

## INTRODUCTION TO OPENACC JSC ADVANCED CUDA COURSE 2023

23 June 2023 | Andreas Herten | Forschungszentrum Jülich



#### **Outline**

OpenACC
History
OpenMP
Modus Operandi
OpenACC's Models

OpenACC Workflow Identify Parallelism Parallelize Loops parallel loops **Nisght Systems** kernels **Data Transfers GPU Memory Spaces Portability** Clause: copy **Nsight Systems Data Locality** Analyse Flow data enter data

Routines

OpenACC by Example

Other Directives
Clause: gang
Conclusions
List of Tasks



## **OpenACC Mission Statement**

[...] OpenACC [is] for writing parallel programs in C, C++, and Fortran that run identified regions in parallel on multicore CPUs or attached accelerators.

[...] a model for parallel programming that is portable across operating systems and various types of multicore CPUs and accelerators.

- OpenACC API Documentation ⚠, openacc.org



## **OpenACC History**

- 2011 OpenACC 1.0 specification is released at SC11 \( \begin{aligned} \ NVIDIA, Cray, PGI, CAPS \end{aligned} \)
- 2013 OpenACC 2.0: More functionality, portability 🔀
- 2015 OpenACC 2.5: Enhancements, clarifications
- 2017 OpenACC 2.6: Deep copy, ... 🔄
- 2019 OpenACC 3.0: Newer C++, more lambdas, ... 🔁 🖹
- 2022 OpenACC 3.3: Gang Parallelism, Nested Loops, ... 🖾 🖹
  - Run as a non-profit organization, OpenACC.org
  - Members from industry and academia
  - ightarrow https://www.openacc.org/ (see also: Best practice guide oxinesigne)

## OpenACC-enabled Applications

- ANSYS Fluent
- Gaussian
- VASP
- COSMO
- GTC
- SOMA
- ٠...



## **Open**{MP↔ACC}

#### **Everything's connected**

- OpenACC modeled after OpenMP ...
- ... but specific for accelerators
- OpenMP 4.0/4.5: Offloading; compiler support improving (Clang, XL, GCC, ...)
- OpenACC more descriptive, OpenMP more prescriptive
- OpenMP 5.0: Descriptive directive loop
- Same basic principle: Fork/join model

Master thread launches parallel child threads; merge after execution

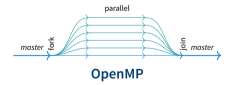


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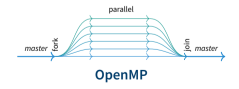


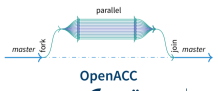
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**Modus Operandi** 

OpenACC

## **OpenACC Acceleration Workflow**

Three-step program

- 1 Annotate code with directives, indicating parallelism
- OpenACC-capable compiler generates accelerator-specific code
- 3 \$uccess





pragmatic

Compiler directives state intend to compiler

```
C/C++
#pragma acc kernels
for (int i = 0; i < 23; i++)
// ...</pre>
```

#### **Fortran**

```
!$acc kernels
do i = 1, 24
! ...
!$acc end kernels
```

- Ignored by compiler which does not understand OpenACC
- OpenACC: Compiler directives, library routines, environment variables
- Portable across host systems and accelerator architectures





- Trust compiler to generate intended parallelism; always check status output!
- No need to know details of accelerator; leave it to expert compiler engineers Tuning possible
- ullet One code can target different accelerators: GPUs, CPUs o Portability



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Compiler	Targets	Languages	oss	Free	Comment
NVIDIA HPC SDK	NVIDIA GPU, CPU	C, C++, Fortran	No	Yes	Best performance
GCC	NVIDIA GPU, AMD GPU	C, C++, Fortran	Yes	Yes	
Clang/LLVM	CPU, NVIDIA GPU	C, C++. Fortran	Yes	Yes	Via Clang OpenMP backend



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## 2 Compiler

#### Flags and options

OpenACC compiler support: activate with compile flag

```
NVHPC nvc -acc
```

- -acc=gpu|multicore Target GPU or CPU
- -acc=gpu -gpu=cc80 Generate Ampere-compatible code
- -gpu=cc80,lineinfo Add source code correlation into binary
  - -gpu=managed Use unified memory
  - -Minfo=accel Print acceleration info

#### GCC gcc -fopenacc

- -fopenacc-dim=geom Use geom configuration for threads
- -foffload="-lm -03" Provide flags to offload compiler
  - -fopt-info-omp Print acceleration info





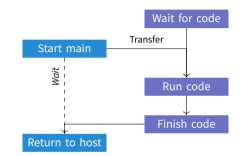
# Expose Parallelism Measure Compile

- Serial to parallel: fast
- Serial to fast parallel: more time needed
- Start simple  $\rightarrow$  refine
- Expose more and more parallelism
- **⇒** Productivity
  - Because of generality: Sometimes not last bit of hardware performance accessible
  - But: Use OpenACC together with other accelerator-targeting techniques (CUDA, libraries, ...)

## **OpenACC Accelerator Model**

For computation and memory spaces

- Main program executes on host
- Device code is transferred to accelerator
- Execution on accelerator is started
- Host waits until return (except: async)
- Two separate memory spaces; data transfers back and forth
  - Transfers hidden from programmer
  - Memories not coherent!
  - Compiler helps; GPU runtime helps







## A Glimpse of OpenACC

```
#pragma acc data copy(x[0:N],y[0:N])
#pragma acc parallel loop
{
    for (int i=0; i<N; i++) {
        x[i] = 1.0;
        y[i] = 2.0;
    }
    for (int i=0; i<N; i++) {
        y[i] = i*x[i]+y[i];
    }
}</pre>
```

```
!$acc data copy(x(1:N),y(1:N))
!$acc parallel loop
   do i = 1. N
       x(i) = 1.0
       v(i) = 2.0
   end do
   do i = 1. N
       v(i) = i*x(i)+v(i):
   end do
!$acc end parallel loop
```

