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Introduction to Alpaka Programming

Kernel Execution

Automatically select a valid kernel launch configuration

```
Vec<Dim, Idx> const globalThreadExtent = vectorValue;
Vec<Dim, Idx> const elementsPerThread = vectorValue;
auto autoWorkDiv = getValidWorkDiv<Acc>(
    device,
    globalThreadExtent, elementsPerThread,
    false,
    GridBlockExtentSubDivRestrictions::Unrestricted);
```

Manually set a kernel launch configuration

Instantiate a kernel and create a task that will run it (does not launch it yet)

Kernel kernel{argumentsForConstructor};
auto taskRunKernel = createTaskKernel<Acc>(workDiv, kernel, parameters);





















alpaka - Abstraction Library for Parallel Kernel Acceleration

Alpaka is...

- A parallel programming library: Accelerate your code by exploiting your hardware's parallelism!
- An abstraction library independent of hardware ecosystem: Create portable code that runs on CPUs and GPUs!
- Free & open-source software







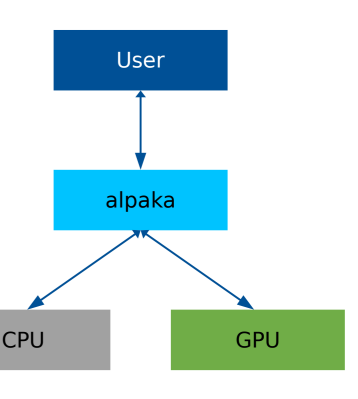
alpaka's purpose

Without alpaka

- Hardware ecosystem is heterogenous and multiple hardware types commonly used (CPUs, GPUs, ...)
- Platforms not inter-operable → parallel programs not easily portable

alpaka: one API to rule them all

- **Abstraction** (not hiding!) of the underlying hardware, compiler and OS platforms
- Easy change of the backend
 - Code needs only minor adjustments to support different accelerators
- Easy indexing of threads in kernels
- Easy setup of the type of parallelism (Block sizes in grid, Thread sizes in block...)
- **Heterogenous Programming**: Using different backends in a synchronized manner

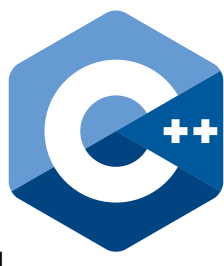






Programming with alpaka

- C++ only!
- Header-only library: No additional runtime dependency introduced
- Modern library: alpaka is written entirely in C++17
- Supports a wide range of modern C++ compilers (g++, clang++, Apple LLVM, MS Visual Studio)
- Portable across operating systems: Linux, macOS, Windows are supported
- Alpaka directly uses vendor API's. For example: Alpaka cuda backend uses Cuda API directly etc.







alpaka is free software (MPL 2.0). Find us on GitHub!

Our GitHub organization: https://www.github.com/alpaka-group

- Contains all alpaka-related projects, documentation, samples, ...
- New contributors welcome!

The library: https://www.github.com/alpaka-group/alpaka

- Full source code
- Issue tracker
- Installation instructions
- Small examples

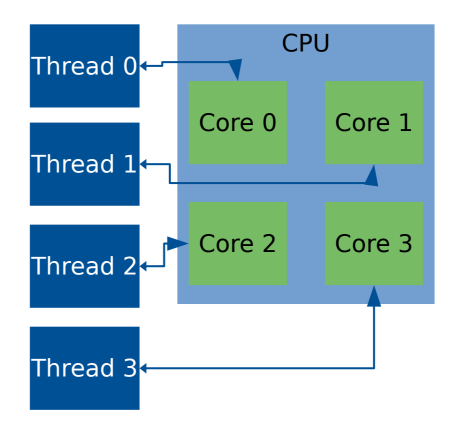






Basics: Thread mapping on CPUs

- CPU consists of multiple cores
 - Because of simultaneous multithreading there can be more logical than physical cores!
- alpaka Threads are executed by CPU cores.
- Single thread per block. Single block per core.
- Multiple elements per thread.



