Master Course – High-Performance Computing

Parallel Programming with OpenMP Part I

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Outline

- Introduction into OpenMP
- Programming and Execution Model
 - Parallel regions: team of threads
 - Syntax
 - Data environment (part 1)
 - Environment variables
 - Runtime library routines
 - Exercise 1: Parallel region / library calls / privat & shared variables
- Worksharing directives
 - Which thread executes which statement or operation?
 - Synchronization constructs, e.g., critical regions
 - Nesting and Binding
 - Exercise 2: Pi
- Data environment and combined constructs
 - Private and shared variables, Reduction clause
 - Combined parallel worksharing directives
 - Exercise 3: Pi with reduction clause and combined constructs
 - Exercise 4: Heat
- Summary of OpenMP API
- OpenMP Pitfalls & Optimization Problems

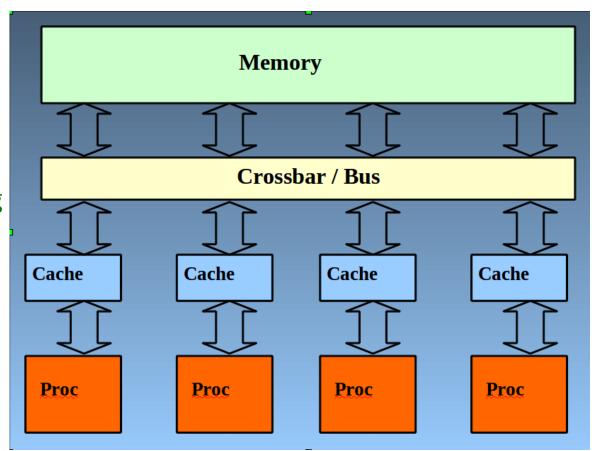
Multiprocessor System with Shared Memory

OpenMP

is a

parallel programming model

for



Shared-memory multiprocessors

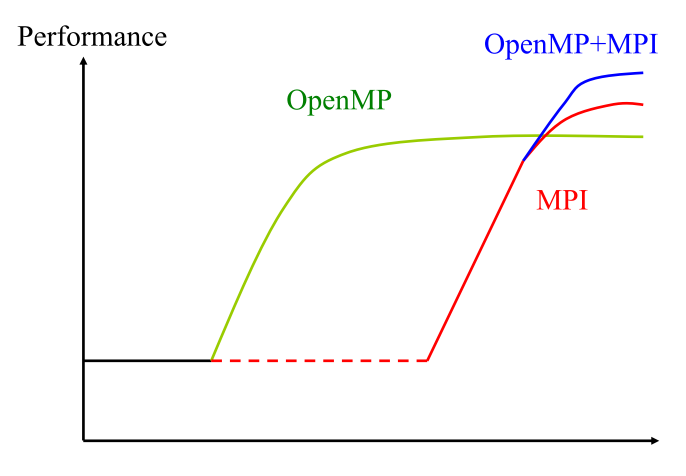
OpenMP Overview: What is OpenMP?

- OpenMP is a standard programming model for shared memory parallel programming
- Portable across all shared-memory architectures
- It allows incremental parallelization
- Compiler based extensions to existing programming language
 - mainly by directives
 - a few library routines
- Fortran and C/C++ binding
- OpenMP is a standard

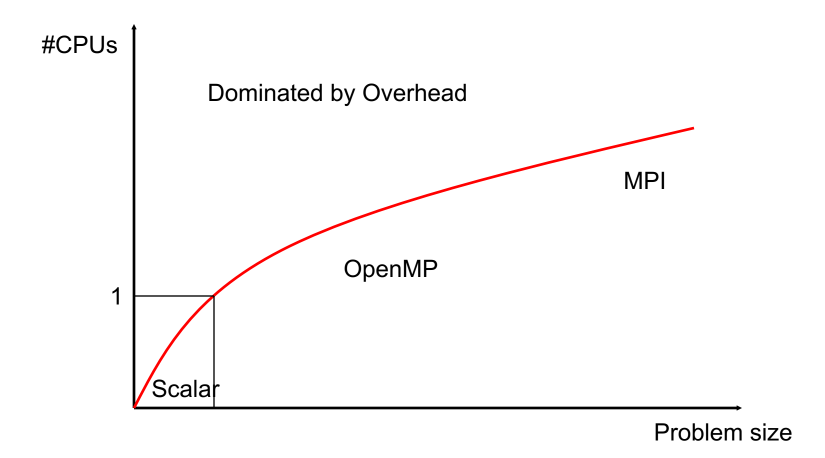
Parallel Computing: Effective Standards for Portable programming

