# N-WAYS GPU BOOTCAMP OPENACC



### OPENACC

#### What to expect?

- Basic introduction to OpenACC directives
- HPC SDK Usage
- Portability across Multicore and GPU



# OpenACC is...

a directives-based

### parallel programming model designed for

performance and portability.

Add Simple Compiler Directive main() <serial code> **#pragma acc kernels** <parallel code> OpenACC



#### **GAUSSIAN 16**



of our efforts.

E3SM

Image coachesy Alabitis

#### **ANSYS FLUENT**

We've effectively used OpenACC for heterogeneous computing in ANSYS Fluent with impressive performance. We're now applying this work to more of our models and new platforms.

#### NUMECA FINE/Open



Porting our unstructured C++ CFD solver FINE/Open to GPUs using OpenACC would have been impossible two or three years ago, but OpenACC has developed enough that we're now getting some really good results.

were able to achieve the acceleration needed for

#### **IBM-CFD**

GPUs.



OpenACC can prove to be a handy loof for tional engineers and researchers to CED, we have obtained order of magnitude improve the overall scalability of the code

#### VASP

For VASP, OpenACC is the way forward for GPU acceleration. Performance is similar and in some cases better than CLIDA C and OpenACC dramatically decreases GPU development and maintenance efforts. We're excited to collaborate with NVIDIA and PGI as an early adopter of CUDA Unified Memory.



Using OpenACC, we've GPUaccelerated the Synopsys TCAD Sentaurus Device EMW simulator to speed up optical simulations of image sensors. GPUs are key to improving simulation throughput in the design of advanced image sensors.



Our team has been evaluating OpenACC as a pathway to performance portability for the Model for Prediction (MPAS) atmospheric model. Using this approach on the MPAS dynamical core, we have achieved performance on a single P100 GPU equivalent to 2.7 dual socketed Intel Xeon nodes on our new Chevenne supercomputer





MAS



Adding OpenACC into MAS has given us the ability to migrate medium-sized simulations from a multi node CPU cluster to a single multi-GPU server. The implementation yielded a portable single-source code for both CPU and GPU runs. Future work will add OpenACC to the remaining model features, enabling GPU-accelerated realistic solar storm modeling.

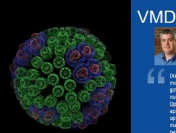
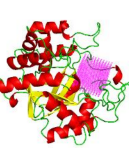


Image courteen Oak Ridge National Laboratory





SANJEEVINI

In an academic environment

provides a great platform for

entire computational task.

maintenance and speedup of existing

computational scientists to accomplish

efforts or manpower in speeding up the

both tasks without involving a lot of

codes is a tedious task. OpenACC

The CAAR project provided us with

early access to Summit hardware and

access to PGI compiler experts. Both

of these were critical to our success.

PGI's OpenACC support remains the

much more intrusive programming

model approaches.

best available and is competitive with

55

approach to getting at least some speedup out of these second-tier routines. In many cases it's completely sufficient because with the current algorithms, GPU performance is bandwidth-bound.









and effort in learning to program



components of our legacy codes to GPU. Especially the routines involving search algorithm and matrix solvers have been well-accelerated to

**OpenACC** 

More Science, Less Programming



CUDA Fortran gives us the full performance potential of the CUDA programming model and NVIDIA GPUs. White leveraging the potential of explicit data movement, ISCUF KERNELS directives give us productivity and source code maintainability. It's the best of both worlds











