



# Alpaka Hackathon Section-II: Heat Equation Solution

PLASMA-PEPSC Workshop on Alpaka and OPENPMD

October 18, 2024





# Workshop Schedule

### Section - I

- Introduction: What is alpaka, where it is used?
- 2 Hands on 1: Installing alpaka and running an example (LUMI)
- Parallel programming concepts and portable parallelization by alpaka
  - Grid Structure and WorkDivision
  - Data Parallelism
  - Indexing
- 4 Hands on 2: HelloIndex kernel which prints indexes

### Section - II

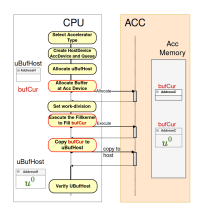
- Memory management for 1D and 2D data
- Filling buffers in parallel
- Hands on 3: Kernel to fill initial conditions of heat equation
- 4 Heat Equation
- 5 Preparing stencil kernel
- 6 Hands on 4: Heat Equation stencil kernel
- Programming features and data-structures of alpaka
- **B** Usability and Optimization
  - Using alpaka mdspan for easier indexing Hands on
  - Domain Decomposition Hands on 6 (Day2)
  - Using async queues for performance increase Hands on 7 (Dav2)
  - Using shared memory for performance increase Hands on 8 (Day2)





# Steps of Filling a Buffer in Parallel

- Select the accelerator
- Create host-device, acc-device and the queue
- 3 Allocate host and device memory
- Decide how to parallelize: set work-division
- Decide where will the parallel and non-parallel parts of the code run
- 6 Create the kernel instance and execute kernel
- Copy the result from Acc (e.g GPU) back to the host buffer







### Allocate memory at Host and at Device

### Define number of dim and index type

```
using Dim = alpaka::DimInt<2u>; // Number of dim: 2 as a type
using Idx = std::size_t; // Index type of the threads and buffers
```

#### Define domain and halo extents

```
// alpaka::Vec is a static array similar to std::array.
// Dim is a compile—time constant, which is 2.
// Create a static array of size Dim.

constexpr alpaka::Vec < Dim, Idx > numNodes {64, 64};
constexpr alpaka::Vec < Dim, Idx > haloSize {2, 2};
constexpr alpaka::Vec < Dim, Idx > extent = numNodes + haloSize;
```

### Allocate memories at host and accelerator

```
// Allocate memory for host—buffer
auto uBufHost = alpaka::allocBuf <double, Idx>(devHost, extent);

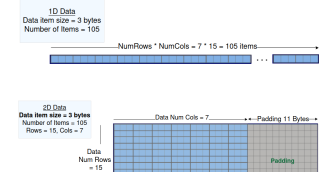
// Allocate memory for accelerator buffer
auto uBufAcc = alpaka::allocBuf <double, Idx>(devAcc, extent);
```





# Allocated area at the memory

Let's assume that 105 item with 3-byte each will be allocated to pass to the kernel. The pitch value (actually the row-pitch) depends on the GPU or CPU type.



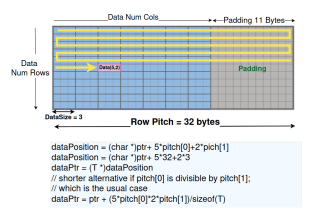
Row Pitch = 32 bytes





### How to access data given the pointer and the pitch?

How to access data at index (5,2) given the pointer ptr and pitch? pitch = {32bytes,3 bytes} as {row-pitch, datasize}







# Passing multi dimensional buffer to the kernel

Pass 3 variables for a buffer: pointer, "row-pitch", and datasize Multi-dimensional memory allocated in memory uses aligned rows. Hence, if a pointer of a 2D buffer is passed to the kernel as a pointer; 2 additional values pitch and item data-size should also be passed.

```
// Signature of function operator of the Kernel
template typename TAcc, typename TDim, typename TIdx>
ALPAKA_FN_ACC auto operator()(
TAcc const& acc,
double* const bufData,
// 2 variables row-pitch and data-type size
alpaka::Vec<TDim, TIdx> const pitch,
double dx,
double dy) const -> void
```