!\$acc end data

# OpenACC by Example

## **Parallelization Workflow**

Member of the Helmholtz Association

Identify available parallelism

Parallelize loops with OpenACC

Optimize data locality

Optimize loop performance



#### Algorithmic description

- Example for acceleration: Jacobi solver
- Iterative solver, converges to correct value
- Each iteration step: compute average of neighboring points
- Example: 2D Poisson equation:  $\nabla^2 A(x,y) = B(x,y)$



$$A_{k+1}(i,j) = -\frac{1}{4} \left( B(i,j) - (A_k(i-1,j) + A_k(i,j+1), +A_k(i+1,j) + A_k(i,j-1)) \right)$$





```
while ( error > tol && iter < iter_max ) {
    error = 0.0:
    for (int ix = ix start; ix < ix end; ix++) {
        for (int iv = iv start: iv < iv end: iv++) {</pre>
            Anew[iy*nx+ix] = -0.25 * (rhs[iy*nx+ix] -
                ( A[iv*nx+ix+1] + A[iv*nx+ix-1]
               + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]);
            error = fmaxr(error. fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
    }}
    for (int iv = iv start: iv < iv end: iv++) {
        for( int ix = ix start; ix < ix end; ix++ ) {
            A[iv*nx+ix] = Anew[iv*nx+ix]:
    }}
    for (int ix = ix start; ix < ix end; ix++) {
            A[0*nx+ix] = A[(nv-2)*nx+ix]:
            A[(ny-1)*nx+ix] = A[1*nx+ix];
    // same for iv
    iter++:
```

#### Source code

```
while ( error > tol && iter < iter_max ) {●
    error = 0.0:
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            error = fmaxr(error. fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
    }}
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Iterate until converged

```
Iterate until converged
while ( error > tol && iter < iter_max ) {●
    error = 0.0:
                                                                              Iterate across
    for (int ix = ix start; ix < ix end; ix++) {
        for (int iy = iy start; iy < iy end; iy++) {
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```
Iterate until converged
while ( error > tol && iter < iter_max ) {●
    error = 0.0:
                                                                               Iterate across
    for (int ix = ix_start; ix < ix_end; ix++) {</pre>
        for (int iy = iy start; iy < iy end; iy++) {
            Anew[iy*nx+ix] = -0.25 * (rhs[iy*nx+ix] -)
                ( A[iv*nx+ix+1] + A[iv*nx+ix-1]
                                                                           Calculate new value
               + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]);
            error = fmaxr(error. fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
    }}
    for (int iv = iv start: iv < iv end: iv++) {
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               + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]);
            error = fmaxr(error, fabsr(Anew[iy*nx+ix]-A[iy*nx+ix]));
    }}
                                                                             Accumulate error
    for (int iv = iv start: iv < iv end: iv++) {
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                                                                            Calculate new value
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    }}
                                                                              Accumulate error
    for (int iv = iv start: iv < iv end: iv++) {
        for( int ix = ix start; ix < ix end; ix++ ) {</pre>
            A[iv*nx+ix] = Anew[iv*nx+ix]:
                                                                             Swap input/output
    }}
    for (int ix = ix start; ix < ix end; ix++) {
            A[0*nx+ix] = A[(nv-2)*nx+ix]:
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Iterate until converged
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                                                                              Swap input/output
    }}
    for (int ix = ix start; ix < ix end; ix++) \{
            A[0*nx+ix] = A[(nv-2)*nx+ix]:
            A[(ny-1)*nx+ix] = A[1*nx+ix];
                                                                         Set boundary conditions
    // same for iv
    iter++;
                                            23 June 2023
                                                        Slide 16174
```

#### **Parallelization Workflow**

Identify available parallelism

Parallelize loops with OpenACC

Optimize data locality

Optimize loop performance



## **Profiling**

**Profile** 

[...] premature optimization is the root of all evil.

- Donald Knuth [3]
- Investigate hot spots of your program!
- $\rightarrow$  Profile!
  - Many tools, many levels: perf, PAPI, Score-P, Intel Advisor, NVIDIA profilers, ...
  - Here: Examples from GCC



## **Profiling**

**Profile** 

```
[...] premature optimization is the root of all evil.

Yet we should not pass up our [optimization] opportunities [...]

- Donald Knuth [3]
```

- Investigate hot spots of your program!
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  - Many tools, many levels: perf, PAPI, Score-P, Intel Advisor, NVIDIA profilers, ...
  - Here: Examples from GCC



## **Profile of Application**

```
. . .
$ gcc -g -pg -DUSE DOUBLE -c -o poisson2d reference.o poisson2d reference.c
  gcc -g -pg -DUSE DOUBLE -lm poisson2d reference.o poisson2d.c -o poisson2d
$ gprof -p -l ./poisson2d gmon.out
  Flat profile:
  Each sample counts as 0.01 seconds.
       cumulative self
                                    self
                                          total
   time
         seconds seconds calls Ts/call Ts/call
                                                   name
   46.29
             1.28
                 1.28
                                                   main (poisson2d.c:107 @ 40135c)
   30.01 2.11 0.83
                                                   main (poisson2d.c:108 @ 4013cd)
   12.66 2.46
                 0.35
                                                   main (poisson2d.c:109 @ 401458)
   6.15 2.63
                     0.17
                                                   main (poisson2d.c:107 @ 401421)
```

- Very simple here: All in main
- Lines 107, 108, 109: within the inner grid loop
- Good position to start! Let's study this further in independency analysis

## **Code Independency Analysis**

#### Independence is key

Member of the Helmholtz Association

```
while ( error > tol δδ iter < iter max ) {
    error = 0.0:
    for (int ix = ix start: ix < ix end: ix++) {
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    }}
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            A[iy*nx+ix] = Anew[iy*nx+ix];
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    for (int ix = ix start; ix < ix end; ix++) {
            A[0*nx+ix] = A[(nv-2)*nx+ix]:
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    // same for iv
    iter++:
```

Slide 20174

## **Code Independency Analysis**

