





Hands-On Session Using OpenACC in MPAS-A

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In this notebook, we explore the GPU enabled MPAS-A (Model Prediction Across Scales-Atmosphere) to apply techniques learned from MiniWeather and implementing OpenACC to develop for GPUs.

- Review of exercises from prior OpenACC/MiniWeather sessions Part 1 and Part 2
- MPAS-Atmosphere model overview
- Managing GPU data in large software projects
- Assessing performance of extracted GPU kernels in MPAS-A

Head to the NCAR JupyterHub portal and **start a JupyterHub session on Casper login** (or batch nodes using 1 CPU, no GPUs) and open the notebook in XXXX. Be sure to clone (if needed) and update/pull the NCAR GPU_workshop directory.

```
# Use the JupyterHub GitHub GUI on the left panel or the below shell commands git clone git@github.com:NCAR/GPU_workshop.git git pull
```

Workshop Etiquette

- Please mute yourself and turn off video during the session.
- Questions may be submitted in the chat and will be answered when appropriate. You
 may also raise your hand, unmute, and ask questions during Q&A at the end of the
 presentation.
- By participating, you are agreeing to UCAR's Code of Conduct
- Recordings & other material will be archived & shared publicly.
- Feel free to follow up with the GPU workshop team via Slack or submit support requests to support.ucar.edu
 - Office Hours: Asynchronous support via Slack or schedule a time with an organizer

Notebook Setup

Set the PROJECT code to a currently active project, ie UCIS0004 for the GPU workshop, and QUEUE to the appropriate routing queue depending on if during a live workshop session (gpuworkshop), during weekday 8am to 5:30pm MT (gpudev), or all other times (casper). Due to limited shared GPU resources, please use GPU_TYPE=gp100 during the workshop. Otherwise, set GPU_TYPE=v100 (required for gpudev) for independent work. See Casper queue documentation for more info.

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```
export PROJECT=UCIS0004
export QUEUE=gpudev
export GPU_TYPE=v100
```

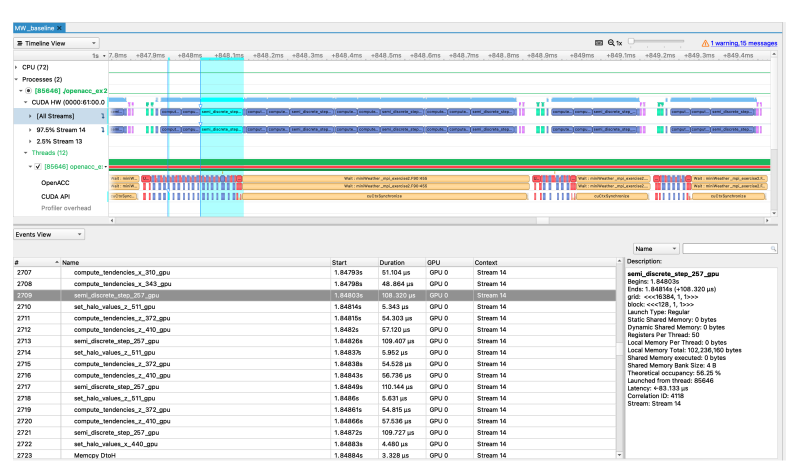
Review of MiniWeather Performance Optimization

At the end of last session, it was suggested to use async and predominantly collapse clauses to achieve optimal performance in MiniWeather kernels. Using NX=1024 and NZ=512, the most expensive kernel in terms of compute time was at Line 231 in the semi_discrete_step subroutine, with NVCOMPILER_ACC_TIME statistics highlighted below:

```
/glade/u/home/dhoward/GPU_workshop/05_DirectivesOpenACC/fortran/miniWeather_mpi_exercise2.F90 # Source file with OpenACC kernel code
semi_discrete_step NVIDIA devicenum=0 # Name of subroutine from which kernel is launched
time(us): 62,147
257: compute region reached 924 times # Specific line number for GPU kernel and number times
reached/launched
257: kernel launched 924 times
grid: [16384] block: [128] # Arrangement of gang/worker/vector in terms of grids
and blocks
device time(us): total=62,147 max=70 min=66 avg=67 # Timing statistics of the GPU kernel
elapsed time(us): total=76,527 max=87 min=80 avg=82 # Timing statistics of the CPU call (less accurate with
asynchronous execution)
257: data region reached 1848 times
```

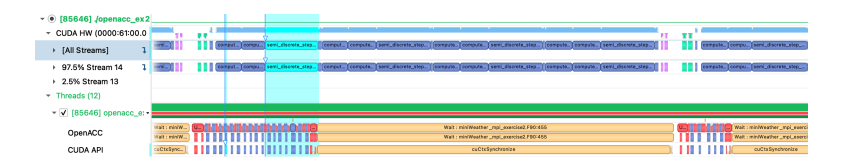
The arrangement of **gang/worker/vector** units is provided by **grid:** [NUM_GANGS] and **block:** [VECTOR_LENGTH x NUM_WORKERS]. The number of workers was 1 in the previous case so is omitted.

Running this version with the NVIDIA NSight Systems Profiler (discussed in later session), we can get a visual representation of the model runtime. You can download and view this profile using the NVIDIA NSight Systems client by downloading (SHIFT + RIGHT-CLICK) MW_baseline.nsys-rep in this folder.



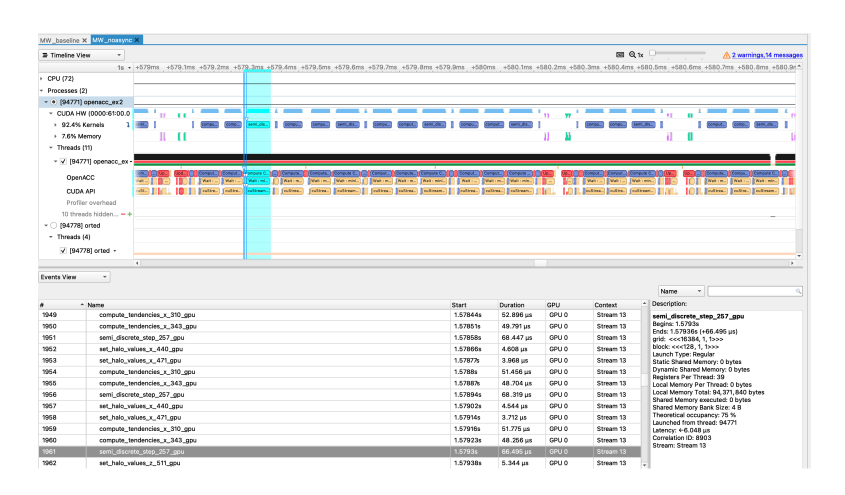
This timeline shows the kernels running on the GPU runtime in the upper **blue** compute kernels, **pink** device to host transfers, and **teal** host to device transfers segments. The lower segments show the CPU runtime in **blue** compute kernel launches, **red** data directives/regions, and beige **wait/synchronize** sections.

The bright blue highlights the most expensive GPU kernel in the semi_discrete_step
subroutine with the associated launch call from the CPU highlighted earlier in the timeline.



Since we used async, the GPU kernels run right after one another without any kernel launch/exit costs.

If we did not use async, the profile would look like this (MW_noasync.qdrep) and time would be lost as the CPU waits between every kernel launch/exit.



MiniWeather - Testing different kernel launch configurations and clauses

Recall the final exercise of the prior MiniWeather session where we experimented with various launch configurations in the miniWeather_mpi_exercise2.F90 source file for specific kernels.

Were you able to achieve any significant speed-up?

The next panels shows statistical results from some launch configuration experiments using parameters _NX=1024 , _NZ=512 , and _SIM_TIME=10 and different clauses in place of *** for the semi discrete step subroutine kernel. Note that NUM VARS=4.

