.12.2.4 set_boundaries()

Set values to be used along the simulation domain boundaries.

```
Indexing is row-major, i.e. A[y][x], so bc = [[y_{lo}, y_{hi}], [x_{lo}, x_{hi}]].
```

Definition at line 30 of file openacc_boundaries.c.

Here is the caller graph for this function:

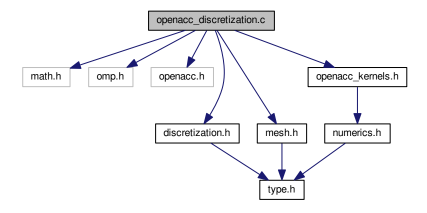


4.13 openacc_discretization.c File Reference

Implementation of boundary condition functions with OpenACC threading.

```
#include <math.h>
#include <omp.h>
#include <openacc.h>
#include "discretization.h"
#include "mesh.h"
#include "openacc_kernels.h"
```

Include dependency graph for openacc_discretization.c:



Functions

void convolution_kernel (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)
 Tiled convolution algorithm for execution on the GPU.

void diffusion_kernel (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, int nx, int ny, int nm, fp_t D, fp_t dt)

Vector addition algorithm for execution on the GPU.

- void compute_convolution (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)

 Perform the convolution of the mask matrix with the composition matrix.
- void solve_diffusion_equation (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm, fp_t bc[2][2], fp_t D, fp_t dt, fp_t *elapsed, struct Stopwatch *sw, int checks)

Update the scalar composition field using old and Laplacian values.

void check_solution (fp_t **conc_new, fp_t **conc_lap, int nx, int ny, fp_t dx, fp_t dy, int nm, fp_t elapsed, fp_t D, fp_t bc[2][2], fp_t *rss)

Compare numerical and analytical solutions of the diffusion equation.

4.13.1 Detailed Description

Implementation of boundary condition functions with OpenACC threading.

4.13.2 Function Documentation

4.13.2.1 check_solution()

```
void check_solution (
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    fp_t dx,
    fp_t dy,
    int nm,
    fp_t elapsed,
    fp_t D,
    fp_t bc[2][2],
    fp_t * rss )
```

Compare numerical and analytical solutions of the diffusion equation.

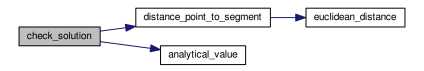
Returns

Residual sum of squares (RSS), normalized to the domain size.

Overwrites *conc_lap*, into which the point-wise RSS is written. Normalized RSS is then computed as the sum of the point-wise values.

Definition at line 106 of file openacc_discretization.c.

Here is the call graph for this function:



Here is the caller graph for this function:



4.13.2.2 compute_convolution()

Perform the convolution of the mask matrix with the composition matrix.

If the convolution mask is the Laplacian stencil, the convolution evaluates the discrete Laplacian of the composition field. Other masks are possible, for example the Sobel filters for edge detection. This function is general purpose: as long as the dimensions nx, ny, and nm are properly specified, the convolution will be correctly computed.

Definition at line 68 of file openacc_discretization.c.

Here is the call graph for this function:



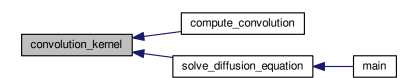
4.13.2.3 convolution_kernel()

```
void convolution_kernel (
    fp_t ** conc_old,
    fp_t ** conc_lap,
    fp_t ** mask_lap,
    int nx,
    int ny,
    int nm )
```

Tiled convolution algorithm for execution on the GPU.

Definition at line 32 of file openacc_discretization.c.

Here is the caller graph for this function:



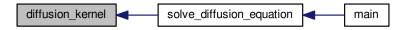
4.13.2.4 diffusion_kernel()

```
void diffusion_kernel (
    fp_t ** conc_old,
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    int nm,
    fp_t D,
    fp_t dt )
```

Vector addition algorithm for execution on the GPU.

Definition at line 52 of file openacc_discretization.c.

Here is the caller graph for this function:

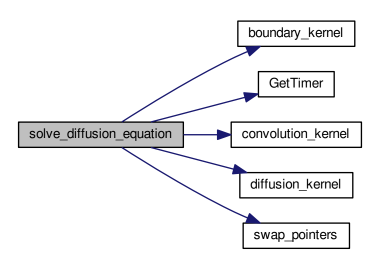


4.13.2.5 solve_diffusion_equation()

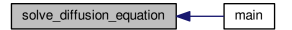
```
void solve_diffusion_equation (
    fp_t ** conc_old,
    fp_t ** conc_new,
    fp_t ** conc_lap,
    fp_t ** mask_lap,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2],
    fp_t dt,
    fp_t * elapsed,
    struct Stopwatch * sw,
    int checks )
```

Update the scalar composition field using old and Laplacian values.

