



# Developing Stencil Code using the YASK Framework

This tutorial applies to YASK version 3.04 (and later versions in most cases)

Chuck Yount  
Intel Americas, Inc.

March 30, 2020  
© 2019-2020 Intel Corporation

# NOTICES AND DISCLAIMERS

Intel technologies' features and benefits depend on system configuration and may require enabled hardware, software or service activation. Performance varies depending on system configuration.

No computer system can be absolutely secure.

Tests document performance of components on a particular test, in specific systems. Differences in hardware, software, or configuration will affect actual performance. For more complete information about performance and benchmark results, visit <http://www.intel.com/benchmarks>.

Software and workloads used in performance tests may have been optimized for performance only on Intel microprocessors. Performance tests, such as SYSmark and MobileMark, are measured using specific computer systems, components, software, operations and functions. Any change to any of those factors may cause the results to vary. You should consult other information and performance tests to assist you in fully evaluating your contemplated purchases, including the performance of that product when combined with other products. For more complete information visit <http://www.intel.com/benchmarks>.

Intel® Advanced Vector Extensions (Intel® AVX)\* provides higher throughput to certain processor operations. Due to varying processor power characteristics, utilizing AVX instructions may cause a) some parts to operate at less than the rated frequency and b) some parts with Intel® Turbo Boost Technology 2.0 to not achieve any or maximum turbo frequencies. Performance varies depending on hardware, software, and system configuration and you can learn more at <http://www.intel.com/go/turbo>.

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Cost reduction scenarios described are intended as examples of how a given Intel-based product, in the specified circumstances and configurations, may affect future costs and provide cost savings. Circumstances will vary. Intel does not guarantee any costs or cost reduction.

Intel does not control or audit third-party benchmark data or the web sites referenced in this document. You should visit the referenced web site and confirm whether referenced data are accurate. Intel, the Intel logo, and Intel Xeon are trademarks of Intel Corporation in the U.S. and/or other countries.

\*Other names and brands may be claimed as property of others.

© 2018 Intel Corporation.

# Outline

## Introduction

- What is YASK?
- Motivation and example YASK application
- Scope of this presentation

## Basic features and usage

- Download, build, and test
- High-level flow
- Writing a simple stencil in the DSL
- Building the YASK library
- Testing and tuning your stencil
- Multi-node usage via MPI

## Using the APIs

- Access, terms, and basic usage
- Advanced APIs and exceptions

## Advanced stencils and tuning

- Multiple stencils
- Boundary regions (sub-domains)
- Temporal conditions
- Vector folding
- Cluster, bundles, and stages
- FP precision and prefetching
- 3 layers of temporal tiling
- Nested OpenMP in sub-blocks
- More on the auto-tuners

## Wrap-up

- Current work
- Further reading and call to action

# INTRODUCTION

# YASK: “Yet Another Stencil Kit”

YASK is a software *framework* for the rapid development of HPC stencil-based applications

- Stencil: an iterative kernel that updates elements in one or more N-dimensional vars using a fixed pattern of computation on neighboring elements
- Fundamental algorithm in many scientific simulations, e.g., finite-difference-method (FDM) approximations of differential equations describing various physical phenomena

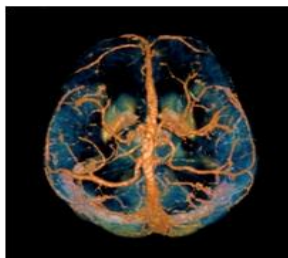
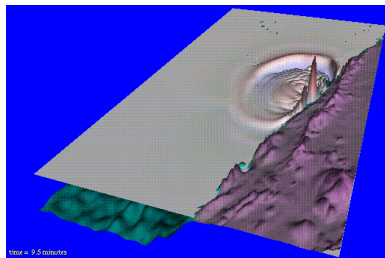
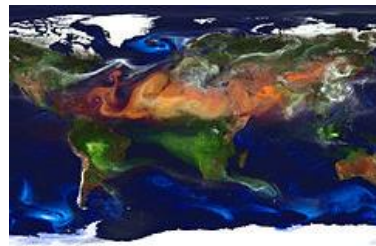


Image Processing



Seismic Modeling



Weather Simulation

Fun fact: prior to version 3, “YASK” stood for “Yet Another Stencil Kernel”

Images from <https://commons.wikimedia.org>

# Technical and business motivation

## Rapid Development

- Stencils in YASK are coded in a simple DSL (domain-specific language)
  - YASK programmer only needs to describe *what* to do, not *how* to do it
  - YASK compiler generates high-perf code from the DSL description
  - Can easily and quickly change the stencil and generate new code
  - Supports arbitrary dimensions, complex stencils, boundary conditions, and more
- Can easily and quickly try different tuning features and parameters without recoding
  - Many complex optimization techniques are available immediately
  - Supports cluster scaling, spatial and temporal tiling, vector folding, and more
- Generated code compiles into a library with documented C++ and Python\* APIs to facilitate integration into real HPC applications

## Performance Portability

- Can re-target stencil for different Intel® platforms by generating multiple libraries from single DSL description
- Future YASK features and supported platforms can be leveraged immediately without recoding

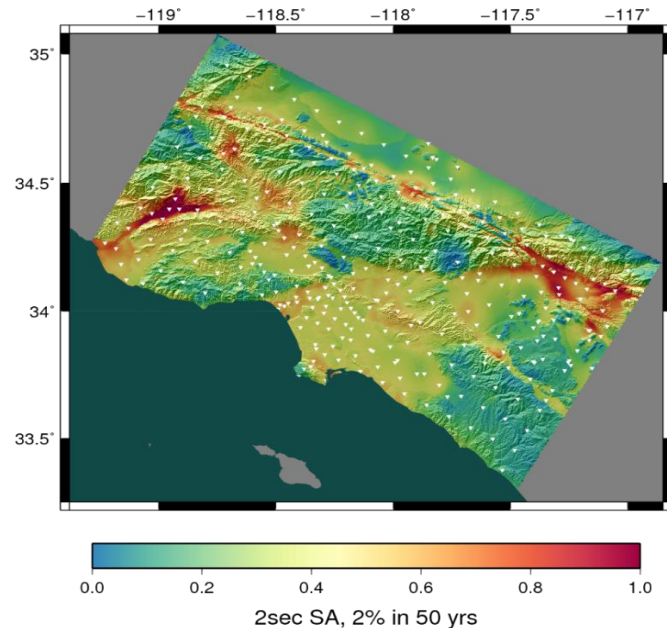
# Example application of YASK

## AWP-ODC: Anelastic Wave Propagation- Olsen, Day, Cui

- Software that simulates seismic wave propagation after a fault rupture
- Widely used by the Southern California Earthquake Center (SCEC) community

## AWP-ODC-OS

- First ever open source release in 2016 (BSD-2 license), including port to Intel Xeon Phi processor, under development by San Diego Supercomputer Center (SDSC) at Univ. of CA, San Diego (UCSD)
- Demonstrated on >9000 nodes of Cori supercomputer



- CyberShake Study 15.4 hazard map for 336 sites around Southern California
- Warm colors represent areas of high hazard

Content on this slide courtesy of UCSD

# Scope of this presentation

## Goals

- Give a practical introduction to using YASK—a tutorial
- Provide an overview of the major YASK performance techniques
- Deliver enough information for someone already familiar with the application of stencil codes to start using YASK

## Non-goals

- Not providing a tutorial on
  - Finite-difference methods, seismic modeling, brain imaging, etc.
  - C++, Linux\*, OpenMP\*, MPI\*, github\*, etc.
  - Intel® instruction sets, microprocessor architecture, etc.
- Not explaining in-depth how the YASK performance techniques work
- Not providing performance data on all the various trade-offs
- Not describing the internal software architecture
- Not a reference manual for the APIs



# BASIC FEATURES AND USAGE

# Download, build, and test

## Code access

- Download from Intel's github\* project
  - `git clone https://github.com/intel/yask`
  - MIT open-source license
- Builds and runs are made from the top-level directory: `cd yask`

## Pre-requisites

- See `README.md` for complete list
- Only supported OS is Linux (no specific distribution recommended)
- Intel® C++ compiler needed for performance
  - Install “Intel® Parallel Studio XE Cluster Edition for Linux”
  - Binary built from gcc\* is consistently lower performance
- Common utilities (gcc, perl, awk, etc.)

Watch for this  
symbol indicating  
steps to try



# Example 1: Iso3dfd stencil

## Description

- Isotropic 3D finite-difference code found in seismic-imaging software used by energy-exploration companies to predict the location of oil and gas deposits
  - Simple stencil with only one updated variable



## Recipes for building and running

- Makefile and run script will automatically determine architecture
  - `make clean; make stencil=iso3dfd`
  - `bin/yask.sh -stencil iso3dfd -g 1024`
- The `-g 1024` option sets the global-domain size to a cube, 1024 elements on a side

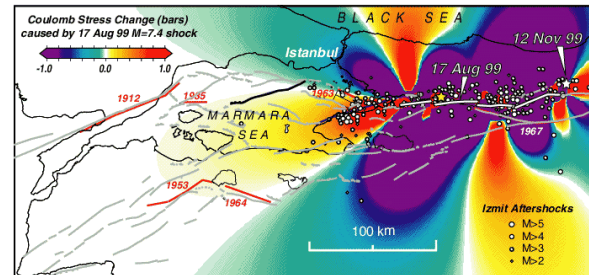


Image from <https://commons.wikimedia.org/wiki/File:PlatformHolly.jpg>. Public domain--U.S. DoE.

# Example 2: AWP stencil

## Description

- Primary compute kernel for earthquake simulator described in the introduction
  - More complex problem that consists of 26 vars in a staggered-grid formulation



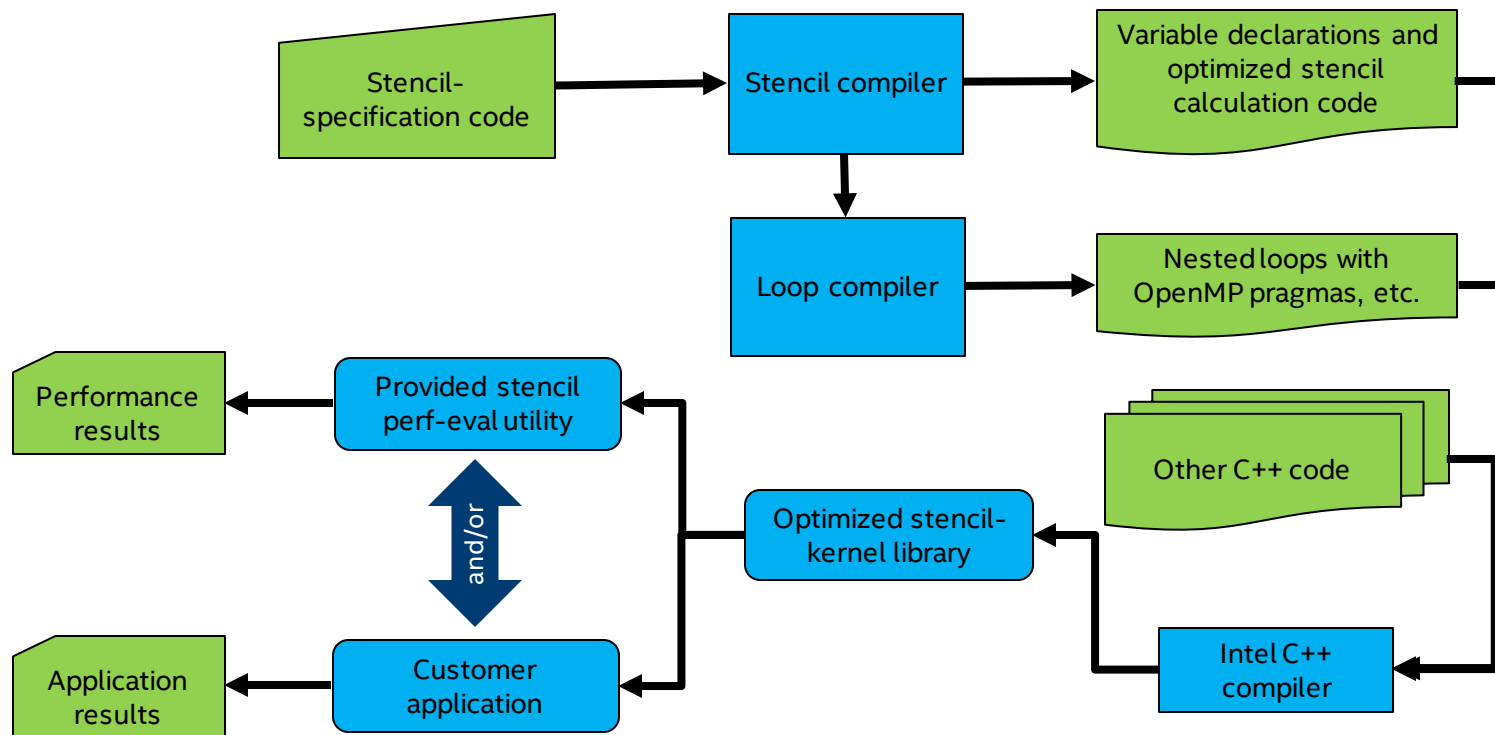
## Recipes for building and running

- Makefile and run script will automatically determine architecture
  - `make clean; make stencil=awp`
  - `bin/yask.sh -stencil awp -gx 1024 -gy 1024 -gz 128`
- The `-g*` options set the global-domain size to specific sizes in each dimension

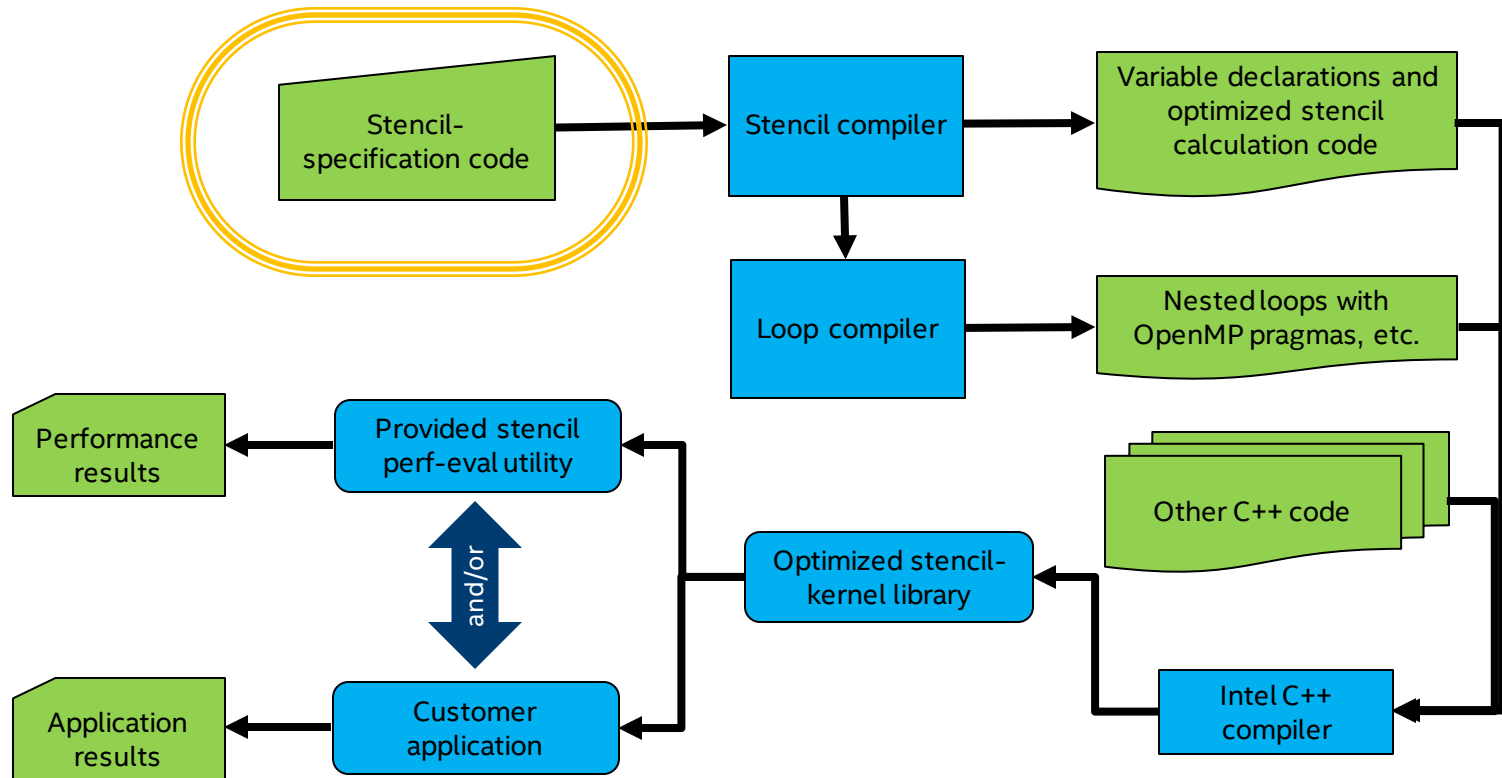


Image from [https://commons.wikimedia.org/wiki/File:Izmit\\_11-12-99.gif](https://commons.wikimedia.org/wiki/File:Izmit_11-12-99.gif). Public domain--U.S.G.S.

