USING OPENMP* EFFECTIVELY ON THETA

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Overview

This talk is not intended to teach basic OpenMP*, but rather focus on new capabilities and an emphasis on application to the Intel® Xeon Phi x200 processors

- Brief introduction to OpenMP*
- OpenMP* tasking
- Using OpenMP* SIMD instructions
- OpenMP* affinity
  - Pure OpenMP*
  - Hybrid MPI+OpenMP*
What is OpenMP*?

OpenMP* stands for Open Multi-Processing. It provides:

- Standardized directive-based multi-language high-level parallelism.
- Portable and Scalable model for shared-memory parallel programmers.
- Language support for C/C++/FORTRAN.
- Provides APIs and environment variables to control the execution of parallel regions.
- Latest specs and examples are available at http://www.openmp.org/specifications/.
- Supported by LLVM, Visual Studio Compiler, Intel Compiler, GNU GCC and others.
OpenMP* Programming Model

Real world applications are a mix of serial and inherently parallel regions.

OpenMP* provides **Fork-Join Parallelism** as a means to exploit inherent parallelism in an application within a **shared memory architecture**.

- Master thread executes in serial mode until a parallel construct is encountered.
- After the parallel region ends team threads synchronize and terminate, but master continues.
OpenMP* Constructs

**Basic Components**

- **Parallel** - thread creation
  - parallel

- **Work Sharing** - work distribution among threads
  - do, for, sections, single

- **Data Sharing** - variable treatment in parallel regions and serial/parallel transitions
  - shared, private

- **Synchronization** - thread execution coordination
  - critical, atomic, barrier

- **Advanced Functionality**
  - Tasking, SIMD, Affinity, Devices (offload)

**Runtime functions and control**

```c
!$OMP PARALLEL
!$OMP DO
  do i = 1, N
    a(i) = b(i) + c(i);
  end do
!$OMP END PARALLEL
```

```c
#define pragma omp parallel
{
  #pragma omp for
    for(int i = 0; i < N; i++)
    {
      a[i] = b[i] + c[i];
    }
}
```
OPENMP* TASKING CONCEPTS
Some Background

Prior to standard version 3.0, OpenMP* was focused exclusively on Data Parallelism, distributing work over threads executing the same code.

This work sharing model presented some limitations

- A need for a known loop count
- Very limited ability for dynamic scheduling
- Inconvenient for naturally task-parallel problems (dependencies, nesting)

Task parallelism constructs were introduced to complement the already existing set that supported data parallelism

Task parallelism is particularly useful in irregular computing
What is an OpenMP* Task?

From the standard document: "specific instance of executable code and its data environment"

- Explicit task: work generated by the `task` construct
- Implicit task: threads of a parallel region

In this section of the talk I will be only discussing explicit tasks.

By default tasks are deferrable, so the generating thread may execute it immediately or queue it.

```
#pragma omp task
myfunc();

#pragma omp task
for(int i = 0; i < N; i++) { ... }
```
Task Synchronization

Sibling tasks

The `taskwait` construct can be used to wait for deferred task completion at some point in the code.

```c
#pragma omp task
myfunc();

#pragma omp task
for(int i = 0; i < N; i++){ ... }

#pragma omp taskwait
```

Nested tasks

Synchronizing siblings and their descendants requires a `taskgroup`.

```c
#pragma omp taskgroup
{
  #pragma omp task
  myfunc();

  #pragma omp task
  {
    for(int i = 0; i < N; i++){
      #pragma omp task
      nestedfunc();
    }
  }
}
Task Decomposition

Often an application can be decomposed into tasks which can execute simultaneously.

Following the Directed Acyclic Graph (DAG) shown on the right:

- Tasks Alice, Bob and Cy can start executing simultaneously.
- Boss can only be executed after Alice and Bob complete execution.
- BigBoss can only be executed after Cy and Boss complete execution.

