





Introduction to OpenACC

Directive Based GPU Programming Vasileios Karakasis, CSCS May 14–15, 2018

Overview

- Day 1 (Understand the basic concepts of OpenACC)
 - Execution and memory model
 - Basic directives
 - Profiling and debugging
 - Hands-on sessions
- Day 2 (Advanced topics)
 - Asynchronous execution and wait gueues
 - Interoperability with CUDA and MPI
 - Deep copy
 - Hands-on sessions



What is OpenACC?

- Collection of compiler directives for specifying loops and regions to be offloaded from a host CPU to an attached accelerator device
- Host + Accelerator programming model
- High-level representation
- Current specification version: 2.6





Why to use OpenACC?

Because

- I don't care about all the little hardware details, I want my science done.
- I want to run on accelerators, but I still need a fast and readable code.
- I need portability across different accelerator vendors, but also be able to run on the multicore.
- I inhereted a large legacy monolithic codebase, which I don't dare to refactor completely, but I need to get my results faster.
- My code is in Fortran.





OpenACC is not a silver bullet

- A high-level representation is not a panacea.
 - You still need to understand and adapt to the programming model.
- Does not substitute hand-tuning, but can serve as a very good starting point.
- User base not yet as large as of classic OpenMP for multicores, but it is expanding.
 - You may run into compiler bugs or specification ambiguities.





Format of directives

- C/C++
 - #pragma acc directive-name [clause-list] new-line
 - Scope is the following block of code
- Fortran
 - !\$acc directive-name [clause-list] new-line
 - Scope is until !\$acc end directive-name



Programming model

- Host-directed execution
- Compute intensive regions are offloaded to attached accelerator devices
- Host orchestrates the execution on the device
 - Allocations on the device
 - Data transfers
 - Kernel launches
 - Wait for events
 - Etc...





- The device executes parallel or kernel regions
- Parallel region
 - Work-sharing loops
- Kernel region
 - Multiple loops to be executed as multiple kernels
- Levels of parallelim
 - 1. Gang
 - 2. Worker
 - 3. Vector
 - Parallelism levels are decided by the compiler but can be fine-tuned by the user



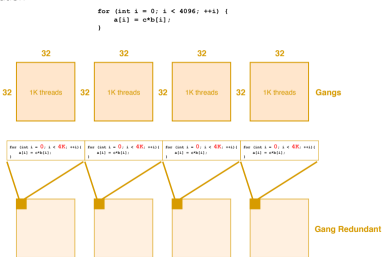
- The device executes parallel or kernel regions
- Parallel region
 - Work-sharing loops
- Kernel region
 - Multiple loops to be executed as multiple kernels
- Levels of parallelim
 - 1. Gang → CUDA block
 - 2. Worker → CUDA warp
 - 3. Vector → CUDA threads
 - Parallelism levels are decided by the compiler but can be fine-tuned by the user
 - Mapping to CUDA blocks/warps/threads is implementation defined



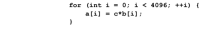


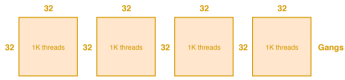
- Gang
 - Gang-redundant (GR)
 - Gang-partioned (GP)
- Worker
 - Worker-single (WS)
 - Worker-partitioned (WP)
- Vector
 - Vector-single (VS)
 - Vector-partitioned (VP)