```
Independence is key
                                                                                    Data dependency
     while ( error > tol && iter < iter_max ) {●
         error = 0.0:
         for (int ix = ix start; ix < ix end; ix++) {
              for (int iv = iv start; iv < iv end; iv++) {</pre>
                  Anew[iv*nx+ix] = -0.25 * (rhs[iv*nx+ix] -
                      ( A[iv*nx+ix+1] + A[iv*nx+ix-1]
                     + A[(iy-1)*nx+ix] + A[(iy+1)*nx+ix]);
                  error = fmaxr(error, fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
         }}
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                  A[iy*nx+ix] = Anew[iy*nx+ix];
         }}
         for (int ix = ix start; ix < ix end; ix++) {</pre>
                  A[0*nx+ix] = A[(nv-2)*nx+ix]:
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         // same for iv
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```

#### **Parallelization Workflow**

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Parallelize loops with OpenACC

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## Parallel Loops: Parallel

#### An important directive

- Programmer identifies block containing parallelism
  - → compiler generates offload code
- Program launch creates gangs of parallel threads on parallel device
- Implicit barrier at end of parallel region
- Each gang executes same code sequentially



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- Each gang executes same code sequentially



# Parallel Loops: Parallel

Clauses

```
Diverse clauses to augment the parallel region
```

```
private(var) A copy of variables var is made for each gang
```

```
firstprivate(var) Same as private, except var will initialized with value from host
```

```
if(cond) Parallel region will execute on accelerator only if cond is true
```

reduction(op:var) Reduction is performed on variable var with operation op; supported:

```
+ * max min ...
```

async[(int)] No implicit barrier at end of parallel region



# Parallel Loops: Loops

Also an important directive

- Programmer identifies loop eligible for parallelization
- Directive must be directly before loop
- Optional: Describe type of parallelism



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# Parallel Loops: Loops

Clauses

# Parallel Loops: Parallel Loops

Maybe the most important directive

- Combined directive: shortcut
   Because its used so often
- Any clause that is allowed on parallel or loop allowed
- Restriction: May not appear in body of another parallel region

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- Restriction: May not appear in body of another parallel region

```
✓ OpenACC: parallel loop

#pragma acc parallel loop [clause, [, clause] ...]
```

# **Parallel Loops Example**

```
double sum = 0.0;
#pragma acc parallel loop
for (int i=0; i<N; i++) {
    x[i] = 1.0;
    y[i] = 2.0;
}

#pragma acc parallel loop reduction(+:sum)
for (int i=0; i<N; i++) {
    y[i] = i*x[i]+y[i];
    sum+=y[i];
}</pre>
```

# Parallel Loops Example

```
double sum = 0.0;
                                                  sum = 0.0
#pragma acc parallel loop
                                                  !$acc parallel loop
for (int i=0; i<N; i++) {
                                                  do i = 1. N
                                                      x(i) = 1.0
    x[i] = 1.0:
   v[i] = 2.0:
                                                      v(i) = 2.0
                                                  end do
                                                  !$acc end parallel loop
#pragma acc parallel loop reduction(+:sum)
                                                  !$acc parallel loop reduction(+:sum)
for (int i=0; i<N; i++) {
                                                  do i = 1. N
    y[i] = i*x[i]+y[i];
                                                      v(i) = i * x(i) + v(i)
                                                                                          Kernel 2
    sum+=v[i];
                                                      sum+=v(i)
                                                  end do
                                                  !$acc end parallel loop
```



#### TASK 2

#### Add parallelism

- Add OpenACC parallelism to main double loop in Jacobi solver source code
- → Congratulations, you are a GPU developer!

#### Task 2: A First Parallel Loop

- Change to Task2/ directory
- Compile: make
- Submit parallel run to the batch system:make run

Adapt the srun call and run with other number of iterations, matrix sizes

#### Fortran

- All tasks available in Fortran: exercises/Fortran/Task2/
- Fortran *much* faster than C
- Slides follow C results
- Fortran: No command line options parsed

#### Source Code

```
#pragma acc parallel loop reduction(max:error)
110
    for (int ix = ix start; ix < ix end; ix++)
111
112
      for (int iy = iy start; iy < iy end; iy++)
113
114
          Anew[iv*nx+ix] = -0.25 * (rhs[iv*nx+ix])
115
                                     (A[iv*nx+ix+1] + A[iv*nx+ix-1]
116
                                     + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]):
117
            error = fmaxr( error. fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
118
119
120
```

#### **Compilation result**

```
$ make
nvc -DUSE DOUBLE -Minfo=accel -O1 -acc=gpu -gpu=managed poisson2d.c poisson2d reference.o
  -o poisson2d
 poisson2d reference.o -o poisson2d
poisson2d.c:
main.
    106. Generating Tesla code
        110. #pragma acc loop gang, vector(128) /* blockIdx.x threadIdx.x */
             Generating reduction(max:error)
        112, #pragma acc loop seq
    106. Generating implicit copvin(A[:]) [if not already present]
         Generating implicit copy(error) [if not already present]
         Generating implicit copyin(rhs[:]) [if not already present]
    112, Complex loop carried dependence of Anew-> prevents parallelization
         Loop carried dependence of Anew-> prevents parallelization
```

Run result

```
$ make run
srun --gres=gpu:1 --time 0:10:00 --pty ./poisson2d
Jacobi relaxation calculation: max 500 iterations on 2048 x 2048 mesh
Calculate reference solution and time with serial CPU execution.
    0. 0.249999
  100. 0.249760
  200. 0....
Calculate current execution.
    0. 0.249999
  100, 0.249760
  200. 0....
2048x2048: Ref: 105.6753 s, This: 14.0692 s, speedup:
                                                        7.51
```