■ Simple Alternative: Pass an alpaka mdspan object

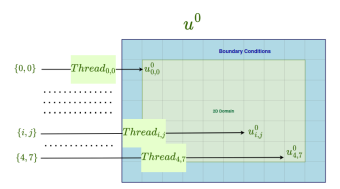
```
template < typename TAcc, typename TDim, typename TIdx, typename TMdSpan > ALPAKA_FN_ACC auto operator()(
TAcc const& acc,
TMdSpan uAccMdSpan
...) const -> void
```





### The Kernel to Initialize Heat Values

Calculate and set initial heat values, the  $u^0$  matrix, by running a grid of threads.





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### The Kernel to Initialize Heat Values

The InitializeBufferKernel fills the given buffer at the accelerator device (e.g GPU) Prepare kernel to set initial heat values

- Thread Index: Find thread index in the kernel to be used as index to set 2D buffer.
- Initial Condition at the point: Find analytically the heat value at the point which has coordinates equal to the 2D thread index.
- Memory Adress in Buffer: Calculate the corresponding memory adress in buffer using thread index. Take into account the row-pitch and data-size
- Set Value at the Adress: Set the data at the memory position to the calculated initial condition.





### Hands-On Session

# Hands-on Session3: Filling an accelerator buffer paralelly





# The Heat Equation

■ The heat equation models the Heat Diffusion over time in a given medium.

$$\frac{\partial u(x, y, t)}{\partial t} = \alpha \left( \frac{\partial^2 u(x, y, t)}{\partial x^2} + \frac{\partial^2 u(x, y, t)}{\partial y^2} \right)$$

Difference approximations for Time and Spatial Derivatives:

$$\left. \frac{\partial u(x,y,t)}{\partial t} \right|_{t=t^n} \approx \left. \frac{u_{i,j}^{n+1} - u_{i,j}^n}{\Delta t} \right. \qquad \left. \frac{\partial^2 u(x,y,t)}{\partial x^2} \right|_{x=x_i,y=y_i} \approx \left. \frac{u_{i+1,j}^n - 2u_{i,j}^n + u_{i-1,j}^n}{\Delta x^2} \right.$$

Resulting difference equation:

$$u_{i,j}^{n+1} = u_{i,j}^{n} + \alpha \Delta t \left( \frac{u_{i+1,j}^{n} - 2u_{i,j}^{n} + u_{i-1,j}^{n}}{\Delta x^{2}} + \frac{u_{i,j+1}^{n} - 2u_{i,j}^{n} + u_{i,j-1}^{n}}{\Delta y^{2}} \right)$$







# The Heat Equation- Cont.

■ The difference equation:

$$u_{i,j}^{n+1} = u_{i,j}^{n} + \alpha \Delta t \left( \frac{u_{i+1,j}^{n} - 2u_{i,j}^{n} + u_{i-1,j}^{n}}{\Delta x^{2}} + \frac{u_{i,j+1}^{n} - 2u_{i,j}^{n} + u_{i,j-1}^{n}}{\Delta y^{2}} \right)$$

■ Substitute:  $\alpha=1$ ,  $r_X=\frac{\Delta t}{\Delta x^2}$ ,  $r_Y=\frac{\Delta t}{\Delta y^2}$ 

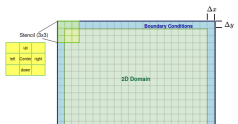
Then  $u_{i,j}^{n+1}$  is:

$$u_{i,j}^{n+1} = u_{i,j}^{n} + r\chi \left( u_{i+1,j}^{n} - 2u_{i,j}^{n} + u_{i-1,j}^{n} \right) + r\gamma \left( u_{i,j+1}^{n} - 2u_{i,j}^{n} + u_{i,j-1}^{n} \right)$$