1. Using worker/vector/seq on each loop respectively, the profiler shows grid: [1] block: [32x4]. Why is this arrangement the least performant?

MiniWeather Kernel L231, semi_discrete_step	Total Device Time (μs)
BaseLine (on V100) - collapse(3) auto vector_length(128)	62,936
clause - gang/worker/vector on each loop resepctively	852,859
<pre>clause - worker/vector/seq (Move NUM_VARS innermost, seq)</pre>	2,271,059
clause - gang/vector/seq (Move NUM_VARS innermost, seq)	72,584

- 1. Did you find any better configurations for this or other kernels in MiniWeather? Explain why it performed better.
- 2. Do you trust the compiler to make relatively optimal choices with minimal direction?

MiniWeather Kernel L231, semi_discrete_step	Total Device Time (μs)
BaseLine (on V100) - collapse(3) auto vector_length(128)	62,936
<pre>clause - collapse(3) vector_length(32)</pre>	100,797
<pre>clause - collapse(3) vector_length(64)</pre>	63,010
<pre>clause - collapse(3) vector_length(256)</pre>	62,990
<pre>clause - collapse(3) vector_length(512)</pre>	63,032
<pre>clause - collapse(3) vector_length(1024)</pre>	66,458

- 1. For tile(), why do you think the (32,1,NUM_VARS=4) clause was closest to the most performant?
- 2. Can you infer the condition that causes the tile() clause to produce incorrect results? Hint: What is the max warp size?

MiniWeather Kernel L231, semi_discrete_step	Total Device Time (μs)
BaseLine (on V100) - collapse(3) auto vector_length(128)	62,936
<pre>clause - tile(32,32,NUM_VARS) INCORRECT</pre>	26,992
clause - tile(32,32,1)	73,476
<pre>clause - tile(32,8,NUM_VARS)</pre>	77,124
<pre>clause - tile(32,1,NUM_VARS)</pre>	65,040
clause - tile(1024,1,1)	67,393
<pre>clause - tile(128,1,NUM_VARS)</pre>	66,421
<pre>clause - tile(128,2,NUM_VARS)</pre>	74,295
clause - tile(128,4,NUM_VARS) INCORRECT	35,999
clause - tile(*,*,*) -> 32,4,32	150,374

MPAS-Atmosphere Overview



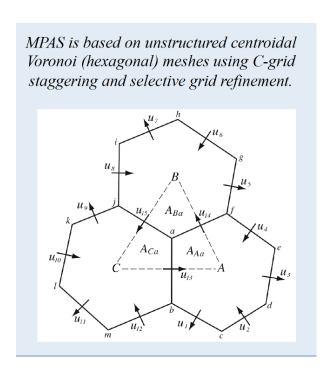


We will now look at a real world production model MPAS (Model Prediction Across Scales), specifically the GPU version of the atmosphere core MPAS-A and how this model leveraged OpenACC to refactor towards GPU devices.

So far, **only the v6.x Atmosphere core has been ported to GPUs** and is freely available to review via their website and the stable v6.x or v7.x develop-openacc branches on GitHub. Some work has also been done on the MPAS-Ocean core given this presentation by PhD student Ashwath Venkataraman.

If you'd like a more complete overview of MPAS, how to run the model, and research applications, see the 2021 MPAS Virtual tutorial page or the upcoming 2022 joint WRF/MPAS workshop.

