
OpenACC for GPU: an introduction

Olga Abramkina, Rémy Dubois, Thibaut Véry

Jun 02, 2023

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Directives

OpenMP since specification 4.5 includes support for offloading to accelerators like GPUs. It uses directives to do so (just like for CPU).

A directive has the following structure:

```
 Sentinel  Name      Clause(option, ...) ...
C/C++: #pragma omp target teams map(from: array) private(var) ...
Fortran: !$omp target teams map(from: array) private(var) ...
```

If we break it down, we have these elements:

- The sentinel is special instruction for the compiler. It tells it that what follows has to be interpreted as OpenACC
- The directive is the action to do. In the example, *target* is the way to open a region that will be offloaded to the GPU
- The clauses are “options” of the directive. In the example we want to copy some data on the GPU.
- The clause arguments give more details for the clause. In the example, we give the name of the variables to be copied

Compiling with NVIDIA compiler

To enable OpenMP GPU offloading you need to activate the compilation options `-mp=gpu -gpu=<gpu, opts>`. For example to compile for NVIDIA V100:

```
nvfortran -mp=gpu -gpu=cc70 -o test test.f90
```

GPU offloading

With OpenMP the offloading is realized with the `omp target` directive. By itself, the directive will only offload the computation and do not activate parallelism. It is similar to the `acc serial compute` construct in OpenACC since only one GPU thread is running.

With OpenMP the developer has to activate manually the parallelism.

Here is an example on how to create a GPU kernel:

```
!$omp target
...
!$omp end target
```

Now that we run on the GPU we have to create the threads.

Thread creation on the GPU

Teams

OpenMP `target teams` directive creates several groups of threads that will be able to work in parallel.

OpenACC for GPU: an introduction

With OpenACC it would correspond to the `gang` level of parallelism.

```
!$omp target teams
...
!$omp end target
```

By default the teams will work in replicated mode meaning that they will perform exactly the same things. If you want to share the iterations of a loop between the threads of the teams you have to use the `teams distribute` directive.

```
!$omp target
  !$omp teams distribute
    do i=0, sys_size
      ...
    enddo
!$omp end target
```

This will split the iterations of the loop among the teams. Each team will have a contiguous set of iterations.

It starting to be interesting but we do not yet take advantage of the full power of the GPU.

More threads with `omp parallel`

With the `omp parallel` directive inside a `omp teams` region we create the threads that will be used inside the team.

```
!$omp target teams distribute parallel
do i=1, sys_size
  ...
enddo
```

In this case the threads generated inside the teams will work in replicated mode. If we want to further split the work among those threads we have to add the `omp do` (Fortran) or `omp for` (C/C++) directive.

```
!$omp target teams distribute parallel do
do i=1, sys_size
  ...
enddo
```

With OpenACC it would correspond to the `worker` level of parallelism.

Let's vectorize with `omp simd`

The last level of parallelism we can leverage with OpenMP is the SIMD vectorization. It is done with the `omp simd` directive:

```
!$omp target teams distribute parallel do simd
do i=1, sys_size
  ...
enddo
```

Note for NVIDIA compilers

The `omp simd` construct is not supported for GPU. Currently, the `parallel` directive creates the threads that should be created with `simd`. Since the directive is just ignored, we recommend that you write it for portability reasons.

collapse clause

The `collapse` clause enables to merge all the iterations of several associated loops into a single large iteration loop. The number of loops that will be merged is indicated as an integer argument to this clause and should be greater than 1.

```
!$omp target teams distribute parallel do simd collapse(3)
do k = 1, nz
    do j = 1, ny
        do i = 1, nx
            ...
        enddo
    enddo
enddo
```

Up to now, we will recommend you to use the `collapse` clause as much as you can with OpenMP target in order to achieve good performance.

Example

```
%%idrrun --options "-mp=gpu -gpu=cc70 -Minfo=all"
!! examples_openmp/Fortran/basic_offloading.f90
program basic_offloading
    use iso_fortran_env, only : real64
    implicit none

    integer :: i, sys_size
    real(kind=real64), allocatable, dimension(:) :: array

    sys_size = 100000000

    allocate(array(sys_size))

    !$omp target teams distribute parallel do simd
    do i=1, sys_size
        array(i) = real(i)
    enddo

    print *, "array(42) = ", array(42)
end program basic_offloading
```

Reductions

Reductions should be performed when a memory location is updated by several threads concurrently, and usually prior to its previous value.

This can be performed by using the `reduction` clause of the target construct. This clause will create a private copy of the variables and initialize them as a function of the requested reduction operation. Once you reach the end of the kernel,

the original variable will be updated with a combination of all the private copies.

The syntax is:

```
!$omp target parallel do reduction(operation:variable_list)
...

```

The available operations are:

- +, -
- -
- &, |, ^, &&, ||

Limitation

The reductions are now only supported for the 2 following combined constructs:

- `omp target parallel for`
- `omp target teams distribute parallel for`

Data management

Implicit behavior

If not specified in a `data map` structure, variables will be mapped implicitly at the entry of one kernel with a default action depending on the type of the variable.

Scalars will be map as `firstprivate`, i.e. every thread will have its own private copy that will be initialized with the value that the scalar have on the CPU before the kernel.

Arrays will be shared in memory between threads and are implicitly mapped as if you specified `map (tofrom:)`.

Pointers will be private by default.