By regrouping the terms related to  $u_{i,j}^n$ , the equation can be rewritten as:

$$u_{i,j}^{n+1} = u_{i,j}^{n} \left(1 - 2r_{X} - 2r_{Y}\right) + r_{X} \left(u_{i+1,j}^{n} + u_{i-1,j}^{n}\right) + r_{Y} \left(u_{i,j+1}^{n} + u_{i,j-1}^{n}\right)$$

$$S = \begin{pmatrix} 0 & r_Y & 0 \\ r_X & 1 - 2r_X - 2r_Y & r_X \\ 0 & r_Y & 0 \end{pmatrix}$$

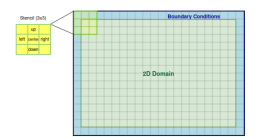






### Parallel Heat Equation Solution

- Data Parallelism: Each point on the grid can be updated independently based on its neighbors, enabling parallel computation.
- Stencil Operations: Stencil is a core computational pattern in PDE solvers. Updates a grid point in time
  using its immediate neighbors (left, right, up, down) according to the difference equation. A 5-point stencil
  is needed.



- Halo Region for BC: A layer of grid cells surrounding the problem domain for Boundary Conditions.
  - Facilitates stencil operations at the boundaries of subdomains.



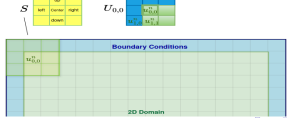


# Calculation of $u_{i,j}^{n+1}$ from $u_{i,j}^n$

- Each kernel execution by alpaka calculates  $u_{i,j}^{n+1}$  using  $u_{i,j}^{n}$
- Each heat point is separately calculated by a thread using Frobenious Inner Product (FIP)
- The Frobenius Inner Product between matrix S and matrix  $U_{i,j}$  is:

$$u_{i,j}^{n+1} = \langle S, U_{i,j}^n \rangle_F = \sum_{m=1}^M \sum_{k=1}^K s_{m,k} u_{m,k}$$

■ S and  $U_{0,0}^n$  is used by a thread to calculate  $u_{0,0}^{n+1}$  using FIP







■ Another thread calculates  $u_{0,1}^{n+1}$  using S and  $U_{0,1}^n$ 

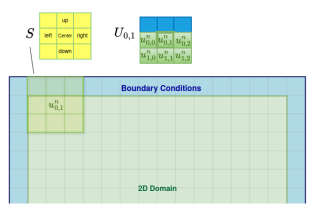
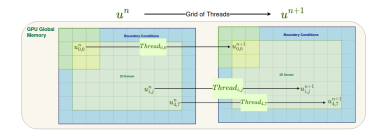


Figure: Second thread calculates  $u_{0,1}^{n+1}$  using





### Stencil Kernel Execution by a grid of threads



- Stencil kernel will update only core nodes not the border
- The workdiv for stencil kernel can be calculated by setting gridthread extent to nodes domain
- The workdiv for the borders kernel would need extended extents, because halo is going to updated as well





# Main Simulation Loop: Leveraging Parallelism

### Initialization:

- Define the "host device" and "accelerator device". The "Host" and "Device" in short.
- Set initial conditions and boundary conditions.
- Allocate data buffers to host and device.
- Copy data from host to device buffer to pass to the kernel.
- Define parallelisation strategy (determine block size).

### Simulation Loop:

- Step 1: Execute StencilKernel to compute next values.
- Step 2: Apply boundary conditions using BoundaryKernel.
- **Step 3:** Swap buffers for the next iteration so that calculated  $u_{i,j}^{n+1}$  becomes the  $u_{i,j}^{n}$  for the next step.

### ■ Parallel Efficiency:

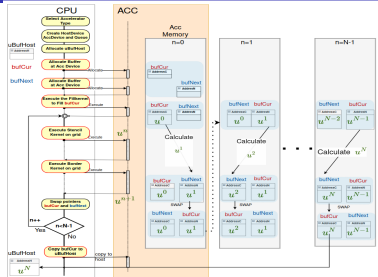
- Subdomains are processed in parallel, with halos ensuring data consistency and correct boundary conditions.
- Optimization: Shared memory optimizes memory access within each block using chunks of data.