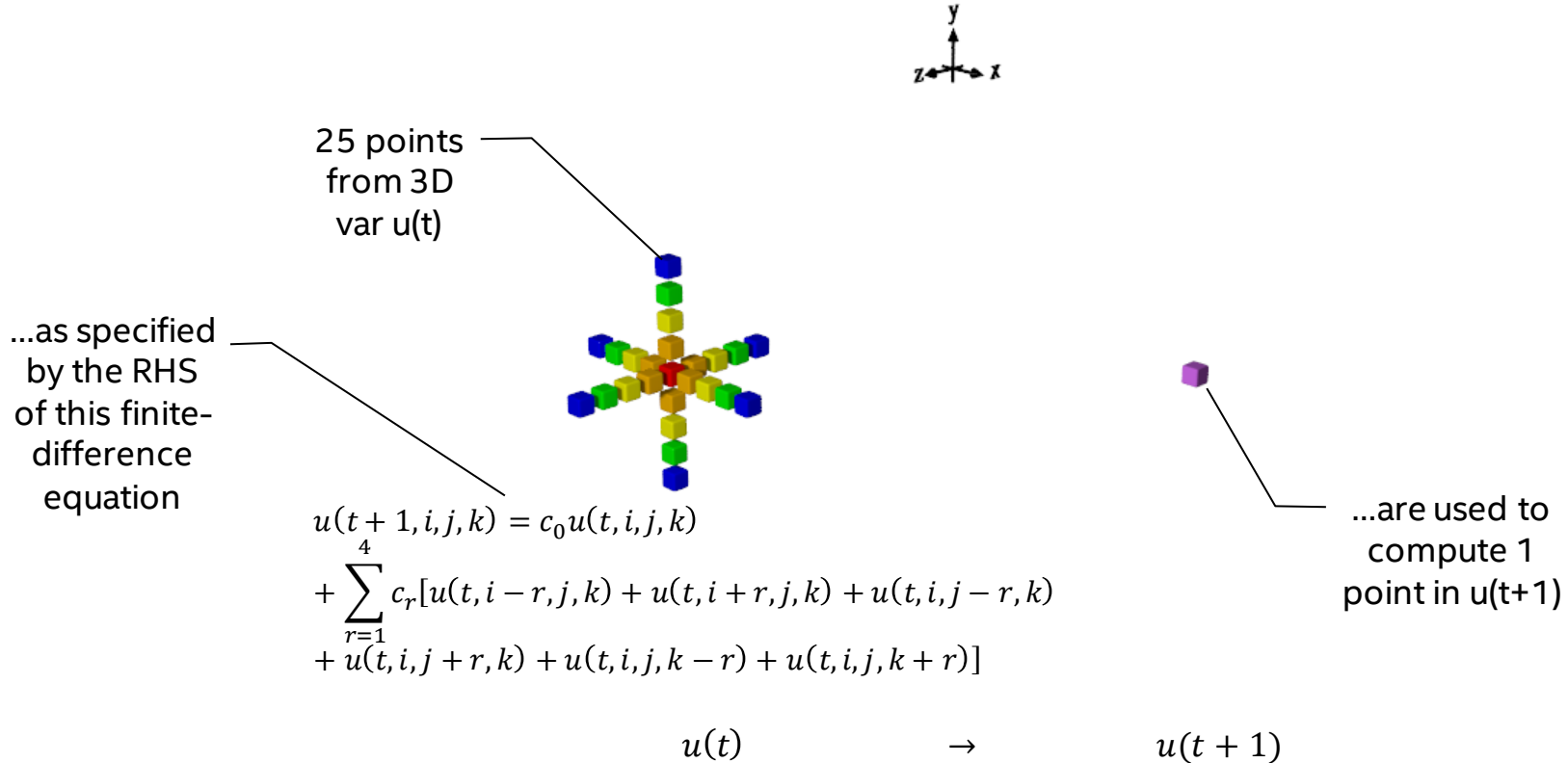
# High-level tool-chain flow



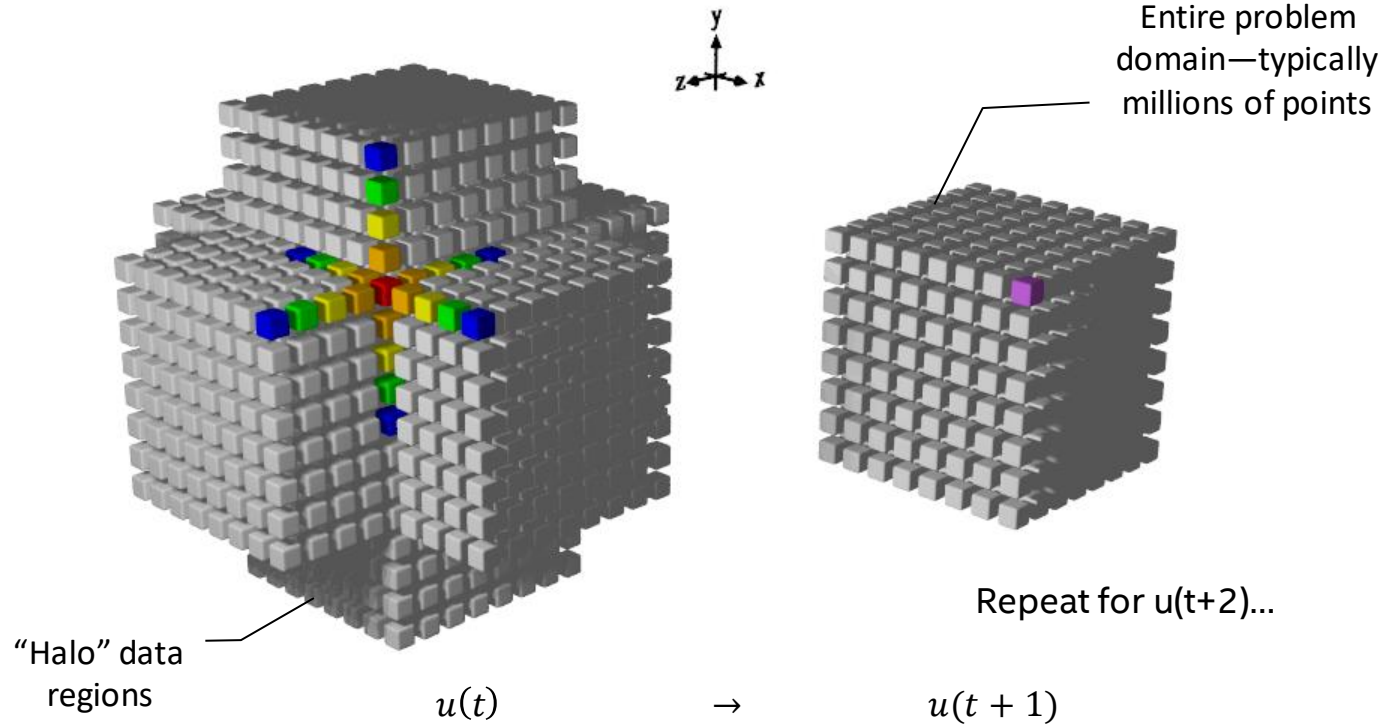
# Stencil specification



# Example simple 25-point 3-D stencil



# Stencil will be applied over entire problem domain





# Example stencil DSL code

Mathematical description of finite-difference approximation implemented in DSL code

$$u(t+1, i, j, k) = c_0 u(t, i, j, k) + \sum_{r=1}^4 c_r [u(t, i-r, j, k) + u(t, i+r, j, k) + u(t, i, j-r, k) + u(t, i, j+r, k) + u(t, i, j, k-r) + u(t, i, j, k+r)]$$

```
#include "yask_compiler_utility_api.hpp"
using namespace yask;
```

```
class MyStencil : public yc_solution_with_radius_base {
```

```
public:
```

```
MyStencil(int radius=4) :
```

```
yc_solution_with_radius_base("my_stencil", radius) { }
```

```
virtual void define() {
```

```
auto t = new_step_index("t");
```

```
auto x = new_domain_index("x");
```

```
auto y = new_domain_index("y");
```

```
auto z = new_domain_index("z");
```

```
yc_var_proxy u("U", get_soln(), { t, x, y, z });
```

```
auto i = new_misc_index("i");
```

```
yc_var_proxy c("C", get_soln(), { i });
```

```
auto nu = c(0) * u(t, x, y, z);
```

```
for (int r = 1; r <= get_radius(); r++)
```

```
nu += c(r) * (u(t, x-r, y, z) + u(t, x+r, y, z) +
```

```
u(t, x, y-r, z) + u(t, x, y+r, z) +
```

```
u(t, x, y, z-r) + u(t, x, y, z+r));
```

```
u(t+1, x, y, z) EQUALS nu;
```

```
 }
```

```
};
```

```
static MyStencil MyStencil_instance;
```

Derive new class from  
yc\_solution\_base (or  
yc\_solution\_with\_radius\_base to  
get radius parameter)

Overload define() method

Declare 1 time and 3 space  
indices and 4D "u" var

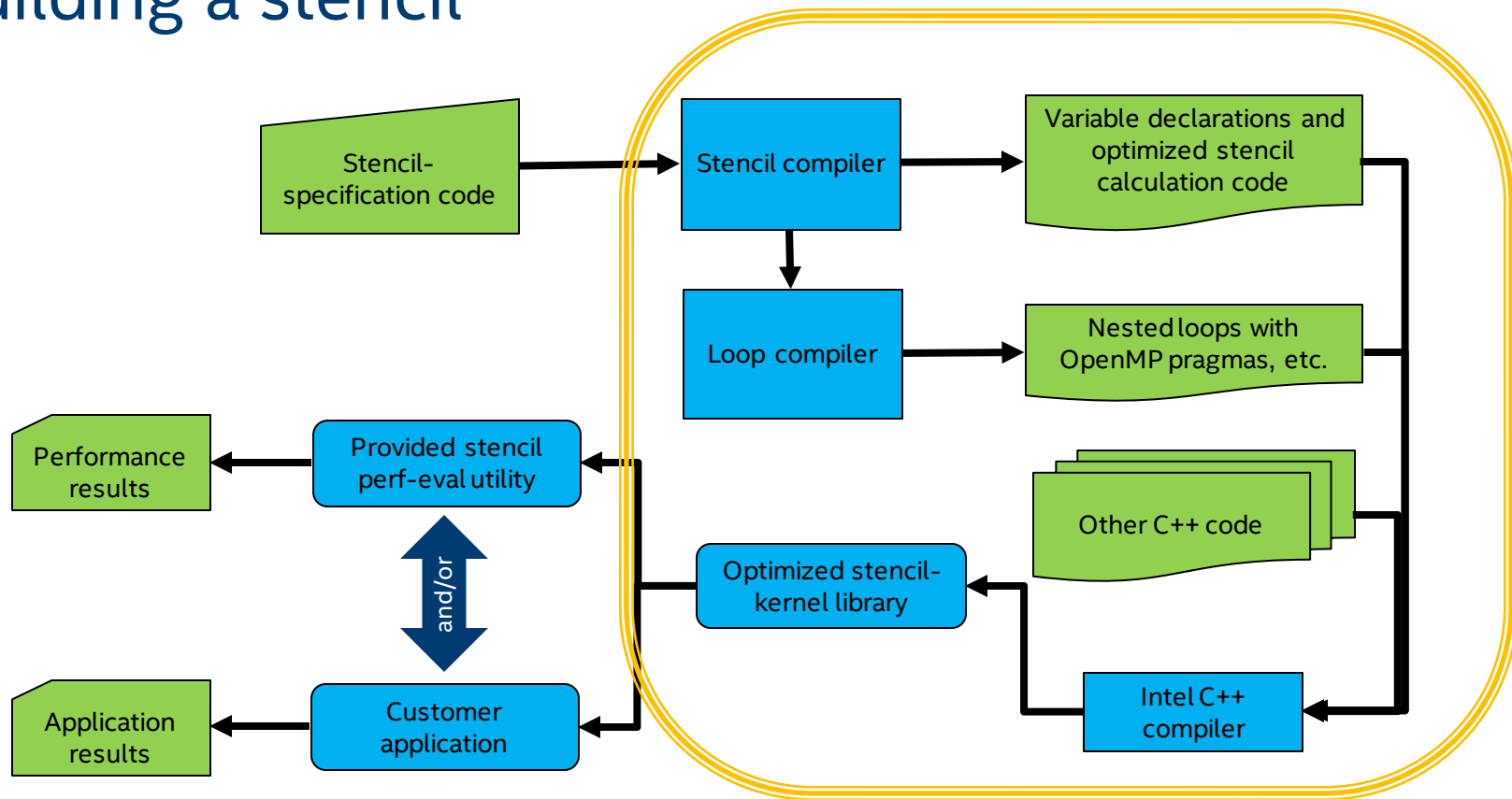
Declare misc index and 1D  
"c" array for coefficients

Write expression for value  
at step t+1

Use EQUALS operator to define  
u(t+1)—this is not an  
assignment!

