# OP2 User's Manual (phase 1)

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#### 1 Introduction

OP2 is an API with associated libraries and preprocessors to generate parallel executables for applications on unstructured grids. The initial API is for C, but FORTRAN 77 will also be supported.

The key concept behind OP2 is that unstructured grids can be described by a number of sets. Depending on the application, these sets might be of nodes, edges, triangular faces, quadrilateral faces, cells of a variety of types, far-field boundary nodes, wall boundary faces, etc. Associated with these sets are both data (e.g. coordinate data at nodes) and mappings to other sets (e.g. edge mapping to the two nodes at each end of the edge). All of the numerically-intensive operations can then be described as a loop over all members of a set, carrying out some operations on data associated directly with the set or with another set through a mapping.

OP2 makes the important restriction that the order in which the function is applied to the members of the set must not affect the final result. This allows the parallel implementation to choose its own ordering to achieve maximum parallel efficiency. Two other restrictions are that the sets and maps are static (i.e. they do not change) and the operands in the set operations are not referenced through a double level of mapping indirection (i.e. through a mapping to another set which in turn uses another mapping to data in a third set).

OP2 currently enables users to write a single program which can be built into three different executables for different platforms:

- single-threaded on a CPU
- parallelised using CUDA for NVIDIA GPUs
- multi-threaded using OpenMP for multicore x86 systems

In the longer-term there will be support for AVX vectorisation for x86 CPUs, and OpenCL for both CPUs and GPUS.

There will also be support for distributed-memory MPI parallelisation in combination with any of the above. This will require parallel file I/O and so there will be routines to handle file I/O for the main datasets, as well as routines to handle terminal I/O.

#### 2 Overview

A computational project can be viewed as involving three steps:

- writing the program
- debugging the program, often using a small testcase
- running the program on increasingly large applications

With OP2 we want to simplify the first two tasks, while providing as much performance as possible for the third one.

To achieve the high performance for large applications, a preprocessor is needed to generate the CUDA code for GPUs or OpenMP code for multicore x86 systems. However, to keep the initial development simple, the single-threaded executable does not use any special tools; the user's main code is simply linked to a set of library routines, most of which do little more than error-checking to assist the debugging process by checking the correctness of the user's program. Note that this single-threaded version will not execute efficiently. The preprocessor is needed to generate efficient OpenMP code for x86 systems.

Figure 1 shows the build process for a single thread CPU executable. The user's main program (in this case jac.cpp) uses the OP header file op\_seq.h and is linked to the OP routines in op\_seq.c using g++, perhaps controlled by a Makefile.

Figure 2 shows the build process for the corresponding CUDA executable. The preprocessor parses the user's main program and produces a modified main program and a CUDA file which includes a separate file for each of the kernel functions. These are then compiled and linked to the OP routines in op\_lib.cu using g++ and the NVIDIA CUDA compiler nvcc, again perhaps controlled by a Makefile. As well as the header file op\_seq.h which is included by the user's main code jac.cpp, there is a header file op\_datatypes.h which is included by all of the files in the CUDA implementation, and by op\_seq.h.

Figure 3 shows the OpenMP build process which is very similar to the CUDA process except that it uses \*.cpp files produced by the preprocessor instead of \*.cu files.

In looking at the API specification, users may think it is a little verbose in places. e.g. users have to re-supply information about the datatype of the datasets being used in a parallel loop. This is a deliberate choice to simplify the task of the preprocessor, and therefore hopefully reduce the chance for errors. It is also motivated by the thought that "programming is easy; it's debugging which is difficult". i.e. writing code isn't time-consuming, it's correcting it which takes the time. Therefore, it's not unreasonable to ask the programmer to supply redundant information, but be assured that the preprocessor or library will check that all redundant information is self-consistent. If you declare a dataset as being of type OP\_DOUBLE and later say that it is of type OP\_FLOAT this will be flagged up as an error at run-time, both in the single-threaded library and in the CUDA library.