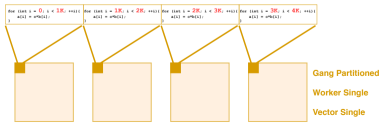






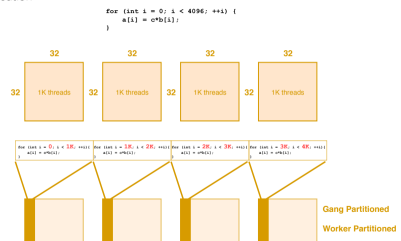






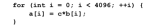


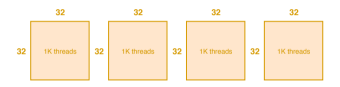
Modes of execution

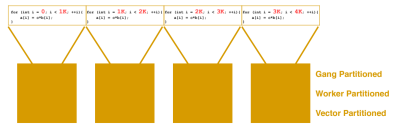




Vector Single









!\$acc end kernels

The kernels construct

Multiple loops inside kernels construct !\$acc kernels !GR mode do i = 1, N !compiler decides on the partitioning (GP/WP/VP modes) y(i) = y(i) + a*x(i) enddo do i = 1, N !compiler decides on the partitioning (GP/WP/VP modes) y(i) = b*y(i) + a*x(i) enddo

- Compiler will try to deduce parallelism
- Loops are launched as different GPU kernels





The parallel construct

```
Parallel construct

! *acc parallel
    do i = 1, N
        ! loop executed in GR mode
        y(i) = y(i) + a*x(i)
    enddo
    ! *acc loop
    do i = 1, N
        ! compiler decides on the partitioning (GP/WP/VP modes)
        y(i) = b*y(i) + a*x(i)
    enddo

! *acc end parallel
```

- No automatic parallelism deduction → parallel loops must be specified explicitly
- Implicit gang barrier at the end of parallel





Work-sharing loops

- C/C++: #pragmaacc loop
 - Applies to the immediately following for loop
- Fortran: !\$acc loop
 - Applies to the immediately following do loop
- Loop will be automatically striped and assigned to different threads
 - Use the independent clause to force striping
- Convenience syntax combines parallel/kernels and loop constructs
 - #pragmaacc parallel loop
 - #pragmaacc kernels loop
 - !\$acc parallel loop
 - !\$acc kernels loop





Work-sharing loops – the collapse clause

Collapse loops !\$acc loop collapse(2) do i = 1, Ndo j = 1, NA(i,j) = coeff*B(i,j)enddo enddo

- OpenACC vs. OpenMP
 - OpenACC: apply the loop directive to the following N loops and possibly collapse their iteration spaces if independent
 - OpenMP: Collapse the iteration spaces of the following N loops



Controlling parallelism

- Amount of parallelism at the kernels and parallel level
 - num_gangs(...), num_workers(...), vector_length(...)
- At the loop level
 - gang, worker, vector

100 thread blocks with 128 threads each

```
!$acc parallel num_gangs(100), vector_length(128)
    !$acc loop gang, vector
   do i = 1, n
       v(i) = v(i) + a*x(i)
    enddo
!$acc end parallel
```



Variable scoping

- Allowed in the parallel directive only
- By default, if outside of a code block, variables are shared in global memory
- private: A copy of the variable is placed in each gang (CUDA block)
- firstprivate: Same as private but initialized from the host value



Variable scoping

- Allowed in the parallel directive only
- By default, if outside of a code block, variables are shared in global memory
- private: A copy of the variable is placed in each gang (CUDA block)
- firstprivate: Same as private but initialized from the host value

Implicit scoping:

- (C/C++/Fortran) Loop variables are private to the thread that executes the loop
- (C/C++ only) Scope of variables declared inside a parallel block depends on the current execution mode:
 - Vector-partitioned mode → private to the thread
 - Worker-partitioned. Vector-single mode → private to the worker
 - Worker-single mode → private to the gang





Reduction operations

- #pragmaacc parallel reduction(<op>:<var>) - e.g., #pragmaacc parallel reduction(+:sum)
- #pragmaacc loop reduction(<op>:<var>)
- var must be scalar.
- var is copied and default initialized within each gang
- Intermediate results from each gang are combined and made available outside the parallel region
- Complex numbers are also supported
- Operators: +, *, max, min, &, |, %, &&, | |



Calling functions from parallel regions

- #pragma acc routine {gang | worker | vector | seq}
 - Just before the function declaration or definition
- !\$acc routine {gang | worker | vector | sea}
 - In the specification part of the subroutine
- Parallelism level of the routine.
 - gang: must be called from GR context
 - worker: must be called from WS context
 - vector: must be called from VS context
 - seq: must be called from sequential context





Where is my data?