You can see the effect of this implicit behavior with the example below:

```
%%idrrun --options "-mp=gpu -gpu=cc70 -Minfo=all"
! examples_openmp/Fortran/Implicit_behavior.f90
program Implicit_behavior
  use iso_fortran_env, only : INT32, REAL64
  implicit none

  real    (kind=REAL64), dimension(:) , allocatable :: Array
  integer(kind=INT32 )                           :: nx, i, scalar

  nx = 10
  allocate(Array(nx))

  scalar = 1000
  !$omp target teams distribute parallel do simd
  do i = 1, nx
```

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```

    Array(i) = scalar + i
enddo

print *, Array

scalar = -1000
!$omp target teams distribute parallel do simd
do i = 1, nx
    Array(i) = scalar + i
enddo

print *, Array

deallocate(Array)
end program Implicit_behavior

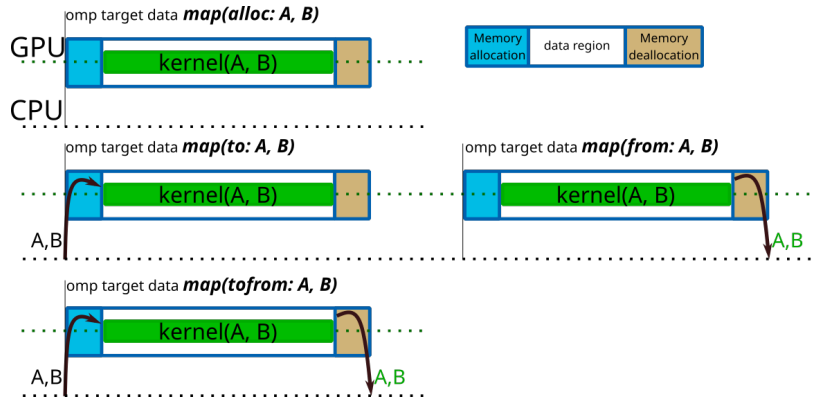
```

Relying only on the implicit behavior can lead to performance degradation as data transfers are performed back and forth at every kernels. This should be avoid by using data regions.

You can define a specific action to perform at the entry and/or the exit of a kernel for a variable or a set of variable with the map clause of the target construct.

The available options are:

- alloc to create the memory space of the variables without prior data transfer.
- to to create the memory space of the variables and transfer the values from CPU to GPU at the entry of the kernel.
- from to create the memory space of the variables and transfer the values from GPU to CPU at the exit of the kernel.
- tofrom to create the memory space of the variables and transfer the values from CPU to GPU at the entry of the kernel, then from GPU to CPU at the exit.



The syntax is:

```

!$omp target map(to:variable1, variable2)
...
!$omp end target

```

It is also possible to modify the status of the variable manually with the private and firstprivate clauses of the target construct or by setting a default mapping that we will see later.

```
!$omp target private(variable1,variable2) firstprivate(variable3)
...
    ! variable1 and variable2 will have independent memory allocations for each
    ↪threads
        ! variable2 will have independent memory allocations for each threads and will be
        ↪initialized with the CPU value
 !$omp end target
```

Structured data region

To run the kernels on GPU, the data should be allocated on the device and eventually the original values should be transferred from the CPU to the GPU. You will also have to retrieve some of the data back from the GPU to the CPU in order to store your results. This can be performed within the same program unit by using the `target data` construct.

If you don't use data regions, implicit copies of the variables will be performed at each entry and exit of every kernels. This implies transfers through the PCIe that could be avoided and thus non-optimal performances.

This construct maps the variable to the device, but only for the extent of the region. The `map` clause enables you to decide which action will be performed on the gpu. These actions could be `alloc`, `to`, `from`, `tofrom`.

You can retrieve the values that were stored on the GPU with `from` and `tofrom` clauses

You can inform the GPU of the original CPU values with the clauses `to` and `tofrom`.

If you use the `alloc` or `from` clause, the initial value on the device is undetermined.

The syntax is:

```
real :: A(nx,ny), B(nx,ny)

 !$omp target data map(tofrom:A,B)
 ...
 !$omp end target data
```

Persistent data (`enter data / exit data`)

If you want to allocate the memory of some variables on the device at a given point of your program but it is not possible to free the memory within the same scope of the program, you can then use the `enter data` and `exit data` constructs.

`enter data` will enable you to allocate or allocate and initialize the variables on the GPU with the `map(alloc:variable_list)` and `map(to:variable_list)` clauses respectively.

`exit data` will enable you to free the memory from the device, resp. free the memory after retrieving the data, with the `map(delete:variable_list)`, resp. `map(from:variable_list)`.

These 2 constructs are not tied to each other, such as one `enter data` construct mapping several variables can lead to several `exit data` constructs in different portions of the code as long as 2 `exit data` are not referring to the same variable in this example.

The syntax is:

```

subroutine some_function_somewhere()
    real :: A(nx,ny), B(nx,ny)

    !$omp target enter data map(to:A)
    !$omp target enter data map(alloc:B)
    ...

end subroutine some_function_somewhere

subroutine some_function_elsewhere_or_maybe_the_same_as_before()
    ...
    !$omp target exit data map(delete:A,B)
end subroutine some_function_elsewhere_or_maybe_the_same_as_before

```

Manual data transfers

When you want to update the values of a given variable, or a set of variables, either on the GPU or on the CPU, you can use the `target update` construct in order to avoid doing it by closing a data structure.

The `to` clause will update the GPU.

The `from` clause will update the CPU.

```
!$omp target update to(variable1,variable2)
```

`defaultmap` clause

You can modify the default mapping for the data transfer upon kernels or data structures with the `defaultmap` clause of the `target` and `target data` constructs.

The new implicit behavior can be specified as `alloc`, `to`, `from`, `tofrom`, `default`, `none`, `firstprivate` or `present` and should be applied to a variable category. Variable categories are:

- scalar
- aggregate (corresponding to arrays and structures in C/C++ and to derived types in Fortran)
- allocatable (only for Fortran arrays that are dynamically allocated)
- pointers

If you specify the implicit behavior as `none`, you should then map explicitly all variables.