Definition at line 78 of file openacc_discretization.c.



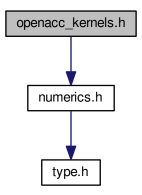
Here is the caller graph for this function:



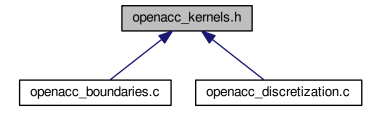
4.14 openacc_kernels.h File Reference

Declaration of functions to execute on the GPU (OpenACC kernels)

#include "numerics.h"
Include dependency graph for openacc_kernels.h:



This graph shows which files directly or indirectly include this file:



Functions

```
    void boundary_kernel (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2])
```

Boundary condition kernel for execution on the GPU.

void convolution_kernel (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)
 Tiled convolution algorithm for execution on the GPU.

void diffusion_kernel (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, int nx, int ny, int nm, fp_t D, fp_t dt)

Vector addition algorithm for execution on the GPU.

4.14.1 Detailed Description

Declaration of functions to execute on the GPU (OpenACC kernels)

4.14.2 Function Documentation

4.14.2.1 boundary_kernel()

Boundary condition kernel for execution on the GPU.

Definition at line 62 of file openacc_boundaries.c.



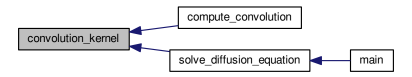
4.14.2.2 convolution_kernel()

```
void convolution_kernel (
    fp_t ** conc_old,
    fp_t ** conc_lap,
    fp_t ** mask_lap,
    int nx,
    int ny,
    int nm )
```

Tiled convolution algorithm for execution on the GPU.

Definition at line 32 of file openacc_discretization.c.

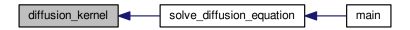
Here is the caller graph for this function:



4.14.2.3 diffusion_kernel()

Vector addition algorithm for execution on the GPU.

Definition at line 52 of file openacc_discretization.c.

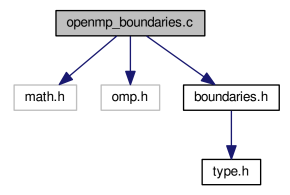


4.15 openmp_boundaries.c File Reference

Implementation of boundary condition functions with OpenMP threading.

```
#include <math.h>
#include <omp.h>
#include "boundaries.h"
```

Include dependency graph for openmp_boundaries.c:



Functions

- void set_boundaries (fp_t bc[2][2])
 - Set values to be used along the simulation domain boundaries.
- void apply_initial_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2])

 Initialize flat composition field with fixed boundary conditions.
- void apply_boundary_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2]) Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

4.15.1 Detailed Description

Implementation of boundary condition functions with OpenMP threading.

4.15.2 Function Documentation

4.15.2.1 apply_boundary_conditions()

```
void apply_boundary_conditions (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Definition at line 62 of file openmp_boundaries.c.

4.15.2.2 apply_initial_conditions()

```
void apply_initial_conditions (
    fp_t ** conc_old,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Initialize flat composition field with fixed boundary conditions.

The boundary conditions are fixed values of c_{hi} along the lower-left half and upper-right half walls, no flux everywhere else, with an initial values of c_{lo} everywhere. These conditions represent a carburizing process, with partial exposure (rather than the entire left and right walls) to produce an inhomogeneous workload and highlight numerical errors at the boundaries.

Definition at line 39 of file openmp boundaries.c.

4.15.2.3 set_boundaries()

Set values to be used along the simulation domain boundaries.

Indexing is row-major, i.e. A[y][x], so $bc = [[y_{lo}, y_{hi}], [x_{lo}, x_{hi}]]$.

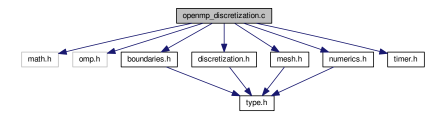
Definition at line 30 of file openmp_boundaries.c.