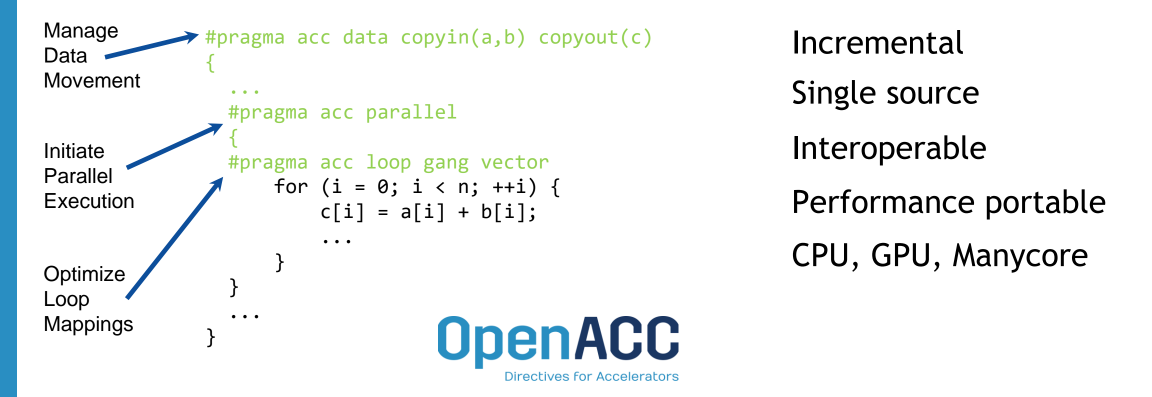
COSMO

develop for GPU-based hardware while retaining a single source for almost all the COSMO physics code

"

mage courtesy: NCAR

# **OpenACC** Directives





# OPENACC SYNTAX

### Syntax for using OpenACC directives in code

C/C++			
<pre>#pragma <code></code></pre>	acc	directive	clauses

Fortran
---------

!\$acc directive clauses
<code>

A *pragma* in C/C++ gives instructions to the compiler on how to compile the code. Compilers that do not understand a particular pragma can freely ignore it.

A *directive* in Fortran is a specially formatted comment that likewise instructions the compiler in it compilation of the code and can be freely ignored.

"acc" informs the compiler that what will come is an OpenACC directive

*Directives* are commands in OpenACC for altering our code.

Clauses are specifiers or additions to directives.