```c
a = alice();
b = bob();
s = boss(a,b);
c = cy();
printf( "%f\n", bigboss(s,c) );
```
Parallel Execution of Tasks

```
#pragma omp parallel
{
    #pragma omp single
    {
        #pragma omp task
        a = alice();
        #pragma omp task
        b = bob();
        #pragma omp task
        c = cy();
    }
}

s = boss(a, b);
printf ( "%f\n", bigboss(s,c) );
```

Start parallel region, forking N threads

Use a single thread to generate the tasks

Each independent code section may be defined as a task

Once generated each task may be performed by any available thread in the parallel region.
Task Generation and Execution

1. Threads are spawned from master

2. Work queue is generated by single thread

3. Tasks in queue are assigned to threads and executed

4. Process continues until queue is empty (or sync point)
Better Scheduling with Depend Clause

```c
#pragma omp parallel
{
    #pragma omp single
    {
        #pragma omp taskgroup
        {
            #pragma omp task depend(out:a)
            a = alice();
            #pragma omp task depend(out:b)
            b = bob();
            #pragma omp task depend(out:c)
            c = cy();
            #pragma omp task depend(in:a,b) depend(out:s)
            s = boss(a, b);
            #pragma omp task depend(in:s,c)
            printf ( "%f\n", bigboss(s,c) );
        }
    }
}
```

The `depend` clause allows to specify dependencies among tasks:

```
depend(<in|out|inout>:<variables>)
```

Based on dependencies `boss()` can start executing once `alice()` and `bob()` are done.

Using the `depend` clause it is possible to execute `cy` and `boss` simultaneously.

The `taskgroup` directive creates an implicit synchronization point, but it is optional in this example.
Parallelize Recursions

```c
void merge_sort_openmp(int a[], int tmp[], int first, int last) {
    if (first < last) {
        int middle = (first + last + 1) / 2;
        if (last - first < 5000) {
            merge_sort(a, tmp, first, middle - 1);
            merge_sort(a, tmp, middle, last);
        } else {
            #pragma omp task
            merge_sort_openmp(a, tmp, first, middle - 1);
            #pragma omp task
            merge_sort_openmp(a, tmp, middle, last);
            #pragma omp taskwait
        }
        merge(a, tmp, first, middle, last);
    }
}
```

Merge sort is common recursive algorithm

- Its recursive nature used to pose a challenge in terms of expressing the parallelism.

- OpenMP* Tasking helps express the parallelism in recursive calls as shown below.

- Explicit taskwait synchronization forces a wait until all sibling tasks complete execution.

- Merging phase can't start until all the tasks spawned above have completed.
Other Interesting Tasking Tidbits

Tasks can be stopped and continued (at scheduling points). By default tasks are tied so they can only be continued by the same thread that started them (hot cache). This behavior can be overridden with the \texttt{untied} clause

\begin{verbatim}
#pragma omp task untied
\end{verbatim}

You may introduce your own scheduling points using the \texttt{taskyield} directive

\begin{verbatim}
#pragma omp taskyield
\end{verbatim}

The \texttt{taskloop} directive may be used to schedule loop iterations as independent tasks with a single generator (Intel® Compiler version 18+)

\begin{verbatim}
#pragma omp taskloop [\texttt{[grainsize|numtask]} [\texttt{untied}] [\texttt{nogroups}] [\texttt{priority}]]
\end{verbatim}

\begin{verbatim}
for( i = 0; i < N; i++){ ...}
\end{verbatim}
Tasking Summary

Introduced to enable task-parallelism in shared memory architectures
Mostly used in irregular computing
Tasks are typically generated by a single thread
Dependencies can be specified to improve scheduling efficiency
Untied task generators can ensure progress
First-private is default data-sharing attribute
Shared variables remain shared
VECTORIZATION WITH OPENMP* SIMD
OpenMP* SIMD

A few critical capabilities were introduced in OpenMP* with the standard specification 4.0 (not an exhaustive list!)

- **Target Constructs**: Accelerator support
- **Task Groups/Dependencies**: Runtime task dependencies & synchronization
  - **SIMD**: fine grained data level parallelism
  - **Affinity**: Pinning workers to cores/HW threads

Refinements to SIMD were also introduced in specification 4.5

SIMD is of critical importance on Theta due to the 512bit width of the KNL processors

Affinity is also of critical importance with 256 threads per socket
The OpenMP* SIMD directive