```

real, dimension(:), allocatable :: A
real :: B
!$omp target data defaultmap(firstprivate:scalar) defaultmap(tofrom:allocatable)
...
!$omp end target data

```

Modular programming

Functions that are call inside a kernel should be executed on the accelerator. You should use the `declare target` construt to inform the compiler that it should produce such an executable. Syntax should be:

```
subroutine my_routine(...)  
!$omp declare target  
...  
end subroutine my_routine
```

If the function and the line from which the function is called are not within the same program unit, you should add a named `declare target` construct within the program unit containing the call.

```
subroutine another_routine  
!$omp declare target (my_routine)  
  
!$omp target teams  
call my_routine()  
!$omp end target teams  
  
end subroutine another_routine
```

Exercise

```
%%idrrun --options "-mp=gpu -gpu=cc70 -Minfo=all"  
! examples_openmp/Fortran/Modular_programming_mean_value_exercise.f90  
module calcul  
    use iso_fortran_env, only : INT32, REAL64  
    contains  
        subroutine rand_init(array,n)  
            real (kind=REAL64), dimension(1,n), intent(inout) :: array  
            integer(kind=INT32 ), intent(in)                 :: n  
            real (kind=REAL64)                            :: rand_val  
            integer(kind=INT32)                           :: i  
  
            call srand(12345900)  
            do i = 1, n  
                call random_number(rand_val)  
                array(1,i) = 2.0_real64*(rand_val-0.5_real64)  
            enddo  
        end subroutine rand_init  
  
        subroutine iterate(array, array_size, cell_size)  
            real (kind=REAL64), dimension(1:array_size,1), intent(inout) :: array  
            integer(kind=INT32 ), intent(in)                 :: array_<br>size, cell_size  
            real (kind=REAL64)                            :: local_<br>mean  
            integer(kind=INT32 )                           :: i  
  
            do i = cell_size/2, array_size-cell_size/2  
                local_mean = mean_value(array(i+1-cell_size/2:i+cell_size/2,1), cell_<br>size)  
                if (local_mean .lt. 0.0_real64) then  
                    array(i,1) = array(i,1) + 0.1  
                end if  
            end do  
        end subroutine iterate
```

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```

        else
            array(i,1) = array(i,1) - 0.1
        endif
    enddo
end subroutine iterate

function mean_value(t, n)
    real (kind=REAL64), dimension(n,1), intent(inout) :: t
    integer(kind=INT32 ), intent(in)                  :: n
    real (kind=REAL64)                                :: mean_value
    integer(kind=INT32 )                                :: i
    mean_value = 0.0_real64

    do i = 1, n
        mean_value = mean_value + t(i,1)
    enddo
    mean_value = mean_value / dble(n)
end function mean_value
end module calcul
program modular_programming
use calcul
implicit none

real (kind=REAL64), dimension(:, :, ), allocatable :: table
real (kind=REAL64), dimension(:, ), allocatable :: mean_values
integer(kind=INT32 )                                :: nx, ny, cell_size, i

nx = 1000000
ny = 3000
allocate(table(nx,ny), mean_values(ny))
table(:, :) = 0.0_real64
call rand_init(table(1,:),ny)
cell_size = 32
do i = 2, ny
    call iterate(table(:,i), nx, cell_size)
enddo

do i = 1, ny
    mean_values(i) = mean_value(table(:,i), nx)
enddo

do i = 1, 10
    write(0,"(a18,i5,a1,f20.8)") "Mean value of row ",i,"=",mean_values(i)
enddo

do i = ny-10, ny
    write(0,"(a18,i5,a1,f20.8)") "Mean value of row ",i,"=",mean_values(i)
enddo

deallocate(table, mean_values)
end program modular_programming

```

Solution

```

%%idrrun --options "-mp=gpu -gpu=cc70 -Minfo=all"
! examples_openmp/Fortran/Modular_programming_mean_value_solution.f90
module calcul
    use iso_fortran_env, only : INT32, REAL64
contains
    subroutine rand_init(array,n)
        real    (kind=REAL64), dimension(1,n), intent(inout) :: array
        integer(kind=INT32 ), intent(in)                      :: n
        real    (kind=REAL64)                                :: rand_val
        integer(kind=INT32 )                                :: i

        call srand(12345900)
        do i = 1, n
            call random_number(rand_val)
            array(1,i) = 2.0_real64*(rand_val-0.5_real64)
        enddo
    end subroutine rand_init

    subroutine iterate(array, array_size, cell_size)
        !$omp declare target
        real    (kind=REAL64), dimension(1:array_size,1), intent(inout) :: array
        integer(kind=INT32 ), intent(in)                      :: array_
        real    (kind=REAL64)                                :: local_
        integer(kind=INT32 )                                :: i

        do i = cell_size/2, array_size-cell_size/2
            local_mean = mean_value(array(i+1-cell_size/2:i+cell_size/2,1), cell_
            size)
            if (local_mean .lt. 0.0_real64) then
                array(i,1) = array(i,1) + 0.1
            else
                array(i,1) = array(i,1) - 0.1
            endif
        enddo
    end subroutine iterate

    function mean_value(t, n)
        !$omp declare target
        real    (kind=REAL64), dimension(n,1), intent(inout) :: t
        integer(kind=INT32 ), intent(in)                      :: n
        real    (kind=REAL64)                                :: mean_value
        integer(kind=INT32 )                                :: i
        mean_value = 0.0_real64

        do i = 1, n
            mean_value = mean_value + t(i,1)
        enddo
        mean_value = mean_value / dble(n)
    end function mean_value
end module calcul
program modular_programming
    use calcul
    implicit none

```

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```

real    (kind=REAL64), dimension(:, :, ), allocatable :: table
real    (kind=REAL64), dimension(:) , allocatable :: mean_values
integer(kind=INT32 )                                :: nx, ny, cell_size, i

nx = 1000000
ny =     3000
allocate(table(nx,ny), mean_values(ny))
table(:, :) = 0.0_real64
call rand_init(table(1,:),ny)
 !$omp target enter data map(to:table)
cell_size = 32
 !$omp target teams distribute parallel do simd
do i = 2, ny
    call iterate(table(:,i), nx, cell_size)
enddo