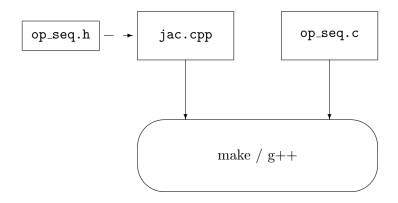


Figure 1: Sequential code build process

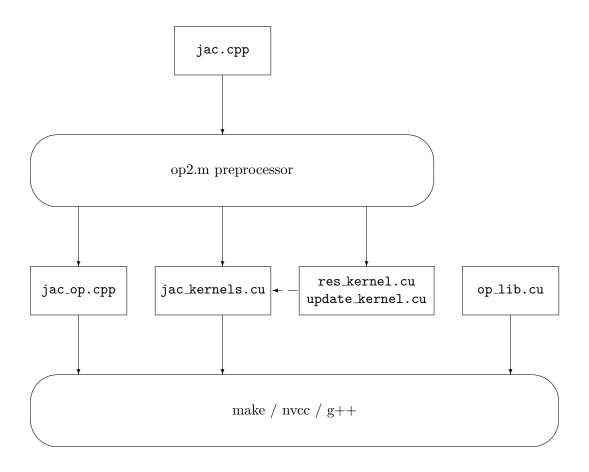


Figure 2: CUDA code build process

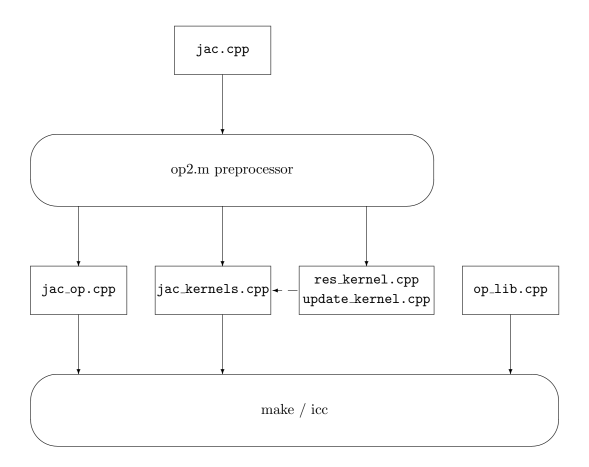


Figure 3: OpenMP code build process

#### 3 Initialisation and termination routines

#### op\_init(int argc, char \*\*argv)

This routine must be called before all other OP routines.

#### op\_decl\_set(int size, op\_set \*set, char \*name)

This routine declares information about a set.

size number of elements in the set

set output OP set ID

name a name used for output diagnostics

# op\_decl\_map(op\_set from, op\_set to, int dim, int \*imap, op\_map \*map, char \*name)

This routine declares information about a mapping from one set to another.

from set pointed from

to set pointed to

dim number of mappings per element

imap input mapping table

map output OP mapping ID

name a name used for output diagnostics

#### op\_decl\_const(int dim, char \*type, T \*dat, char \*name)

This routine declares constant data with global scope to be used in user's kernel functions. Note: in sequential version, it is the user's responsibility to define the appropriate global variable.

dim dimension of data (i.e. array size)

at present this must be a literal constant (i.e. a number not a variable); this restriction will be removed in the future but a literal constant will

remain more efficient

type datatype, either intrinsic ("float", "double", "int", "uint", "ll", "ull" or

"bool") or user-defined

dat input data of type T (checked for consistency with type at run-time)

name global name to be used in user's kernel functions:

a scalar variable if dim=1, otherwise an array of size dim

# op\_decl\_dat(op\_set set, int dim, char \*type, T \*dat, op\_dat \*data, char \*name)

This routine declares information about data associated with a set.

set set

dim dimension of dataset (number of items per set element)

at present this must be a literal constant (i.e. a number not a variable); this restriction will be removed in the future but a literal constant will

remain more efficient

type datatype, either intrinsic or user-defined

dat input data of type T (checked for consistency with type at run-time)

dat output OP dataset ID

name a name used for output diagnostics

#### op\_fetch\_data(op\_dat dat)

This routine transfers data from the GPU back to the CPU.

dat OP dataset ID – data is put back into original input array

#### op\_diagnostic\_output()

This routine prints out various useful bits of diagnostic info about sets, mappings and datasets

#### op\_exit()

This routine must be called last to cleanly terminate the OP computation.