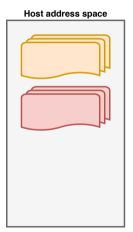
- The host and the device have separate address spaces
 - Data management between the host and the device is the programmer's responsibility
 - You must make sure that all the necessary data for a computation is available on the accelerator before entering the compute region
 - You must make sure to transfer the processed data back to the host if needed



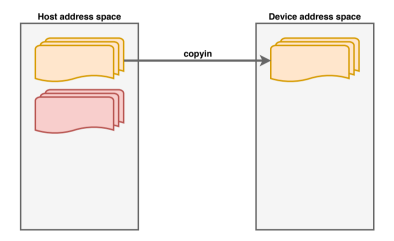
Where is my data?

- The host and the device have separate address spaces
 - Data management between the host and the device is the programmer's responsibility
 - You must make sure that all the necessary data for a computation is available on the accelerator before entering the compute region
 - You must make sure to transfer the processed data back to the host if needed
- But there can be some exceptions:
 - The "device" might be the multicore → no need for data management
 - Some compilers may infer automatically the necessary data transfers
 - Nvidia Pascal GPUs provide efficient support for a unified memory view between the host and the accelerator

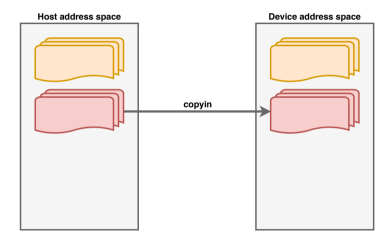




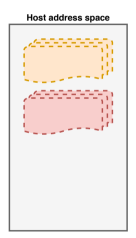


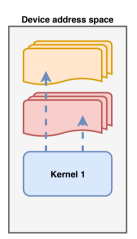


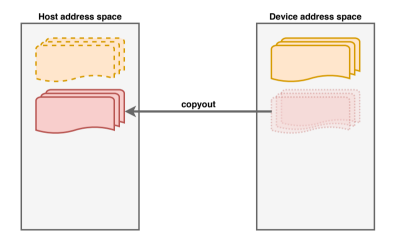






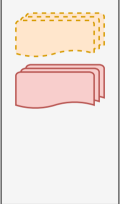




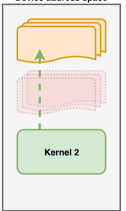


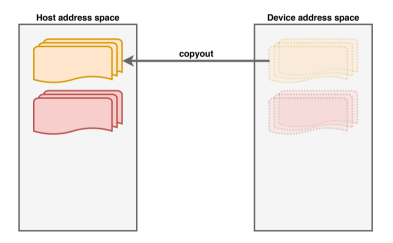






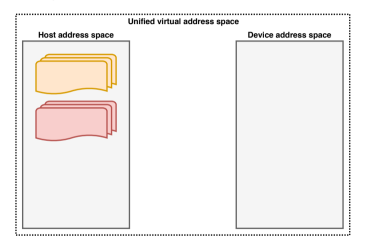
Device address space





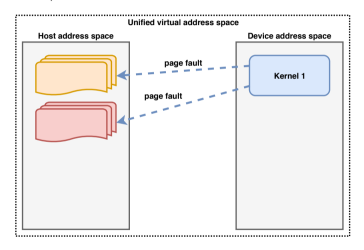


Unified memory address space



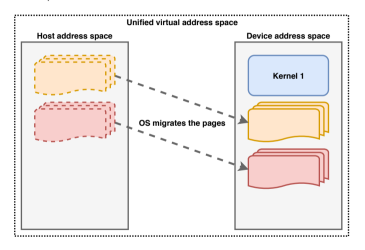


Unified memory address space





Unified memory address space





Directives accepting data clauses

Data clauses may appear in the following directives:

- Compute directives:
 - #pragmaacc kernels
 - #pragmaacc parallel
- Data directives:
 - #pragmaacc data
 - #pragmaacc enter data
 - #pragmaacc exit data
 - #pragmaacc declare
 - #pragmaacc update