Register this stencil with the YASK  
compiler utility

# Building a stencil



# Building your example stencil

## Build your stencil and the YASK compiler

- Put the code from a couple of slides back into a new file `src/stencils/MyStencil.cpp`
  - If you really don't want to copy the example stencil during this tutorial, just substitute `iso3dfd` for `my_stencil` in the example commands from here on
  - If you want to write your own stencil, see the examples in `src/stencils` and the full DSL documentation for the YASK Compiler as discussed in the “Using the APIs” section later in this tutorial
- *Optional:* rebuild and run the YASK compiler manually
  - Build the compiler: `make -j compiler`
  - Run the compiler: `bin/yask_compiler.exe -h`
    - “`my_stencil`” should be listed as a valid parameter to the `-stencil` option
    - Lots of other options shown; most of these are passed automatically from the `Makefile`

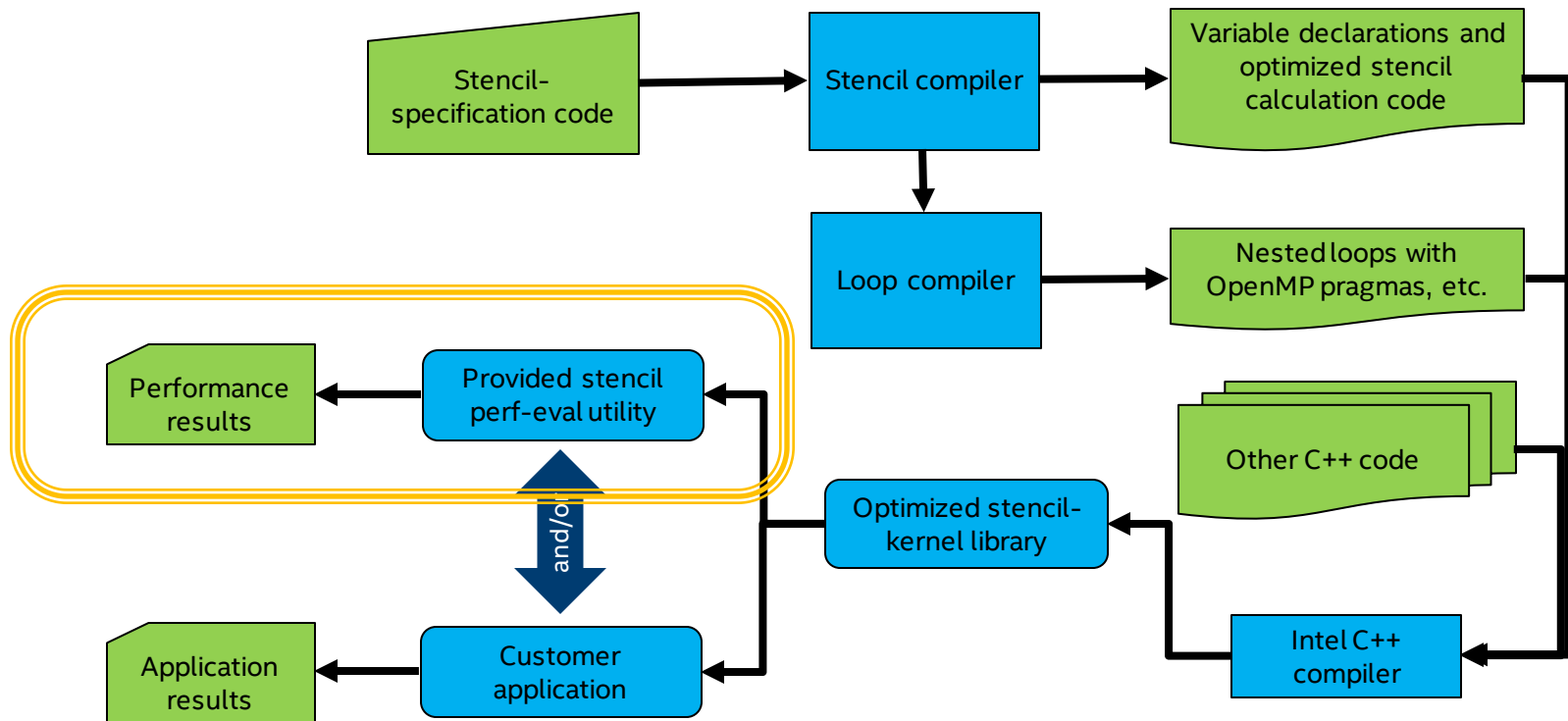


## Build the YASK kernel

- Build the kernel: `make -j stencil=my_stencil radius=4`
  - Targets the instruction-set architecture of your current platform by default
    - If you want to cross-compile, specify `arch=isa-code`, e.g., `kn1, avx2`
  - Uses all the default parameters for static options such as data layout and prefetching
  - Dynamic options such as problem size are specified at run-time
- If you skipped the step of building the compiler above, it will be done here for you automatically



# Running a stencil



# Running your example stencil on one node

## Run the provided test utility

- `bin/yask.sh -stencil my_stencil -g 512 -no-pre_auto_tune`
  - Problem size is specified as 512 in the 3 spatial dimensions (512<sup>3</sup> points) using `-g` option (“g” for global-domain)
  - Other parameters are set to their defaults, except the auto-tuner is disabled
  - The utility prints lots of stats about the stencil, memory usage, performance, etc.
  - Runs 3 trials for about 10 sec each and reports the best and median time and throughput (rate) stats
  - A log file is kept in the “logs” directory, which includes some compilation and platform information for posterity



## Use the auto-tuner

- `bin/yask.sh -stencil my_stencil -g 512`
  - As above, but with the block-size auto-tuner enabled (by default)
    - The block size determines the amount of work done by each OpenMP “parallel for” iteration
    - The utility prints each block size that the auto-tuner tries and its measured performance
  - Compare performance of the two runs



## If your node has more than one socket, try an optional communication method

- `bin/yask.sh -stencil my_stencil -g 512 -use_shm`
  - As above, but uses shared-memory buffers to speed up inter-socket communication
  - This is not the default setting because not all nodes are configured to allow enough shared memory for large problems
    - You will get an out-of-memory error (or sometimes MPI will just exit) when trying to allocate memory if this is the case



# Tuning the size parameters

## Block size

- From the log file of the last run, find the best block size found by the auto-tuner
  - For example, `auto-tuner: best-block-size: t=1 * x=96 * y=64 * z=128`
- Specify this manually to bypass the auto-tuner, e.g.,
  - `bin/yask.sh -stencil my_stencil -g 512 -no-pre_auto_tune \`  
`-bx 96 -by 64 -bz 128`
  - You should get similar performance
  - Play around with block size to see its effect
  - *Contribution opportunity:* github ticket #159 to save and reuse the settings automatically



## General usage of size-parameter options

- Most domain-dimension sizes can be specified in two ways
  - Spatial dims set separately as we did with block size, e.g., `-bx`, `-by`, `-bz`
  - Leave off the dimension names to set all sizes as we did with domain size, e.g., `-g`
- Play around with various problem sizes to see the effect on memory and performance



# Getting a list of options

The YASK test utility is composed of a script driver and a binary

- Print script options: `bin/yask.sh -h`
  - There is only a short list of options here
  - These are just the ones that control how the binary is launched from the script
- Print binary options: `bin/yask.sh -stencil my_stencil -help`
  - Need to specify the stencil because there are different binaries for each
    - If you don't specify a stencil, the script will show the ones that have been compiled
  - *Warning:* the long list might be overwhelming; we'll point out the most important ones in this tutorial 😊



# Scaling out to multiple nodes

## Domain decomposition

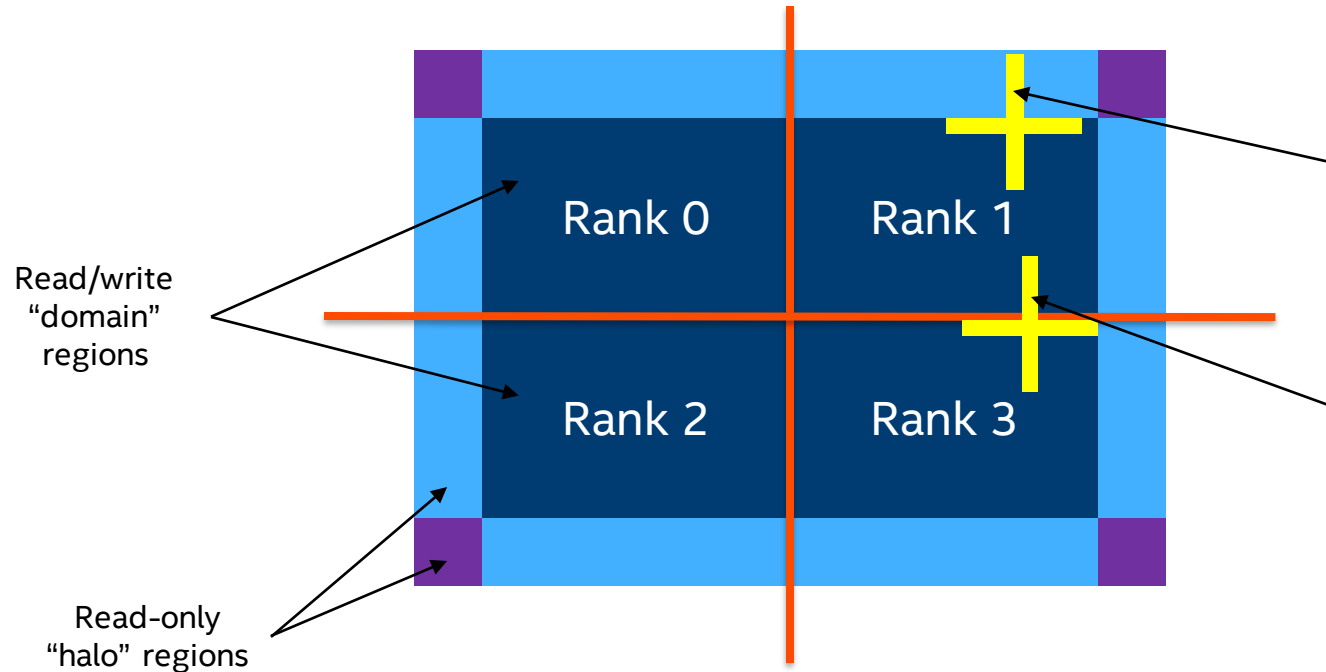
- Many HPC problem sizes won't fit in the DRAM memory of one node
- Most HPC problems can be divided spatially across multiple nodes

## YASK implementation

- Exchanges required inter-node data via two-sided MPI\* communication
- *Important:* strong vs. weak HPC scaling
  - Global-domain size parameters ( $-g$ ) apply to the overall problem size
    - This implements a *strong-scaling* model, where the overall problem size remains constant regardless of the number of nodes
  - Local-domain size parameters ( $-l$ ) apply to the size on each MPI rank
    - This implements a *weak-scaling* model, where the overall problem size increases with the number of nodes



# Conceptual view of 2D, 2×2 rank domains



Due to the read-radius of stencils, halo data must be read to calculate some points in the domain.

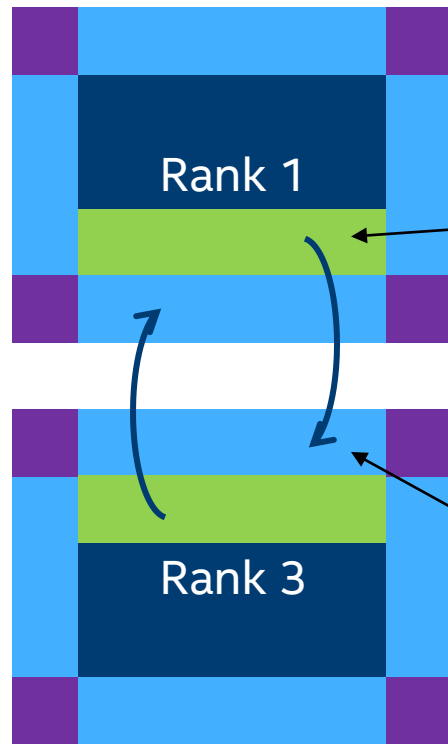
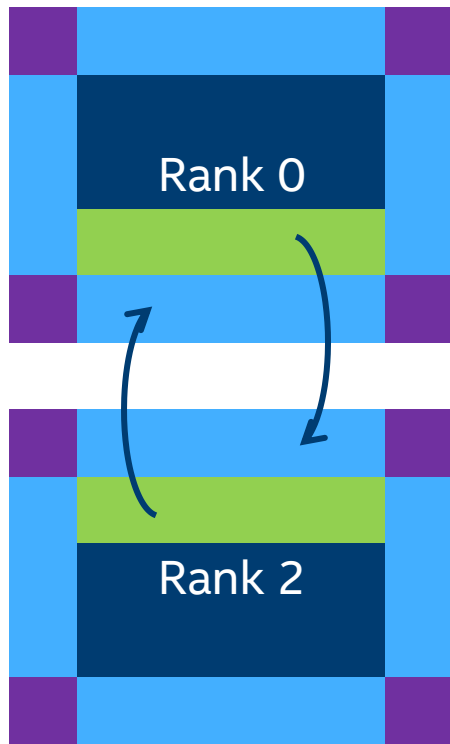
Similarly, at rank boundaries, each rank needs to read data from its neighbors.

This data changes every time-step and must be exchanged between ranks every time-step to keep the data consistent.

# Schematic of 2D, 2×2 rank y-edge halo exchanges

Halos are exchanged at beginning of run and after each time-step.

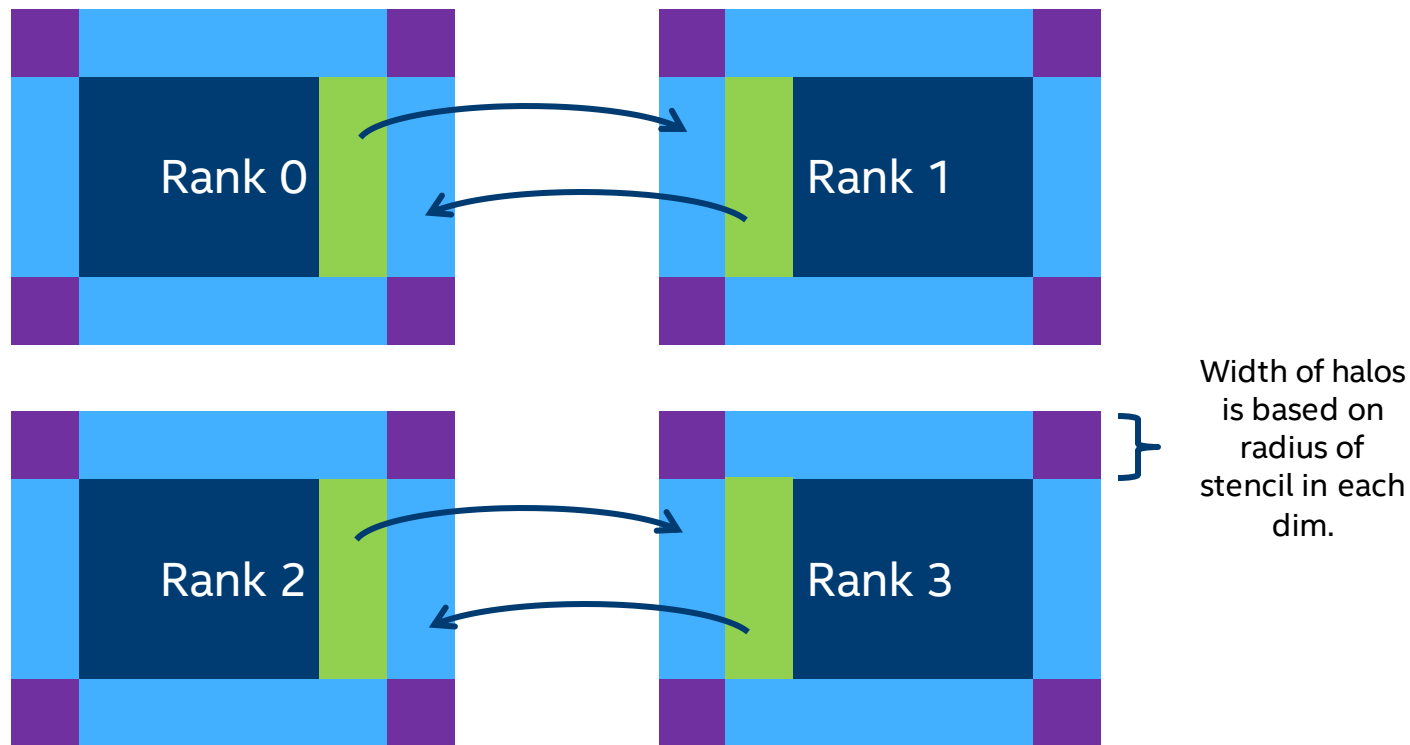
Can get more complex when there are multiple dependent stencils.