4.16 openmp_discretization.c File Reference

Implementation of boundary condition functions with OpenMP threading.

```
#include <math.h>
#include <omp.h>
#include "boundaries.h"
#include "discretization.h"
#include "mesh.h"
#include "numerics.h"
#include "timer.h"
```

Include dependency graph for openmp_discretization.c:



Functions

- void compute_convolution (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)

 Perform the convolution of the mask matrix with the composition matrix.
- void solve_diffusion_equation (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm, fp_t bc[2][2], fp_t D, fp_t dt, fp_t *elapsed, struct Stopwatch *sw, int checks)

Update the scalar composition field using old and Laplacian values.

void check_solution (fp_t **conc_new, fp_t **conc_lap, int nx, int ny, fp_t dx, fp_t dy, int nm, fp_t elapsed, fp_t D, fp_t bc[2][2], fp_t *rss)

Compare numerical and analytical solutions of the diffusion equation.

4.16.1 Detailed Description

Implementation of boundary condition functions with OpenMP threading.

4.16.2 Function Documentation

4.16.2.1 check_solution()

```
void check_solution (
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    fp_t dx,
    fp_t dy,
    int nm,
    fp_t elapsed,
    fp_t D,
    fp_t * rss )
```

Compare numerical and analytical solutions of the diffusion equation.

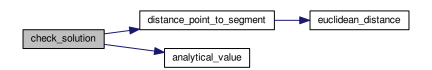
Returns

Residual sum of squares (RSS), normalized to the domain size.

Overwrites *conc_lap*, into which the point-wise RSS is written. Normalized RSS is then computed as the sum of the point-wise values.

Definition at line 84 of file openmp_discretization.c.

Here is the call graph for this function:



4.16.2.2 compute_convolution()

Perform the convolution of the mask matrix with the composition matrix.

If the convolution mask is the Laplacian stencil, the convolution evaluates the discrete Laplacian of the composition field. Other masks are possible, for example the Sobel filters for edge detection. This function is general purpose: as long as the dimensions nx, ny, and nm are properly specified, the convolution will be correctly computed.

Definition at line 33 of file openmp_discretization.c.

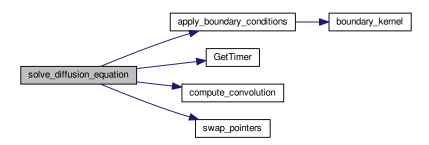
Here is the caller graph for this function:



4.16.2.3 solve_diffusion_equation()

Update the scalar composition field using old and Laplacian values.

Definition at line 56 of file openmp_discretization.c.

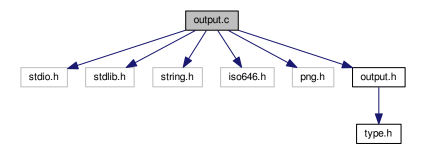


4.17 output.c File Reference

Implementation of file output functions for diffusion benchmarks.

```
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#include <iso646.h>
#include <png.h>
#include "output.h"
```

Include dependency graph for output.c:



Functions

• void param_parser (int argc, char *argv[], int *nx, int *ny, int *nm, int *code, fp_t *dx, fp_t *dy, fp_t *D, fp_t *linStab, int *steps, int *checks)

Read parameters from file specified on the command line.

• void print_progress (const int step, const int steps)

Prints timestamps and a 20-point progress bar to stdout.

void write_csv (fp_t **conc, int nx, int ny, fp_t dx, fp_t dy, int step)

Writes scalar composition field to diffusion.??????csv.

void write_png (fp_t **conc, int nx, int ny, int step)

Writes scalar composition field to diffusion.???????.png.

4.17.1 Detailed Description

Implementation of file output functions for diffusion benchmarks.

4.17.2 Function Documentation

4.17.2.1 param_parser()

```
void param_parser (
    int argc,
    char * argv[],
    int * nx,
    int * ny,
    int * nm,
    int * code,
    fp_t * dx,
    fp_t * D,
    fp_t * linStab,
    int * steps,
    int * checks )
```

Read parameters from file specified on the command line.

Definition at line 32 of file output.c.

Here is the caller graph for this function:



4.17.2.2 print_progress()

Prints timestamps and a 20-point progress bar to stdout.

Call inside the timestepping loop, near the top, e.g.

```
for (int step=0; step<steps; step++) {
    print_progress(step, steps);
    take_a_step();
    elapsed += dt;
}</pre>
```

Definition at line 123 of file output.c.

Here is the caller graph for this function:



4.17.2.3 write_csv()

Writes scalar composition field to diffusion.??????csv.

Definition at line 148 of file output.c.