### Validation





### Complete Heat Equation solution







### The Stencil Kernel

### What kind of parallelization needed to calculate $u_{i,j}^{n+1}$ using $u_{i,j}^{n}$

- StencilKernel needs a Work-division to work on domain of all nodes (without halo)
- Boundary kernel needs a Workdivision which covers nodes + halo region
- Boundary kernel will only use threads corresponding to halo region

#### Coding StencilKernel

- Input: Check the size of the input buffer, it should include halo region as well
- Thread Index: Find thread index in the kernel. This index will be used as the center of 3x3 stencil.
- Memory Adress in Buffer: Calculate the corresponding memory adress in buffer using thread index. Take into account pitch and data-size
- Calculate new heat value: Calculate  $u_{i,j}^{n+1}$  using Frobenious Inner Product of 3x3 matrices
- Set Value at the Adress: Set the data at the memory adress.

```
struct StencilKernel
{
    template < typename TAcc, typename TDim, typename TIdx>

    ALPAKA_FN_ACC auto operator()(
        TAcc const& acc,
        double const* const uCurrBuf, double* const uNextBuf,
        alpaka::Vec<TDim, TIdx> const chunkSize,
        alpaka::Vec<TDim, TIdx> const pitchCurr, alpaka::Vec<TDim, TIdx> const pitchNext,
        double const dx,double const dy, double const dt) const -> void
    {
        ...
}
```





### Hands-On Session

# Hands-on Session4: Stencil Kernel to Calculate $u^{n+1}$





# alpaka Basics

# Setting up the stage to run kernels

- Selecting the accelerator and host devices
- Allocating and setting host and accelerator device memory
- 3 Alpaka Vector, Buffer and View?
- Passing data to the accelarator
- 5 WorkDiv
- 6 Define Queue





### Accelerator, Device and Host

### Define number of dim and index type

```
using Dim = alpaka::DimInt<2u>; // Number of dim: 2 as a type
using Idx = std::size_t; // Index type of the threads and buffers
```

### Define the accelerator

```
// AccGpuCudaRt, AccGpuHipRt, AccCpuThreads, AccCpuSerial,
// AccCpuOmp2Threads, AccCpuOmp2Blocks, AccCpuTbbBlocks
using Acc = alpaka::AccGpuHipRt<Dim, Idx>;
using DevAcc = alpaka::Dev<Acc>;
```

### Select a device from platform of Acc

```
auto const platform = alpaka::Platform<Acc>{};
auto const devAcc = alpaka::getDevByIdx(platform, 0);
```

### Select a host and hostype to allocate memory for data

```
// Get the host device for allocating memory on the host.
auto const platformHost = alpaka::PlatformCpu{};
auto const devHost = alpaka::getDevByIdx(platformHost, 0);
// Host device type is needed, still not known
sing DevHost = alpaka::DevCou:
```





### What is Accelerator

Accelerator hides hardware specifics behind alpaka's abstract API

- On Host: Accelerator is a type. A Meta-parameter for choosing correct physical device and dependent types
- using Acc = acc::AccGpuHipRt < Dim, Idx>;
  - Inside Kernel: Accelerator is a variable. Contains thread state, provides access to alpaka's device-side API
    - The Accelerator provides the means to access to the indices

```
// get thread index on the grid
auto gridThreadIdx = alpaka::getIdx<Grid, Threads>(acc);
// get block index on the grid
auto gridBlockIdx = alpaka::getIdx<Grid, Blocks>(acc);
```

 The Accelerator gives access to alpaka's shared memory (for threads inside the same block)

■ Enables synchronization on the block level

```
1 // synchronize all threads within the block
2 alpaka::syncBlockThreads(acc);
```





### What is alpaka Buffer, Vector and View

- alpaka::Buf is multi-dimensional dynamic (runtime sized) container.
  - Contains memory adress, extent, datatype and the device that memory belongs to!
  - Since buffer already knows the it's device and extent; device to device copy is easy in alpaka
  - Supports [] operator but not [][].

```
// Allocate buffers
auto bufCpu = alpaka::allocBuf<float, Idx>(devCpu, extent);
auto bufGpu = alpaka::allocBuf<float, Idx>(devGpu, extent);
....
// Copy buffer from CPU to GPU - destination comes first
alpaka::memcopy(gpuQueue, bufGpu, bufCpu);
// cuda way: cudaMemcpy(b_d, b_host, sizeof(float)*N, cudaMemcpyHostToDevice)
```

alpaka::Vec is a static 1D array.

```
alpaka::Vec<SizeOfArrayAsType,DataT> myVec;
```

 alpaka::View is a non-owning view to an already allocated memory, so that it can be used in alpaka::memcpy