# **Nsight Systems**

**NVIDIA's Application Profiler** 

- Profiler for GPU applications
- CLI and GUI (timeline view)
- Sister tool: Nsight Compute (kernel profiler)
- More: tomorrow in dedicated session



## Profile of Jacobi

#### With nsys

```
$ make profile
srun --gres=gpu:1 --time 0:10:00 --pty nsys nyprof ./poisson2d 10
CUDA APT Statistics:
 Time(%) Total Time (ns) Num Calls Average Minimum Maximum
                                                                   Name
   90.9 160.407.572 30 5.346.919.1 1.780 25.648.117 cuStreamSynchronize
CUDA Kernel Statistics:
 Time(%) Total Time (ns) Instances Average
                                          Minimum Maximum
                                                                  Name
  100.0 158,686,617 10 15,868,661.7 14,525,819 25,652,783 main_106_gpu
               25.120
                                  2,512.0 2,304 3,680 main_106_gpu__red
    0.0
                           10
```

## **Profile of Jacobi**

With nsys

```
. . .
                     Only one function is parallelized!
Let's do the rest!
$ make profile
srun --gres=gpu:1 --time 0:10:00 --ptv nsys nynre
CUDA API Statistics:
 Time(%) Total Time
    90.9
             160.407
CUDA Kernel Statistics
 Time(%)
         Total Time (ray Instances
                                     Average
                                                 Minimum
                                                            Maximum
                                                                            Name
   100.0
             158,686,617
                                10 15,868,661.7 14,525,819 25,652,783 main 106 gpu
     0.0
                  25,120
                                10
                                        2,512.0
                                                     2,304
                                                                       main 106 gpu red
```

## More Parallelism: Kernels

#### More freedom for compiler

- Kernels directive: second way to expose parallelism
- Region may contain parallelism
- Compiler determines parallelization opportunities
- → More freedom for compiler
  - Rest: Same as for parallel

```
✓ OpenACC: kernels
```

#pragma acc kernels [clause, [, clause] ...]



# **Kernels Example**

```
double sum = 0.0;
#pragma acc kernels
{
    for (int i=0; i<N; i++) {
        x[i] = 1.0;
        y[i] = 2.0;
}
for (int i=0; i<N; i++) {
        y[i] = i*x[i]+y[i];
        sum+=y[i];
}</pre>
```

Kernels created here



# kernels vs. parallel

• Both approaches equally valid; can perform equally well



# kernels vs. parallel

- Both approaches equally valid; can perform equally well
- kernels
  - Compiler performs parallel analysis
  - Can cover large area of code with single directive
  - Gives compiler additional leeway
- parallel
  - Requires parallel analysis by programmer
  - Will also parallelize what compiler may miss
  - More explicit
  - Similar to OpenMP



# kernels vs. parallel

- Both approaches equally valid; can perform equally well
- kernels
  - Compiler performs parallel analysis
  - Can cover large area of code with single directive
  - Gives compiler additional leeway
- parallel
  - Requires parallel analysis by programmer
  - Will also parallelize what compiler may miss
  - More explicit
  - Similar to OpenMP
- Both regions may not contain other kernels/parallel regions
- No braunching into or out
- Program must not depend on order of evaluation of clauses
- At most: One if clause





#### Add more parallelism

- Add OpenACC parallelism to other loops of while (L:123 L:141)
- Use either kernels or parallel
- Do they perform equally well?

#### Task 3: More Parallel Loops

- Change to Task3/ directory
- Compile: make Study the compiler output!
- Submit parallel run to the batch system: make run
- ? What's your speed-up?



#### **Source Code**

```
while ( error > tol && iter < iter max ) {
   error = 0.0:
   #pragma acc parallel loop reduction(max:error)
   for (int ix = ix_start; ix < ix_end; ix++) {
        for (int iv = iv start; iv < iv end; iv++) {
            Anew[iv*nx+ix] = -0.25 * (rhs[iv*nx+ix] -
                ( A[iv*nx+ix+1] + A[iv*nx+ix-1]
               + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]):
            error = fmaxr(error, fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
    }}
   #pragma acc parallel loop
   for (int iv = iv start: iv < iv end: iv++) {
       for( int ix = ix start: ix < ix end: ix++ ) {
           A[iv*nx+ix] = Anew[iv*nx+ix]:
    }}
   #pragma acc parallel loop
    for (int ix = ix start: ix < ix end: ix++) {
           A[0*nx+ix] = A[(ny-2)*nx+ix]:
           A[(nv-1)*nx+ix] = A[1*nx+ix]:
   // same for iv
   iter++:
```



#### **Compilation result**

```
$ make
nvc -c -DUSE DOUBLE -Minfo=accel -O1 -accegpu -gpu=managed poisson2d reference.c -o poisson2d reference.o
nvc -DUSE DOUBLE -Minfo=accel -01 -acc=gpu -gpu=managed poisson2d.c poisson2d reference.o -o poisson2d
poisson2d.c:
main:
    106. Generating Tesla code
        110, #pragma acc loop gang, vector(128) /* blockIdx.x threadIdx.x */
             Generating reduction(max:error)
        112, #pragma acc loop seq
    106. . . .
    118. Generating Tesla code
        123. #pragma acc loop gang /* blockIdx.x */
        125. #pragma acc loop vector(128) /* threadIdx.x */
    118. Generating implicit copyin(Anew[:]) [if not already present]
         Generating implicit copyout(A[:]) [if not already present]
    125, Loop is paral...
```



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Run result

```
$ make run
srun --gres=gpu:1 --time 0:10:00 --pty ./poisson2d
Jacobi relaxation calculation: max 500 iterations on 2048 x 2048 mesh
Calculate reference solution and time with serial CPU execution.
    0. 0.249999
  100. 0.249760
  200. 0....
Calculate current execution.
    0. 0.249999
  100, 0.249760
  200. 0....
2048x2048: Ref: 105.4636 s, This: 0.3448 s, speedup: 305.86
```



Run result

```
. . .
$ make run
srun --gres=gpu:1 --time 0:10:00 --pty ./poisson2d
Jacobi relaxation calculation: max 500 iterations on 2048 x 2048 mesh
Calculate reference solution and time with
                                                       execution.
    0. 0.249999
  100. 0.249760
  200, 0....
Calculate current execution.
    0. 0.249999
  100. 0.249760
  200. 0....
2048x2048: Ref: 105.4636 s, This: 0.3448 s, speedup:
                                                        305.86
```



# Data Transfers

OpenACC by Example

## **Automatic Data Transfers**

- Up to now: We did not care about data transfers
- Compiler and runtime care
- Magic keyword: -gpu=managed
- Only feature of (recent) NVIDIA GPUs!