Here is the caller graph for this function:



4.17.2.4 write_png()

Writes scalar composition field to diffusion.??????ngg.

Definition at line 182 of file output.c.

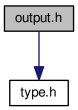
Here is the caller graph for this function:



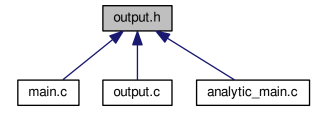
4.18 output.h File Reference

Declaration of output function prototypes for diffusion benchmarks.

#include "type.h"
Include dependency graph for output.h:



This graph shows which files directly or indirectly include this file:



Functions

void param_parser (int argc, char *argv[], int *nx, int *ny, int *nm, int *code, fp_t *dx, fp_t *dy, fp_t *D, fp_t *linStab, int *steps, int *checks)

Read parameters from file specified on the command line.

• void print_progress (const int step, const int steps)

Prints timestamps and a 20-point progress bar to stdout.

void write_csv (fp_t **conc, int nx, int ny, fp_t dx, fp_t dy, int step)

Writes scalar composition field to diffusion.???????csv.

void write_png (fp_t **conc, int nx, int ny, int step)

Writes scalar composition field to diffusion.??????ng.

4.18.1 Detailed Description

Declaration of output function prototypes for diffusion benchmarks.

4.18.2 Function Documentation

4.18.2.1 param_parser()

```
void param_parser (
    int argc,
    char * argv[],
    int * nx,
    int * ny,
    int * code,
    fp_t * dx,
    fp_t * D,
    fp_t * linStab,
    int * steps,
    int * checks )
```

Read parameters from file specified on the command line.

Definition at line 32 of file output.c.



4.18.2.2 print_progress()

Prints timestamps and a 20-point progress bar to stdout.

Call inside the timestepping loop, near the top, e.g.

```
for (int step=0; step<steps; step++) {
    print_progress(step, steps);
    take_a_step();
    elapsed += dt;
}</pre>
```

Definition at line 123 of file output.c.

Here is the caller graph for this function:



4.18.2.3 write_csv()

Writes scalar composition field to diffusion.??????csv.

Definition at line 148 of file output.c.



4.18.2.4 write_png()

Writes scalar composition field to diffusion.??????ng.

Definition at line 182 of file output.c.

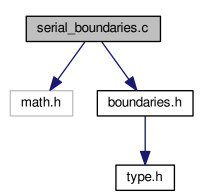
Here is the caller graph for this function:



4.19 serial_boundaries.c File Reference

Implementation of boundary condition functions without threading.

```
#include <math.h>
#include "boundaries.h"
Include dependency graph for serial_boundaries.c:
```



Functions

• void set_boundaries (fp_t bc[2][2])

Set values to be used along the simulation domain boundaries.

void apply_initial_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2])

Initialize flat composition field with fixed boundary conditions.

```
• void apply_boundary_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2]) Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.
```

4.19.1 Detailed Description

Implementation of boundary condition functions without threading.

4.19.2 Function Documentation

4.19.2.1 apply_boundary_conditions()

```
void apply_boundary_conditions (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Definition at line 54 of file serial boundaries.c.

4.19.2.2 apply_initial_conditions()

Initialize flat composition field with fixed boundary conditions.

The boundary conditions are fixed values of c_{hi} along the lower-left half and upper-right half walls, no flux everywhere else, with an initial values of c_{lo} everywhere. These conditions represent a carburizing process, with partial exposure (rather than the entire left and right walls) to produce an inhomogeneous workload and highlight numerical errors at the boundaries.

Definition at line 37 of file serial_boundaries.c.

4.19.2.3 set_boundaries()

Set values to be used along the simulation domain boundaries.

Indexing is row-major, i.e. A[y][x], so $bc = [[y_{lo}, y_{hi}], [x_{lo}, x_{hi}]]$.

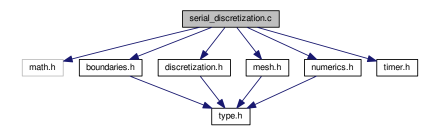
Definition at line 28 of file serial boundaries.c.

4.20 serial discretization.c File Reference

Implementation of boundary condition functions without threading.