Data calculated in rank 1 at time-step  $n...$

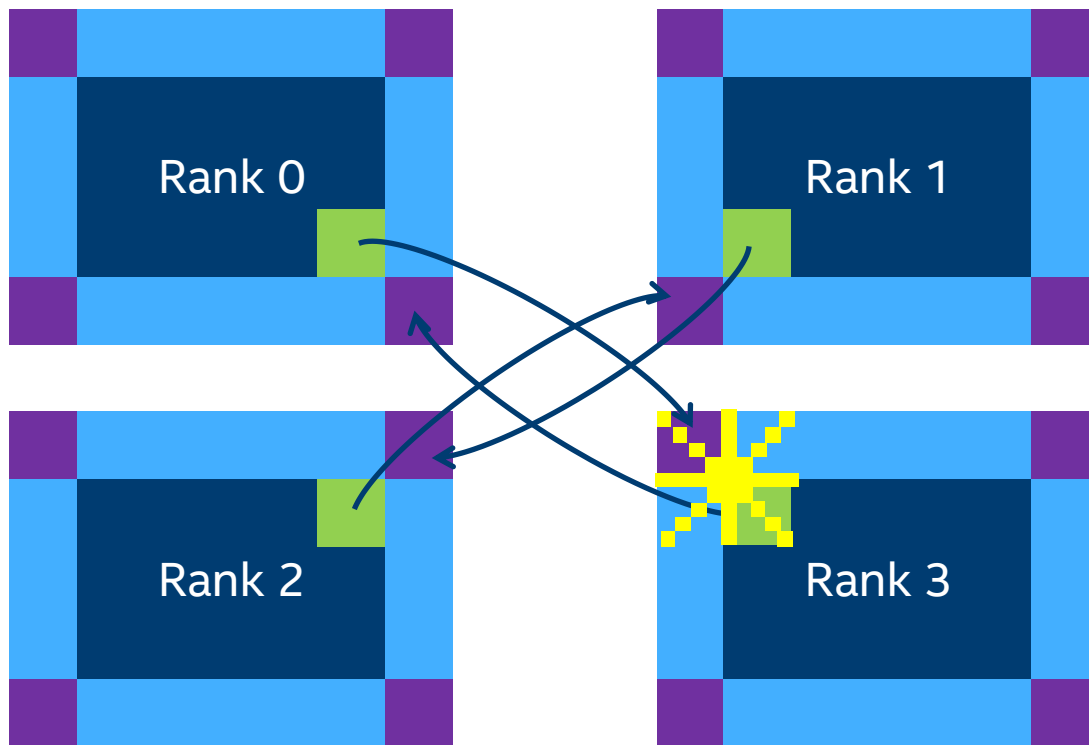
...is needed by rank 3 at time-step  $n+1$

# Schematic of 2D, 2×2 rank x-edge halo exchanges



# Schematic of 2D, 2×2 rank corner exchanges

Corner exchanges are only needed for halos that have diagonal points.




For 3D stencils, exchanges may be needed along faces, edges, and corners of the rectangular solids.


This concept can be extended to any number of dims.

# Running on multiple nodes

## Running the test utility with an MPI command

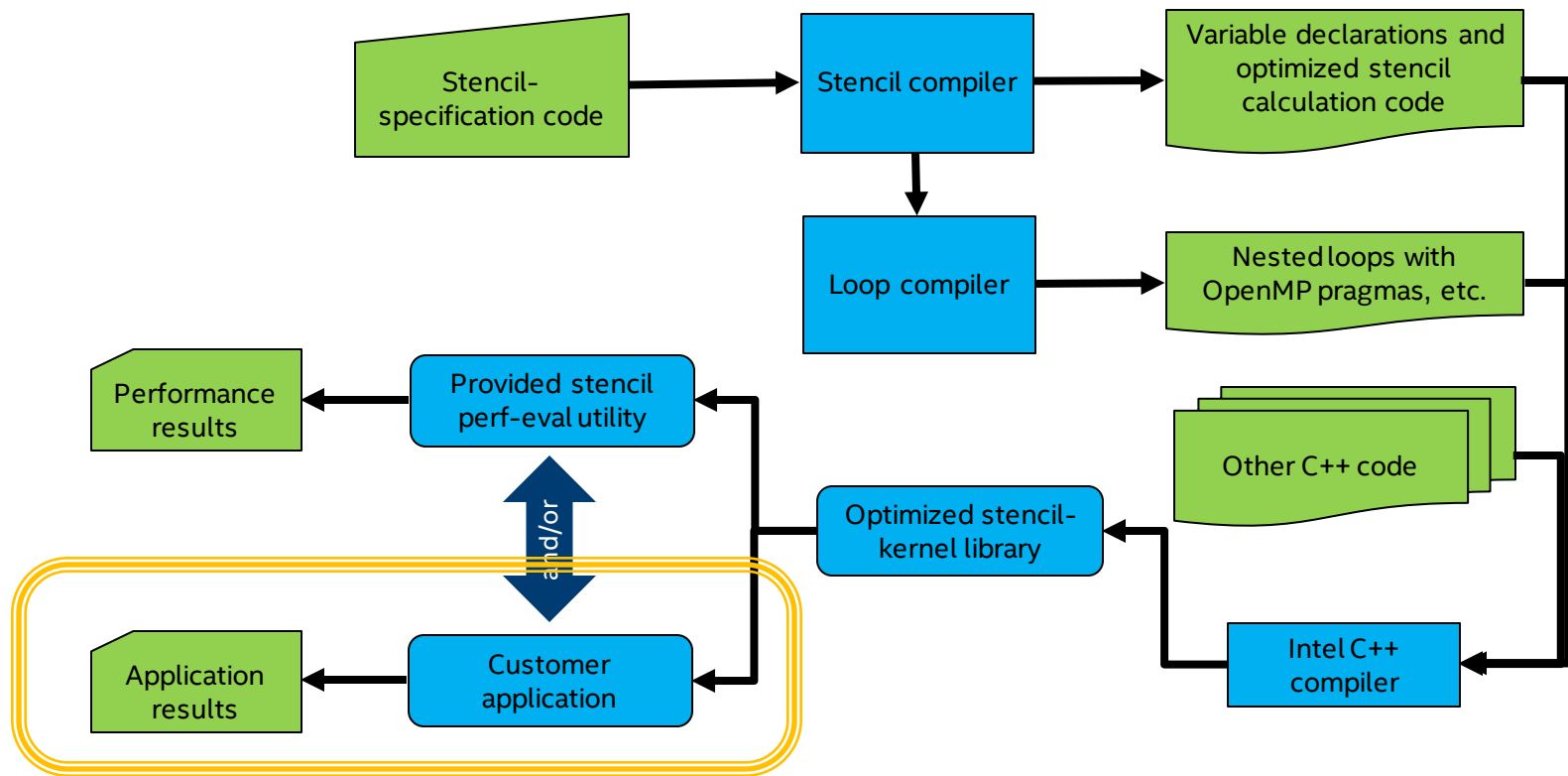
- `bin/yask.sh -mpi_cmd 'mpirun -f $PBS_NODEFILE -ppn 2' -stencil my_stencil -g 1024 -no-pre_auto_tune`
  - This example uses `$PBS_NODEFILE` host file created by IBM Platform LSF\*
    - Use the technique for your scheduler, explicit `-hosts` option, or whatever is used on your cluster
    - *Important:* specify how many processes to run on each node, e.g., with `-ppn`, usually corresponding to the number of NUMA nodes
  - Use the `-nr*` options to control the topology of the MPI ranks
    - Example: `-nr 2` specifies 2 ranks in each spatial dimension
    - Example: `-nrx 2 -nry 4 -nrz 8` specifies 2 ranks in the 'x' dimension, 4 in the 'y', and 8 in the 'z', for 64 total ranks
  - Auto-tuning is turned off in this example, but it can also be used across multiple nodes
  - You can also specify `-use_shm`: it will automatically be applied only between ranks that share a virtual address space

## Implementation optimization

- For typical configurations, YASK overlaps communication with [most] computation using this sequence:
  1. Only the data needed by another rank is calculated on each rank for a given time-step
  2. Asynchronous (non-blocking) MPI receive and send requests are initiated
  3. The remaining data on each rank is calculated while the MPI data is in-flight
  4. The MPI requests are completed before the next time-step is begun
- You can disable this feature via `-no-overlap_comms` to measure the impact

# USING THE APIS

# Making a stencil application



# API Overview

## Purpose

- Define the DSL (Domain-Specific Language) used by the YASK compiler
- Facilitate inclusion of stencil code generated by YASK into real applications

## Target languages

- C++ natively
- Python\* interface generated via SWIG\*

## Design principles

- Consistent interface
- Stable interface with backward-compatibility maintained except in rare cases
- Documented
- Hidden implementation via C++ pure-virtual classes in most cases
- “Factory” pattern: use small number of factory objects to create more objects



# API documentation

From <https://github.com/intel/yask>, follow the “API documentation” link in the Overview section of `README.md`

There are four main tabs

- **Main Page** contains an overview of the YASK workflow and API sets
  - Read this first
- **Modules** lists the API sets
  - This is the best place to start looking for the API you want
  - **YASK Compiler** APIs are used to define stencil solutions, i.e., the “DSL”
    - Objects in the compiler API start with “`yc_`”
  - **YASK Kernel** APIs are used to create your own application from a YASK library
    - Objects in the kernel API start with “`yk_`”
  - **YASK Common Utilities** contains some common utility classes like output streams
    - Objects here start with “`yask_`”
- **Classes** lists all the C++ classes in alphabetical order
- **Files** lists the C++ header files

# Access to the APIs

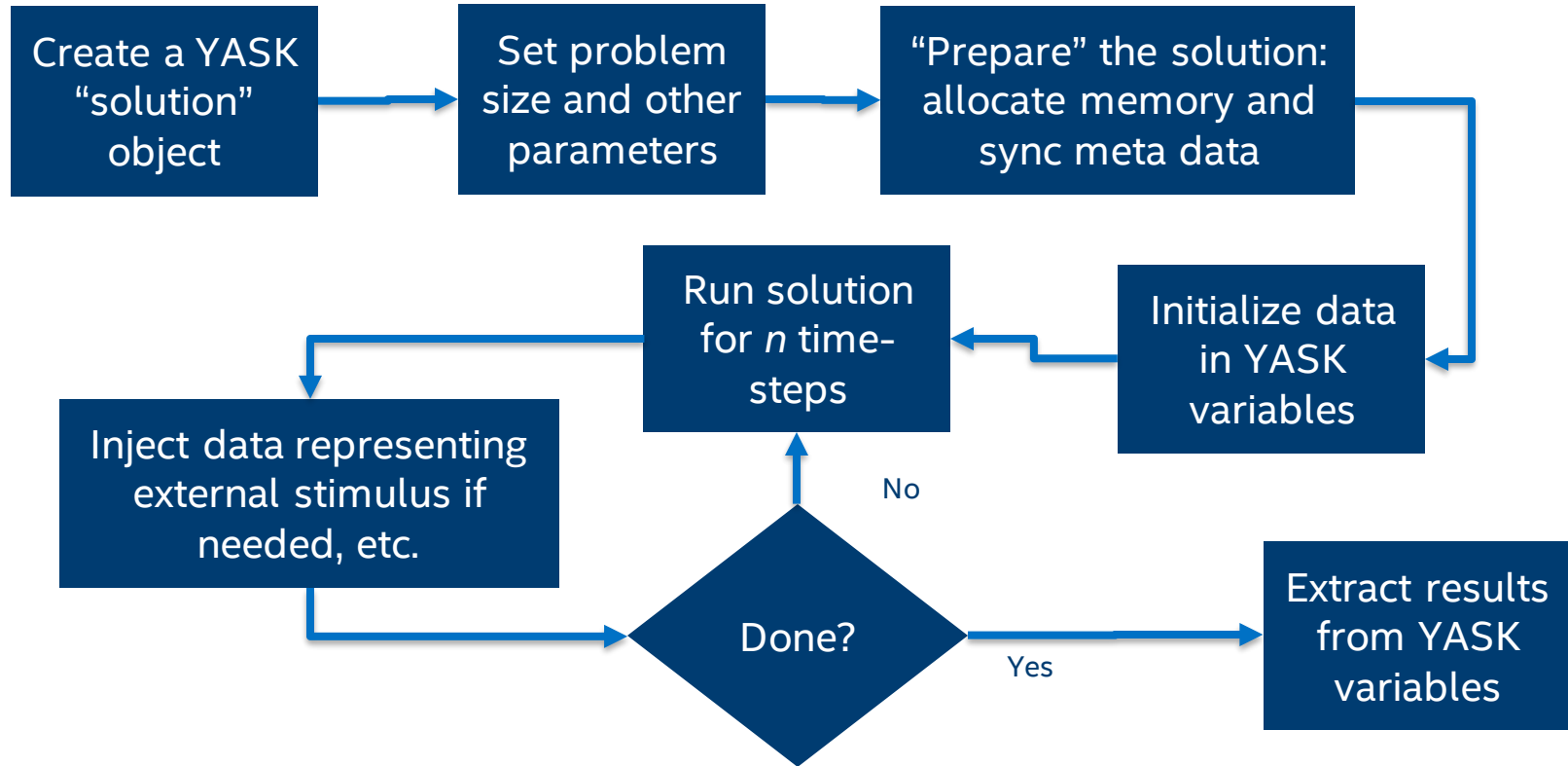
## C++

- The standard method of writing YASK stencils uses the compiler API library
- When you build a kernel as in the previous examples, you have implicitly made a kernel API library
  - A shared object is created in the `lib` directory, one for each stencil and architecture
  - The performance-test utility we've been using is just a wrapper around the library
- The header files are in the `include` directory
  - `#include "yask_kernel_api.hpp"` in your application
  - See `src/kernel/tests/yask_kernel_api_test.cpp` for an example

## Build the APIs for Python\* only if needed

- Download src code as before: `git clone https://github.com/intel/yask`
  - See `README.md` for SWIG\* version requirement
- From `yask` directory, run `make -j api`
- Python APIs are *not* documented separately
- Performance-wise, the stencil calculations will be the same speed as in C++, but data access is slower