Location, location

At the Beginning CPU and GPU memory very distinct, own addresses

CPU
CPU Memory

Scheduler Interconnect L2 DRAM

Location, location

At the Beginning CPU and GPU memory very distinct, own addresses

CUDA 4.0 Unified Virtual Addressing: pointer from same address pool, but data copy manual

Scheduler Interconnect L2 DRAM

CPU

**CPU Memory** 

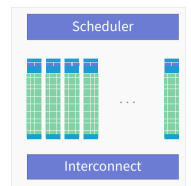
Unified
Virtual

Location, location

At the Beginning CPU and GPU memory very distinct, own addresses

CUDA 4.0 Unified Virtual Addressing: pointer from same address pool, but data copy manual

CUDA 6.0 Unified Memory\*: Data copy by driver, but whole data at once



L2

Unified Memorv

**CPU** 

Location, location, location

At the Beginning CPU and GPU memory very distinct, own addresses

CUDA 4.0 Unified Virtual Addressing: pointer from same address pool, but data copy manual

CUDA 6.0 Unified Memory\*: Data copy by driver, but whole data at once

CUDA 8.0 Unified Memory (truly): Data copy by driver, page faults on-demand initiate data migrations (Pascal)

Scheduler Interconnect

L2

Unified Memory

**CPU** 

Location, location, location

At the Beginning CPU and GPU memory very distinct, own addresses

CUDA 4.0 Unified Virtual Addressing: pointer from same address pool, but data copy manual

CUDA 6.0 Unified Memory\*: Data copy by driver, but whole data at once

CUDA 8.0 Unified Memory (truly): Data copy by driver, page faults on-demand initiate data migrations (Pascal)

Future Address Translation Service (POWER); Heterogeneous Memory Management (Linux)

Scheduler ...

Interconnect

L2

Unified Memory

# **Portability**

- Managed memory: Very productive feature
- Manual transfers: Fine-grained control, possibly faster, portability
- → Code should also be fast without -gpu=managed!
  - Let's remove it from compile flags!



### **Portability**

- Managed memory: Very productive feature
- Manual transfers: Fine-grained control, possibly faster, portability
- → Code should also be fast without -gpu=managed!
  - Let's remove it from compile flags!

```
$ make
nvc -c -DUSE_DOUBLE -Minfo=accel -O1 -acc=gpu poisson2d_reference.c -o
    poisson2d_reference.o
nvc -DUSE_DOUBLE -Minfo=accel -O1 -acc=gpu poisson2d.c poisson2d_reference.o -o poisson2d
poisson2d.c:
NVC++-S-0155-Compiler failed to translate accelerator region (see -Minfo messages): Could
    not find allocated-variable index for symbol - rhs (poisson2d.c: 106)
...
NVC++-F-0704-Compilation aborted due to previous errors. (poisson2d.c)
NVC++/x86-64 Linux 21.9-0: compilation aborted
```

Compiler implicitly created copy clauses to copy data to device

```
106, Generating implicit copyin(A[:],rhs[:]) [if not already present]

Generating implicit copy(error) [if not already present]
```

• It couldn't determine length of copied data ...but before: no problem – Unified Memory!



Compiler implicitly created copy clauses to copy data to device

```
106, Generating implicit copyin(A[:],rhs[:]) [if not already present]
Generating implicit copy(error) [if not already present]
```

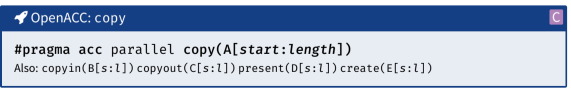
- It couldn't determine length of copied data ...but before: no problem Unified Memory!
- Now: Problem! We need to give that information! (Fortran: can often be determined by compiler)



Compiler implicitly created copy clauses to copy data to device

```
106, Generating implicit copyin(A[:],rhs[:]) [if not already present]
Generating implicit copy(error) [if not already present]
```

- It couldn't determine length of copied data ...but before: no problem Unified Memory!
- Now: Problem! We need to give that information! (Fortran: can often be determined by compiler)

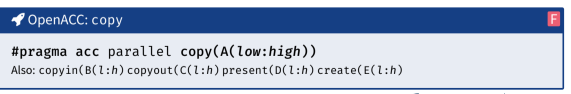




Compiler implicitly created copy clauses to copy data to device

```
106, Generating implicit copyin(A[:],rhs[:]) [if not already present]
Generating implicit copy(error) [if not already present]
```

- It couldn't determine length of copied data ...but before: no problem Unified Memory!
- Now: Problem! We need to give that information! (Fortran: can often be determined by compiler)





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#### Get that data!

Add copy clause to parallel regions

#### Task 4: Data Copies

- Change to Task4/ directory
- Work on TODOs
- Compile: make
- Submit parallel run to the batch system: make run
- ? What's your speed-up?