 !$omp target teams distribute parallel do simd map(from:mean_values)
do i = 1, ny
    mean_values(i) = mean_value(table(:,i), nx)
enddo

do i = 1, 10
    write(0,"(a18,i5,a1,f20.8)") "Mean value of row ",i,"=",mean_values(i)
enddo

do i = ny-10, ny
    write(0,"(a18,i5,a1,f20.8)") "Mean value of row ",i,"=",mean_values(i)
enddo

 !$omp target exit data map(delete:table)
deallocate(table, mean_values)
end program modular_programming

```

Using multiple GPUs with OpenMP

If you have multiple accelerators available, you can select the one on which you run the kernels with the `device` clause of the `target` construct. It includes both `target data` constructs and `target teams/parallel` constructs.

You should give an integer that refers to the gpu number (starting from 0) to the `device` clause, such as :

```

call mpi_comm_rank(MPI_COMM_WORLD, rank, code)
num_gpus = omp_get_num_devices()
my_gpu   = mod(my_rank, num_gpus)
 !$omp target data map(...) device(my_gpu)
 ...
 !$omp end target data

```

Exercise

In this exercise, you should bring on the gpu the MPI version of the generation of the Mandelbrot set on the gpu with OpenMP and by using multiple devices.

```
%%idrrun --cliopts "2000 1000" -m 4 -g 4 --options "-mp=gpu -gpu=cc70 -Minfo=all"
! examples_openmp/Fortran/mandelbrot_mpi_exercise.f90
program mandelbrot_mpi
    use MPI
    implicit none
    real, parameter :: min_re = -2.0, max_re = 1.0
    real, parameter :: min_im = -1.0, max_im = 1.0
    integer :: first, last, width, height
    integer :: num_elements
    real :: step_w, step_h
    integer :: numarg, i, length, j, first_elem, last_elem
    integer :: rest_eucli, local_height
    integer :: rank, nb_procs, code
    character(len=:), allocatable :: arg1, arg2
    integer (kind=1), allocatable :: picture(:)
    real :: x, y

    numarg = command_argument_count()
    if (numarg .ne. 2) then
        write(0,*) "Error, you should provide 2 arguments of integer kind : width and"
        length
        stop
    endif
    call get_command_argument(1, LENGTH=length)
    allocate(character(len=length) :: arg1)
    call get_command_argument(1, VALUE=arg1)
    read(arg1,'(i10)') width
    call get_command_argument(2, LENGTH=length)
    allocate(character(len=length) :: arg2)
    call get_command_argument(2, VALUE=arg2)
    read(arg2,'(i10)') height
    step_w = 1.0 / real(width)
    step_h = 1.0 / real(height)

    call mpi_init(code)
    call mpi_comm_rank(MPI_COMM_WORLD, rank, code)
    call mpi_comm_size(MPI_COMM_WORLD, nb_procs, code)

    local_height = height / nb_procs
    first = 0
    last = local_height
    rest_eucli = mod(height, nb_procs)

    if ((rank .eq. 0) .and. (rank .lt. rest_eucli)) last = last + 1

    if (rank .gt. 0) then
        do i = 1, rank
            first = first + local_height
            last = last + local_height
            if (rank .lt. rest_eucli) then
                first = first + 1
                last = last + 1
            endif
        enddo
    endif

    if (rank .lt. rest_eucli) local_height = local_height + 1
```

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```

num_elements = local_height * width

write(unit=*,fmt="(a9,i3,a18,i8,a3,i8,a5,i10,a9)") "I am rank",rank, &
" and my range is [",first," ",last,"[ ie ",num_elements," elements"

allocate(picture(first*width:last*width))

do i=first,last-1
    do j=0,width-1
        x = min_re + j * step_w * (max_re - min_re)
        y = min_im + i * step_h * (max_im - min_im)
        picture(i*width+j) = mandelbrot_iterations(x,y)
    enddo
enddo

call output()
deallocate(picture)

call mpi_finalize(code)

contains

subroutine output
    integer                      :: fh
    integer(kind=MPI_OFFSET_KIND) :: woffset

    woffset=first*width
    call MPI_File_open(MPI_COMM_WORLD,"mandel.gray",MPI_MODE_WRONLY+MPI_MODE_-
CREATE,MPI_INFO_NULL,fh,code)
    call MPI_File_write_at(fh,woffset,picture,num_elements,MPI_INTEGER1,MPI_-
STATUS_IGNORE,code);
    call MPI_File_close(fh,code)
end subroutine output
integer(kind=1) function mandelbrot_iterations(x,y)
    integer, parameter           :: max_iter = 127
    real, intent(in)             :: x,y
    real                          :: z1,z2,z1_old,z2_old

    z1 = 0.0
    z2 = 0.0
    mandelbrot_iterations = 0
    do while (((z1*z1+z2*z2) .le. 4) .and. (mandelbrot_iterations .lt. max_-
iter))
        z1_old = z1
        z2_old = z2
        z1 = z1_old*z1_old - z2_old*z2_old + x
        z2 = 2.0*z1_old*z2_old + y
        mandelbrot_iterations = mandelbrot_iterations + 1
    enddo
end function mandelbrot_iterations
end program mandelbrot_mpi

```

```

from idrcomp import show_gray
show_gray("mandel.gray", 2000, 1000)