# 4 Parallel execution routine

As an example, the parallel loop syntax when the user's kernel function has 3 arguments, with the third being a local constant or global reduction array, is:

op\_par\_loop\_3(void (\*kernel)(T0 \*, T1 \*, T2 \*), char \*name, op\_set set, op\_dat arg0, int idx0, op\_map map0, int dim0, char \*typ0, op\_access acc0, op\_dat arg1, int idx1, op\_map map1, int dim1, char \*typ1, op\_access acc1, T2 \*arg2, int idx2, op\_map map2, int dim2, char \*typ2, op\_access acc2)

kernel	user's kernel function with 3 arguments of arbitrary type (this is only used for the single-threaded CPU build)
name	name of kernel function, used for output diagnostics
set	OP set ID, giving set over which the parallel computation is performed
arg	OP dataset ID, or pointer to constant or global reduction array
idx	index of mapping to be used (-1 $\equiv$ no mapping indirection)
map	OP mapping ID (OP_ID for identity mapping, i.e. no mapping indirection, OP_GBL for constant or global reduction array)
dim	dataset dimension (redundant info, checked at run-time for consistency)
	at present this must be a literal constant (i.e. a number not a variable); this restriction will be removed in the future but a literal constant will remain more efficient
typ	dataset datatype (redundant info, checked at run-time for consistency)
acc	access type:  OP_READ: read-only  OP_WRITE: write-only, but without potential data conflict  OP_RW: read and write, but without potential data conflict  OP_INC: increment, or global reduction to compute a sum  OP_MAX: global reduction to compute a maximum
	OP_MIN: global reduction to compute a minimum

In this example, kernel is a function with 3 arguments of arbitrary type which performs a calculation for a single set element. This will get converted by a preprocessor into a routine called by the CUDA kernel function. The preprocessor will also take the specification of the arguments and turn this into the CUDA kernel function which loads in indirect data (i.e. data addressed indirectly through a mapping) from the device main memory into the

shared storage, then calls the converted kernel function for each element for each line in the above specification. Indirect data is incremented in shared memory (with thread coloring to avoid possible data conflicts) before being updated at the end of the CUDA kernel call.

The restriction that OP\_WRITE and OP\_RW access must not have any potential data conflict means that two different elements of the set cannot through a mapping indirection reference the same elements of the dataset.

Furthermore, with OP\_WRITE the user's kernel function must set the value of all DIM components of the dataset. If the user's kernel function does not set all of them, the access should be specified to be OP\_RW since the kernel function needs to read in the old values of the components which are not being modified.

Different numbers of arguments are handled similarly by routines with names of the form op\_par\_loop\_n where n is the number of arguments. Each argument can be either a dataset or a local constant or global reduction array, following the syntax shown above.

## 5 User-defined datatypes

If the user defines a new datatype mytype then this must be included in a header file along with

• a type-checking routine:

```
inline int type_error(const mytype *,const char *type)
{return strcmp(type, "mytype");}
```

which is used at run-time to check the consistency of the user's type declarations in input arguments.

• a "zero element" declaration of the form:

```
#define ZERO_mytype 0;
```

as well as an appropriate overloaded addition operator if there is any OP\_INC access to the datatype. The zero element and overloaded addition have to be such that 0 + x = x where x represents any element of the user's datatype and 0 represents the declared zero element.

• an overloaded implementation of the inequality operators < and > if there are any OP\_MIN, OP\_MAX accesses to the datatype.

In addition, the user must specify the name of the new header file using the environment variable OP\_USER\_DATATYPES so that this header file is included into the OP2 header file op\_datatypes.h.