#### **Data Copies**

#### **Compiler Output**

```
$ make
nyc -DUSE DOUBLE -Minfo=accel -O1 -acc=gpu poisson2d.c poisson2d reference.o -o poisson2d
poisson2d.c:
main:
    106. Generating copy(A[:ny*nx].rhs[:ny*nx]) [if not already present]
         Generating implicit copy(error) [if not already present]
         Generating Tesla code
        110, #pragma acc loop gang, vector(128) /* blockIdx.x threadIdx.x */
             Generating reduction(max:error)
        112. #pragma acc loop seg
    106. Generating copy(Anew[:ny*nx]) [if not already present]
    112. Complex loop carried dependence of Anew-> prevents parallelization
         Loop carried dependence of Anew-> prevents parallelization
         Loop carried backward dependence of Anew-> prevents vectorization
```



#### **Data Copies**

#### **Run Result**

```
. . .
$ make run
srun --partition=gpus --gres=gpu:1 ./poisson2d
Jacobi relaxation calculation: max 500 iterations on 2048 x 2048 mesh
Calculate reference solution and time with serial CPU execution.
    0. 0.249999
  100. 0.249760
  200, 0...
Calculate current execution.
    0. 0.249999
  100. 0.249760
  200. 0...
2048x2048: Ref: 89.8862 s, This: 22.8402 s, speedup:
                                                            3.94
```



#### **Data Copies**

#### **Run Result**

```
. . .
$ make run
srun --partition=gpus --gres=gpu:1
                                     ./poisson2d
Jacobi relaxation calculation: max 500
                                                        48 x 2048 mesh
                                      Slower?!
Why?
Calculate reference solution and tim
                                                        ecution.
    0. 0.249999
  100. 0.249760
  200, 0...
Calculate current execution.
    0. 0.249999
  100. 0.249760
  200. 0...
2048x2048: Ref: 89.8862 s, This: 22.8402 s, speedup:
                                                           3.94
```



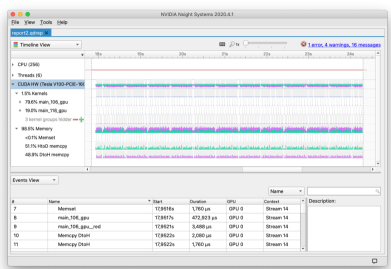
# **Nsight Systems**

- Let's check again with profiler!
- This time: GUI of Nsight Systems with timeline



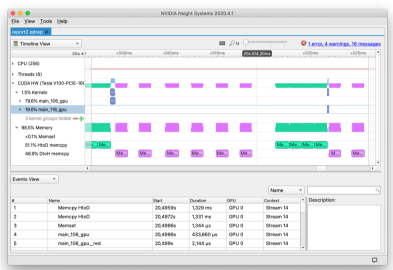
# **Nsight Systems**

#### Overview



### **Nsight Systems**

#### Zoom to kernels



#### **Parallelization Workflow**

Identify available parallelism

Parallelize loops with OpenACC

**Optimize data locality** 

Optimize loop performance



#### In code

```
while (error > tol && iter < iter_max) {
    error = 0.0;</pre>
```

#### A, Anew resident on **host**

#### #pragma acc parallel loop

```
iter++
}
```



```
In code
```

```
while (error > tol && iter < iter_max) {
    error = 0.0;</pre>
```

A, Anew resident on **host** 

сору

#pragma acc parallel loop

A, Anew resident on **device** 

```
iter++
```



```
In code
```

```
while (error > tol && iter < iter max) {
    error = 0.0:
```

A, Anew resident on host

copy

#pragma acc parallel loop

A, Anew resident on device

```
for (int ix = ix_start; ix < ix_end;</pre>
\hookrightarrow ix++) {
     for (int iy = iy_start; iy < iy_end;</pre>
     \hookrightarrow iv++) {
     // ...
}}
```

A. Anew resident on **device** 

```
iter++
```



```
In code
```

```
while (error > tol && iter < iter max) {
    error = 0.0:
```

A, Anew resident on **host** 

copy

#pragma acc parallel loop

A, Anew resident on **device** 

```
for (int ix = ix_start; ix < ix_end;</pre>
\hookrightarrow ix++) {
     for (int iy = iy_start; iy < iy_end;</pre>
     \hookrightarrow iv++) {
     // ...
}}
```

A. Anew resident on **device** 

A, Anew resident on host 4

```
iter++
```



```
In code
```

```
while (error > tol && iter < iter max) {
    error = 0.0:
```

A, Anew resident on host

copy

#pragma acc parallel loop

A, Anew resident on **device** 

```
for (int ix = ix_start; ix < ix_end;</pre>
\hookrightarrow ix++) {
     for (int iy = iy_start; iy < iy_end;</pre>
     \hookrightarrow iv++) {
     // ...
```

}}

A, Anew resident on device

A, Anew resident on **host** 

```
iter++
```

```
In code
 while (error > tol && iter < iter max) {
     error = 0.0:
  A, Anew resident on host
```

Copies are done

between **each** loop and in **each** iteration!

A, Anew resident on **host** 

```
iter++
```

#pragma acc parallel loop

A, Anew resident on **device** 

```
for (int ix = ix_start; ix < ix_end;</pre>
\hookrightarrow ix++) {
     for (int iy = iy_start; iy < iy_end;</pre>
     \hookrightarrow iv++) {
     // ...
}}
```

A. Anew resident on **device** 



copy

**Summary** 

- By now, whole algorithm is using GPU
- At beginning of while loop, data copied to device; at end of loop, copied by to host
- Depending on type of parallel regions in while loop: Data copied in between regions as well



**Summary** 

- By now, whole algorithm is using GPU
- At beginning of while loop, data copied to device; at end of loop, copied by to host
- Depending on type of parallel regions in while loop: Data copied in between regions as well
- Slow! Data copies are expensive!