```
#include <math.h>
#include "boundaries.h"
#include "discretization.h"
#include "mesh.h"
#include "numerics.h"
#include "timer.h"
```

Include dependency graph for serial_discretization.c:



Functions

- void compute_convolution (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)

 Perform the convolution of the mask matrix with the composition matrix.
- void solve_diffusion_equation (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm, fp_t bc[2][2], fp_t D, fp_t dt, fp_t *elapsed, struct Stopwatch *sw, int checks)

Update the scalar composition field using old and Laplacian values.

void check_solution (fp_t **conc_new, fp_t **conc_lap, int nx, int ny, fp_t dx, fp_t dy, int nm, fp_t elapsed, fp_t D, fp_t bc[2][2], fp_t *rss)

Compare numerical and analytical solutions of the diffusion equation.

4.20.1 Detailed Description

Implementation of boundary condition functions without threading.

4.20.2 Function Documentation

4.20.2.1 check_solution()

```
void check_solution (
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    fp_t dx,
    fp_t dy,
    int nm,
    fp_t elapsed,
    fp_t D,
    fp_t * rss )
```

Compare numerical and analytical solutions of the diffusion equation.

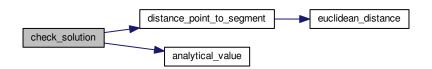
Returns

Residual sum of squares (RSS), normalized to the domain size.

Overwrites *conc_lap*, into which the point-wise RSS is written. Normalized RSS is then computed as the sum of the point-wise values.

Definition at line 78 of file serial_discretization.c.

Here is the call graph for this function:



4.20.2.2 compute_convolution()

Perform the convolution of the mask matrix with the composition matrix.

If the convolution mask is the Laplacian stencil, the convolution evaluates the discrete Laplacian of the composition field. Other masks are possible, for example the Sobel filters for edge detection. This function is general purpose: as long as the dimensions nx, ny, and nm are properly specified, the convolution will be correctly computed.

Definition at line 32 of file serial_discretization.c.

Here is the caller graph for this function:

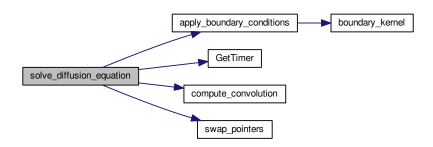


4.20.2.3 solve_diffusion_equation()

Update the scalar composition field using old and Laplacian values.

Definition at line 51 of file serial_discretization.c.

Here is the call graph for this function:

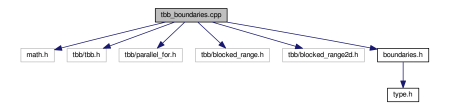


4.21 tbb_boundaries.cpp File Reference

Implementation of boundary condition functions with TBB threading.

```
#include <math.h>
#include <tbb/tbb.h>
#include <tbb/parallel_for.h>
#include <tbb/blocked_range.h>
#include <tbb/blocked_range2d.h>
#include "boundaries.h"
```

Include dependency graph for tbb_boundaries.cpp:



Functions

• void set_boundaries (fp_t bc[2][2])

Set values to be used along the simulation domain boundaries.

- $\bullet \ \ void\ apply_initial_conditions\ (fp_t\ **conc,\ int\ nx,\ int\ ny,\ int\ nm,\ fp_t\ bc[2][2])$
 - Initialize flat composition field with fixed boundary conditions.
- void apply_boundary_conditions (fp_t **conc, int nx, int ny, int nm, fp_t bc[2][2]) Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

4.21.1 Detailed Description

Implementation of boundary condition functions with TBB threading.

4.21.2 Function Documentation

4.21.2.1 apply_boundary_conditions()

```
void apply_boundary_conditions (
    fp_t ** conc,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Set fixed value (c_{hi}) along left and bottom, zero-flux elsewhere.

Definition at line 77 of file tbb_boundaries.cpp.

4.21.2.2 apply_initial_conditions()

```
void apply_initial_conditions (
    fp_t ** conc_old,
    int nx,
    int ny,
    int nm,
    fp_t bc[2][2] )
```

Initialize flat composition field with fixed boundary conditions.

The boundary conditions are fixed values of c_{hi} along the lower-left half and upper-right half walls, no flux everywhere else, with an initial values of c_{lo} everywhere. These conditions represent a carburizing process, with partial exposure (rather than the entire left and right walls) to produce an inhomogeneous workload and highlight numerical errors at the boundaries.

Definition at line 41 of file tbb boundaries.cpp.

4.21.2.3 set_boundaries()

Set values to be used along the simulation domain boundaries.

Indexing is row-major, i.e. A[y][x], so $bc = [[y_{lo}, y_{hi}], [x_{lo}, x_{hi}]]$.