```

Solution

CONTENTS

```
%%idrrun --cliopts "2000 1000" -m 4 -g 4 --options "-mp=gpu -gpu=cc70 -Minfo=all"
! examples_openmp/Fortran/mandelbrot_mpi_solution.f90
program mandelbrot_mpi
    use MPI
    implicit none
    real, parameter :: min_re = -2.0, max_re = 1.0
    real, parameter :: min_im = -1.0, max_im = 1.0
    integer :: first, last, width, height
    integer :: num_elements
    real :: step_w, step_h
    integer :: numarg, i, length, j, first_elem, last_elem
    integer :: rest_eucli, local_height
    integer :: rank, nb_procs, code
    character(len=:), allocatable :: arg1, arg2
    integer (kind=1), allocatable :: picture(:)
    real :: x, y

    numarg = command_argument_count()
    if (numarg .ne. 2) then
        write(0,*) "Error, you should provide 2 arguments of integer kind : width and"
        length
        stop
    endif
    call get_command_argument(1,LENGTH=length)
    allocate(character(len=length) :: arg1)
    call get_command_argument(1,VALUE=arg1)
    read(arg1,'(i10)') width
    call get_command_argument(2,LENGTH=length)
    allocate(character(len=length) :: arg2)
    call get_command_argument(2,VALUE=arg2)
    read(arg2,'(i10)') height
    step_w = 1.0 / real(width)
    step_h = 1.0 / real(height)

    call mpi_init(code)
    call mpi_comm_rank(MPI_COMM_WORLD, rank, code)
    call mpi_comm_size(MPI_COMM_WORLD, nb_procs, code)

    local_height = height / nb_procs
    first = 0
    last = local_height
    rest_eucli = mod(height, nb_procs)

    if ((rank .eq. 0) .and. (rank .lt. rest_eucli)) last = last + 1

    if (rank .gt. 0) then
        do i = 1, rank
            first = first + local_height
            last = last + local_height
            if (rank .lt. rest_eucli) then
                first = first + 1
                last = last + 1
            endif
        enddo
    endif

    if (rank .lt. rest_eucli) local_height = local_height + 1
```

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```

num_elements = local_height * width

write(unit=*,fmt="(a9,i3,a18,i8,a3,i8,a5,i10,a9)") "I am rank",rank, &
" and my range is [",first," ,",last,"[ ie ",num_elements," elements"

allocate(picture(first*width:last*width))
 !$omp target data map(tofrom:picture) device(rank)
 !$omp target teams distribute parallel do simd collapse(2) device(rank)
 do i=first,last-1
   do j=0,width-1
     x = min_re + j * step_w * (max_re - min_re)
     y = min_im + i * step_h * (max_im - min_im)
     picture(i*width+j) = mandelbrot_iterations(x,y)
   enddo
 enddo
 !$omp end target data
 call output()
 deallocate(picture)

call mpi_finalize(code)

contains
 subroutine output
   integer :: fh
   integer(kind=MPI_OFFSET_KIND) :: woffset

   woffset=first*width
   call MPI_File_open(MPI_COMM_WORLD, "mandel.gray", MPI_MODE_WRONLY+MPI_MODE_<CREATE,MPI_INFO_NULL,fh,code)
   call MPI_File_write_at(fh,woffset,picture,num_elements,MPI_INTEGER1,MPI_<STATUS_IGNORE,code);
   call MPI_File_close(fh,code)
 end subroutine output
 integer(kind=1) function mandelbrot_iterations(x,y)
 !$omp declare target
 integer, parameter :: max_iter = 127
 real, intent(in) :: x,y
 real :: z1,z2,z1_old,z2_old

 z1 = 0.0
 z2 = 0.0
 mandelbrot_iterations = 0
 do while (((z1*z1+z2*z2) .le. 4) .and. (mandelbrot_iterations .lt. max_<iter))
   z1_old = z1
   z2_old = z2
   z1 = z1_old*z1_old - z2_old*z2_old + x
   z2 = 2.0*z1_old*z2_old + y
   mandelbrot_iterations = mandelbrot_iterations + 1
 enddo
 end function mandelbrot_iterations
end program mandelbrot_mpi

```

```

from idrcomp import show_gray
show_gray("mandel.gray", 2000, 1000)

```

Using NV-link with OpenMP target

You can specify to the accelerator the pointer to a given data structure already present on the device that should be used with `use_device_addr` clause of the `data` construct.

Exercise

As an exercise, you can complete the following MPI code that measures the bandwidth between the GPUs:

1. Add directives to create the buffers on the GPU
2. Measure the effective bandwidth between GPUs by adding the directives necessary to transfer data from one GPU to another one in the following cases:
 - Not using NVLink
 - Using NVLink

We have a bug for MPI in the notebooks and you need to save the file before running the next cell. It is a good way to practice manual building! Please add the correct extension for the language you are running.

```
%%writefile MultiGPU_mpi_exercise.<extension>
! examples_openmp/Fortran/MultiGPU_mpi_exercise.f90
! you should add ` --option "-cpp" ` as argument to the idrrun command
program MultiGPU_exercice
    use ISO_FORTRAN_ENV, only : INT32, REAL64
    use mpi
    use openacc
    implicit none
    real    (kind=REAL64), dimension(:), allocatable :: send_buffer, receive_buffer
    real    (kind=REAL64)                                :: start, finish , data_volume
    integer(kind=INT32 ), parameter                      :: system_size = 2e8/8
    integer                                         :: comm_size, my_rank, code, reps,
    i, j, k
    integer                                         :: num_gpus, my_gpu
    integer(kind=acc_device_kind)                      :: device_type
    integer, dimension(MPI_STATUS_SIZE)                :: mpi_stat

    ! Useful for OpenMPI and GPU DIRECT
    call initialisation_openacc()

    ! MPI stuff
    reps = 5
    data_volume = dble(reps*system_size)*8*1024_real64**(-3.0)

    call MPI_Init(code)
    call MPI_Comm_size(MPI_COMM_WORLD, comm_size, code)
    call MPI_Comm_rank(MPI_COMM_WORLD, my_rank, code)
    allocate(send_buffer(system_size), receive_buffer(system_size))