#### **Structured Data Regions**

- Defines region of code in which data remains on device
- Data is shared among all kernels in region
- Explicit data transfers

#### 🕜 OpenACC: data

```
#pragma acc data [clause, [, clause] ...]
```



Clauses

#### Clauses to augment the data regions

create(var) Allocates memory of var on GPU

present(var) Data of var is not copies automatically to GPU but considered present

### **Data Region Example**

```
#pragma acc data copyout(v[0:N]) create(x[0:N])
double sum = 0.0:
#pragma acc parallel loop
for (int i=0; i<N; i++) {
   x[i] = 1.0;
   v[i] = 2.0:
#pragma acc parallel loop
for (int i=0; i<N; i++) {
   v[i] = i*x[i]+v[i]:
```

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```
!\$acc data copyout(v(1:N)) create(x(1,N))
sum = 0.0:
!$acc parallel loop
do i = 1. N
    x(i) = 1.0
    v(i) = 2.0
end do
!$acc end parallel loop
!$acc parallel loop
do i = 1, N
    v(i) = i * x(i) + v(i)
end do
!$acc end parallel loop
!$acc end data
```

Slide 56174

# **Data Regions II**

#### **Unstructured Data Regions**

- Define data regions, but not for structured block
- Clauses executed at the very position the directive encountered
- Closest to cudaMemcpy()
- Still, explicit data transfers

#### ✓ OpenACC: enter data

```
#pragma acc enter data [clause, [, clause] ...]
#pragma acc exit data [clause, [, clause] ...]
```



TASK

More parallelism, Data locality

• Add data regions such that all data resides on device during iterations

#### Task 5: Data Region

- Change to Task5/ directory
- Work on TODOs
- Compile: make
- Submit parallel run to the batch system: make run
- ? What's your speed-up?



#### Parallel Jacobi II

#### Source Code

```
#pragma acc data copv(A[0:nx*nv]) copvin(rhs[0:nx*nv]) create(Anew[0:nx*nv])
105
106
      while ( error > tol && iter < iter max )
107
108
          error = 0.0:
109
          // Jacobi kernel
110
111
          #pragma acc parallel loop reduction(max:error)
112
          for (int ix = ix_start; ix < ix_end; ix++)</pre>
113
114
              for (int iv = iv start: iv < iv end: iv++)
115
116
                   Anew[iv*nx+ix] = -0.25 * (rhs[iv*nx+ix] - (A[iv*nx+ix+1] + A[iv*nx+ix-1])
117
                                                           + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]));
                   error = fmaxr( error, fabsr(Anew[iv*nx+ix]-A[iv*nx+ix])):
118
119
120
121
122
          // A <-> Anew
123
          #pragma acc parallel loop
124
          for (int iy = iy start; iy < iy_end; iy++)
125
          // ...
126
```

**Compiler Output** 

```
$ make
nvc -DUSE DOUBLE -Minfo=accel -O1 -acc=gpu poisson2d.c poisson2d reference.o -o poisson2d
poisson2d.c:
main:
    105. Generating create(Anew[:nv*nx]) [if not already present]
         Generating copy(A[:nv*nx]) [if not already present]
         Generating copyin(rhs[:ny*nx]) [if not already present]
    107. Generating Tesla code
        111, #pragma acc loop gang, vector(128) /* blockIdx.x threadIdx.x */
             Generating reduction(max:error)
        113. #pragma acc loop seg
```

#### **Run Result**

```
. . .
$ make run
srun --pty ./poisson2d
Jacobi relaxation calculation: max 500 iterations on 2048 x 2048 mesh
Calculate reference solution and time with serial CPU execution.
    0. 0.249999
  100. 0.249760
  200, 0...
Calculate current execution.
    0. 0.249999
  100, 0.249760
  200. 0...
2048x2048: Ref: 94.3213 s, This: 0.3506 s, speedup: 269.05
```



#### **Run Result**

```
$ make run
srun --pty ./poisson2d
Jacobi relaxation Calculation: 2048 x 20
Calculate reference solution and time CA
     0 0.250000
   100 0.002396
   200 0 . . .
GPU execution.
       0.250000
   100 0.002396
   200 0...
2048 x 2048: 1 GPU: 0.1570s, 1 CPU cores: 3.5955s, speedup: 22.90
```

23 June 2023

#### **Parallelization Workflow**

Identify available parallelism

Parallelize loops with OpenACC

Optimize data locality

Optimize loop performance



#### **Loop Performance Optimization**

**Opportunities** 

To be discussed in other sessions!



# Routines

OpenACC by Example

#### **Accelerated Routines**

- Enable functions/sub-routines for acceleration
- Make routine callable from device (CUDA: \_\_device\_\_)
- Needed for refactoring, modular designs, ...
- Position
  - C At declaration and implementation; immediately before See next slide Fortran Within specification part sub-routine



#### **Routine Details**

Clauses to Directive gang worker vector seq Type of parallelism used inside of routine



#### **Routine Details**

Clauses to Directive
gang worker vector seq Type of parallelism used inside of routine
(name) Second version of directive

- Make named routine accelerated
- Applies to function within current scope with name name
- To be inserted before definition of named function



## **Routine Details**

#### **Clauses to Directive**

gang worker vector seq Type of parallelism used inside of routine
 (name) Second version of directive

- Make named routine accelerated
- Applies to function within current scope with name name
- To be inserted before definition of named function

#### bind(func) Bind routine to func device function

```
#pragma acc routine bind(func_dev)
void func(float * ) {}
void func(float * A) {A[0] = 2;}
#pragma acc routine
void func_dev(float * A) {A[0] = 23;}
int main() {
   float * A = (float*) malloc(1*sizeof(float));
   func(A) // A[0] == 2
   #pragma acc parallel
   func(A) // A[0] == 23
```

## Routine



#### Getting some routine!

- Extract the inner part of the double for-loop into a dedicated routine called inner\_loop()
- C: error needs to be passed by reference!

#### Task 6: Routine

- Change to Task6/ directory
- Work on TODOs
- Compile: make
- Submit parallel run to the batch system: make run
- Fortran: Why did it get slower?!



## **Jacobi Routine**

#### **Source Code**