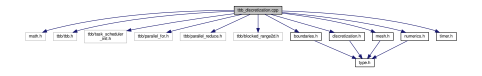
Definition at line 32 of file tbb_boundaries.cpp.

4.22 tbb_discretization.cpp File Reference

Implementation of boundary condition functions with TBB threading.

```
#include <math.h>
#include <tbb/tbb.h>
#include <tbb/task_scheduler_init.h>
#include <tbb/parallel_for.h>
#include <tbb/parallel_reduce.h>
#include <tbb/blocked_range2d.h>
#include "boundaries.h"
#include "discretization.h"
#include "mesh.h"
#include "numerics.h"
#include "timer.h"
```

Include dependency graph for tbb_discretization.cpp:



Functions

- void compute_convolution (fp_t **conc_old, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm)

 Perform the convolution of the mask matrix with the composition matrix.
- void solve_diffusion_equation (fp_t **conc_old, fp_t **conc_new, fp_t **conc_lap, fp_t **mask_lap, int nx, int ny, int nm, fp_t bc[2][2], fp_t D, fp_t dt, fp_t *elapsed, struct Stopwatch *sw, int checks)

Update the scalar composition field using old and Laplacian values.

void check_solution (fp_t **conc_new, fp_t **conc_lap, int nx, int ny, fp_t dx, fp_t dy, int nm, fp_t elapsed, fp_t D, fp_t bc[2][2], fp_t *rss)

Compare numerical and analytical solutions of the diffusion equation.

4.22.1 Detailed Description

Implementation of boundary condition functions with TBB threading.

4.22.2 Function Documentation

4.22.2.1 check_solution()

```
void check_solution (
    fp_t ** conc_new,
    fp_t ** conc_lap,
    int nx,
    int ny,
    fp_t dx,
    fp_t dy,
    int nm,
    fp_t elapsed,
    fp_t D,
    fp_t bc[2][2],
    fp_t * rss )
```

Compare numerical and analytical solutions of the diffusion equation.

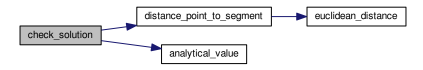
Returns

Residual sum of squares (RSS), normalized to the domain size.

Overwrites *conc_lap*, into which the point-wise RSS is written. Normalized RSS is then computed as the sum of the point-wise values.

Definition at line 91 of file tbb_discretization.cpp.

Here is the call graph for this function:



4.22.2.2 compute_convolution()

Perform the convolution of the mask matrix with the composition matrix.

If the convolution mask is the Laplacian stencil, the convolution evaluates the discrete Laplacian of the composition field. Other masks are possible, for example the Sobel filters for edge detection. This function is general purpose: as long as the dimensions nx, ny, and nm are properly specified, the convolution will be correctly computed.

Definition at line 37 of file tbb_discretization.cpp.

Here is the caller graph for this function:

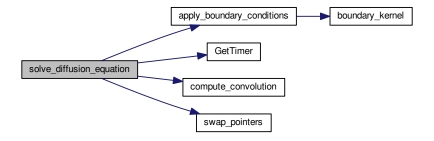


4.22.2.3 solve_diffusion_equation()

Update the scalar composition field using old and Laplacian values.

Definition at line 58 of file tbb_discretization.cpp.

Here is the call graph for this function:

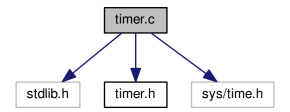


4.23 timer.c File Reference

High-resolution cross-platform machine time reader.

```
#include <stdlib.h>
#include "timer.h"
#include <sys/time.h>
```

Include dependency graph for timer.c:



Functions

• void StartTimer ()

Set CPU frequency and begin timing.

• double GetTimer ()

Return elapsed time in seconds.

Variables

• struct timeval timerStart

4.23.1 Detailed Description

High-resolution cross-platform machine time reader.

Author

NVIDIA

4.23.2 Function Documentation

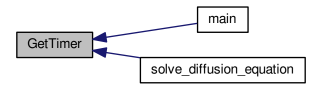
4.23.2.1 GetTimer()

```
double GetTimer ( )
```

Return elapsed time in seconds.

Definition at line 64 of file timer.c.

Here is the caller graph for this function:



4.23.2.2 StartTimer()

```
void StartTimer ( )
```

Set CPU frequency and begin timing.

Definition at line 48 of file timer.c.

Here is the caller graph for this function:



4.23.3 Variable Documentation

4.23.3.1 timerStart

struct timeval timerStart

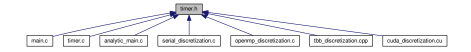
Platform-dependent data type of hardware time value

Definition at line 45 of file timer.c.