- Thread Libraries
 - Win32 API
 - POSIX threads
- Compiler Directives
 - OpenMP portable shared memory parallelism
- Our focus

- OpenACC, OpenCL
- Global Address Space Languages
 - Unified Parallel C
 - Titanium
 - -X10
 - Chapel
- Message Passing Libraries
 - MPI



Time/Effort



Multithreading versus Multi-Processing

- Multiple Processes (Heavyweight Process model)
 - traditional UNIX process model
 - interprocess communication techniques supported by OS: shared memory, sockets, file IO, memory map
 - Higher overhead associated with process creation and destruction
- Multiple Threads (Lightweight Process model, LWP)
 - thread concept: independent flow of control within one process with its own context: stack, register set
 - process data and opened files are shared
 - lower overhead of thread creation and destruction
 - shared address space
 - Auto-Parallelization, OpenMP, Explicit Multithreading using P-Threads
- **Hybrid Models** (e.g. MPI + OpenMP)

• 1997: OpenMP Version 1.0 for Fortran



- de facto standard for shared memory programming
- now available for all SMP systems
- replaces proprietary parallelization directives and in many cases the explicit programming of [p]threads
- 1998: OpenMP V1.0 for C and C++
- 1999: OpenMP V1.1 for Fortran (error corrections, explanations)
- 2000: OpenMP V2.0 for Fortran (support of Fortran90 modules)
- 2001: OpenMP V2.0 for C and C++
- 2008: OpenMP V3.0 for Fortran/ C and C++
- 2014: OpenMP V4.0 for Fortran/C and C++

OpenMP - Information

- The OpenMP Architecture Review Board (ARB) Fortran and C Application Program Interfaces (APIs): www.openmp.org
- NCSA online course on OpenMP:
 http://www.citutor.org/login.php?course=24
 (you need to register for a free account, very useful)
- OpenMP tutorial from Lawrence Livermore National Laboratory: https://computing.llnl.gov/tutorials/openMP/
- Book: Rohit Chandra, et.al. "Parallel Programming in OpenMP" Morgan Kaufmann, ISBN 1-55860-671-8

 SEARCH INSIDE!™

Parallel Programming

OpenMP - Overview:

CSOMP FLUSH #pragma omp critical C\$OMP THREADPRIVATE (/ABC/) CALL OMP SET NUM THREADS (10) call omp test lock(jlok) C\$OMP parallel do shared(a, b, c) OpenMP: An API for Writing Multi-threaded Application A set of compiler directives and library routines for C\$OM parallel application programmers Makes it easy to create multi-threaded (MT) programs C\$C in Fortran, C and C++ Standardizes last 15 years of SMP practice #pragma omp parallel for private(A, B) !\$OMP BARRIER C\$OMP PARALLEL COPYIN(/blk/) C\$OMP DO lastprivate(XX) Nthrds = OMP GET NUM PROCS() omp set lock(lck)

OpenMP Overview: How is OpenMP typically used?

- OpenMP is usually used to parallelize loops:
 - Find your most time consuming loops.
 - Split them up between threads.