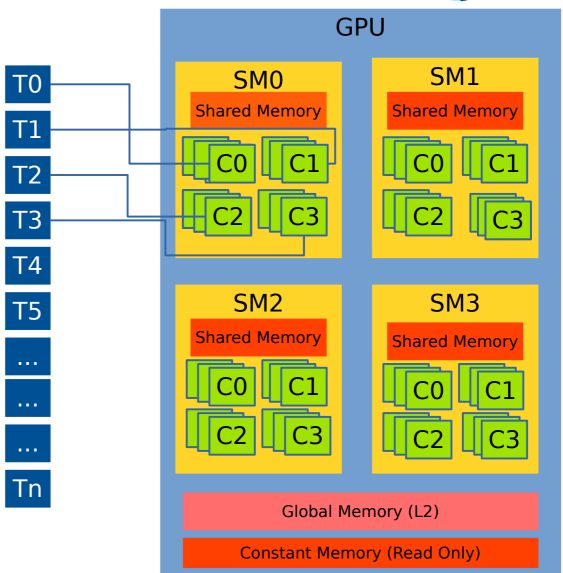




Matching threads to cores in GPUs:

How distribute threads between SMs while matching each thread to a core?

- Main determinants of mapping threads to the SMs:
 - Number of cores SM.
 - Register size in each SM,
 - Shared memory usage of each thread,
 - Threads per SM, Threads per Block, Blocks per Device
- Memory latencies: Global Memory and Constant memory has different latencies.
- Memory sizes: Size of shared memory used by threads in a block or blocks assigned to an SM

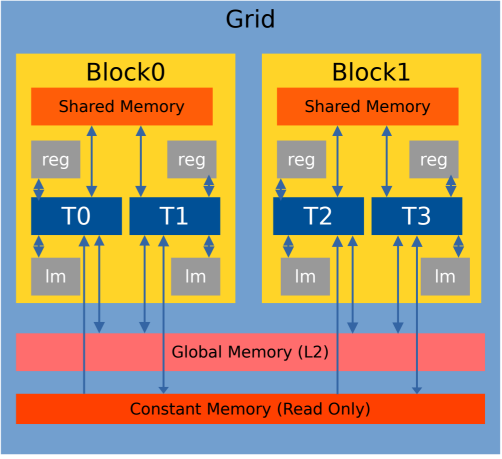






Alpaka proposes and validates WorkDivision!

- WorkDivision data structure consists:
 - Number of Blocks per grid
 - Number of Threads per block
 - Elements per thread
- Alpaka validates and proposes correct parallelisation strategies to map threads to SMs

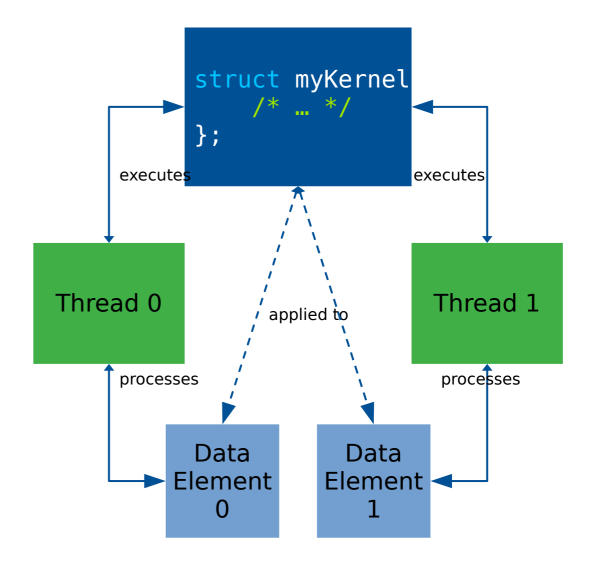






Basics:Threads, Kernels and Indexing

- A Kernel is executed by a number of Threads
- Threads are executing the same algorithm for different data elements
- **Indexing:** Distributing the data to be processed among threads by mapping data indexes to thread indices in the kernel code!
- A Kernel **defines** an algorithm
- A Thread **applies** an algorithm. **Uses** the data part determined by it's thread index.







What is an Alpaka Kernel?

- Contains the algorithm
- Written on per-data-element basis
- alpaka Kernels are functors (function-like C++ structs / classes)
- operator() is annotated with ALPAKA FN ACC specifier
- operator() must return void
- operator() must be const

```
struct HelloWorldKernel {
   template <typename Acc>
   ALPAKA_FN_ACC void operator()(Acc const & acc) const {
        using namespace alpaka;
        uint32 t threadIdx = getIdx<Grid, Threads>(acc)[0];
        printf("Hello, World from alpaka thread %u!\n", threadIdx);
```