Data clauses

- create(a[0:n]): Allocate array a on device
- copyin(a[0:n]): Copy array a to device
- copyout (a[0:n]): Copy array a from device
- copy(a[0:n]): Copy array a to and from device
- present (a): Inform OpenACC runtime that array a is on device
- delete(a): Deallocate array a from device (exit data only)

Not for the acc update directive



The acc data directive

- Defines a scoped data region
 - Data will be copied in at entry of the region and copied out at exit
 - A structural reference count is associated with each memory region that appears in the data clauses
- C/C++: #pragma acc data [data clauses]
 - The next block of code is a data region
- Fortran: !\$acc data [data clauses]
 - Defines a data region until !\$acc end data is encountered



The acc enter/exit data directives

- Defines an unscoped data region
 - Data will be resident on the device until a corresponding exit data directive is found
 - Useful for managing data on the device across compilation units
 - A dynamic reference count is associated with each memory region that appears in the data clauses

C/C++:

- #pragma acc enter data [data clauses]
- #pragma acc exit data [data clauses]

Fortran:

- !\$acc enter data [data clauses]
- !\acc exit data [data clauses]



The acc declare directive

- Functions, subroutines and programs define implicit data regions
- The acc declare directive is used in variable declarations for making them available on the device during the lifetime of the implicit data region
- Useful for copying global variables to the device

- C/C++: #pragma acc declare [data clauses]
- Fortran: !\$acc declare [data clauses]





The acc update directive

 May be used during the lifetime of device data for updating the copies on either host or the device

- #pragmaacc update host(<var-list>)
 - Update host copy with corresponding data from the device

- #pragmaacc update device(<var-list>)
 - Update device copy with corresponding data from the host



Array ranges

Data clauses may accept as arguments

- Whole arrays
 - C/C++: You must specify bounds for dynamically allocated arrays
 - #pragmaacc data copyin(a[0:n])
 - But #pragmaacc data present(a) is acceptable: a's bounds can be inferred by the runtime
 - Fortran: array shape information is already embedded in the data type
 - !\$acc data copyin(a)
- Array subranges
 - #pragmaacc data copyin(a[2:n-2])



Synchronization directives

- Atomic operations
 - #pragmaacc atomic [atomic-clause]
 - !\$acc atomic [atomic-clause]
 - Atomic clauses: read, write, update and capture
 - Example of "capturing" a value:
 - v = x++:
- No global barriers → cannot be implemented due to hardware restrictions
- No equivalent of __syncthreads()





Leverage the unified memory

- Virtual address space shared between CPU and GPU
- The CUDA driver and the hardware take care of the page migration
- Introduced with the Kepler architecture and CUDA 6, but is significantly improved with Pascal





Leverage the unified memory

- Virtual address space shared between CPU and GPU
- The CUDA driver and the hardware take care of the page migration
- Introduced with the Kepler architecture and CUDA 6, but is significantly improved with Pascal

- You could completely omit the data management in OpenACC!
- Supported by the PGI compiler using the -ta=tesla:managed option





Hands-on exercises

General information

The initial course material is available on Github and it will be update during the course:

- git clone https://github.com/vkarak/openacc-training.git
- git pull origin master to get the latest version

Directory structure:

- practicals/: The hands-on exercises
- scripts/: Set up scripts to make your life easier
- slides/: Slides of the course
- ci/: Continuous integration tests for the exercises (ask me offline if interested)



Hands-on exercises

General information

- grep TODO *.{cpp,f90,f03}
- Both Cray/PGI compilers are supported, unless otherwise stated
 - Suggest using PGI for the advanced examples
- source <repodir>/scripts/setup_pgi.sh → will make available PGI 18.4
- module load craype-accel-nvidia60 for loading CUDA and set the target architecture to the GPU
- make
 - For Cray compiler you may use make VERBOSE=1 to get diagnostics information



Exercise 1 – AXPY

- practicals/axpy/axpy_openacc.{cpp,f90}
- Run as: srun --reserv=openacc -Cgpu ./axpy.openacc [ARRAY_SIZE]
 - ARRAY_SIZE is power of 2, default is 16
- Try with different sizes. Does the GPU outperform the CPU version?