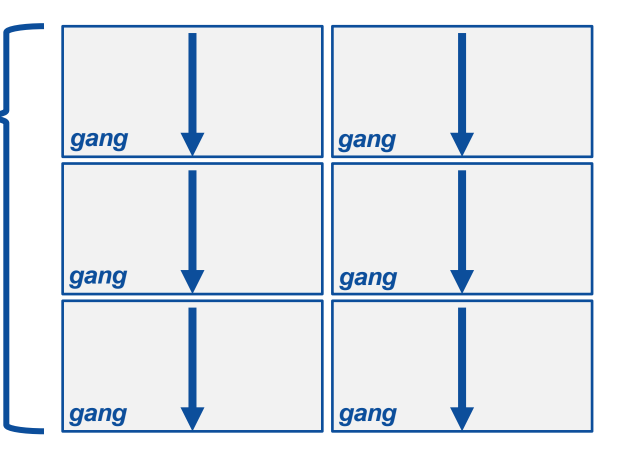
OpenAC

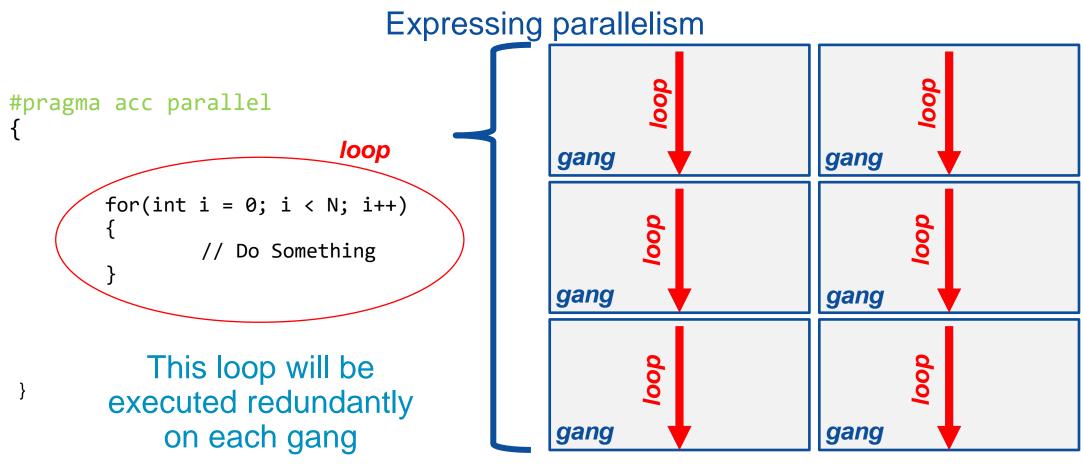
### Expressing parallelism



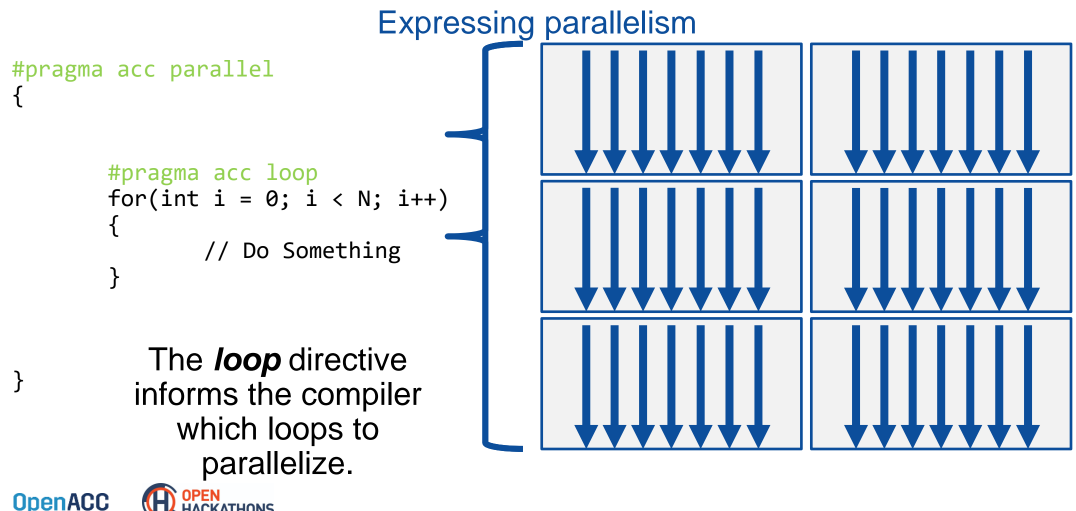
OpenACC

When encountering the *parallel* directive, the compiler will generate *1 or more parallel gangs*, which execute redundantly.









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#### Parallelizing a single loop

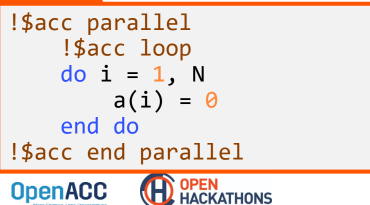
#### C/C++

```
#pragma acc parallel
```

# Use a **parallel** directive to mark a region of code where you want parallel execution to occur

This parallel region is marked by curly braces in C/C++ or a start and end directive in Fortran

#### Fortran



The **loop** directive is used to instruct the compiler to parallelize the iterations of the next loop to run across the parallel gangs

#### Combined directive for parallelising a single loop

C/C++		
<pre>#pragma for(int</pre>	acc i =	<pre>parallel loop 0; j &lt; N; i++) = 0;</pre>

#### Fortran

This pattern is so common that you can do all of this in a single line of code

In this example, the parallel loop directive applies to the next loop

This directive both marks the region for parallel execution and distributes the iterations of the loop.