```c
#pragma omp simd [clause]
for(int i = 0; i < N; i++)
{
    ...
}

!$omp simd [clause]
do i = 1, N
    ...
end do
!$omp end simd
```

WARNING: The compiler ignores dependencies when using the `simd` directive.

Multiple clauses available

- safelen(length)
- simdlen(length)
- linear(list[:linear-step])
- aligned(list[:alignment])
- private(list)
- lastprivate(list)
- reduction(op: list)
- collapse(n)
Details and Limitations

Do/For-loop has to be in “canonical loop form” (see OpenMP 4.0 API:2.6)

safelen(n) : The compiler can assume a vectorization for a vector of length of n to be safe

simdlen(n) : Preferred vector length

linear(var:step) : For every iteration of the original scalar loop var is incremented by step. Therefore it will be incremented by step * vector_length for the vectorized loop.

aligned(var:base) : Assert that var is aligned to base bytes; (default is architecture specific alignment)
SIMD Example

This example instructs the compiler to ignore data dependencies, asserts array alignment, and indirectly mitigates the control flow dependence.

OpenMP* SIMD must be enabled at compilation time with either `-qopenmp` or `-qopenmp-simd` flags

```c
#pragma omp simd safelen(32) aligned(a:64, b:64)
for(int i = 0; i < N; i++)
{
    a[i] = (a[i] > 1.0) ? a[i]*b[i] : a[i+off]*b[i];
}
```
SIMD Enabled Functions

Applying the `declare simd` construct to a function creates one or more versions of the function that can process multiple arguments using SIMD instructions from a single invocation from a SIMD loop.

### Multiple clause options

- `simdlen(length)`
- `linear(list[:linear-step])`
- `aligned(list[:alignment])`
- `uniform(list)`
- `inbranch`
- `notinbranch`

```plaintext
#pragma omp declare simd [clause]
double work(double *a, double *b, int off);

function work(a, b, off)
!$omp declare simd [clause]
implicit none
integer :: off
double precision :: a(*), b(*)
... end function
```
SIMD Enabled Function Example

```c
#pragma omp declare simd simdlen(16) notinbranch uniform(a, b, off)
double work( double *a, double *b, int i, int off )
{
    return (a[i] > 1.0) ? a[i]*b[i] : a[i + off]*b[i];
}

void vec2( double *a, double *b, int off, int len )
{
    #pragma omp simd safelen(64) aligned(a:64, b:64)
    for( int i = 0; i < len; i++ )
    {
        a[i] = work( a, b, i, off );
    }
}
```
SIMD + Threads

By combining syntax we can both parallelize and vectorize a loop:

```c
#pragma omp parallel for simd [clause]

 !$omp parallel do simd [clause]
```

Where the clauses are those valid for either a do/for directive or a simd directive.

Loop will distributed among threads using chunks that are multiples of the vector size
**SIMD on Theta**

Ensure `safelen` and `simdlen` are compatible with AVX512

- Minimum of 8 for double precision
- Minimum of 16 for single precision

Use the `processor` clause

- Extension introduced in Intel Compiler version 17
- Use `processor(mic_avx512)` to target KNL

Remember not all vector operations are equally effective

- Alignment (array + accesses)
- Strided access (gather/scatter operations reduce performance)
- Masking (enables conditional execution, but at a cost)
AFFINITY CONTROL WITH OPENMP*
Thread Affinity in OpenMP*

OpenMP* 4.0 introduces the concept of Places and Policies

- Set of threads running on one or more processors
- Places can be defined by the user
- Predefined places available: threads, cores, sockets
- Predefined policies: spread, close, master

And means to control these settings

- Environment variables OMP_PLACES and OMP_PROC_BIND
- Clause proc_bind for parallel regions

Optimal settings depend on application and workload
Pure OpenMP* on Theta

For pure OpenMP* based codes the most effective way to set affinity is to disable affinity in aprun and then use OpenMP settings to bind threads.

Disabling affinity with aprun is simple:

```sh
$ aprun -n 1 -N 1 -cc none ./exe
```

Now threads can be pinned to specific hardware resources using the OMP_PLACES and OMP_PROC_BIND environmental variables.

Rich set of options with lots of flexibility and configuration granularity, but a few simple setups cover the vast majority of production cases.
Pinning Step 1: OMP_PLACES

Two levels of granularity. You may specify a policy:

\[
\text{OMP\_PLACES} = \text{<policy>}
\]

Where policy may be

- sockets: threads are allowed to float on sockets (multiple cores)
- cores: threads are allowed to float on cores (multiple logical processors)
- threads: threads are bound to specific logical processors

Or you may specify a list:

\[
\text{OMP\_PLACES} = \{\text{lower\_bound: length: stride}\}:\text{repeat:increment}
\]
Pinning Step 2: OMP_PROC_BIND

To specify how threads are bound within the defined places use:

```plaintext
OMP_PROC_BIND=<policy>
```

Where `policy` must be chosen from:

- **close**: threads paced consecutively, as near to the master place as possible
- **spread**: threads spread equally on hardware to use most resources
- **master**: threads placed on master place to enhance locality

Note that specifying `master` could lead to heavy oversubscription of hardware resources, depending on the defined places.

It is possible to print out your pinning specification as interpreted by OpenMP* using `OMP_DISPLAY_ENV=true`
Some examples

```plaintext
OMP_NUM_THREADS=4; OMP_PLACES="{0:4:2}"  
Bound to [0] [2] [4] [6]

OMP_NUM_THREADS=4; OMP_PLACES=threads; OMP_PROC_BIND=close  
Bound to [0] [64] [128] [192]

OMP_NUM_THREADS=4; OMP_PLACES=threads; OMP_PROC_BIND=spread  
Bound to [0] [16] [32] [48]

OMP_NUM_THREADS=4; OMP_PLACES=cores; OMP_PROC_BIND=spread  
Bound to [0,64,128,192] [16, 80, 144, 208] [32, 96, 160, 224] [48, 112, 176, 240]
```
Hybrid MPI + OpenMP*

When using hybrid applications aprun must be configured to create pinning ranges for each MPI task, and then OpenMP variables may be set to control thread pinning within each rank processor range. Example: 4 MPI tasks, 16, 8 nodes

```bash
export OMP_NUM_THREADS=16
export OMP_PLACES=cores;
export OMP_PROC_BIND=spread
aprun -n 32 -N 4 -cc depth -d 64 -j 4 ./exe
```

<table>
<thead>
<tr>
<th>Task 0</th>
<th>Thread 1</th>
<th>...</th>
<th>Thread 15</th>
</tr>
</thead>
<tbody>
<tr>
<td>[0, 64, 128, 192]</td>
<td>[1, 65, 129, 193]</td>
<td>...</td>
<td>[15, 79, 143, 207]</td>
</tr>
<tr>
<td>Task 1</td>
<td>[16, 80, 144, 208]</td>
<td>[17, 81, 145, 209]</td>
<td>...</td>
</tr>
<tr>
<td>Task 2</td>
<td>[32, 96, 160, 224]</td>
<td>[33, 97, 161, 225]</td>
<td>...</td>
</tr>
<tr>
<td>Task 3</td>
<td>[48, 112, 176, 240]</td>
<td>[49, 113, 177, 241]</td>
<td>...</td>
</tr>
</tbody>
</table>

*Other names and brands may be claimed as the property of others.
NUMA considerations

Locality

- Local memory accesses reduce latency.
- Use Linux first touch policy to your advantage by initializing data in an OpenMP* loop in the same way that it will be used later.

MCDRAM

- Provides higher bandwidth
- Important to make a conscious choice if running on flat mode

If running on flat mode you may use `numactl` to attach to the numa node 1 (MCDRAM):

```
aprun -n <ntot> -N <ppn> numactl --membind=1 ./exe
aprun -n <ntot> -N <ppn> numactl --preferred=1 ./exe
```
Recommended settings for Theta

The following setup is recommended for jobs using up to 4 threads per core

```
OMP_PLACES=cores

OMP_PROC_BIND=spread

aprun -n <totalTasks> -N <tasksPerNode> -cc depth -d 256/<tasksPerNode> -j 4
```

If using multiple threads per core you may want to test the effect of changing the default wait policy to passive:

```
OMP_WAIT_POLICY=passive
```