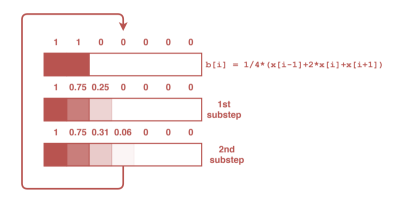


Exercise 2 - Dot product

- practicals/basics/dot_openacc.{cpp,f90}
- Run as: srun --reserv=openacc -Cgpu ./dot.openacc [ARRAY_SIZE]
 - ARRAY_SIZE is power of 2, default is 2
- Try with different sizes. Does the GPU outperform the CPU version?



Exercise 3 – 1D blur kernel





Exercise 3 - 1D blur kernel

- practicals/basics/blur_openacc.{cpp,f90}
- Run as:

```
srun --reserv=openacc -Cgpu ./blur.openacc [ARRAY_SIZE]
```

- ARRAY_SIZE is power of 2, default is 20
- Offload to GPU the loops of the naive kernel; why is it so slow?



Exercise 3 – 1D blur kernel

- practicals/basics/blur_openacc.{cpp,f90}
- Run as:

```
srun --reserv=openacc -Cgpu ./blur.openacc [ARRAY_SIZE]
```

- ARRAY_SIZE is power of 2, default is 20
- Offload to GPU the loops of the naive kernel; why is it so slow?
- Moving data to and from the device is slow (≈7–8 GB/s per direction)
- Avoid unnecessary data movement in the nocopies kernel
 - Move the necessary data to GPU early enough and keep it there as long as possible
 - Update host copies using #pragma acc update directive if needed



Exercise 4 – Experiment with the unified memory

- Remove all the data directives and data clauses
- Compile the blur_twice_naive kernel with -Mcuda=managed
- How does it compare to the manual data management in terms of performance?
- Can you explain the performance difference?





Interoperability with MPI and CUDA

- 1. Call an optimised library function that expects data on the device, e.g., cuBLAS
- Let optimised MPI implementations do RDMA between remote devices' memory
- 3. Manual data management with CUDA, but parallelisation with OpenACC
 - The safest way to manipulate pointers on the device





Interoperability with MPI and CUDA

- Call an optimised library function that expects data on the device, e.g., cuBLAS
- Let optimised MPI implementations do RDMA between remote devices' memory
- 3. Manual data management with CUDA, but parallelisation with OpenACC
 - The safest way to manipulate pointers on the device

Scenarios (1) and (2)

```
#pragmaacc host_data use_device(<varlist>)
```





Interoperability with MPI and CUDA

- Call an optimised library function that expects data on the device, e.g., cuBLAS
- 2. Let optimised MPI implementations do RDMA between remote devices' memory
- 3. Manual data management with CUDA, but parallelisation with OpenACC
 - The safest way to manipulate pointers on the device

Scenarios (1) and (2)

#pragmaacc host_data use_device(<varlist>)

Scenario (3)

Use the deviceptr (<ptrlist>) clause with parallel, kernels and data





2D diffusion example

Source code:

- diffusion2d_omp.{cpp,f90}: our baseline code
 - Single node OpenMP version for the CPU
- diffusion2d_openacc.{cpp,f90}
 - Single node OpenACC version
- diffusion2d_openacc_mpi.{cpp,f90}
 - MPI+OpenACC version
 - If OPENACC_DATA is undefined, data management is performed by CUDA



Calling cuBLAS methods

Source code:

topics/openacc/practicals/gemm/gemm.cpp

Steps:

- Compile with 'make CPPFLAGS=' to get also the naive implementation → too slow!
- 2. Offload the GEMM method to the GPU using OpenACC
- 3. Make use of cuBLAS GEMM through OpenACC