Why Alpaka - 1: Easy Indexing of Threads and Data

- Direct calculation of the index of a thread with respec to a grid or block origin in the kernel.
- Mapping the thread indexes to less dimensional space.

```
truct HelloWorldKernel
   template<tvpename TAcc>
   ALPAKA_FN_ACC auto operator()(TAcc const& acc) const -> void
       using Dim = alpaka::Dim<TAcc>;
       using Idx = alpaka::Idx<TAcc>;
       using Vec = alpaka::Vec<Dim, Idx>;
       using Vec1D = alpaka::Vec<alpaka::DimInt<1u>, Idx>;
       // In the most cases the parallel work distibution depends on the current index of a thread
       // and how many threads exist overall. These information can be obtained by
       // getIdx() and getWorkDiv(). In this example these values are obtained for a global scope.
       Vec const globalThreadIdx = alpaka::getIdx<alpaka::Grid, alpaka::Threads>(acc);
       Vec const globalThreadExtent = alpaka::getWorkDiv<alpaka::Grid, alpaka::Threads>(acc);
       // one dimensional thread index space. We call it
       // linearize the thread index.
       Vec1D const linearizedGlobalThreadIdx = alpaka::mapIdx<1u>(globalThreadIdx, globalThreadExtent);
       // Each thread prints a hello world to the terminal together with the global index of the thread in
       // order [z][y][x] where the last index is the fast one.
       printf(
           "[z:%u, y:%u, x:%u][linear:%u] Hello World\n",
           static_cast<unsigned>(globalThreadIdx[@u]),
          static_cast<unsigned>(globalThreadIdx[1u]),
           static_cast<unsigned>(globalThreadIdx[2u]),
           static_cast<unsigned>(linearizedGlobalThreadIdx[0u]));
```





Obtaining the indices in Kernel

- alpaka provides several API functions for obtaining indices:
 - Index of Thread on the Grid: alpaka::getIdx<alpaka::Grid, alpaka::Threads>(acc)[dim];
 - Index of Thread on a Block: alpaka::getIdx<alpaka::Block, alpaka::Threads>(acc)[dim];
 - Index of Block on the Grid: alpaka::getIdx<alpaka::Grid, alpaka::Blocks>(acc)[dim];
- You can also obtain the extents of the Grid or the Blocks:
 - Number of Threads on the Grid: alpaka::getWorkDiv<alpaka::Grid, alpaka::Threads>(acc) [dim];
 - Number of Threads on a Block: alpaka::getWorkDiv<alpaka::Block, alpaka:Threads>(acc) [dim];
 - Number of Blocks on the Grid: alpaka::getWorkDiv<alpaka::Grid, alpaka::Blocks>(acc) [dim];





Why Alpaka-2:

Easy definition of Type of Parallelism by WorkDivision

- Determines the number of kernel instantiations
- Determines the type of parallelism
 - Dimensions of a grid in terms of blocks,
 - Dimensions of a block in terms of threads
 - Elements per thread

```
// Define the work division
// The workdiv is divided in three levels of parallelization:
// - grid-blocks:
                      The number of blocks in the grid
// - block-threads:
                      The number of threads per block (parallel, synchronizable).
// - thread-elements: The number of elements per thread (sequential, not synchronizable)
                       Each kernel has to execute its elements sequentially.
using Vec = alpaka::Vec<Dim, Idx>;
auto const elementsPerThread = Vec::all(static_cast<Idx>(1));
auto const threadsPerGrid = Vec{4, 2, 4};
using WorkDiv = alpaka::WorkDivMembers<Dim, Idx>;
WorkDiv const workDiv = alpaka::getValidWorkDiv<Acc>( devAcc, threadsPerGrid,
    elementsPerThread, false, alpaka::GridBlockExtentSubDivRestrictions::Unrestricted);
// Instantiate the kernel function object
HelloWorldKernel helloWorldKernel;
// Run the kernel
// To execute the kernel, you have to provide the
// work division as well as the additional kernel function parameters.
alpaka::exec<Acc>(queue,workDiv,helloWorldKernel/* put kernel arguments here */);
```





Easy definition of Type of Parallelism: Preparing the Host for 2D Grid

• Go the top of main() and enable 2D dimensionality on the Host:

```
using Dim = dim::DimInt<2>;
```

• Further down in main(), set up a 2D Thread hierarchy:

```
auto blocksPerGrid = alpaka::Vec<Dim, Idx>{2u, 4u};
auto threadsPerBlock = alpaka::Vec<Dim, Idx>{1u, 1u};
auto elementsPerThread = alpaka::Vec<Dim, Idx>{1u, 1u};
```