When applied to a loop with a data dependency, parallel loop may produce incorrect results



# BUILD AND RUN THE CODE



## NVIDIA HPC SDK

- Comprehensive suite of compilers, libraries, and tools used to GPU accelerate HPC modeling and simulation application
- The NVIDIA HPC SDK includes the new NVIDIA HPC compiler supporting OpenACC C and Fortran
  - The command to compile C code is 'nvc'
  - The command to compile C++ code is 'nvc++'
  - The command to compile Fortran code is 'nvfortran'

nvc –fast –Minfo=accel –ta=tesla:managed main.c

OpenACC

nvfortran –fast –Minfo=accel –ta=tesla:managed main.f90

## **BUILDING THE CODE**

-Minfo shows more details

\$ nvc -fast -ta=multicore -Minfo=accel laplace2d\_uvm.c
main:

- 63, Generating Multicore code
  - 64, #pragma acc loop gang
- 64, Accelerator restriction: size of the GPU copy of Anew, A is unknown Generating reduction(max:error)
- 66, Loop is parallelizable



# RDF

### Pseudo Code - C

for (int frame=0;frame<nconf;frame++) {</pre>

```
for(int id1=0;id1<numatm;id1++) {</pre>
```

```
for(int id2=0;id2<numatm;id2++) {
    dx=d_x[]-d_x[];
    dy=d_y[]-d_y[];
    dz=d_z[]-d_z[];
    r=sqrtf(dx*dx+dy*dy+dz*dz);</pre>
```

```
if (r<cut) {
    ig2=(int)(r/del);
    d_g2[ig2] = d_g2[ig2] +1;
}</pre>
```

OpenACC

Across Frames

• Find Distance

• Reduction

# RDF

### Pseudo Code - C

for (int frame=0;frame<nconf;frame++) {
 #pragma acc parallel loop</pre>

```
for(int id1=0;id1<numatm;id1++) {</pre>
```

for(int id2=0;id2<numatm;id2++) {
 dx=d\_x[]-d\_x[];
 dy=d\_y[]-d\_y[];
 dz=d\_z[]-d\_z[];
 r=sqrtf(dx\*dx+dy\*dy+dz\*dz);</pre>

if (r<cut) {
 ig2=(int)(r/del);
 #pragma acc atomic
 d\_g2[ig2] = d\_g2[ig2] +1;</pre>

THONS

**OpenACC** 

Parallel loop construct

• Atomic construct

### RDF Pseudo Code - Fortran

```
do iconf=1,nframes
     if (mod(iconf,1).eq.0) print*,iconf
     !$acc parallel loop
     do i=1,natoms
       do j=1,natoms
         dx=x(iconf,i)-x(iconf,j)
         dy=y(iconf,i)-y(iconf,j)
         dz=z(iconf,i)-z(iconf,j)
         if(r<cut)then
           !$acc atomic
           g(ind)=g(ind)+1.0d0
         endif
       enddo
     enddo
   enddo
```

• Parallel Loop construct

#### Atomic Construct



### REFERENCES

https://www.openacc.org/get-started

https://developer.nvidia.com/hpc-sdk



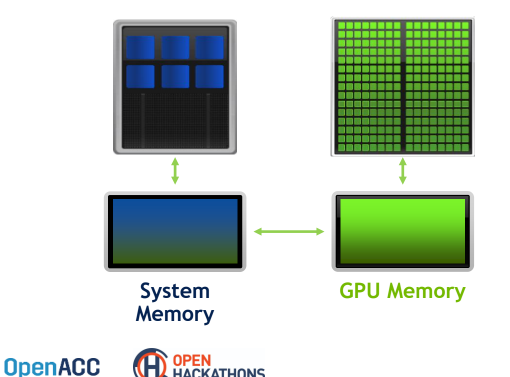
# Memory management



# CUDA UNIFIED MEMORY

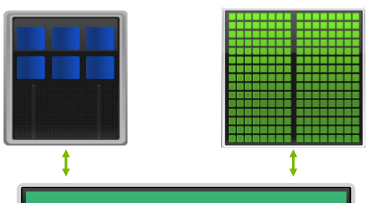
Simplified Developer Effort

Without Managed Memory



Commonly referred to as *"managed memory."* 

#### With Managed Memory



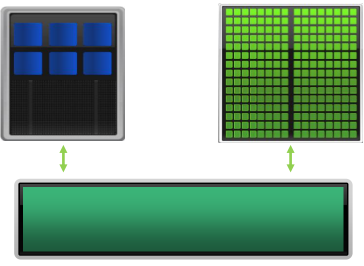
CPU and GPU memories are combined into a single, shared pool