# What is alpaka::Queue

- alpaka::Queue is "a queue of tasks".
- Used for sycnhronization of tasks like memcpy or kernel-execution
- Queue is always FIFO, everything is sequential (in-order) inside the alpaka::queue
- If the Queue is non-blocking the caller(host) is not blocked
- Different non-blocking queues can run in parallel
- Within a single queue accelerator back-ends can be mixed (used in interleaves)

```
using QueueProperty = alpaka::NonBlocking;
// Create queue
using QueueAcc = alpaka::Queue<Acc, QueueProperty>;
QueueAcc computeQueue{devAcc};
// Copy host -> device, use the queue
alpaka::memcpy(computeQueue, uCurrBufAcc, uBufHost);
alpaka::wait(computeQueue); // Not needed, we have single queue
// Create kernel instance
StencilKernel
stencilKernel</pre
```





# Multiple Queues

Queues are used for synchronization



Copying and running synchronously

```
using QueueProperty = alpaka::NonBlocking;
using QueueGpu = alpaka::Queue<AccGpu, QueueProperty>;

StencilKernel<sharedMemSize> stencilKernel;
auto queueGpu1 = QueueGpu{devGpu};
auto queueGpu2 = QueueGpu{devGpu};
alpaka::memcpy(queueGpu1, uCurrBufAcc, uBufHost);
alpaka::wait(QueueGpu1);
alpaka::wait(QueueGpu2, uCurrBufAcc2, uBufHost2);
// Execute kernel using queue
alpaka::exec<Acc>(queueGpu1, workDiv_manual, stencilKernel...)
alpaka::wait(QueueGpu1);
alpaka::wait(QueueGpu2);
alpaka::wait(QueueGpu2);
```





### Queue Operations and Tasks

 Device-side related operations (kernels, memory operations) can be wrapped in tasks.

Tasks are executed by enqueue() function.

■ Wait can be used to make queue finish all tasks enqueued

```
// wait until all tasks have completed
alpaka::wait(queueA);
// block queueA until otherQueue has completed
alpaka::wait(queueA, otherQueue);
```

- Queues can be checked for completion of all tasks
- bool done = alpaka::empty(myQueue);

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# Queue Operations and Events

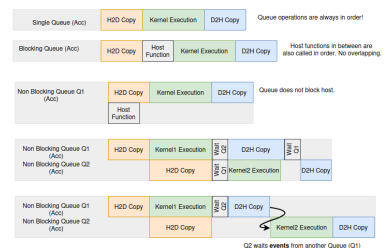
For easier synchronization, alpaka Events can be inserted before, after or between Tasks:

```
auto myEvent = alpaka::Event<alpaka::Queue>(myDev);
alpaka::enqueue(queueA, myEvent);
alpaka::wait(queueB, myEvent);
// queueB will only resume after queueA reached myEvent
```





## Some sychronization scenarios



ato events from another educat (ex)





# Kernel execution and copying results back

### Create the queue

```
// Create queue,
// queue is needed for kernel execution and copies to/from accelerator
alpaka::Queue<Acc, alpaka::NonBlocking> queue{devAcc};
```

■ Execute the kernel using the queue, the workdiv and kernel arguments:

```
alpaka::exec<Acc>(queue, workDiv, initBufferKernel, uBufAcc.data(),
pitchCurrAcc, dx, dy);
```

Copy the filled buffer back to the host

```
// Copy device -> host
// Since buffers know their corresponding devices (host or acc) memcopy does
not need any device variable
alpaka::memcpy(queue, uBufHost, uBufAcc);
alpaka::wait(queue);
```