Split-up this loop between multiple threads

```
#include "omp.h"
void main()
{
    double Res[1000];
    for( int i=0; i<1000; i++)
    {
        do_huge_comp(Res[i]);
    }
}</pre>
#include "omp.h"
void main()
{
    double Res[1000];
    #pragma omp parallel for
    for(int i=0;i<1000;i++)
    {
        do_huge_comp(Res[i]);
    }
}
```

Sequential Program

Parallel Program

Outline — Programming and Execution Model

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Components - Environment, Variables, Directives, Runtime

```
environment variables
#!/bin/tcsh
                                         directives
# Shell-Script
                                         (special comment lines)
gcc -fopenmp test.c
setenv OMP NUM THREADS 4
                                             runtime library
a.out
/* Source file test.c */
#include <stdio.h>
                                                  me: 0
#include <omp.h>
int main(void)
                                                  me: 3
                                                  me: 2
   #pragma omp parallel
                                                  me: 1
   printf("me: %d\n",omp_get thread num());
```

OpenMP Overview: How do threads interact?

- OpenMP is a shared memory model.
 - Threads communicate by sharing variables.
- Unintended sharing of data causes race conditions:
 - <u>race condition</u>: when the program's outcome changes as the threads are scheduled differently.
- To control race conditions
 - Use synchronization to protect data conflicts.
- Synchronization is expensive so:
 - Change how data is accessed to minimize the need for synchronization.

OpenMP: Structured blocks

- Most OpenMP constructs apply to structured blocks
- Structured block: a block with one point of entry at the top and one point of exit at the bottom.
- The only "branches" allowed are exit() in C/C++.

```
#pragma omp parallel
{
  int id = omp_get_thread_num();
  more:
    res(id) = do_big_job(id);
  if( conv(res(id) ) goto more;
}
printf(" All done \n");
```

A structured block

```
if(go_now()) goto more;
#pragma omp parallel
{
  int id = omp_get_thread_num();
  more:
    res(id) = do_big_job(id);
  if( conv(res(id) ) goto done;
  go to more;
}
done:
  if(!really_done()) goto more;
```

Not a structured block

OpenMP: Structured blocks / Conditional Compilation

• In C/C++: a block is a single statement or a group of statements between brackets {}

```
#pragma omp parallel
{
   id = omp_thread_num();
   res(id) =
   lots_of_work(id);
}
```

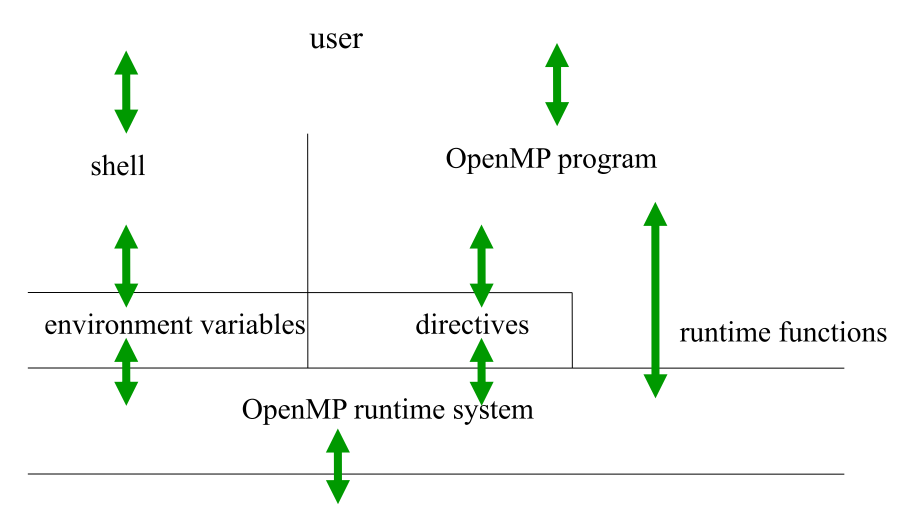
```
#pragma omp for
for(I=0; I<N; I++)
{
    res[I] = big_calc(I);
    A[I] = B[I] + res[I];
}</pre>
```

• Conditional Compilation

```
#ifdef _OPENMP
   iam=omp_get_thread_num();
#endif
```