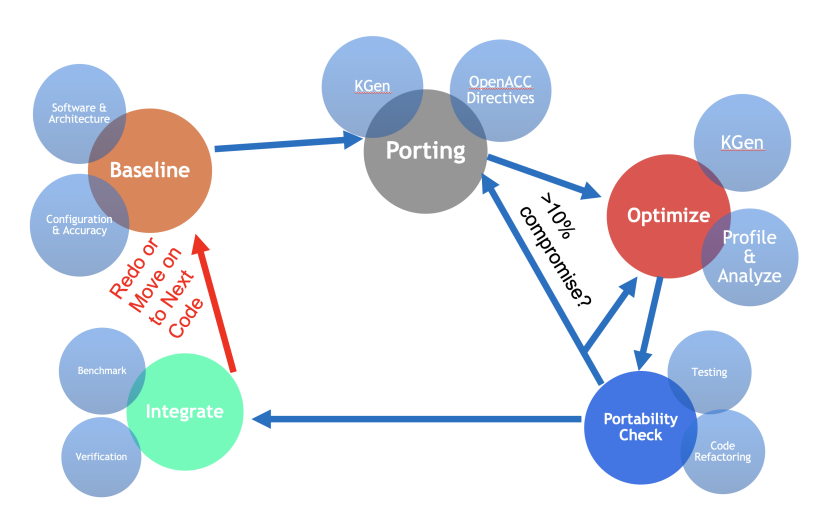
- Fully compressible non-hydrostatic equations written in flux form
- Split-Explicit timestepping via 3rd Order Runge-Kutta, see AMS Paper Klemp,
 Skamarock, and Dudhia



The MPAS-A kernels we will focus on computes coefficients for vertically implicit gravity-wave/acoustic computations needed for each Runge-Kutta timestep. The previously linked paper, specifically section 2 and the appendix, covers this in depth with a broader overview given in the 2021 tutorial Time Integration presentation.

However, understanding the numerical physics at play is not required to port well written code to GPUs.

Development Process of MPAS-A



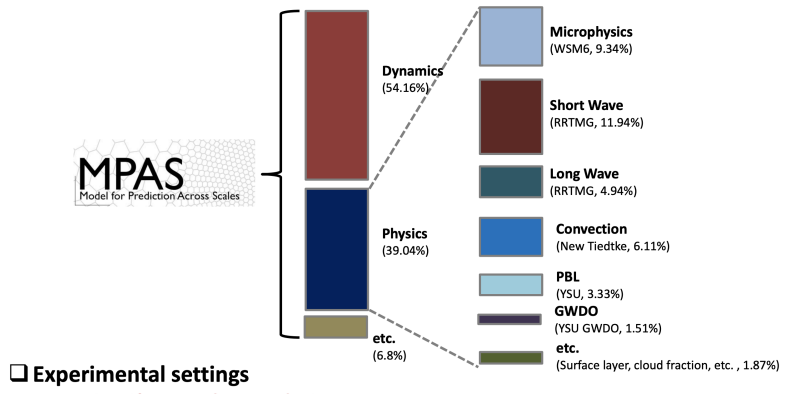
Courtesy of Raghu Raj Kumar, NVIDIA

Identifying an established iterative process for GPU development ahead of work performed significantly eases development cost and increases success outcomes.

- 1. **Establish a baseline**, ensure working and accurate configuration with target hardware and external software.
- 2. **Port the code**, using incremental addition of OpenACC, perhaps using tools for kernel extraction like KGen (Fortran only) to allow separation of concerns.
 - See KGen Guide if interested
- 3. **Optimize computationally expensive kernels** individually via an analysis and profiling iterative process.
- 4. **Check portability expectations** are met and that code satisfies both CPU and GPU unit tests.
 - Look for and eliminate any GPU anti-patterns such as linked lists data structures or global memory variables which may cause excessive data movement.
 - Repeat Steps 2-4 as needed.
- 5. **Integrate changes into benchmarks and verification suite**, utilizing version control and ideally a continuous integration process.

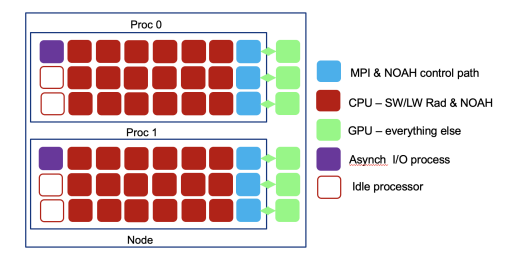
MPAS-A Performance Baseline

Getting an accurate baseline helps inform where to dedicate development effort. This can be measured using internal timing metrics or your preferred CPU profiler (like TAU, Arm Forge Map, *gprof*, etc), to **identify hotspots** in the code.

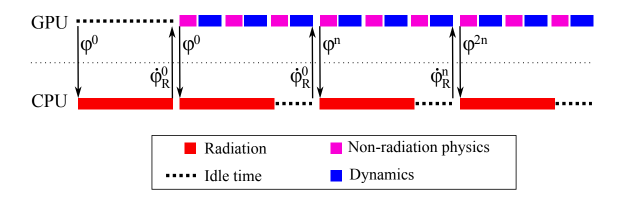


- Quasi-uniform 60-km resolution (163,842 cells)
- Δt=180 sec
- 41 vertical layers

Specific dynamics/physics schemes were prioritized for GPU while some set for CPU.



A lagged computation of radiation was established in order to utilize idle CPUs. Requires manual tuning of load balancing between number of CPU and GPU tasks.



Managing GPU Data in MPAS-A

Recall that using !\$acc kernels ... and similar directives will generate lists of variables needed to manage data movement for each compute region.