# Setting WorkDiv using getValidWorkDiv function

### Let alpaka calculate work division for you:

```
// All kernel inputs are needed because work—division depends on the kernel
// Create kernel instance
InitializeBufferKernel initBufferKernel;
// Elements per thread needed to determine work—div
constexpr alpaka::Vec<Dim, Idx> elemPerThreaddi, 1};
// Bundle the extent and elements per thread
alpaka::KernelCfg<Acc> const kernelCfg = {extent, elemPerThread};
// Determine the work—div using kernel and kernel arguments
auto workDiv = alpaka::getValidWorkDiv(kernelCfg, devAcc, initBufferKernel,
uBuffacc,data(), pitchCurrAcc, dx. dv):
```





# Setting WorkDiv Manually: Set 3 vectors of workdiv

```
// Set Dim and Index type
using Idx = uint32_t;
using Dim2D = alpaka::DimInt<2u>; // 2 as a type
alpaka::Vec<Dim2D, Idx> gridBlockExtent{M,N}; // 2D grid
alpaka::Vec<Dim2D, Idx> blockThreadExtent{32,32}; // 2D block
alpaka::Vec<Dim2D, Idx> elementExtentPerThread{1,1};
// MxN blocks each has 32x32 threads, each level is 2D
alpaka::WorkDivMembers<Dim2D, Idx> workdiv2D{gridBlockExtent,
blockThreadExtent, elementExtentPerThread};
```

```
using DimlD = alpaka::DimInt<l>;//Set number of dims to 1
using VecID = alpaka::VecKDhmLD, Idx>;//Define alias
auto workDivID = alpaka::WorkDivMembers(VecID[M], VecID[Au], VecID[1u]);
// alternatively
using DimlD = alpaka::DimInt<a>;//Set number of dims to 3
using VecID = alpaka::VecKDhmlD, Idx>; //Define alias
auto workDiv3D = alpaka::WorkDivMembers(Vec3D[1,1,M], Vec3D[1,1,4u], Vec3D[1,1,1u]);
```





# Optimization and usability

# alpaka Usability and Optimization Features

- Use alpaka mdspan to set, get, pass buffers easily (Hands-On 5)
- 2 Use Domain Decomposition: Divide the domain in chunks (Hands-On 6)
- Use 2 asynch queues for performance increase (Hands-On 7)
- 4 Use shared memory for performance increase (Hands-On 8)





# alpaka::experimental::mdspan

### Mdspan a multi-dimensional and non-owning view

- Part of C++23 standard. Can be used with C++17.
- Consists pointer, pitch and data size
- Has member functions to get/set data and to get extents

### Mdspan Installation

- Set alpaka\_USE\_MDSPAN cmake variable to FETCH while installing alpaka
- Alternatively, set alpaka\_USE\_MDSPAN cmake variable to FETCH while configuring example if it is not already set while installation

```
// in build directory
cmake -Dalpaka_USE_MDSPAN=FETCH ..
```

### Passing mdspan to kernel

```
// Host code: Allocate device memory
auto bufDevA = alpaka::allocBuf<DataType, Idx>(devAcc, extentA);
// Create mdspan views for device buffers using alpaka::experimental::getMdSpan
auto mdDevA = alpaka::experimental::getMdSpan(bufDevA);
// Execute the kernel
alpaka::exec<Acc>(queue, workDiv, kernel, mdDevA, mdDevB, mdDevC);
```





### Kernel using mdspan instead of data pointer and pitch info

### Accessing data at host or at accelerator

```
struct MatrixAddKernel
         template < typename TAcc, typename TMdSpan >
         ALPAKA FN ACC void operator()(TAcc const& acc. TMdSpan A. TMdSpan B. TMdSpan
               C) const
         ł
             auto const i = alpaka::getIdx<alpaka::Grid, alpaka::Threads>(acc)[0];
             auto const j = alpaka::getIdx<alpaka::Grid, alpaka::Threads>(acc)[1];
             if(i < C.extent(0) && j < C.extent(1))
                      C(i, i) = A(i, i) + B(i, i):
         } };
    struct StencilKernel
        template < typename TAcc, typename TDim, typename TIdx, typename TMdSpan>
        ALPAKA_FN_ACC auto operator()(
            TAcc const& acc.
            TMdSpan uCurrBuf,
            TMdSpan uNextBuf,
            alpaka::Vec < TDim, TIdx > const chunkSize,
            double const dx.
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            double const dy,
            double const dt) const -> void
        f ....
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        } };
```