```
#pragma acc routine
42
    void inner loop(int ix, int nx, int iy start, int iy end, real * A, real * Anew, real * rhs,
43

    real * error) {

44
        #pragma acc loop
        for (int iy = iy start; iy < iy end; iy++)</pre>
            Anew[iy*nx+ix] = -0.25 * (rhs[iy*nx+ix] - (A[iy*nx+ix+1] + A[iy*nx+ix-1])
47
                                                     + A[(iv-1)*nx+ix] + A[(iv+1)*nx+ix]):
48
            *error = fmaxr( *error. fabsr(Anew[iv*nx+ix]-A[iv*nx+ix]));
49
50
51
```



## **Jacobi Routine**

#### **Compiler Output**



# Other Directives

# **Further Keywords**

#### **Directives**

```
serial Serial GPU Region
```

wait Wait for any async operation

atomic Atomically access data (no interference of concurrent accesses)

cache Fetch data to GPU caches

declare Make data live on GPU for implicit region directly after variable declaration

update Update device data

shutdown Shutdown connection to GPU



# **Further Keywords**

# **Directives** serial Serial GPU Region wait Wait for any async operation atomic Atomically access data (no interference of concurrent accesses) cache Fetch data to GPU caches declare Make data live on GPU for implicit region directly after variable declaration update Update device data

shut down Shutdown connection to GPU

#### **Clauses**

gang worker vector Type of parallelism collapse Combine tightly-nested loops tile Split loop into two loops (first)private Create thread-private data (and init) attach Reference counting for data pointers async Schedule operation

asynchronously



# **Further Keywords**

# **Directives** serial Serial GPU Region wait Wait for any async operation atomic Atomically access data (no interference of concurrent accesses) cache Fetch data to GPU caches declare Make data live on GPU for implicit region directly after variable declaration update Update device data

shut down Shutdown connection to GPU

#### **Clauses**

attach Reference counting for data

pointers

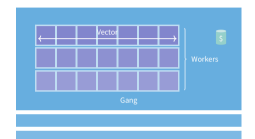
async Schedule operation asynchronously



# **Launch Configuration**

Specify number of threads and blocks

- 3 clauses for changing distribution of group of threads (clauses of parallel region (parallel, kernels))
- Presence of keyword: Distribute using this level
- Optional size: Control size of parallel entity



#### → OpenACC: gang worker vector

#pragma acc parallel loop gang worker vector
Size: num\_gangs(n), num\_workers(n), vector\_length(n)



# Conclusions

## **Conclusions**

- OpenACC directives and clauses#pragma acc parallel loop copyin(A[0:N]) reduction(max:err) vector
- Start easy, optimize from there; express as much parallelism as possible
- Optimize data for locality, prevent unnecessary movements
- OpenACC is interoperable to other GPU programming models



## **Conclusions**

- OpenACC directives and clauses#pragma acc parallel loop copyin(A[0:N]) reduction(max:err) vector
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Appendix List of Tasks Glossary References



## **List of Tasks**

Task 2: A First Parallel Loop

48Task 3: More Parallel Loops

60Task 4: Data Copies

78Task 5: Data Region

98Task 6: Routine

110



# Glossary I

- AMD Manufacturer of CPUs and GPUs. 11, 12, 13, 14
- Ampere GPU architecture from NVIDIA (announced 2019). 15
  - CUDA Computing platform for GPUs from NVIDIA. Provides, among others, CUDA C/C++. 16, 67, 68, 69, 70, 71
    - GCC The GNU Compiler Collection, the collection of open source compilers, among others for C and Fortran. 15, 30, 31
  - LLVM An open Source compiler infrastructure, providing, among others, Clang for C. 11, 12, 13, 14
- NVHPC NVIDIA HPC SDK; Collection of GPU-capable compilers and libraries. Formerly known as PGI.. 15



# Glossary II

- NVIDIA US technology company creating GPUs. 4, 11, 12, 13, 14, 52, 66, 123, 124, 125
- OpenACC Directive-based programming, primarily for many-core machines. 2, 4, 5, 6, 7, 8, 9, 10, 15, 16, 17, 18, 19, 20, 29, 35, 36, 37, 38, 40, 41, 43, 44, 45, 48, 55, 60, 65, 76, 77, 85, 94, 97, 103, 105, 106, 117, 119, 120
- OpenMP Directive-based programming, primarily for multi-threaded machines. 2, 5, 6, 7, 11, 12, 13, 14, 57, 58, 59
  - PAPI The Performance API, a C/C++ API for querying performance counters. 30, 31
  - Pascal GPU architecture from NVIDIA (announced 2016). 67, 68, 69, 70, 71
    - perf Part of the Linux kernel which facilitates access to performance counters; comes with command line utilities. 30, 31



# **Glossary III**

- PGI Compiler creators. Formerly *The Portland Group, Inc.*; since 2013 part of NVIDIA. 123
- POWER CPU architecture from IBM, earlier: PowerPC. See also POWER8. 67, 68, 69, 70, 71, 125
- POWER8 Version 8 of IBM's POWER processor, available also within the OpenPOWER Foundation. 125
  - CPU Central Processing Unit. 11, 12, 13, 14, 67, 68, 69, 70, 71, 123, 125
  - GPU Graphics Processing Unit. 2, 11, 12, 13, 14, 17, 48, 52, 66, 67, 68, 69, 70, 71, 92, 93, 95, 119, 120, 123, 124

### References I

[3] Donald E. Knuth. "Structured Programming with Go to Statements." In: ACM Comput. Surv. 6.4 (Dec. 1974), pp. 261–301. ISSN: 0360-0300. DOI: 10.1145/356635.356640. URL: http://doi.acm.org/10.1145/356635.356640 (pages 30, 31).



# **References: Images, Graphics**

- [1] Bill Jelen. SpaceX Falcon Heavy Launch. Freely available at Unsplash. URL: https://unsplash.com/photos/lDEMa5dPcNo.
- [2] Setyo Ari Wibowo. *Ask*. URL: https://thenounproject.com/term/ask/1221810.