4.24 timer.h File Reference

Declaration of timer function prototypes for diffusion benchmarks.

This graph shows which files directly or indirectly include this file:



Functions

- void StartTimer ()
 - Set CPU frequency and begin timing.
- double GetTimer ()

Return elapsed time in seconds.

4.24.1 Detailed Description

Declaration of timer function prototypes for diffusion benchmarks.

4.24.2 Function Documentation

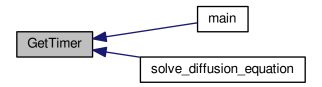
4.24.2.1 GetTimer()

```
double GetTimer ( )
```

Return elapsed time in seconds.

Definition at line 64 of file timer.c.

Here is the caller graph for this function:



4.24.2.2 StartTimer()

```
void StartTimer ( )
```

Set CPU frequency and begin timing.

Definition at line 48 of file timer.c.

Here is the caller graph for this function:



4.25 type.h File Reference

Definition of scalar data type and Doxygen diffusion group.

This graph shows which files directly or indirectly include this file:



Classes

• struct Stopwatch

Typedefs

typedef double fp_t

4.25.1 Detailed Description

Definition of scalar data type and Doxygen diffusion group.

4.25.2 Typedef Documentation

4.25.2.1 fp_t

typedef double fp_t

Specify the basic data type to achieve the desired accuracy in floating-point arithmetic: float for single-precision, double for double-precision. This choice propagates throughout the code, and may significantly affect runtime on GPU hardware.

Definition at line 36 of file type.h.

Index

analytic_main.c, 5	d_bc, 12
main, 5	set_boundaries, 11
solve_diffusion_equation, 6	cuda_discretization.cu, 12
analytical_value	check_solution, 13
numerics.c, 33	compute_convolution, 14
numerics.h, 39	convolution_kernel, 14
apply_boundary_conditions	d_mask, 16
boundaries.h, 8	diffusion_kernel, 15
cuda_boundaries.cu, 11	solve_diffusion_equation, 15
openacc_boundaries.c, 45	cuda_kernels.cuh, 16
openmp_boundaries.c, 55	boundary_kernel, 18
serial_boundaries.c, 67	convolution_kernel, 18
tbb_boundaries.cpp, 71	d_bc, 19
apply_initial_conditions	d_mask, 20
boundaries.h, 8	diffusion_kernel, 19
cuda_boundaries.cu, 11	MAX_TILE_H, 17
openacc_boundaries.c, 45	MAX TILE W, 18
openmp boundaries.c, 56	:
serial_boundaries.c, 67	d_bc
tbb boundaries.cpp, 71	cuda_boundaries.cu, 12
_ 1117	cuda_kernels.cuh, 19
boundaries.h, 7	d_mask
apply_boundary_conditions, 8	cuda_discretization.cu, 16
apply_initial_conditions, 8	cuda_kernels.cuh, 20
set_boundaries, 9	diffusion_kernel
boundary_kernel	cuda_discretization.cu, 15
cuda_boundaries.cu, 11	cuda_kernels.cuh, 19
cuda_kernels.cuh, 18	openacc_discretization.c, 50
openacc_boundaries.c, 46	openacc_kernels.h, 54
openacc_kernels.h, 53	discretization.h, 20
. –	check_solution, 21
check_solution	compute_convolution, 22
cuda_discretization.cu, 13	solve_diffusion_equation, 22
discretization.h, 21	distance_point_to_segment
openacc_discretization.c, 48	numerics.c, 33
openmp_discretization.c, 57	numerics.h, 40
serial_discretization.c, 69	
tbb_discretization.cpp, 73	euclidean_distance
compute_convolution	numerics.c, 34
cuda_discretization.cu, 14	numerics.h, 40
discretization.h, 22	file
openacc_discretization.c, 49	Stopwatch, 4
openmp_discretization.c, 58	five_point_Laplacian_stencil
serial_discretization.c, 69	numerics.c, 34
tbb_discretization.cpp, 73	
conv	numerics.h, 41
Stopwatch, 4	fp_t
convolution_kernel	type.h, 79
cuda_discretization.cu, 14	free_arrays mesh.c, 26
cuda_kernels.cuh, 18	
openacc_discretization.c, 50	mesh.h, 29
openacc_kernels.h, 53	GetTimer
cuda_boundaries.cu, 10	timer.c, 76
apply_boundary_conditions, 11	timer.h, 77
apply_initial_conditions, 11	, • •
boundary_kernel, 11	MAX_MASK_H