### Hands-On Session

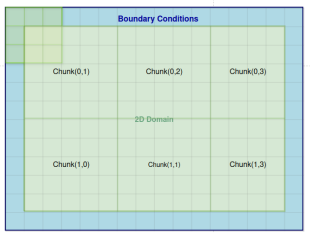
# Hands-on Session5: Use mdspan to pass data to kernel





#### Chunk Definition

**Chunk:** Subdomains needed for latency management of block level parallelisation

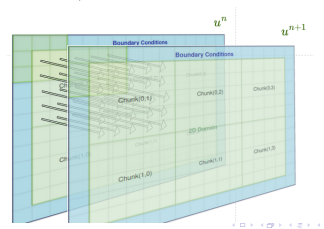






# Calculation by a block of grids of stencil kernel

- $\blacksquare$  Chunking is a Domain Decomposition method
- A block of threads update a chunk of heat data
- A grid of threads updates the whole domain

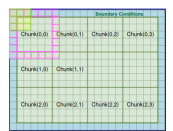


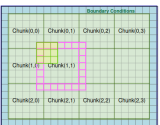




# Chunks in Parallel Grid Computations

- Halo Region around chunk: A layer of grid cells surrounding the subdomains.
- Halo Size: Typically 1 for a 5-point stencil.
- Chunks-size could be larger than the block size.

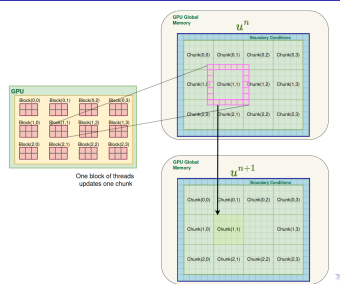








# A block is responsible from a chunk







# Determine WorkDiv for Chunked Solution

#### Set work division fields directly:

```
// Define a workdiv for the shared memory based heat egn solution
    constexpr alpaka::Vec < Dim , Idx > elemPerThread {1, 1};
    // Get max threads that can be run in a block for this kernel
    auto const kernelFunctionAttributes = alpaka::getFunctionAttributes<Acc>(
        devAcc.
        stencilKernel.
        uCurrBufAcc.data(), uNextBufAcc.data(),
        chunkSize.
        pitchCurrAcc, pitchNextAcc,
9
        dx.dv. dt):
    auto const maxThreadsPerBlock = kernelFunctionAttributes.maxThreadsPerBlock;
    auto const threadsPerBlock
        = maxThreadsPerBlock < chunkSize.prod() ? alpaka::Vec < Dim , Idx > {
             maxThreadsPerBlock, 1} : chunkSize:
    alpaka::WorkDivMembers < Dim , Idx > workDiv_manual { numChunks , threadsPerBlock ,
14
         elemPerThreadl:
```





## Hands-On Session

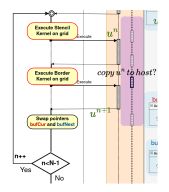
# Hands-on Session6: Optimized Heat Eqn. solution by Domain Decomposition





# Running 2 parallel queues: Additional queue to dump temporary results

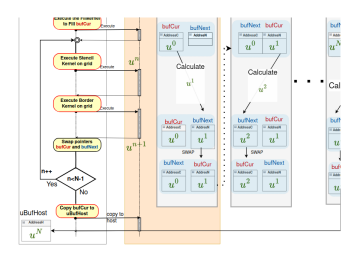
- Create an additional alpaka::queue instance at accelerator to run parallely
- The temporary heat result u<sup>n</sup> will be copied to host from accelerator at the end of each iteration
- Copying can start while the stencil and boundary kernels are running
- In order to run 2 queues paralelly they should be a NonBlocking queues
- The copied heat data will be used to create an animation of images