# Main YASK kernel process steps



# Creating a YASK kernel via the APIs

## Bootstrap

- Declare an object of type `yk_factory`
- Call `yk_factory::new_env()` to create an “environment” object
  - This can be used to provide an existing MPI communicator if desired
- Call `yk_factory::new_solution()` to create a YASK “solution” object
  - The solution object provides methods to configure the problem size, access YASK variables, and run the stencil calculations

## What's with all the output?

- Calling many APIs results in debug output written to a stream
  - By default, this stream is standard output
  - Handy for development, but you probably don't want this in your final application
- Call `yk_solution::set_debug_output()` to change it to a file or discard it
  - Use a `yask_output_factory` to create new output streams
  - NB: this mechanism was used instead of raw C++ streams for Python usage

# Key terms related to domain sizes

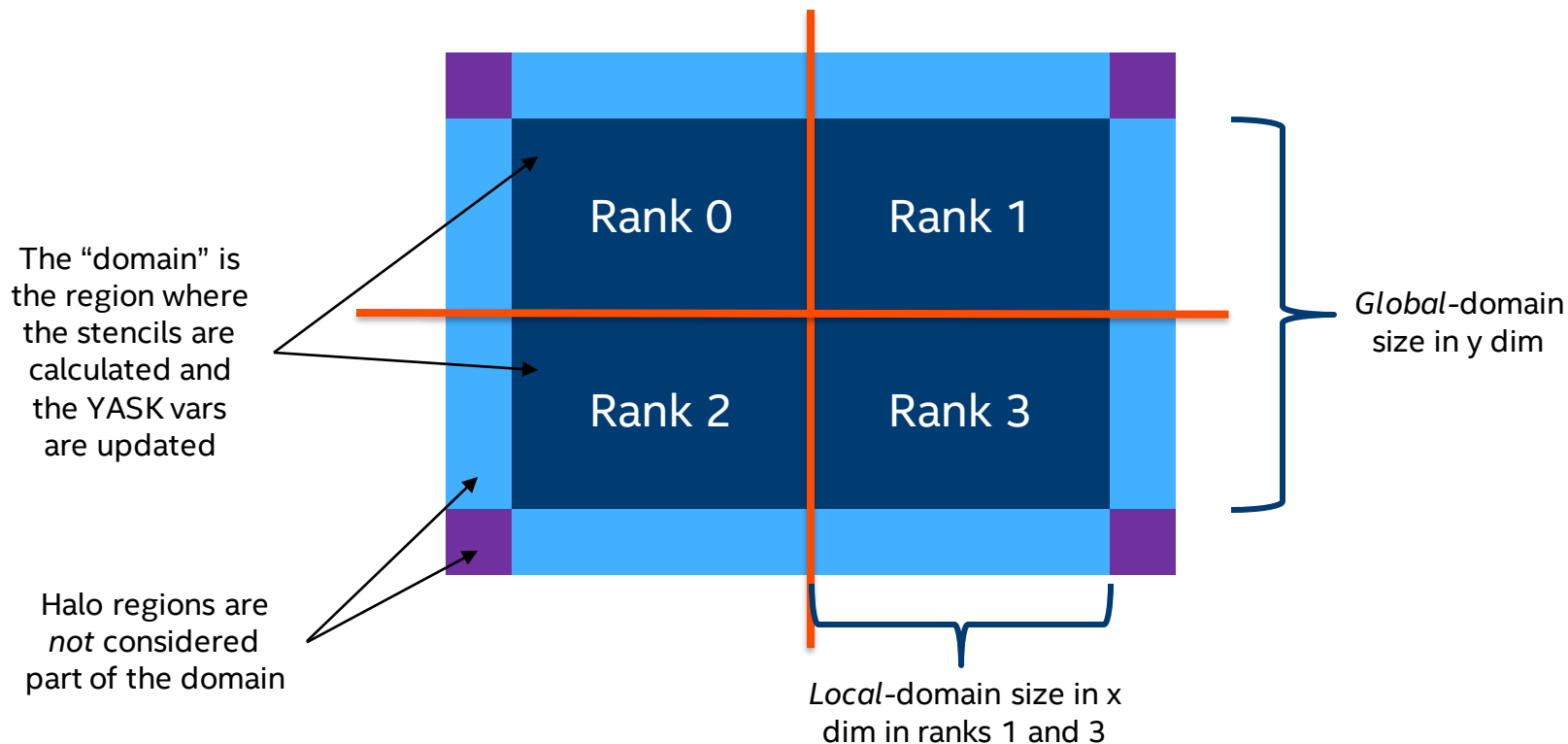
## Overall (global) and rank (local) domain sizes

- As with the test utility, each MPI rank can define the local-domain size or the global-domain size in each dimension
  - The size that is not specified will be calculated automatically
  - All ranks in a given row must have the same rank-domain height, etc.
- Also, each rank can specify the number of ranks in each dimension or let the run-time assign them automatically

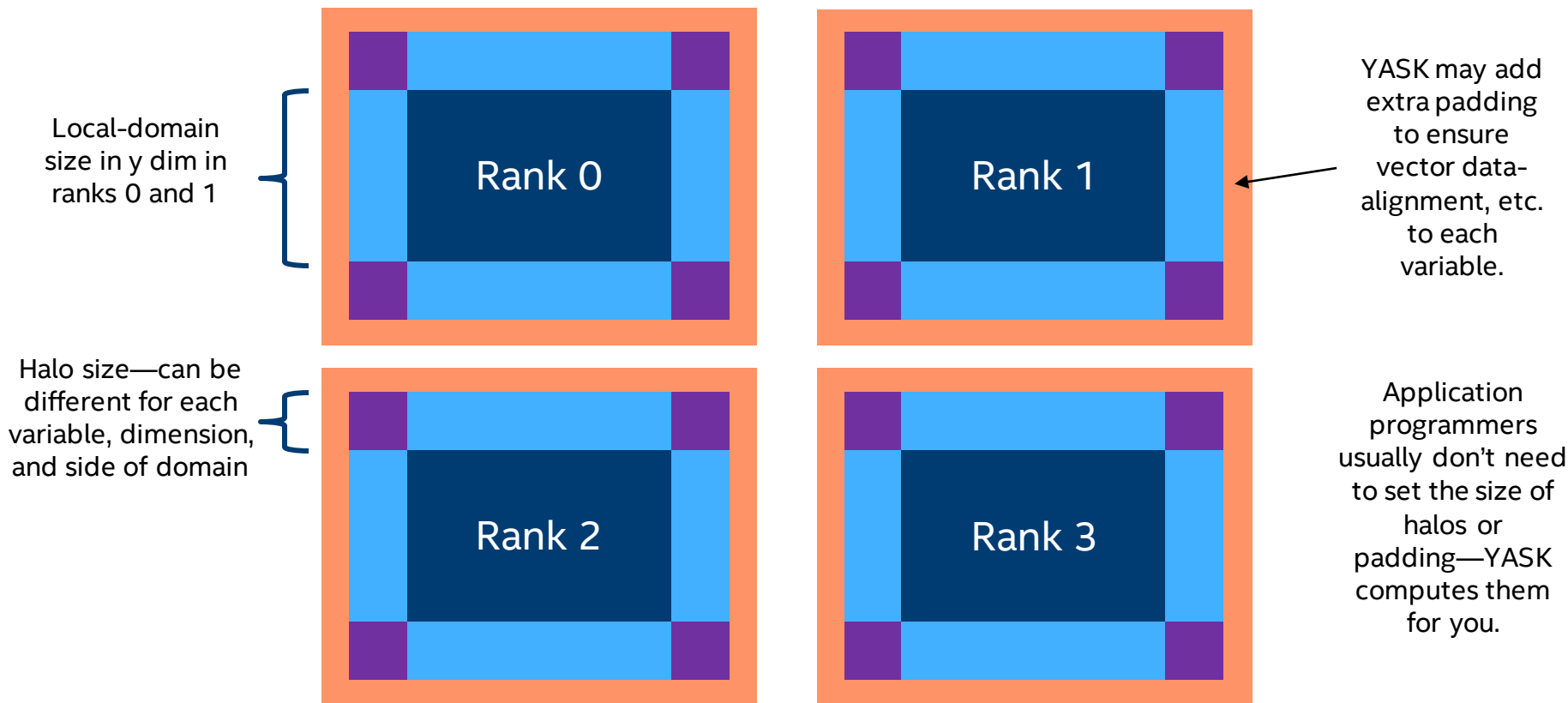
## YASK-variable dimensions and sizes

- When YASK variables are defined, they can have three types of dims
  - **Step dim:** usually “t” for “time”
  - **Domain dim:** usually used for spatial dims like “x” and “y”
    - Generally, can be any domain-size parameter that may be decomposed over ranks
  - **Misc dim:** an index that is known at YASK-compile time like a coefficient index
- The size of the domain will be used when allocating each variable
  - Read the “Detailed Description” section of the `yk_var` API page for an explanation of each size

# Global view of 2D problem size



# Per-rank view of 2D problem sizes



# Prepare the kernel solution

## Set up the problem sizes and ranks

- Call `yk_solution::set_overall_domain_size()` to set the global-domain sizes or `yk_solution::set_rank_domain_size()` to set the local-domain sizes
- Call `yk_solution::set_num_ranks()` to explicitly set the number of ranks if desired
- Call `yk_solution::set_rank_index()` to explicitly set the position of each rank if desired

## Optional: Set other solution parameters

- Call `yk_solution::set_block_size()` to use the best block size you found during the tuning process
- Alternatively, or in addition, you can use the block-size auto-tuner
  - The auto-tuner can be controlled with APIs
  - There are several ways to use the auto-tuner, discussed in the advanced section



# Allocate data and synchronize info across ranks

Call `yk_solution::prepare_solution()`

- Shares MPI positions across ranks or calculates a default position for each rank
  - Since `prepare_solution()` uses MPI calls, it is critical to call it from each rank
- Ensures domain-size consistency across ranks
- Allocates data for each YASK variable based on halo sizes determined by the YASK compiler and the local domain sizes
- Determines MPI buffer requirements and sizes and allocates data for them

# Initialize data

## Access YASK variables

- Get a list via `yk_solution::get_vars()` or find a specific var by name via `yk_solution::get_var()`
- Either one returns pointers to variables of type `yk_var`
  - Used to find all meta data about each var
  - Use `get_*_index()` APIs to determine valid indices
  - Use `idx_t` as a type for indices: `size_t` and `int` are *not* good alternatives.

## Data access

- YASK uses a non-standard tiled data layout (“vector folding” discussed later)
  - Thus, it does not support simple overlay with native row-major or column-major arrays of floating-point numbers
  - However, github issue #147 requests to support this under certain restrictions for easier integration with legacy applications
- There are several APIs for writing one or more FP elements from and to the variables
  - Start with `set_element()`, `set_elements_in_slice()`, and `set_all_elements_same()` in `yk_var`
  - The “slice” versions are threaded for performance and convert to and from row-major layout
    - But you’ll probably want to avoid copying whole vars this way to avoid wasting memory

# YASK-variable indices

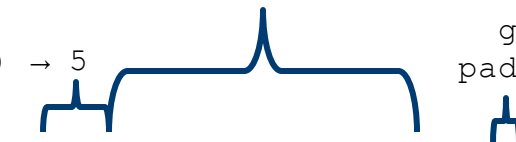
`get_left_halo_size("x") → 5`

`get_rank_domain_size("x") → 100`

`get_right_extra_\npadding_size("x") → 3`

*Important: var  
indices are  
always global!*

*Note: some may  
be negative.*



*Tip: use an API to  
determine each  
index instead of  
adding sizes or  
doing similar math.*

`get_first_rank_\nhalo_index("x") → -5`

`get_first_rank_domain_index("x") → 0`

`get_last_rank_domain_index("x") → 99`

`get_last_rank_halo_index("x") → 104`

`get_last_local_index("x") → 107`

`get_last_rank_\nhalo_index("x") → 204`

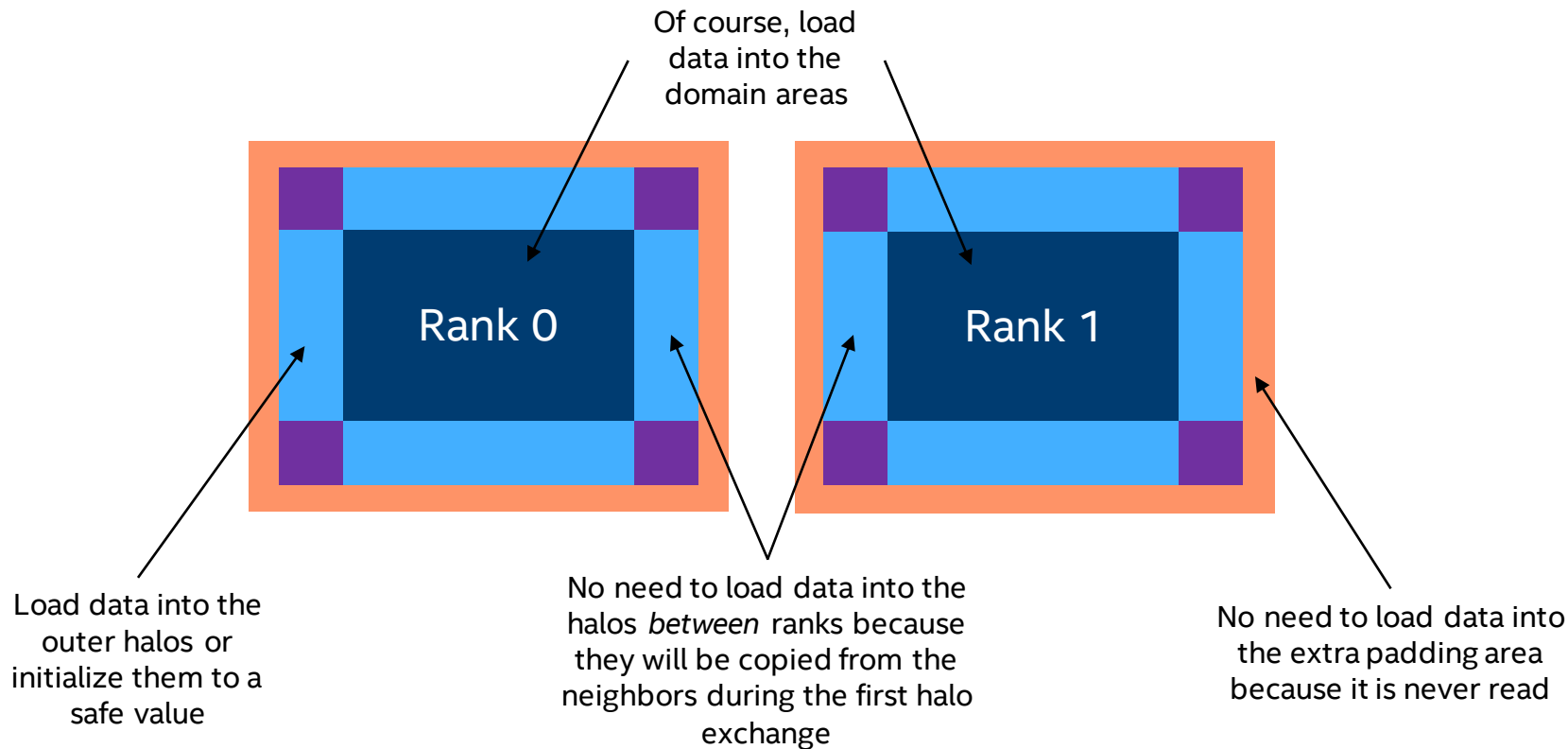
`get_last_rank_domain_index("x") → 199`

`get_first_rank_domain_index("x") → 100`

`get_first_rank_halo_index("x") → 95`

`get_first_local_index("x") → 92`

# Notes on initializing data



# Make the stencil calculations and get results

## Advance the simulation through a series of time-steps

- Call `yk_solution::run_solution()`
  - General form takes first and last time indices, e.g.,
    - `run_solution(0, 9)` runs the first 10 time-steps
    - `run_solution(10, 19)` runs the next 10
- If needed, between time-steps, you can access var data, e.g., for source injection

## Access results

- Use appropriate var methods analogous to the data-writing ones listed earlier
  - Start with `get_element()` and `get_elements_in_slice`
- Call `yk_solution::get_stats()` to collect some performance stats if desired
- Call `yk_solution::end_solution()` when all done to release memory, etc.