# MANAGED MEMORY

### Limitations

- The programmer may be able to get better performance by manually handling data transfers
- Memory allocation/deallocation takes longer with managed memory

#### With Managed Memory



#### Managed Memory



### DATA CLAUSES

**copy**(*list*) Allocates memory on GPU and copies data from host to GPU when entering region and copies data to the host when exiting region.

**Principal use:** For many important data structures in your code, this is a logical default to input, modify and return the data.

**copyin(** *list* ) Allocates memory on GPU and copies data from host to GPU when entering region.

**Principal use** Think of this like an array that you would use as just an input to a subroutine.

copyout ( *list* ) Allocates memory on GPU and copies data to the host when exiting region.

**Principal use:** A result that isn't overwriting the input data structure.

**create**(*list*) Allocates memory on GPU but does not copy.

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## ARRAY SHAPING

Sometimes the compiler needs help understanding the shape of an array

The first number is the start index of the array

In C/C++, the second number is how much data is to be transferred

In Fortran, the second number is the ending index

copy(array[starting\_index:length]) C/C++
copy(array(starting\_index:ending\_index)) Fortran



## ARRAY SHAPING (CONT.)

#### Multi-dimensional Array shaping

copy(array[0:N][0:M])

Both of these examples copy a 2D array to the device

copy(array(1:N, 1:M))

Fortran

### OPENACC DATA DIRECTIVE Definition

The data directive defines a lifetime for data on the device beyond individual loops

During the region data is essentially "owned by" the accelerator

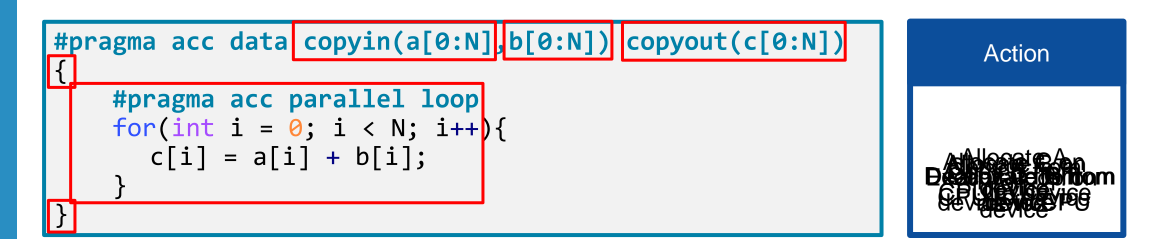
Data clauses express shape and data movement for the region

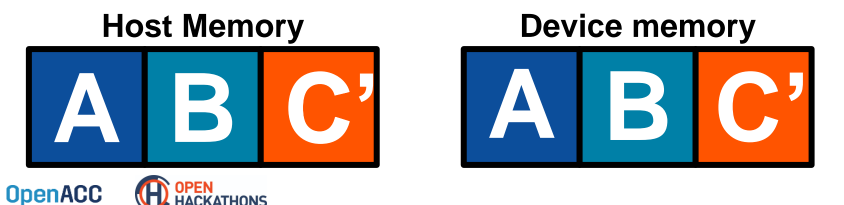
#pragma acc <b>data</b> <i>clauses</i> {
< Sequential and/or Parallel code >
}
!\$acc data clauses

< Sequential and/or Parallel code >

\$acc end data

### STRUCTURED DATA DIRECTIVE Example





## Acknowledgment

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