    ! OpenACC stuff
    #ifdef _OPENACC
    device_type = acc_get_device_type()
    num_gpus = acc_get_num_devices(device_type)
    my_gpu   = mod(my_rank,num_gpus)
```

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```

call acc_set_device_num(my_gpu, device_type)
#endiff

do j = 0, comm_size - 1
    do i = 0, comm_size - 1
        if ( (my_rank .eq. j) .and. (j .ne. i) ) then
            start = MPI_Wtime()
            do k = 1, reps
                call MPI_Send(send_buffer,system_size, MPI_DOUBLE, i, 0, MPI_COMM_
COMM_WORLD, code)
            enddo
        endif
        if ( (my_rank .eq. i) .and. (i .ne. j) ) then
            do k = 1, reps
                call MPI_Recv(receive_buffer, system_size, MPI_DOUBLE, j, 0, MPI_
COMM_WORLD, mpi_stat, code)
            enddo
        endif
        if ( (my_rank .eq. j) .and. (j .ne. i) ) then
            finish = MPI_Wtime()
            write(0,"(a11,i2,a2,i2,a2,f20.8,a5)") "bandwidth ",j,"->",i,": ",data_
volume/(finish-start)," GB/s"
        endif
    enddo
enddo

deallocate(send_buffer, receive_buffer)

call MPI_Finalize(code)

contains

#ifdef _OPENACC
subroutine initialisation_openacc
    use openacc
    implicit none
    type accel_info
        integer :: current_devices
        integer :: total_devices
    end type accel_info

    type(accel_info) :: info
    character(len=6) :: local_rank_env
    integer :: local_rank_env_status, local_rank
! Initialisation of OpenACC
 !$acc init

! Recovery of the local rank of the process via the environment variable
! set by Slurm, as MPI_Comm_rank cannot be used here because this routine
! is used BEFORE the initialisation of MPI
    call get_environment_variable(name="$SLURM_LOCALID", value=local_rank_env,_
status=local_rank_env_status)
    info%total_devices = acc_get_num_devices(acc_get_device_type())
    if (local_rank_env_status == 0) then
        read(local_rank_env, *) local_rank
        ! Definition of the GPU to be used via OpenACC
        call acc_set_device_num(local_rank, acc_get_device_type())
end subroutine
#endif

```

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```

        info%current_devices = local_rank
    else
        print *, "Error : impossible to determine the local rank of the_
→process"
        stop 1
    endif
end subroutine initialisation_openacc
#endif

end program MultiGPU_exercice

```

```

module load nvidia-compilers/21.9 cuda/11.2 openmpi/4.0.5-cuda
# Add compiling here
mpi.....
srun -A for@gpu --gpus-per-node=2 --ntasks-per-node=4 --cpus-per-task=5 ./a.out

```

Solution

We have a bug for MPI in the notebooks and you need to save the file before running the next cell. It is a good way to practice manual building! Please add the correct extension for the language you are running.

```

%%writefile MultiGPU_mpi_exercise.<extension>
! examples_openmp/Fortran/MultiGPU_mpi_solution.f90
! you should add ` --option "-cpp" ` as argument to the idrrun command
program MultiGPU_solution
    use ISO_FORTRAN_ENV, only : INT32, REAL64
    use mpi
    use openacc
    implicit none
    real    (kind=REAL64), dimension(:), allocatable :: send_buffer, receive_buffer
    real    (kind=REAL64)
    integer(kind=INT32 ), parameter
    integer
    ↵ i, j, k
    integer
    integer(kind=acc_device_kind)
    integer, dimension(MPI_STATUS_SIZE)
                                :: start, finish , data_volume
                                :: system_size = 2e8/8
                                :: comm_size, my_rank, code, reps,
                                :: num_gpus, my_gpu
                                :: device_type
                                :: mpi_stat

    ! Useful for OpenMPI and GPU DIRECT
    call initialisation_openacc()

    ! MPI stuff
    reps = 5
    data_volume = dble(reps*system_size)*8*1024_real64**(-3.0)

    call MPI_Init(code)
    call MPI_Comm_size(MPI_COMM_WORLD, comm_size, code)
    call MPI_Comm_rank(MPI_COMM_WORLD, my_rank, code)
    allocate(send_buffer(system_size), receive_buffer(system_size))
    !$omp target enter data map(alloc: send_buffer(1:system_size), receive_
→buffer(1:system_size))

```

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```

! OpenMP target stuff
#ifndef _OPENACC
device_type = acc_get_device_type()
num_gpus = acc_get_num_devices(device_type)
my_gpu   = mod(my_rank, num_gpus)
call acc_set_device_num(my_gpu, device_type)
#endif

do j = 0, comm_size - 1
    do i = 0, comm_size - 1
        if ( (my_rank .eq. j) .and. (j .ne. i) ) then
            start = MPI_Wtime()
            !$omp target data use_device_ptr(send_buffer)
            do k = 1, reps
                call MPI_Send(send_buffer, system_size, MPI_DOUBLE, i, 0, MPI_COMM_
COMM_WORLD, code)
            enddo
            !$omp end target data
        endif
        if ( (my_rank .eq. i) .and. (i .ne. j) ) then
            !$omp target data use_device_ptr(send_buffer)
            do k = 1, reps
                call MPI_Recv(receive_buffer, system_size, MPI_DOUBLE, j, 0, MPI_
COMM_WORLD, mpi_stat, code)
            enddo
            !$omp end target data
        endif
        if ( (my_rank .eq. j) .and. (j .ne. i) ) then
            finish = MPI_Wtime()
            write(0,"(a11,i2,a2,i2,a2,f20.8,a5)") "bandwidth ",j,"->",i,": ",data_
volume/(finish-start)," GB/s"
        endif
    enddo
enddo
 !$omp target exit data map(delete: send_buffer, receive_buffer)
deallocate(send_buffer, receive_buffer)

call MPI_Finalize(code)

contains
#ifndef _OPENACC
subroutine initialisation_openacc
use openacc
implicit none
type accel_info
    integer :: current_devices
    integer :: total_devices
end type accel_info

type(accel_info) :: info
character(len=6) :: local_rank_env
integer          :: local_rank_env_status, local_rank
! Initialisation of OpenACC
 !$acc init

! Recovery of the local rank of the process via the environment variable