```
153, Generating implicit copyin(rdzu(:)) [if not already present]

Generating implicit copyout(cofwr(:,:)) [if not already present]

Generating implicit copyin(p(:,:)) [if not already present]

Generating implicit copyout(cofwz(:,:)) [if not already present]

Generating implicit copyin(fzp(:),t(:,:),zz(:,:),fzm(:),cqw(:,:)) [if not already present]

Generating implicit copyout(coftz(:,:)) [if not already present]
```

These lists can be used and leveraged for your own data directives as GPU development progresses.

```
do iCell=cellSolveStart,cellSolveEnd
  do i=1,nEdgesOnCell(iCell)
      iEdge = edgesOnCell(i,iCell)
      !DIR$ IVDEP
      do k = 2, nVertLevels
         flux = edgesOnCell_sign(i,iCell) * fzm(k) * u_tend(k,iEdge)
         w_tend(k,iCell) = w_tend(k,iCell) - zb_cell(k,i,iCell)
      end do
  end do
   !DIR$ IVDEP
  do k = 2, nVertLevels
      w_{tend}(k,iCell) = (fzm(k) * zz(k,iCell) + fzp(k) * zz(k-1,iCell)
  end do
end do
  !$acc data present(w_tend, &
  !$acc edgesoncell, edgesoncell_sign, fzm, fzp,nedgesoncell, u_tend, &
  !$acc zb3_cell, zb_cell, zz)
  !$acc parallel num_workers(8) vector_length(32)
  !$acc loop gang worker private(iEdge, flux)
  do iCell=cellSolveStart,cellSolveEnd
    do i=1,nEdgesOnCell(iCell)
    iEdge = edgesOnCell(i,iCell)
    !DIR$ IVDEP
       do k = 2, nVertLevels
           flux = edgesOnCell_sign(i,iCell) * fzm(k) * u_tend(k,iEdge)
           w_tend(k,iCell) = w_tend(k,iCell) - zb_cell(k,i,iCell)
        end do
    end do
  !DIR$ IVDEP
    do k = 2, nVertLevels
       w_{tend}(k,iCell) = (fzm(k) * zz(k,iCell) + fzp(k) * zz(k-1,iCell))
    end do
  end do
  !$acc end parallel
  !$acc end data
```