Important Alpaka Structures

- Accelerator provides abstract view of all capable physical devices. AccCpuThreads, AccGpuCudaRt, AccGpuHipRt...
- **Device** represents a single physical device
- Queue enables communication between the host and a single Device
- Platform is a union of Accelerator, Device and Kernel
- Task is a device-side operation (e.g kernel, memory operation)
- Others: **Event**, **Buffer** (dynamic array), **Vector** (static array)





Why Alpaka3: Changing the accelerator with minimal code change

- Accelerator concept is an abstraction of concrete devices and programming models
- The programmer changes the accelerator in just one line of code
- In the background, an entirely different code path for the "new" device is chosen
- Accelerator provides portable access to device-specific functions

```
/* Before the code change */
using Acc = alpaka::AccCpuOmp2Blocks<Dim,</pre>
Idx>;
/* Kernels will run on CPUs */
/* Parallelism provided by OpenMP 2.x */
/* After the code change */
using Acc = alpaka::AccGpuHipRt<Dim, Idx>;
/* Kernels will run on AMD + NVIDIA GPUs */
/* Parallelism provided by HIP */
```





Switching the Accelerator

- alpaka provides a number of pre-defined Accelerators in the acc namespace.
- For GPUs:
 - AccGpuCudaRt for NVIDIA GPUs
 - AccGpuHipRt for AMD and NVIDIA GPUs
- For CPUs
 - AccCpuFibers based on Boost.fiber
 - AccCpu0mp2Blocks based on OpenMP 2.x
 - AccCpu0mp4 based on OpenMP 4.x
 - AccCpuTbbBlocks based on TBB
 - AccCpuThreads based on std::thread

```
Example: CPU accelerator
using Acc =
alpaka::AccCpuOmp2Blocks<Dim, Idx>;
// Example: CUDA GPU accelerator
using Acc = alpaka::AccGpuCudaRt<Dim,</pre>
Idx>;
  Example: HIP GPU accelerator
using Acc = alpaka::AccGpuHipRt<Dim,</pre>
Idx>;
```

al/aka Accelerator Details



- Accelerator chosen by the programmer and hides hardware specifics behind alpaka's abstract API using Acc = acc::AccGpuCudaRt<Dim, Idx>;
- Inside Kernel: 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)

```
// allocate a variable in block shared static memory
auto & mvSharedVar = block::shared::st::allocVar<int, COUNTER >(acc);
// get pointer to the block shared dynamic memory
float * mySharedBuffer = block::shared::dyn::getMem<float>(acc);
```

It also enables synchronization on the block level

```
// synchronize all threads within the block
block::sync::syncBlockThreads(acc);
```

- Internally, the accelerator maps all device-side functions to their native counterparts
 - Device-side functions require the accelerator as first argument: math::sqrt(acc, /* ... */); time::clock(acc); atomic::atomicOp<atomic::op::Or>(acc, /* ... */, hierarchy::Grids); (Atomics)
- On Host: Meta-parameter for choosing correct physical device and dependent types





Physical device information and management by "alpaka Device"

- Each alpaka Device represents a single physical device;
- Contains device information:

```
auto const name = alpaka::getName(myDev);
                                                   // Back-end-defined device name
auto const bytes = alpaka::getMemBytes(myDev);
                                                  // Size of device memory

    auto const free = alpaka::getFreeMemBytes(myDev); // Size of available device

 memory
```

Provides the means for device management:

```
alpaka::reset(myDev);
                                                   // Reset GPU device state
```

Encapsulates back-end device:

```
    auto nativeDevice = alpaka::getDev(myDev);

                                                       // nativeDevice is not portable!
```

al/aka Queue data structure



Queue: Connecting Host and Device

- alpaka Queues enable communication between Host and Device
- Two queue types: blocking and nonblocking
- Blocking queues block the Host until Device-side command returns
- Non-blocking queues return control to Host immediately, Device-side command runs asynchronously

```
// Choose queue behaviour - Blocking or
NonBlocking
using QueueProperty = alpaka::NonBlocking;
// Define queue type
using Queue = alpaka::Queue<Acc,</pre>
QueueProperty>;
// Create queue for communication with
myDev
auto myQueue = Queue{myDev};
```





Queue operations

- Queues execute Tasks (see next slide):
 - alpaka::enqueue(myQueue, taskRunKernel);
- Check for completion:
 - bool done = alpaka::empty(myQueue);
- Wait for completion, Events (see next slide), or other Queues:
 - alpaka::wait(myQueue); // blocks caller until all operations have completed
 - alpaka::wait(myQueue, myEvent); // blocks myQueue until myEvent has been reached
 - alpaka::wait(myQueue, otherQueue); // blocks myQueue until otherQueue's ops have completed