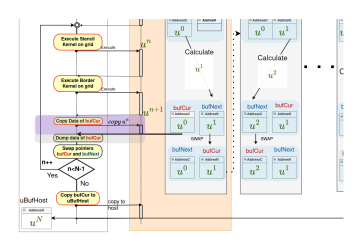
# Current Loop







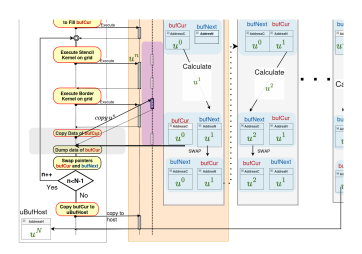
# Copy $u^n$ back at each iteration sequentially







# Copy $u^n$ back at each iteration in parallel by a second Acc Queue







# Hands-On Session

Hands-On Session7: Running 2 parallel queues to dump heat at each step





# Efficient Stencil Application with Shared Memory

#### Shared Memory at GPUs

- A fast, limited-size memory accessible by all threads within a block.
- Used to store data locally in Compute Unit(or SM), reducing the need to access slower global memory.
- Shared Memory allocation can be static or dynamic
  - Static (compile time determined extent)
  - Dynamic (runtime determined extent)
- Filling shared memory is done by the same kernel calculating the stencil
- Threads in a block must synchronize to ensure all data is loaded into shared memory before computation begins.

#### Benefits:

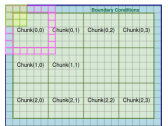
 Reduces memory latency by storing the working set of data (halo + core) in shared memory.





# Shared memory data: chunk with halo

- Halo Region around chunk: A layer of grid cells surrounding the subdomains. In order to use the heat value beside the current chunk
- Halo Size: Typically 1 for a 5-point stencil.
- Chunks might include more than one blocks depending on the blocksize
- Kernel will install the data to shared memory then use the data from shared memory

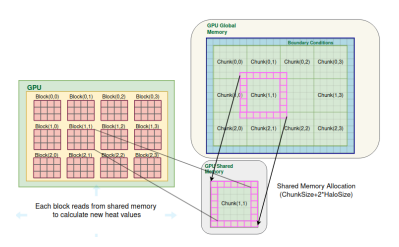








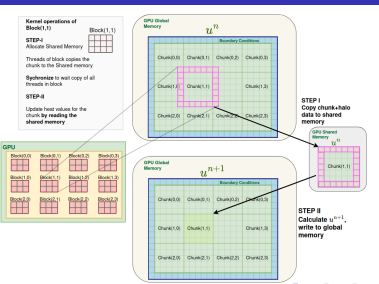
# Updating chunks using shared memory







# Kernel Operations of a Block to Find $u^{n+1}$ for block data







# Steps of Stencil Kernel using shared memory

#### Allocate shared memory inside kernel

```
// Allocate shared memory inside kernel, this will be done only once per
block although it is in the kernel
// Size is determined in compile time and is passed to kernel as a type
autok sdata = alpaka::declareSharedVar<double[T_SharedMemSize1D],
__COUNTER__>(acc);
```

- Calculate thread index
- Fill the shared memory by block of threads
- Wait for shared memory to be filled by all block threads

```
alpaka::syncBlockThreads(acc);
```

- Calculate new heat value using the data from the shared memory
- Set the new heat value





## Hands-On Session

Hands-on Session8: Optimized Heat Eqn. solution by using shared memory





# Conclusion: Parallel Techniques For Solving Heat Equation

#### Kernel Definition

- Kernel to fill a buffer in parallel
- Stencil Kernel for calculating the next set of heat values
- Boundary Kernel

#### Work division

- Getting a valid work division according to accelerator
- Setting work-division manually

#### Allocating and Setting Memory at Host and Accelerator

- Using alpaka::buffer
- Using alpaka::memcpy

#### Alpaka Structures

Accelerator, Device, Queue, Task

#### Optimizations and Usability

- Using alpaka Mdspan
- Domain Decomposition
- Using Multiple Async Queues
- Using GPU's Shared Memory





# End of Alpaka Hackathon. Any Questions?