```
!$acc data copy(w_tend, &
!$acc edgesoncell, edgesoncell_sign, fzm, fzp,nedgesoncell, u_tend, &
!$acc zb3_cell, zb_cell, zz)
!$acc kernel
do iCell=cellSolveStart,cellSolveEnd
  do i=1,nEdgesOnCell(iCell)
     iEdge = edgesOnCell(i,iCell)
      !DIR$ IVDEP
      do k = 2, nVertLevels
        flux = edgesOnCell_sign(i,iCell) * fzm(k) * u_tend(k,iEdge)
         w_tend(k,iCell) = w_tend(k,iCell) - zb_cell(k,i,iCell)
      end do
   end do
   !DIR$ IVDEP
   do k = 2, nVertLevels
     w_{tend}(k,iCell) = (fzm(k) * zz(k,iCell) + fzp(k) * zz(k-1,iCell))
   end do
end do
!$acc end kernel
!$acc end data
```

Given ported kernels, MPAS-A was designed to create CPU and GPU data copies at initialization via !\$acc declare create(...) and copy data at unstructured data regions via !\$acc enter data copyin(...) prior to each kernel call. Then, each kernel would only require a present(...) clause using the prior variable lists. **Reference counters** would mitigate excessive copies.

Any lingering excessive data copies could be identified by profilers and fixed while other required copies for CPU algorithms & I/O were managed by !\$acc update directives.

MPAS-A Kernel Extraction

We will focus on the atm_compute_vert_imp_coefs_work subroutine and kernels as extracted by Supreeth Suresh, TDD/ASAP in CISL. This is the link, Line 2641 to the source subroutine in the full model codebase and in this workshop directory is the the extracted set of kernels mpas_atm_compute_vert_imp_coefs_work.F90.

Assuming data locality is resolved, this extracted kernel simply utilizes **randomized input data** as we will be **focusing on optimizing the performance** of the subroutine's kernels. The kernel is run in a repeating loop so we can get a relatively consistent average of measured performance. A validation tool has not been included at this time but is typically highly recommended.

For large codebases, building and/or using an automated tool like NCAR's KGen for Fortran codes or Kernel Tuner from NL eScience Center for CUDA/OpenCL codes will likely speed up the development/optimization process.

EXERCISE: MPAS-A Kernel Optimization

Open the mpas_atm_compute_vert_imp_coefs_work.F90 source file and convert the !\$acc kernels loops to optimized !\$acc parallel ... compute constructs. Analyze each set of loops and apply appropriate sets of kernel configuration clauses to achieve improved performance. Note: !DIR\$ IVDEP tells compiler to ignore loop dependencies for serial vector SIMD compilations.

You are encouraged to reference the initial attempts at optimization done by the !\$acc kernels directive output during the compilation process. Data management has already been done for you using -gpu=managed and present(var-list) / create(var-list) clauses.

Record results of your optimization experiments on a chosen kernel and try to determine optimal configurations for that kernel. Compare your achieved performance with the original at Line 2641. Work on other kernels as time allows. Note that most kernels may benefit from similar clause specifications since they operate on similar domain sizes/variables.

In []:

module load nvhpc/22.2 &> /dev/null
export _OPENACC=true
make

```
In []:
    module load nvhpc/22.2 &> /dev/null
    export _OPENACC=true
    make

In []:
    qcmd -l select=l:ncpus=l:ngpus=l -l gpu_type=v100 -v NVCOMPILER_ACC_TIME=l -- \
    `pwd`/vert_implicit_coefs.exe
```

In []:
 module load nvhpc/22.2 &> /dev/null
 export _OPENACC=true
 make

In []:
 qcmd -l select=1:ncpus=1:ngpus=1 -l gpu_type=v100 -v NVCOMPILER_ACC_TIME=1 -- \
 `pwd`/vert_implicit_coefs.exe

MPAS-A Kernels L###	Device Time (μs)
BaseLine (on V100) - !\$acc kernels	XX
clause - gang/vector	XX
clause - tile(##,##)	XX
<pre>clause - tile(*,*)</pre>	XX
clause - vector_length(XX)	XX
clause - num_workers(XX)	XX
•••	XX

Final Points

- 1. Plan for and **commit to a defined iterative GPU development process** to remove pain points and manage long term goals of your code project
 - Smaller, validated incremental changes are easier to debug
- 2. Start with descriptive !\$acc kernels then add prescriptive !\$acc parallel ... kernels as needed for expensive kernels
 - !\$acc kernels can still achieve meaningful performance alone
- 3. Understand that the GPU development process takes time and effort but **specific tools/techniques can drastically speed up development time**.

Suggested Resources

- 2021 MPAS Virtual tutorial
- Computers & Geosciences, GPU acceleration of MPAS microphysics WSM6 using OpenACC directives: Performance and verification by J. Kim, J. Kang, and M. Joh (KISTI)
- OpenACC.org and NVIDIA managed GitHub, presentations, and learning materials
 GPU Bootcamps
 - Lab sequence on OpenACC
 - Lab sequence on Profiling Tools with MiniWeather
 - Lab sequence on Various GPU Programming Paradigms (CUDA, OpenACC, stdPar, OpenMP)
 - Lab sequence on Multi-GPU Programming
 - Lab sequences on GPU AI with CFD, PINNs, and Climate models

After this session, we will have three weeks until the next workshop. Order of upcoming sessions will also be adjusted to accomodate availability of a NVIDIA engineer to present on Multi-GPU programming. Look out for upcoming announcements.