```

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```

! set by Slurm, as MPI_Comm_rank cannot be used here because this routine
! is used BEFORE the initialisation of MPI
    call get_environment_variable(name="SLURM_LOCALID", value=local_rank_env,_
→status=local_rank_env_status)
        info%total_devices = acc_get_num_devices(acc_get_device_type())
        if (local_rank_env_status == 0) then
            read(local_rank_env, *) local_rank
            ! Definition of the GPU to be used via OpenACC
            call acc_set_device_num(local_rank, acc_get_device_type())
            info%current_devices = local_rank
        else
            print *, "Error : impossible to determine the local rank of the_
→process"
            stop 1
        endif
    end subroutine initialisation_openacc
#endif

end program MultiGPU_solution

```

```

%%bash
module load nvidia-compilers/21.9 cuda/11.2 openmpi/4.0.5-cuda
# Add compiling here
mpi....
srun -A for@gpu --gpus-per-node=4 --ntasks-per-node=8 --cpus-per-task=5 ./a.out

```

Asynchronism

Concurrent executions within the same stream

An implicit barrier is set at the end of each `target` construct to ensure that the parent task (the task on the host) can not move on until the target task has ended. You can disable this implicit behavior and submit several kernels on the GPU by explicitly adding the `nowait` clause to the `target` construct.

In order to avoid race conditions that could arise from the lack of barrier between kernels, it is possible to specify a scheduling of the kernels based on a dependency mechanism. To do so, you should use the `depend` clause.

Exercise

```

%%idrrun --options "-mp=gpu -gpu=cc70 -Minfo=all"
! examples_openmp/Fortran/async_async_exercise.f90
program prod_mat
    use iso_fortran_env, only : INT32, REAL64
    implicit none
    integer (kind=INT32) :: rank=5000
    real (kind=REAL64), allocatable :: A(:, :, :), B(:, :, :), C(:, :, :)
    integer (kind=INT32) :: i, j, k
    integer (kind=INT32) :: streamA, streamB, streamC

```

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```

streamA = 1
streamB = 2
streamC = 3

call create_mat(A, rank, streamA)
call create_mat(B, rank, streamB)
call create_mat(C, rank, streamC)

call init_mat(A, rank, 3.0_real64 , streamA)
call init_mat(B, rank, 14.0_real64, streamB)
call init_mat(C, rank, 0.0_real64 , streamC)

do j=1, rank
    do k=1, rank
        do i=1, rank
            C(i,j) = C(i,j) + A(i,k)*B(k,j)
        enddo
    enddo
enddo
print *, "Check that this is close to 42.0:", C(12,12)
deallocate(A, B, C)
contains
    subroutine create_mat(mat, rank, stream)
        real      (kind=REAL64), intent(inout), allocatable :: mat(:,:)
        integer(kind=INT32 ), intent(in)                      :: rank, stream
        allocate(mat(rank,rank))
    end subroutine create_mat

    subroutine init_mat(mat, rank, diag, stream)
        real      (kind=REAL64), intent(inout)   :: mat(:,:)
        real      (kind=REAL64), intent(in)       :: diag
        integer (kind=INT32 ), intent(in)       :: rank, stream
        integer (kind=INT32 )                  :: i, j

        do j=1, rank
            do i=1, rank
                mat(i,j) = 0.0_real64
            enddo
        enddo

        do j=1, rank
            mat(j,j) = diag
        enddo
    end subroutine init_mat
end program prod_mat

```

Solution

```

%%idrrun --options "-mp=gpu -gpu=cc70 -Minfo=all"
! examples_openmp/Fortran/async_async_solution.f90
program prod_mat
    use iso_fortran_env, only : INT32, REAL64
    implicit none

```

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```

integer (kind=INT32) :: rank=5000
real (kind=REAL64), allocatable :: A(:,:,:), B(:,:,:)
integer (kind=INT32) :: i, j, k
integer (kind=INT32) :: streamA, streamB, streamC

streamA = 1
streamB = 2
streamC = 3

call create_mat(A, rank, streamA)
call create_mat(B, rank, streamB)
call create_mat(C, rank, streamC)

call init_mat(A, rank, 3.0_real64 , streamA)
call init_mat(B, rank, 14.0_real64, streamB)
call init_mat(C, rank, 0.0_real64 , streamC)

!$omp target teams distribute parallel do simd collapse(3)
do j=1, rank
    do k=1, rank
        do i=1, rank
            C(i,j) = C(i,j) + A(i,k)*B(k,j)
        enddo
    enddo
enddo
!$omp target exit data map(delete:A,B)
!$omp target exit data map(from:C)
print *, "Check that this is close to 42.0:", C(12,12)
deallocate(A, B, C)
contains
    subroutine create_mat(mat, rank, stream)
        real (kind=REAL64), intent(inout), allocatable :: mat(:,:)
        integer(kind=INT32 ), intent(in)                 :: rank, stream
        allocate(mat(rank,rank))
        !$omp target enter data map(alloc:mat) nowait depend(out:mat)
    end subroutine create_mat

    subroutine init_mat(mat, rank, diag, stream)
        real (kind=REAL64), intent(inout)   :: mat(:,:)
        real (kind=REAL64), intent(in)     :: diag
        integer (kind=INT32 ), intent(in)  :: rank, stream
        integer (kind=INT32 )             :: i, j