# 6 Preprocessor

The prototype preprocessor has been written in MATLAB. It is run by the command op2('main')

where main.cpp is the user's main program. It produces as output a modified main program main\_op.cpp, and a new CUDA file main\_kernels.cu which includes one or more files of the form xxx\_kernel.cu containing the CUDA implementations of the user's kernel functions.

If the user's application is split over several files it is run by a command such as op2('main','sub1','sub2','sub3')

where sub1.cpp, sub2.cpp, sub3.cpp are the additional input files which will lead to the generation of output files sub1\_op.cpp, sub2\_op.cpp, sub3\_op.cpp in addition to main\_op.cpp, main\_kernels.cu and the individual kernel files.

The preprocessor cannot currently handle cases in which the same user kernel is used in more than one parallel loop, or when global constant data is set/updated in more than one place within the code. This will be addressed in the future.

## 7 Future changes

There will be a new function **op\_partition** which will re-number all of the elements in each set, to maximise the data reuse within each mini-partition. This is likely to use the same partitioning algorithm which will be employed for the higher-level distributed-memory partitioning for the MPI implementation in phase 2.

## 8 Phase 2 proposal

As explained in the introduction, phase 2 of the OP2 project will handle distributed-memory parallelisation using MPI. Because this links into other work by Leigh Lapworth and others at Rolls-Royce, discussions have begun about how this will be handled within OP2, and this has led to the following proposal.

My starting point is that we anticipate dealing with extremely large datasets and so we need to support parallel file I/O. There also seems to be general agreement that HDF5 has become the *de facto* standard underlying file format, with various other standards like CGNS layered on top.

Originally, my idea was to modify the OP2 set, mapping and dataset declarations so that these were read in by OP2 from a specified HDF5 file using specified keywords. Thus the OP2 library would have been entirely responsible for the parallel file I/O.

However, my new proposal is to adopt a layered approach:

- a minor extension to the existing API, leaving the parallel file I/O to the developer
- an example implementation of the parallel file I/O for HDF files, which some developers may choose to use unaltered, and others may modify to suit their needs

The rationale for this is to allow developers to make the tradeoff between ease-of-use and flexibility. Some will want maximum ease-of-use and are prepared to pay the price of working with HDF5 files with the flat keyword-based hierarchy which we will assume. Others will want the flexibility to manage their data storage in the way they wish, and will accept the additional programming effort this will entail.

In an MPI application, multiple copies of the same program are executed as separate processes, often on different nodes of a compute cluster. Hence, the OP2 declarations will be invoked on each process. The extensions to the existing API are as follows:

- op\_decl\_set: size is the number of elements of the set which will be provided by this MPI process
- op\_decl\_map: imap provides the part of the mapping table which corresponds to its share of the from set
- op\_decl\_dat: dat provides the data which corresponds to its share of set

For example, if an application has 4 processes,  $4 \times 10^6$  nodes and  $16 \times 10^6$  edges, then each process might be responsible for providing  $10^6$  nodes and  $4 \times 10^6$  edges. Process 0 (the one with MPI rank 0) would be responsible for providing the first  $10^6$  nodes, process 1 the next  $10^6$  nodes, and so on, and the same for the edges.

The edge  $\rightarrow$  node mapping tables would still contain the same information as in a single process implementation, but process 0 would provide the first  $4 \times 10^6$  entries, process 1 the next  $4 \times 10^6$  entries, and so on.

This is effectively using a simple contiguous block partitioning of the datasets, but it is very important to note that this will not be used for the parallel computation. OP2 will repartition the datasets (in parallel, probably using parmetis), will re-number the mapping tables as needed (as well as constructing import/export lists for halo data exchange) and will move all data/mappings/datasets to the correct MPI process.

The second layer would look similar to the existing API:

- op\_decl\_set\_hdf5: similar to op\_decl\_set but with size replaced by file which defines the HDF5 file from which size is read using keyword name
- op\_decl\_map\_hdf5: similar to op\_decl\_map but with imap replaced by file from which the mapping table is read using keyword name
- op\_decl\_dat\_hdf5: similar to op\_decl\_dat but with dat replaced by file from which the data is read using keyword name