Setting up Accelerator, Device and Queue

```
// Choose types for dimensionality and indices
using Dim = alpaka::DimInt<1>;
using Idx = std::size t;
// Choose the back-end
using Acc = alpaka::AccGpuHipRt<Dim, Idx>;
// Obtain first device in the HIP GPU list
auto myDev = alpaka::getDevByIdx<Acc>(0u);
// Create non-blocking queue for chosen device
using Queue = alpaka::Queue<Acc, alpaka::NonBlocking>;
auto myQueue = Queue{myDev};
// Done! We can now enqueue device-side operations.
```





Tasks and Events

- Device-side operations (kernels, memory operations) are called Tasks
- Tasks on the same queue are executed in order (FIFO principle)

```
alpaka::enqueue(queueA, task1);
alpaka::enqueue(queueA, task2); // task2 starts after task1 has finished
```

- Order of tasks in different gueues is unspecified
 - alpaka::enqueue(queueA, task1); alpaka::enqueue(queueB, task2); // task2 starts before, after or in parallel to task1
- For easier synchronization, alpaka Events can be inserted before, after or between Tasks:

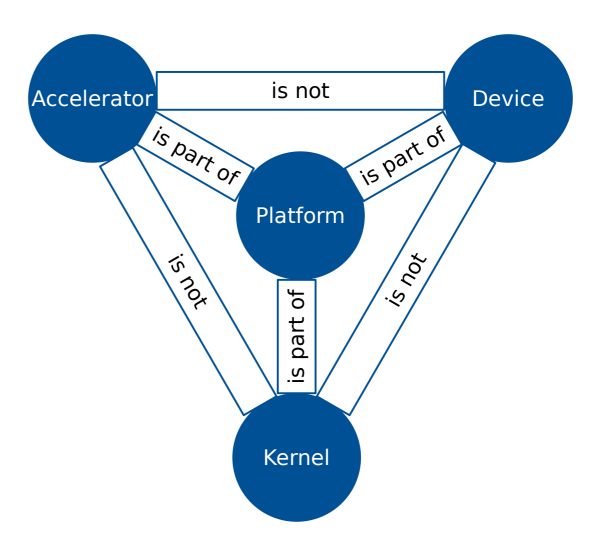
```
auto myEvent = alpaka::Event<alpaka::Queue>(myDev);
alpaka::enqueue(queueA, myEvent);
alpaka::wait(queueB, myEvent); // queueB will only resume after queueA reached myEvent
```





alpaka Platform

- Platform is meta-concept in alpaka
- Union of Accelerator, Device and Kernel functionality
 - Accelerator gives structure to the host side and functionality to the device side
 - · Device gives functionality to the host side
 - Kernels are agnostic of Device details
 - → Portable Kernels







Changing the target platform

```
using namespace alpaka;
using Dim = dim::DimInt<1u>;
using Idx = std::size t;
/*** BEFORE ***/
using Acc = alpaka::AccCpu0mp2Blocks<Dim, Idx>;
/*** AFTER ***/
using Acc = alpaka::AccGpuHipRt<Dim, Idx>;
/* No change required - dependent types and variables are automatically changed */
auto myDev = alpaka::getDevByIdx<Acc>(0u);
using Queue = alpaka::Queue<Acc, queue::NonBlocking>;
auto myQueue = Queue{myDev};
```



Summary of Alpaka Structures

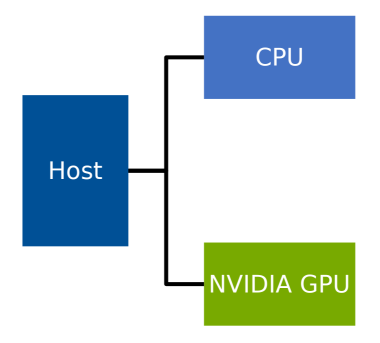
- Accelerator provides abstract view of all capable physical devices
- **Device** represents a single physical device
- **Queue** enables communication between the host and a single Device
- Platform is a union of Accelerator, Device and Kernel
- **Task** is a device-side operation (e.g kernel, memory operation)
- Others: **Event, Buffer** (dynamic array), **Vector** (static array)
- Question: How is portability between back-ends achieved?