# Advanced APIs

## YASK Compiler APIs

- Objects in the compiler API start with “yc\_”
- Very similar functionality to the DSL and the YASK compiler utility
  - Methods to make dims, vars, expressions, spatial and temporal conditions, etc.
  - Also provides operator overloading in C++ and Python to enable readable expressions (except for logical *and*, *or*, and *not* in Python)
- Only needed if you want to write a program to create a YASK stencil
  - However, a planned feature of YASK version 3 will replace the [undocumented] DSL with these APIs (with a conversion utility)

## Misc interesting APIs (not [remotely] exhaustive)

- Call `yk_var::set_numa_preferred()` to set the NUMA node for a given variable—useful for explicit placement in MCDRAM or DDR on Xeon Phi CPUs
- Call `yk_solution::new_var()` to make variables beyond those used in the solution
- Call `yk_solution::apply_command_line_options()` to parse a command-line string—useful for quickly applying options from the test utility in another application
- Call `yk_var::add_to_element()` to atomically update a var element—useful for threaded source injection

# Exceptions

YASK APIs that can trigger errors throw exceptions

- Allows applications to catch and process errors
- Find exception documentation in the **YASK Common Utilities** module tab

## C++

- Throws object of type `yask_exception`
- Call `yask_exception::get_message()` for a human-readable explanation
- See `src/kernel/tests/yask_kernel_api_exception_test.cpp`

## Python

- Throws `RuntimeError` as defined by SWIG\*
- Call `format()` for a human-readable explanation
- See `src/kernel/tests/yask_kernel_api_exception_test.py`

# ADVANCED STENCILS AND TUNING



# Solutions with multiple stencils

## Purpose

- Many modern seismic simulations require multiple variables to be updated, each with a different stencil

## Independent stencils

- *Definition:* for any two stencils, the input of one stencil does *not* depend on the output of the other within the same time-step, and vice-versa
- *Example:*  $x\text{-stress}(t+1)$  depends on  $x\text{-stress}(t)$ , and  $y\text{-stress}(t+1)$  depends on  $y\text{-stress}(t)$ , but  $x\text{-stress}(t+1)$  does not depend on  $y\text{-stress}(t+1)$  or vice-versa
  - See `define_str_TL()` in `src/stencils/SSGElasticStencil.cpp`
  - Thanks to contributor Albert Farres from the Barcelona Supercomputing Center!
- For independent stencils, YASK applies the stencils simultaneously in one pass over the domain

## Dependent stencils

- *Definition:* for any two stencils, the output of one stencil is required for the input of the other stencil in the same time-step (the opposite cannot also be true)
- *Example:*  $x\text{-velocity}(t+1)$  depends on  $x\text{-stress}(t)$ , and  $x\text{-stress}(t+1)$  depends on  $x\text{-velocity}(t+1)$  –see same DSL file
- For dependent stencils, YASK cannot apply the stencils simultaneously and makes subsequent passes over the domain
- YASK automatically determines the dependencies between equations in the DSL and schedules their stencils

# Absorbing boundaries

## Used in wave-propagation simulation

- Simulation domain covers only a finite block of earth, water, etc.
- Simulated waves will erroneously reflect from the arbitrary boundaries
- Artificial absorbing boundary conditions (ABC) or boundary layers are used to reduce these reflections
- There are many techniques and papers covering this field of study

## Two high-level approaches to use in YASK

- Modify the stencil(s) to include simple attenuation factors
  - *Example:* Cerjan sponge layers
  - See `Iso3dfdSpongeStencil` class in `src/stencils/Iso3dfdStencil.cpp`
    - Try modifying your test stencil to use sponge layers and check the impact on performance
  - This stencil illustrates the use of combining 3D and 2D variables to save memory
  - *Tip:* this stencil and its parent class also show the recommended practice of creating functions to return expressions, allowing more flexible composition
- Use *different* stencils in the boundary layers by creating *sub-domains*
  - *Examples:* Higdon BCs, perfectly-matched-layers (PML), and the convolutional form (CPML)



# Sub-domain (spatial) conditions

A sub-domain is a subset of the domain in which a stencil is applied

- Defining stencils in boundaries is a common use-case for sub-domains

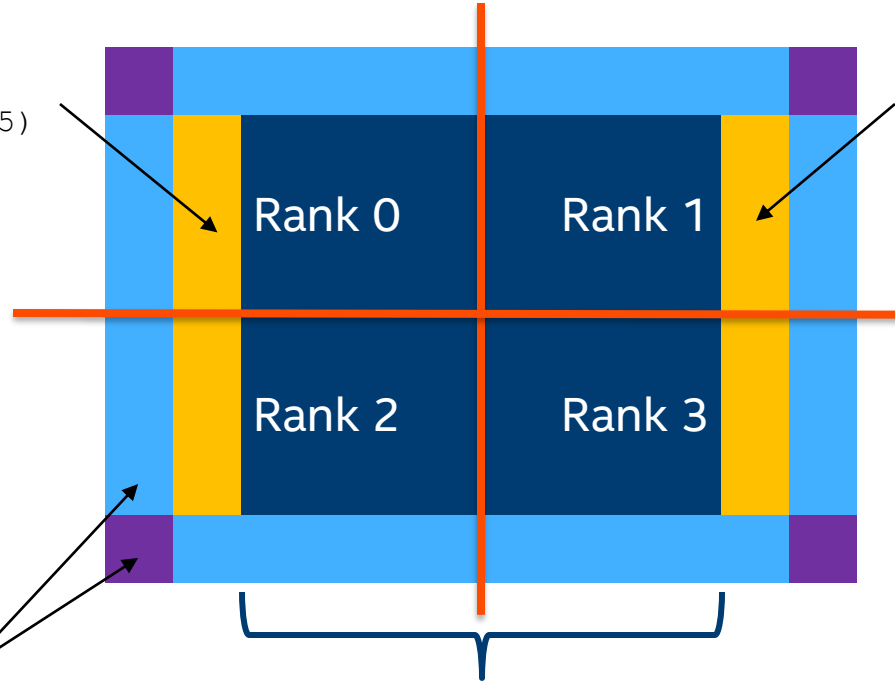
A sub-domain is defined in the DSL by a Boolean expression on the domain indices

- The DSL includes terms for the left-most and right-most indices in the domain
- *Example:* `(x < first_index(x) + 5)` defines a sub-domain on the left side of the domain, 5 elements wide
  - `(x > last_index(x) - 5)` would be the same on the right side
  - `(x >= first_index(x) + 5) && (x <= last_index(x) - 5)` would be used for the interior sub-domain between the left and right boundary sub-domains
- These conditions are written after the equation using the `IF_DOMAIN` operator in the DSL
  - Search for `TestBoundaryStencil` in `src/stencils/TestStencils.cpp` for simple synthetic examples
  - See `src/stencils/FSGElasticStencil.cpp` for a more complex real-world example

# Sub-domains on multiple ranks

Sub-domain described by  
 $(x < \text{first\_index}(x) + 5)$

Sub-domain described by  
 $(x > \text{last\_index}(x) - 5)$



Halo regions are  
*not* considered  
part of any domain  
or sub-domain

Sub-domain described by  
 $(x \geq \text{first\_index}(x) + 5) \ \&\& \ (x \leq \text{last\_index}(x) - 5)$

Sub-domains  
always apply  
across the overall  
problem domain.

(This is one reason  
that domain  
indices are always  
global.)

# Step (temporal) conditions

A step-condition expresses which time-steps are valid for a stencil

- Some stencil applications require special processing on regular intervals
- Wave-field sub-sampling can be implemented with step conditions

Like sub-domain conditions, step conditions restrict stencil application

- Sub-domain expressions can use only the domain indices
- Step-condition expressions can use the step index (usually time) or values in other YASK variables (but not via domain indices)
- *Example:*  $(t \% 8 == 0)$  is true every 8<sup>th</sup> time-step
- These conditions are written after the equation using the `IF_STEP` operator in the DSL
  - Look for `TestStepCondStencil` in `src/stencils/SimpleTestStencils.cpp` for a simple example

# Vector folding (multi-dimensional vectorization)

## Concept

- Store small 2D or 3D block of data into each SIMD vector
- *Pros*: reduces memory loads and memory streams compared to traditional 1D in-line vectorization
- *Cons*: requires non-traditional tiled data layout and additional shift and/or permute operations preceding SIMD arithmetic operations

## Results

- Significant speedup shown on large-radius stencils, especially on Intel® Xeon Phi™ processors
- Works well paired with tiling and other performance techniques

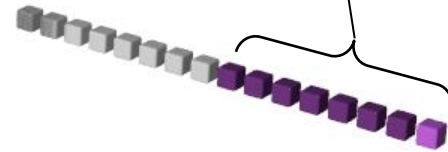
## Implementation

- The YASK compiler automatically generates the proper shift and permute instructions
- The YASK kernel code automatically generates code to store the tiled data and look up elements by index when needed
- Intel® AVX-512 instruction set is needed for efficient permutes, so only enabled when available
- See the paper in the upcoming reading list showing up to  $1.5\times$  speedup from vector-folding

# Traditional 1D Vectorization



Inner 3D loop iterates  
in x direction, i.e., *same  
dimension* as  
vectorization



8 new vectors must  
be read for  $k \pm r$   
points  
(4 for  $k+r$  and 4 for  
 $k-r$  for  $r=1..4$ )

8 new vectors must  
be read for  $j \pm r$   
points

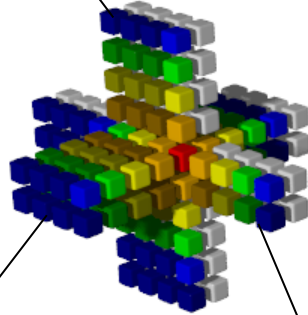
Only 1 new vector  
must be read for  $i \pm r$   
points due to  
overlap along x axis

Total BW cost for  
traditional "in-line" vectors  
= **17** new vector inputs for  
each vector of output  
(some loads will come from  
cache with blocking)

# 2D Vector-Folding

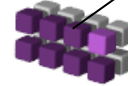


4 new 4x2x1 vectors  
must be read for  $j \pm r$   
points



Only 1 new  
vector must be  
read for  $k \pm r$   
points

2 new vectors  
must be read for  
 $i \pm r$  points



Inner 3D loop  
iterates in z  
direction, i.e.,  
*perpendicular*  
to 2D vector

Total BW cost for 4x2x1 vector  
with z-axis loop= 7 new vector  
inputs for each vector of output  
**(2.4x lower than in-line)**



# Vector-Folding Memory Layout and Code Gen

## 2D “4x2” vector folding

Logical indices in 2D with 8-element SIMD in x and y dimensions

5,1	5,2	5,3	5,4	5,5	5,6	5,7	5,8	5,9	...
4,1	4,2	4,3	4,4	4,5	4,6	4,7	4,8	4,9	...
3,1	3,2	3,3	3,4	3,5	3,6	3,7	3,8	3,9	...
2,1	2,2	2,3	2,4	2,5	2,6	2,7	2,8	2,9	...
1,1	1,2	1,3	1,4	1,5	1,6	1,7	1,8	1,9	...

y

x

Access to elements in custom memory layout encapsulated behind C++ & Python APIs

Load and permute instructions generated automatically by YASK stencil compiler

Layout in memory (1D)

1,1	1,2	1,3	1,4	2,1	2,2	2,3	2,4	1,5	1,6	1,7	1,8	2,5	2,6	2,7	2,8	1,9	...
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

- 2D vector folding layout (8×1)
- Two aligned vectors are colored
- Unaligned read shown with bold borders done by loading aligned vectors and then shuffling the requisite elements via an AVX-512 *permute* instruction

# Vector-folding customization

## Specify the vectorization length in each dimension

- Use the `fold='x=n,y=n,z=n'` argument to the `make` command-line
  - The values are passed to the YASK stencil compiler and used to generate code
  - The fold settings are also included in non-generated code during compilation
  - *Example:* `make fold='x=1,y=2,z=8'` generates code using a  $1 \times 2 \times 8$  fold
  - Try different fold settings on the test stencil and check the impact on performance
    - *Important:* Be sure to run `make clean` before re-compiling when changing compile-time options like vectorization
- The product of the fold lengths should equal the number of SIMD vector elements in the target architecture and FP precision (single or double)
  - *Example:* single-precision FP using 512-bit SIMD contains 16 elements per vector
  - If the fold is not the right size, the compiler will adjust the requested fold using a heuristic algorithm
  - The vector length in any dimension not specified defaults to one (1)
  - The default fold varies depending on dimensionality, architecture, and FP precision
    - Is  $4 \times 4 \times 1$  for a 3D problem using SP FP on Xeon Scalable CPUs, for example



# More compile-time settings

## Floating-point precision

- The default FP size is 4 bytes, which is the norm in seismic modeling
- Build with double-precision by adding `real_bytes=8` to the `make` command

## Prefetching

- Software prefetch instructions can be added by the YASK compiler
  - Use `make ... pfd_l1=n` to set the L1 prefetch distance to  $n$  iterations ahead
  - Use `make ... pfd_l2=n` to set the L2 prefetch distance to  $n$  iterations ahead
  - Set  $n$  to zero (0) to disable generation of any prefetch instructions for a level
- The default prefetch depends on the architecture

## Time allocation

- By default, the YASK compiler tries to determine the minimum number of time-steps that need to be saved in each variable, but you may want to override it
- Override this with `make ... time_alloc=n` (to affect all vars at compile-time)
- Override at run-time with the `yk_var::set_alloc_size()` API

# More compile-time settings

## Order of domain dimensions

- The order of domain dimensions determines
  - The inner-most or “unit-stride” dimension in the memory layout
  - The nesting order of loops for the stencil-calculation code
  - Default vector-folding and default rank layout
- By default, the order of domain dimensions is determined from the variable declarations
  - For example, `new_var("A", {t, x, y, z})` implies domain dimensions are in 'x, y, z' order
  - If vars have different domain-dimension index orders, dimensions are sorted in the order they are seen by the compiler
- You can explicitly set the order of domain dimensions
  - For example, `make ... domain_dims='z, y, x'` creates domain dimensions in 'z, y, x' order
  - This setting overrides the order of the dimensions specified in the variable declarations and affects memory layout, looping order, vector-folding, and rank layout
  - Try different orders on the test stencil and check the performance impact (remember to `make clean`)