        !$omp target teams distribute parallel do simd collapse(2) nowait_
        ~depend(inout:mat)
        do j=1, rank
            do i=1, rank
                mat(i,j) = 0.0_real64
            enddo
        enddo

        !$omp target teams distribute parallel do simd nowait depend(in:mat)
        do j=1, rank
            mat(j,j) = diag
        enddo
    end subroutine init_mat
end program prod_mat

```

OPENMP CHEAT SHEET

1.1 Directive syntax

 Sentinel Name Clause(option, ...) ...
C/C++: #pragma omp target teams map(from: array) private(var) ...
Fortran: !\$omp target teams map(from: array) private(var) ...

If we break it down, we have those elements:

- The sentinel is a special instruction for the compiler. It tells him that what follows has to be interpreted as OpenMP directives
- The directive is the action to do. In the example, *target* is the way to open a parallel region that will be offloaded to the GPU
- The clauses are “options” of the directive. In the example we want to copy some data from the GPU.
- The clause arguments give more details for the clause. In the example, we give the name of the variables to be copied

1.2 Creating kernels

The way to open kernels on the GPU is to use the `omp target` directive with `directive` to create threads.

1.2.1 Creating threads

The threads creation is the job of the developer in OpenMP. The standard defines 3 levels of parallelism:

- `omp teams`: Several groups of threads are created but only the master thread is active.
- `omp parallel`: The other threads of the team are activated.
- `omp simd`: SIMD threads are activated

1.2.2 Work Sharing

Creating threads is not enough to have the full power of the GPU. You have to share work among threads:

- `omp teams distribute`: distribute work among teams
- `omp parallel for/do`: distribute work inside a team

1.2.3 *omp target* Clauses

Clause	effect
<code>private(vars, ...)</code>	Make <i>vars</i> private at <i>team</i> level
<code>firstprivate(vars, ...)</code>	Make <i>vars</i> private at <i>team</i> level and copy the value <i>vars</i> had on the host before
<code>device(dev_num)</code>	Set the device on which to run the kernel

Other clauses might be available. Check the specification and the compiler documentation for full list.

1.2.4 *omp teams* Clauses

Clause	effect
<code>num_teams(#teams)</code>	Set the number of teams for the target region
<code>thread_limit(#threads)</code>	Set the maximum number of threads inside a team
<code>private(vars, ...)</code>	Make <i>vars</i> private at <i>team</i> level
<code>firstprivate(vars, ...)</code>	Make <i>vars</i> private at <i>team</i> level and copy the value <i>vars</i> had on the host before
<code>reduction(op:vars, ...)</code>	Perform a reduction of the variables <i>vars</i> with operation <i>op</i>

Other clauses might be available. Check the specification and the compiler documentation for full list.

1.2.5 *omp parallel* Clauses

Clause	effect
<code>private(vars, ...)</code>	Make <i>vars</i> private at <i>parallel</i> level
<code>firstprivate(vars, ...)</code>	Make <i>vars</i> private at <i>parallel</i> level and copy the value <i>vars</i> had on the host before
<code>reduction(op:vars, ...)</code>	Perform a reduction of the variables <i>vars</i> with operation <i>op</i>

Other clauses might be available. Check the specification and the compiler documentation for full list.

1.2.6 *omp simd* Clauses

Clause	effect
<code>private(vars, ...)</code>	Make <i>vars</i> private at <i>simd</i> level
<code>firstprivate(vars, ...)</code>	Make <i>vars</i> private at <i>simd</i> level and copy the value <i>vars</i> had on the host before
<code>reduction(op:vars, ...)</code>	Perform a reduction of the variables <i>vars</i> with operation <i>op</i>
<code>simdlen(vector_size)</code>	Set the length of the vector

Other clauses might be available. Check the specification and the compiler documentation for full list.

1.3 Combined constructs for loops

It is possible to combine the

1.4 Managing data

1.4.1 Data regions

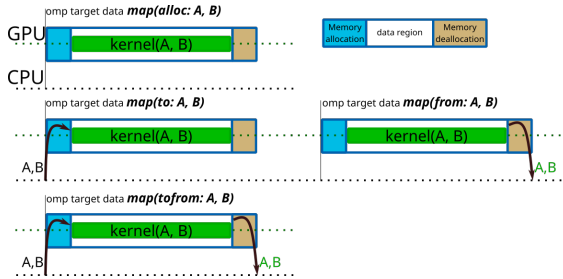
Region	Directive
Program lifetime	omp target enter data & omp target exit data
Structured	omp target data
Kernels	omp target map(...)

1.4.2 Data clauses

To choose the right data clause you need to answer the following questions:

- Does the kernel need the values computed on the host (CPU) beforehand? (Before)
- Are the values computed inside the kernel needed on the host (CPU) afterward? (After)

	Needed after	Not needed after
Needed Before	map(tofrom:var1, ...)	map(to:var2, ...)
Not needed before	map(from:var3, ...)	map(alloc:var4, ...)



1.4.3 Updating data already present on the GPU

It is not possible to update data present on the GPU with the data clauses on a data region. To do so you need to use `omp target update`

omp target update Clauses

- To update CPU with data computed on GPU: `omp target update from(data, ...)`
- To update GPU with data computer on CPU: `omp target update to(data, ...)`

1.5 GPU routines

A routine called from a kernel needs to be inside a `declare target` region.

```
subroutine my_routine(...)  
!$omp declare target  
...  
end subroutine my_routine
```

1.6 Using data on the GPU with GPU aware libraries

To get a pointer to the device memory for a variable you have to use:

- `omp data use_device_ptr(var, ...)` for pointers
- `omp data use_device_addr(var, ...)` for allocatables