82 INDEX

numerics.h, 39	check_solution, 48
MAX MASK W	compute_convolution, 49
numerics.h, 39	convolution_kernel, 50
MAX TILE H	diffusion_kernel, 50
cuda kernels.cuh, 17	solve_diffusion_equation, 51
MAX TILE W	openacc_kernels.h, 52
cuda_kernels.cuh, 18	boundary_kernel, 53
main	convolution_kernel, 53
analytic_main.c, 5	diffusion_kernel, 54
main.c, 24	openmp_boundaries.c, 55
main.c, 24	apply_boundary_conditions, 55
main, 24	apply_bothdary_conditions, 56
	set_boundaries, 56
make_arrays	
mesh.c, 27 mesh.h, 30	openmp_discretization.c, 57
	check_solution, 57
manhattan_distance	compute_convolution, 58
numerics.c, 35	solve_diffusion_equation, 59
numerics.h, 41	output.c, 60
mesh.c, 25	param_parser, 60
free_arrays, 26	print_progress, 61
make_arrays, 27	write_csv, 62
swap_pointers, 27	write_png, 62
swap_pointers_1D, 28	output.h, 63
mesh.h, 29	param_parser, 64
free_arrays, 29	print_progress, 64
make_arrays, 30	write_csv, 65
swap_pointers, 30	write_png, 65
swap_pointers_1D, 31	
	param_parser
nine_point_Laplacian_stencil	output.c, 60
numerics.c, 35	output.h, 64
numerics.h, 42	print_progress
numerics.c, 32	output.c, 61
analytical_value, 33	output.h, 64
distance_point_to_segment, 33	and havedous a 00
euclidean_distance, 34	serial_boundaries.c, 66
five_point_Laplacian_stencil, 34	apply_boundary_conditions, 67
manhattan_distance, 35	apply_initial_conditions, 67
nine_point_Laplacian_stencil, 35	set_boundaries, 67
set_mask, 36	serial_discretization.c, 68
slow_nine_point_Laplacian_stencil, 37	check_solution, 69
numerics.h, 37	compute_convolution, 69
analytical_value, 39	solve_diffusion_equation, 70
distance_point_to_segment, 40	set_boundaries
euclidean_distance, 40	boundaries.h, 9
five_point_Laplacian_stencil, 41	cuda_boundaries.cu, 11
MAX_MASK_H, 39	openacc_boundaries.c, 46
MAX_MASK_W, 39	openmp_boundaries.c, 56
manhattan_distance, 41	serial_boundaries.c, 67
nine_point_Laplacian_stencil, 42	tbb_boundaries.cpp, 72
set_mask, 42	set_mask
slow_nine_point_Laplacian_stencil, 43	numerics.c, 36
, ,	numerics.h, 42
openacc_boundaries.c, 44	slow_nine_point_Laplacian_stencil
apply_boundary_conditions, 45	numerics.c, 37
apply_initial_conditions, 45	numerics.h, 43
boundary_kernel, 46	soln
set_boundaries, 46	Stopwatch, 4
openacc_discretization.c, 47	solve_diffusion_equation

INDEX 83

```
analytic_main.c, 6
     cuda_discretization.cu, 15
     discretization.h, 22
     openacc_discretization.c, 51
     openmp_discretization.c, 59
     serial discretization.c, 70
     tbb_discretization.cpp, 74
StartTimer
     timer.c, 76
     timer.h, 78
step
     Stopwatch, 4
Stopwatch, 3
     conv, 4
     file, 4
     soln, 4
     step, 4
swap_pointers
     mesh.c, 27
     mesh.h, 30
swap_pointers_1D
     mesh.c, 28
     mesh.h, 31
tbb_boundaries.cpp, 71
     apply_boundary_conditions, 71
     apply_initial_conditions, 71
     set_boundaries, 72
tbb_discretization.cpp, 72
     check_solution, 73
     compute convolution, 73
     solve_diffusion_equation, 74
timer.c, 75
     GetTimer, 76
     StartTimer, 76
     timerStart, 77
timer.h, 77
     GetTimer, 77
     StartTimer, 78
timerStart
     timer.c, 77
type.h, 78
     fp_t, 79
write_csv
     output.c, 62
     output.h, 65
write_png
     output.c, 62
     output.h, 65
```