## Misc settings

- Any preprocessor macro can be set by `make ... EXTRA_MACROS='name=value ...'`
- Type `make help` for examples of settings the C++ optimization level and more

# More grouping terms: clusters, bundles, and stages

## Clusters

- The YASK compiler can also unroll the innermost loops a specified number of times, effectively calculating a *cluster* of SIMD vectors in each iteration
- Control via the `cluster='x=n,y=n,z=n'` argument to the make command-line similar to the `fold` argument for vector-folding
  - Try different cluster settings on the test stencil and check the performance impact (remember to `make clean`)
- The default clustering is to not unroll (cluster size of one in each dimension)
- Clustering increases register pressure, making it useful only for small stencils



## Bundles

- A *bundle* is the term used for independent equations that are grouped into one function by the YASK compiler
- A bundle cannot contain equations from different sub-domains
- The compiler eliminates common sub-expressions between stencils in a bundle

## Stages

- A *stage* is the term used for a group of equation bundles that are not inter-dependent
- This can occur when there are equations in different sub-domains that are not dependent on one another
- Two or more stages may be evaluated within one cache block that crosses sub-domain boundaries
- Multiple stages are scheduled in the correct order based on dependencies (e.g., in a staggered-grid problem)

# Temporal wave-front rank tiling

## Goal

- Increase temporal locality in caches at a package level
- Especially useful for large unified cache (such as MCDRAM cache on Xeon Phi)

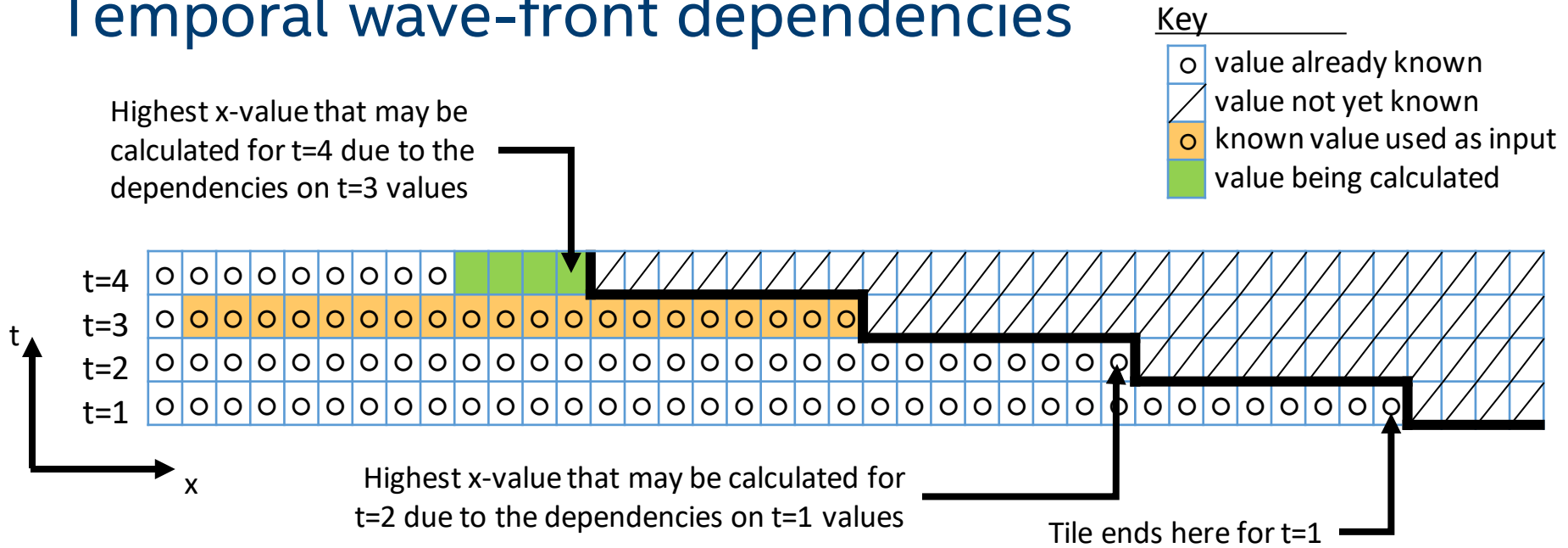
## Technique

- Evaluate  $n$  time-steps in a subset of local rank domain
  - We refer to this subset as a *region* in YASK
- Evaluate each region *sequentially* until entire rank domain is evaluated for  $n$  time-steps

## Increased software complexity

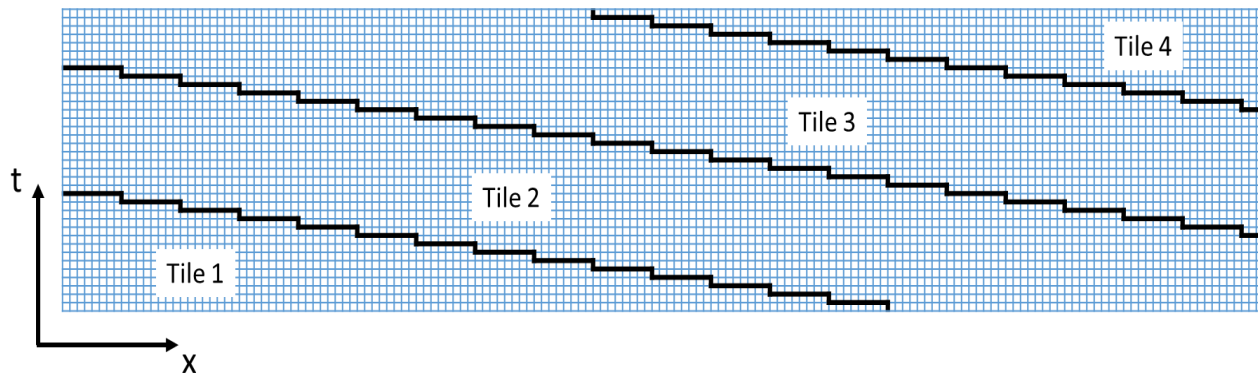
- Must respect mathematical dependencies between time-steps
- Halo exchanges between ranks become more complex
- Any inter-time-step data processing (like source injection) becomes more complex

# Temporal wave-front dependencies



- In the first temporal wave-front tile shown here, the number of values that can be calculated is reduced for each time-step (and/or each stage within a time-step)
- The amount of shift is called the *wave-front* or *temporal angle* and is based on the radius of the largest stencil (shift of 8 shown)

# Covering a temporal range via multiple tiles



- Wave-front tiles are computed *sequentially*, but multiple values within one time-step of a given tile may still be evaluated *concurrently*
- After last tile is complete, YASK variables contain the same data as they would have without temporal tiling
- This concept is directly extended to 2D or 3D stencils by shifting in each spatial dimension
- To handle MPI halo exchanges, some redundant calculations are made between ranks



# Using temporal wave-front rank tiling in YASK

## Another level of tiling hierarchy

- Earlier, we explained how each rank domain is divided into *blocks*, set via `-b`
- Ranks may also be divided into *regions*, where each region is a wave-front tile
- Control the spatial size of a region with `-r`, `-rx`, `-ry`, etc.
- Control the temporal size of a region with `-rt` (assuming your time dim is “t”)
- By default, the spatial size of a region is the size of the rank, and the temporal size is one (1)

## Selecting values

- When using a very large unified cache (e.g., a 16GiB MCDRAM cache on Xeon Phi), set the spatial size to fit within this cache
  - See the paper in the upcoming reading list showing  $\sim 2\times$  speedup on two stencils
- For processors that have much smaller third-level caches (e.g., a Xeon Scalable processor), values are much more critical, and additional performance is more difficult to obtain

# Temporal block tiling

## Goal

- Increase temporal locality in caches at a core level, e.g., level-2 caches

## Technique





- Evaluate  $n$  time-steps in each block
  - Recall that a rank is composed of regions, and a region is composed of blocks
- Evaluate blocks *concurrently* until entire region is evaluated for  $n$  time-steps

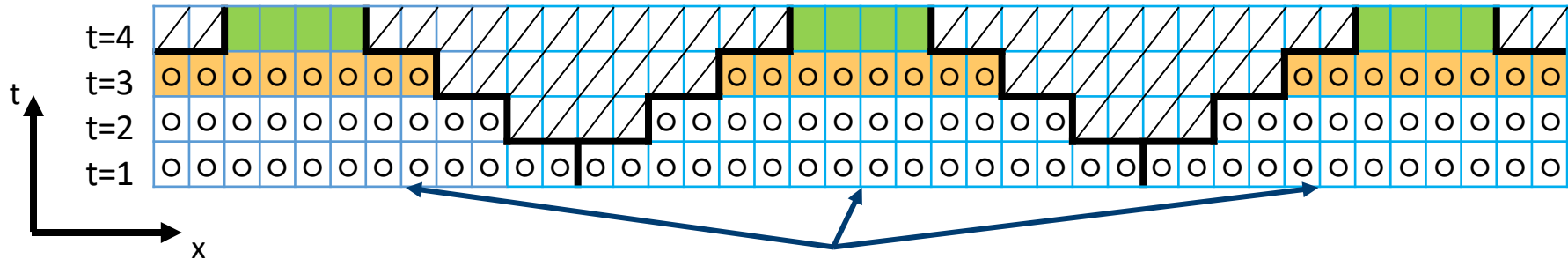
## Increased software complexity (similar to wave-front tiling)

- Must respect mathematical dependencies between time-steps
  - Due to concurrency, shapes of blocks are more complex than with sequential wave-front tiling
- Any inter-time-step data processing (like source injection) becomes more complex

# Temporal block dependencies

Key

	value already known
	value not yet known
	known value used as input
	value being calculated



These three “upward” triangle blocks may be evaluated concurrently.  
When they are complete, the “downward” triangle blocks may be evaluated.

- In one spatial dimension (as shown), this is called triangle or trapezoid tiling (half of diamond tiling)
- Extension into multiple spatial dimensions requires a more complex series of multi-dimensional shapes (polytopes) to tessellate the space (not shown)
- The amount of shift or *temporal angle* is based on the radius of the largest stencil (shift of 2 shown)

# Using temporal block tiling in YASK

## Not another level of tiling hierarchy (yet)

- As before, control the spatial size of a block with `-b`, `-bx`, `-by`, etc.
- Control the temporal size of a block with `-bt` (assuming your time dim is “t”)
- By default, the temporal size of a block is one (1)
- Temporal block tiling occurs *within* wave-front rank tiling, so the *temporal* size of a region defaults to the temporal size of a block
  - But the default *spatial* size of a region is still the spatial size of a rank domain

## Selecting values

- When selecting spatial sizes, consider level-2 cache size, number of YASK variables accessed, the FP-element size, and the number of cores

# Temporal wave-front mini-block tiling

## Goal

- *Observation*: block sizes are used as a unit-of-work for OpenMP\* threads as well as level-2 cache targeting
  - These objectives can conflict when small cache sizes relative to the number of variables calls for small blocks, which can lead to small OpenMP tasks
- Also, it may be beneficial in some cases to combine the concepts of triangle tiling and wave-front tiling across different dimensions
- Goal is to separate the thread and cache block concepts and provide more tiling flexibility

## Technique





- Evaluate  $n$  time-steps in a subset of each block
  - We refer to this subset as a *mini-block* in YASK
- Evaluate each mini-block *sequentially* until entire block is evaluated

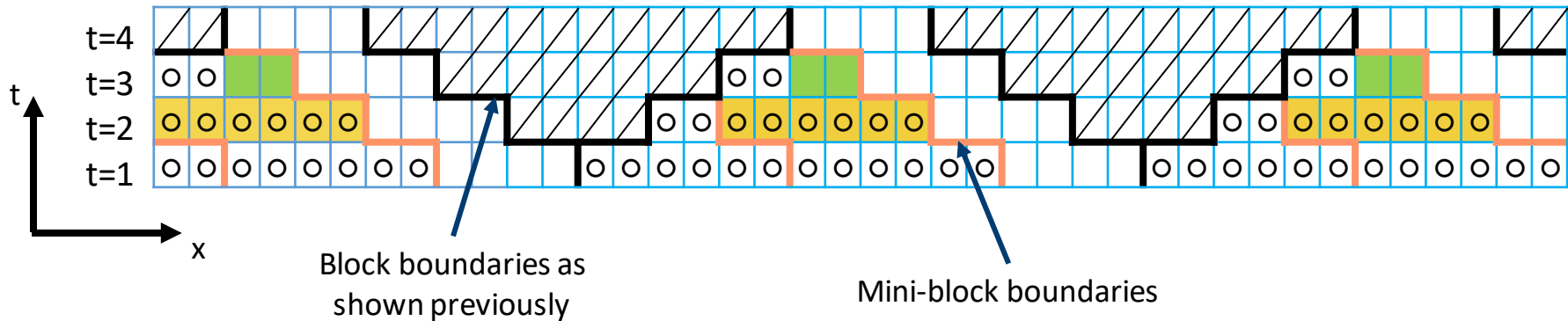
## Increased software complexity (similar to previous tiling)

- Must respect mathematical dependencies between time-steps
  - Similar to sequential wave-front tiling, but within [hyper] triangles of temporal blocks

# Temporal mini-block dependencies

Key

	value already known
	value not yet known
	known value used as input
	value being calculated



- Blocks are evaluated concurrently as before using OpenMP threads
- Mini-blocks are evaluated sequentially within each block
- Mini-blocks use wave-front tiling, similar to wave-front rank tiling, but inside blocks instead of ranks

# Using mini-block tiling in YASK

## Yet another level of tiling hierarchy

- Blocks may be divided into *mini-blocks*, where each mini-block is a wave-front tile
- Control the spatial size of a mini-block with `-mb`, `-mbx`, `-mby`, etc.
- The temporal size of a mini-block is always the same as a block, so it is set implicitly via `-bt`
- By default, the spatial size of a mini-block is the size of a block

## Selecting values

- The size of a mini-block should normally correspond to the size of a level-2 cache, considering the number of variables accessed and the FP-element size
- The block sizes can now be larger, considering the number of cores with thread balancing
- By sizing mini-blocks as wide as blocks in one or two dimensions and/or sizing blocks as wide as regions in some dimensions, interesting special-case scenarios may be created

# Nested OpenMP threads

## Goal

- *Observation:* using hyper-threads (SMT) across blocks effectively reduces the usable size of the level-2 cache available to each thread
- This is particularly impactful on Xeon Phi processors with not only 4 hyper-threads per core but also 2 cores sharing a level-2 cache
- *Goal:* allow multiple threads to use shared caches constructively rather than destructively

## Technique

- Evaluate subsets of each mini-block by threads that share caches
  - We refer to this subset as a *sub-block* in YASK
- Evaluate sub-blocks *concurrently* until a single time-step in a mini-block is evaluated (thus, no temporal sub-block tiling)
- Since blocking already uses OpenMP threads, sub-blocking is implemented with a nested level of OpenMP threading