Programming Heterogeneous Systems

- Real-world scenario: Use all available compute power
- Also real-world scenario: Multiple different hardware types available
- Requirement: Usage of one back-end per hardware platform
- Requirement: Back-ends need to be interoperable







Programming Heterogeneous Systems

Why Alpaka - 4: Using multiple **Platforms Synchronously**

- Alpaka enables easy heterogeneous programming!
- Create one Accelerator per backend
- Acquire at least one Device per Accelerator
- Create one Queue per Device

```
// Define Accelerators
using AccCpu = alpaka::AccCpu0mp2Blocks<Dim, Idx>;
using AccGpu = alpaka::AccGpuCudaRt<Dim, Idx>;
// Acquire Devices
auto devCpu = alpaka::getDevByIdx<AccCpu>(0u);
auto devGpu = alpaka::getDevByIdx<AccGpu>(0u);
// Create Oueues
using QueueProperty = alpaka::NonBlocking;
using QueueCpu = alpaka::Queue<AccCpu, QueueProperty>;
using QueueGpu = alpaka::Queue<AccGpu, QueueProperty>;
auto queueCpu = QueueCpu{devCpu};
auto queueGpu = QueueGpu{devGpu};
```





Programming Heterogeneous Systems

Communication by Buffers

- Buffers are defined and created per Device
- Buffers can be copied between different Devices / Oueues
- Not restricted to a single platform!
- **Restriction**: CPU to GPU copies (and vice versa) require GPU queue

```
// Allocate buffers
auto bufCpu = alpaka::allocBuf<float, Idx>(devCpu, extent);
auto bufGpu = alpaka::allocBuf<float, Idx>(devGpu, extent);
/* Initialization ... */
// Copy buffer from CPU to GPU - destination comes first
alpaka::memcopy(gpuQueue, bufGpu, bufCpu, extent);
// Execute GPU kernel
alpaka::enqueue(qpuQueue, someKernelTask);
// Copy results back to CPU and wait for completion
alpaka::memcopy(qpuQueue, bufCpu, bufGpu, extent);
// Wait for GPU, then execute CPU kernel
alpaka::wait(cpuQueue, gpuQueue);
alpaka::enqueue(cpuQueue, anotherKernelTask);
```





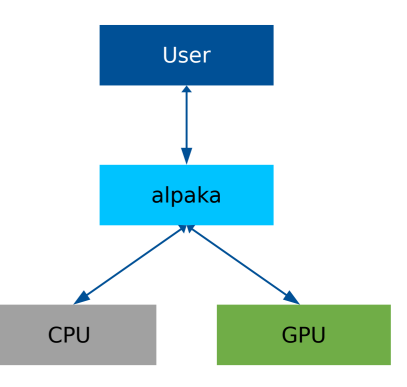
As a summary

Without alpaka

- Multiple hardware types commonly used (CPUs, GPUs, ...)
- Increasingly heterogeneous hardware configurations available
- Platforms not inter-operable → parallel programs not easily portable

alpaka: one API to rule them all

- **Abstraction** (not hiding!) of the underlying hardware & software platforms
 - · AMD, Nvidia, Intel GPUs, CPU
- Easy change of the backend
 - Code needs only minor adjustments to support different accelerators
- Easy indexing of threads in kernels
- Easy setup of the type of parallelism by WorkDivision (Block sizes in grid, Thread sizes in I
- Heterogenous Programming: Using different backends in a synchronized manner







Thank you! If you use alpaka for your research, please cite one of the following publications:

Matthes A., Widera R., Zenker E., Worpitz B., Huebl A., Bussmann M. (2017): Tuning and Optimization for a Variety of Many-Core Architectures Without Changing a Single Line of Implementation Code Using the Alpaka Library. In: Kunkel J., Yokota R., Taufer M., Shalf J. (eds) High Performance Computing. ISC High Performance 2017. Lecture Notes in Computer Science, vol 10524. Springer, Cham, DOI: 10.1007/978-3-319-67630-2 36.

E. Zenker et al., "Alpaka - An Abstraction Library for Parallel Kernel Acceleration", 2016 IEEE International Parallel and Distributed Processing Symposium Workshops (IPDPSW), Chicago, IL, 2016, pp. 631 – 640, DOI: **10.1109/IPDPSW.2016.50**.

Worpitz, B. (2015, September 28). Investigating performance portability of a highly scalable particle-in-cell simulation code on various multi-core architectures. Zenodo. DOI: 10.5281/zenodo.49768.



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