# Using sub-block tiling in YASK

## The lowest level of tiling hierarchy

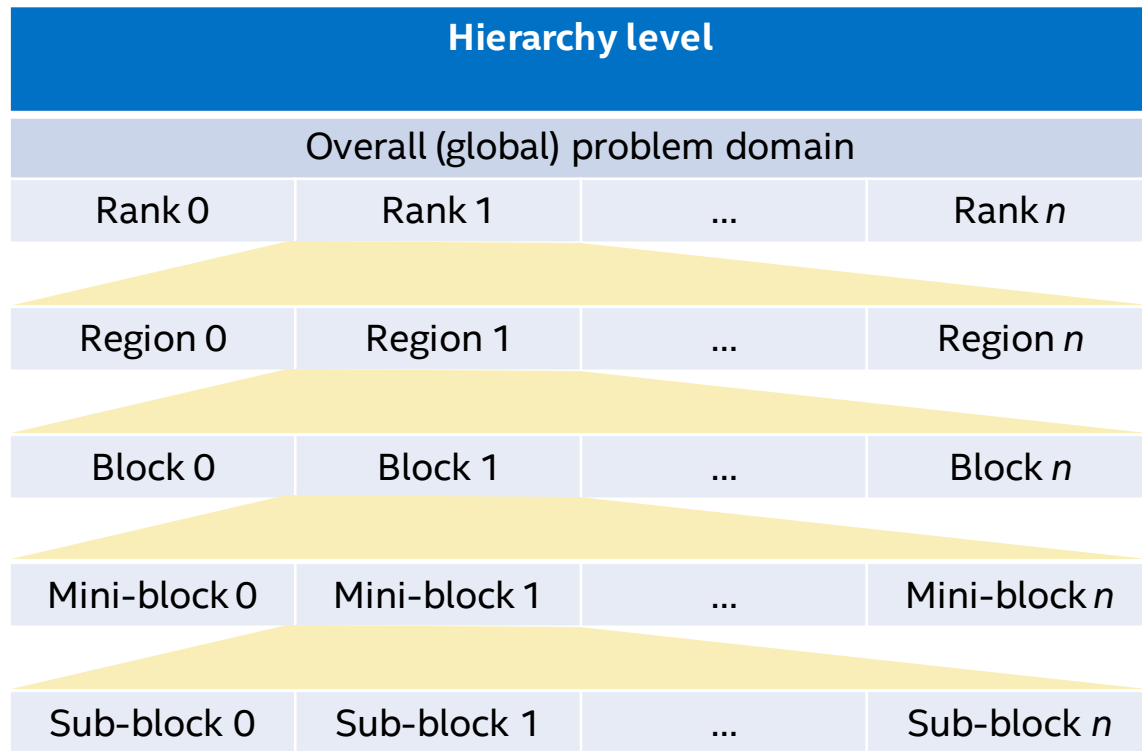
- Mini-blocks may be divided into *sub-blocks*
- Control the spatial size of a sub-block with `-sb`, `-sbx`, `-sby`, etc. (the temporal size is always one)

## Selecting values

- By default, the spatial size of a sub-block depends on the number of threads are used per block
  - If there is one thread per block (no nested OpenMP), the default size of a sub-block is the size of a mini-block
    - In this case, it is recommended to use only one thread per level-2 cache
  - If there are >1 threads per block (nested OpenMP active), the default size of a sub-block is a narrow slab the width of one vector (in the first dim) and the size of a mini-block in the other dims
    - This setting is intended to increase reuse between hyper-threads while keeping threading overhead as low as possible
- Control the number of threads per blocks with `-block_threads`
  - If you do not want to use hyper-threads, also use `-thread_divisor`, which divides the default number of threads
    - On a Xeon CPU, use `-thread_divisor 2 -block_threads 1` to run one thread from each core per block
  - For Xeon Phi, choosing 8 threads per block often works well and is the default
  - For Xeon processors, using both threads is better for some stencils, and using only one is better for others
- Try adjusting threads-per-block and/or sub-block sizes on the test stencil and check performance



# Review of the hierarchy



Target structure	Concurrency
Cluster	
NUMA node	MPI
Pkg cache	Sequential
OMP thread	OMP team
L2 cache	Sequential
OMP thread	Nested OMP

# Combining tile settings

Try the following on a two-socket Intel® Xeon® Scalable CPU

- ```
bin/yask.sh -stencil my_stencil -l 1024 \  
-mbx 32 -mby 32 -mbz 128 \  
-bx 220 -by 256 -bz 256 -bt 12 -trial_steps 36 \  
-block_threads 1 -thread_divisor 2 -no-pre_auto_tune
```
- Mini-block settings selected for a 1MiB L2 cache
  - $32 \times 32 \times 128 \times 4B \times 2 = 1\text{MiB}$  (128 in unit stride; “ $\times 2$ ” for two time-steps in memory)
- Spatial block settings selected for a multiple of number of cores per socket
  - $\text{Ceil}(1024 \div 220) = 5$  in x dimension,  $1024 \div 256 = 4$  in y and z
  - $5 \times 4 \times 4 = 80 = 4 \times 20$  cores (modify block size if your CPU has a different core count)
  - A small multiple (like 4) often works better than the exact count because several blocks per core gives more opportunity for dynamic load balancing
- Temporal block setting (`-bt 12`) chosen experimentally
- Region size not used because there is no very large cache (like MCDRAM on Xeon Phi CPUs)
- Block-threads and thread-divisor chosen to use only one hyper-thread, so sub-blocks are automatically scaled up to mini-block size



# More about the block-size automatic tuner

## Pre-calculation mode

- Runs *before* the desired stencil calculations are done
- Does *not* ensure that calculations are done in the proper order
  - Thus, important to [re]initialize data *after* running in this mode for use in deployed applications
- Intended for benchmarking to tune block-size before running actual time-steps
- Use `[-no]-pre_auto_tune` option to control in provided test utility (default is on)
- Call API `run_auto_tuner_now()` to activate outside in your application

## Intra-calculation mode

- Runs *during* desired stencil calculations
- Maintains proper calculations
- Intended for deployment, esp. on multiple platforms or when all final platforms are not known *a priori*
- Use `[-no]-auto_tune` option to control in provided test utility (default is off)
- Call `reset_auto_tuner()` to turn off or on explicitly
- Call API `is_auto_tuner_enabled()` to determine whether it is [still] running

# Multi-parameter auto-tuner

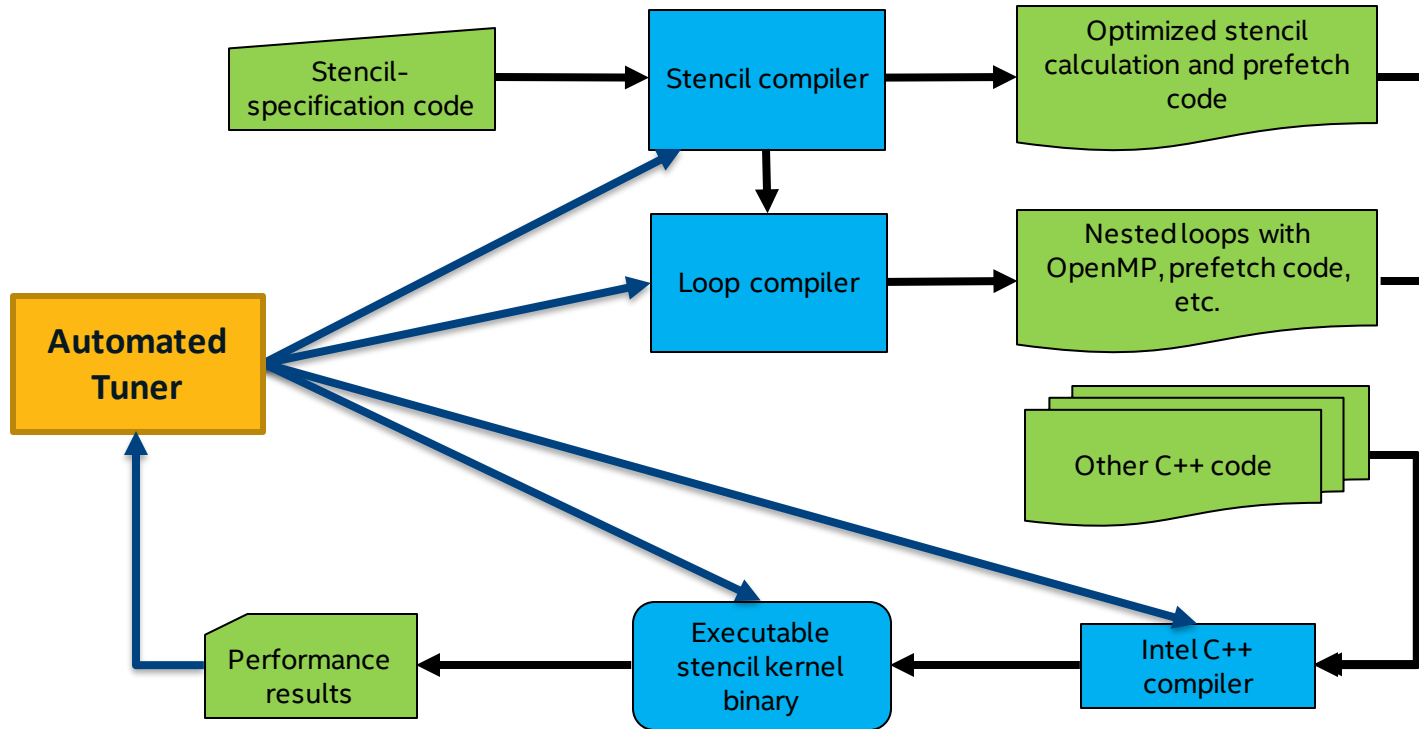
## Goal

- Use a genetic algorithm to explore the design space of many parameters
- Can target compile-time and/or run-time parameters

## Usage

- Run `utils/bin/yask_tuner.pl` to see options
- By default, explores almost every compile-time and run-time parameter
  - Practically, you probably want to limit some of them
  - To fix the problem size, use `-g*` or `-l*` options, e.g., `-g=1024 -gz=128`
  - To search only run-time parameters, use `-noBuild` option
- Tuner will stop when there has been no improvement over 5 generations
- *Tip:* alternate between manual and automated tuning for final optimization
- Currently only works for 3D (spatial) problems with dims named `t`, `x`, `y`, and `z`

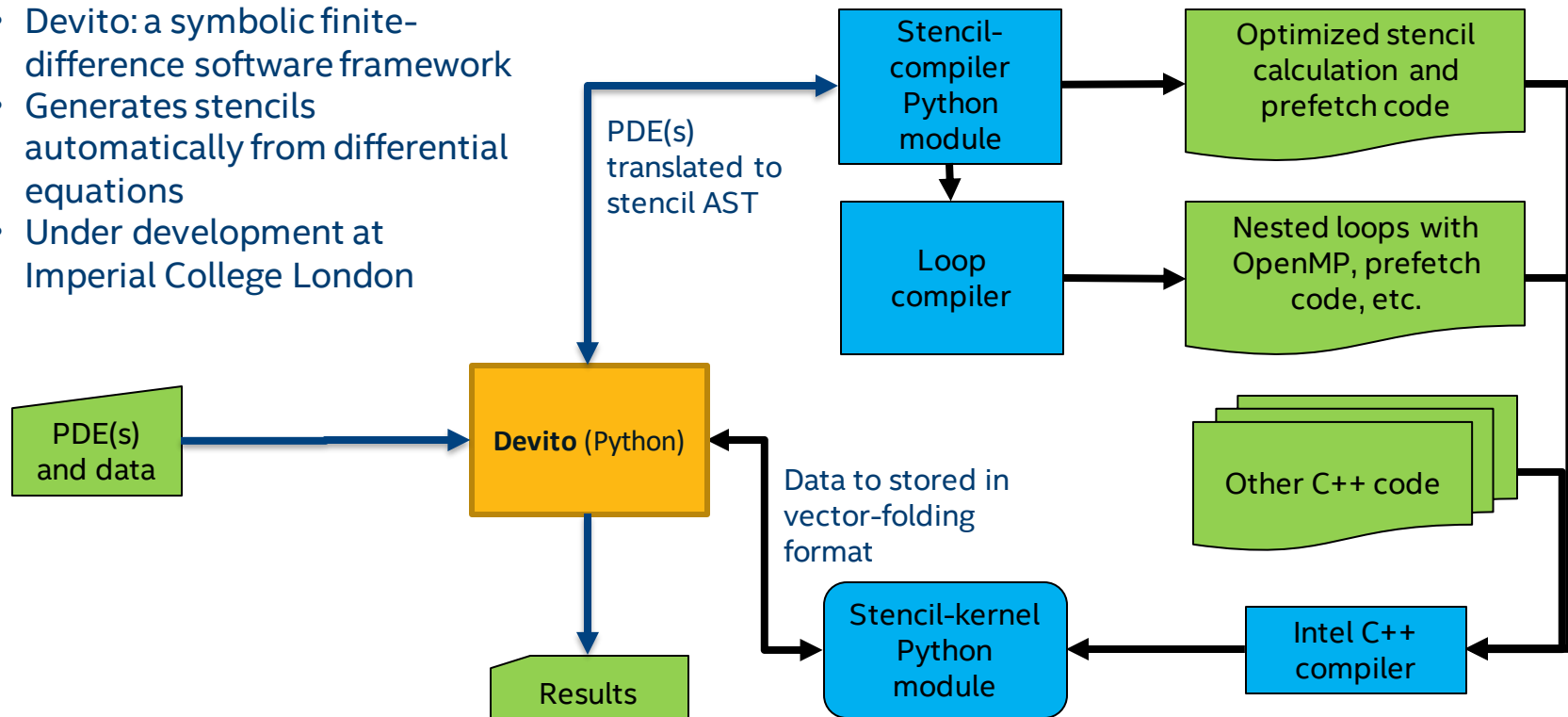
# YASK workflow driven by auto-tuner



# WRAP-UP

# Current work item: Devito\* integration

- Devito: a symbolic finite-difference software framework
- Generates stencils automatically from differential equations
- Under development at Imperial College London





# Read more about YASK features and applications

- “Vector Folding: improving stencil performance via multi-dimensional SIMD-vector representation.” C Yount. *High Performance Computing and Communications (HPCC)*, 2015
- “YASK—Yet Another Stencil Kernel: A Framework for HPC Stencil Code-Generation and Tuning.” C Yount, J Tobin, A Breuer, A Duran. *Domain-Specific Languages and High-Level Frameworks for High Performance*, 2016
- “Effective use of large high-bandwidth memory caches in HPC stencil computation via temporal wave-front tiling.” C Yount, A Duran. *Performance Modeling, Benchmarking and Simulation of High Performance*, 2016
- “Accelerating seismic simulations using the Intel Xeon Phi knights landing processor.” J Tobin, A Breuer, A Heinecke, C Yount, Y Cui. *International Supercomputing Conference*, 139-157, 2017
- “Multi-level spatial and temporal tiling for efficient HPC stencil computation on many-core processors with large shared caches.” C Yount, A Duran, J Tobin. *Future Generation Computer Systems*, 2017
- “Performance Optimization of Fully Anisotropic Elastic Wave Propagation on 2nd Generation Intel® Xeon Phi (TM) Processors.” A Farres, C Rosas, M Hanzich, A Duran, C Yount. *2018 IEEE International Parallel and Distributed Processing Symposium*

# Call to action

Work through this tutorial!

Code your own stencil

- Use it in a real application
- Please feel free to contact the developers with questions
- Contribute your stencil for others to use: create a fork on github\* and submit a pull request
- Please tell the developers about your experience, maybe even co-publish results

Contribute to the project

- See the “issues” database on github\* for the to-do list
- Talk to the developers about something you'd like to work on
  - Probably some good academic projects in there!



<https://github.com/intel/yask>