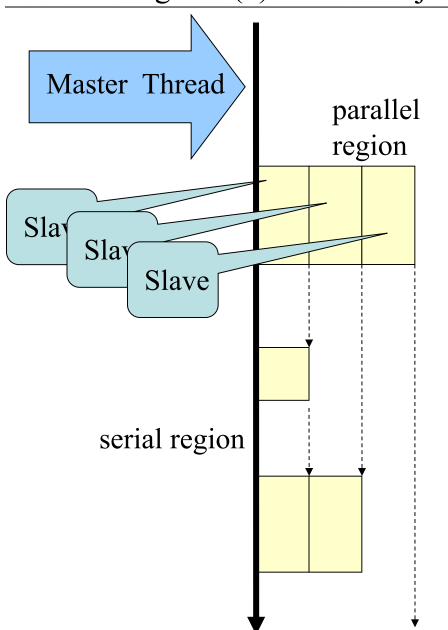
```
/* OpenMP directive */
#pragma omp directive [clause ..]
/* OpenMP directive with
   continuation line */
#pragma omp directive clause \
   clause ...
```

OpenMP Components Diagram



operating system – threads

Parallel Regions (1) - The fork-join concept

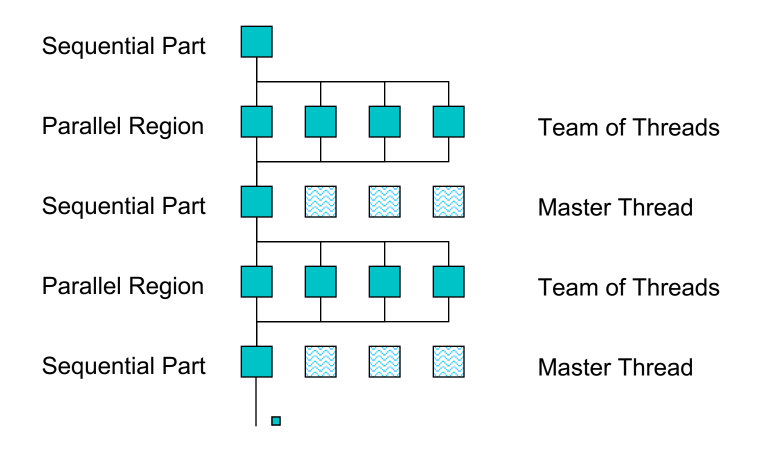


The OpenMP program starts like a serial program: single threaded

In the beginning of the first parallel region the slave threads are started. Together with the master, they form a team.

Between the parallel regions the slave threads are put to sleep.

OpenMP Execution Model

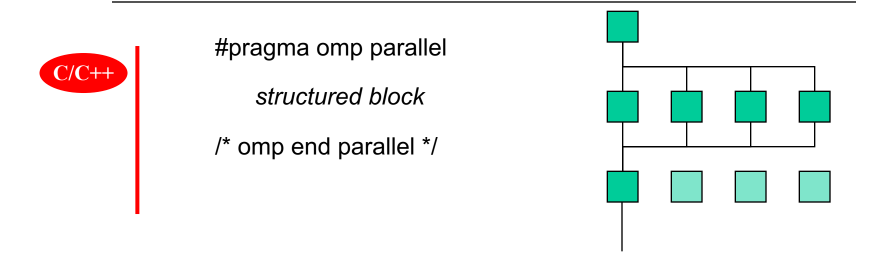


OpenMP: Some syntax details to get us started

- Most of the constructs in OpenMP are compiler directives or pragmas.
 - For C and C++, the pragmas take the form:

```
#pragma omp construct [clause [clause]...]
```

- Include OpenMP file
 - #include "omp.h"



OpenMP Parallel Region Construct Syntax

- C/C++:
 #pragma omp parallel [clause [[,] clause] ...] new-line
 structured-block
- *clause* can be one of the following:
 - private(list)
 - shared(*list*)
 - **...**
- Conditional compilation

```
#ifdef _OPENMP
    printf("%d avail.processors\n",omp_get_num_procs());
#endif
```

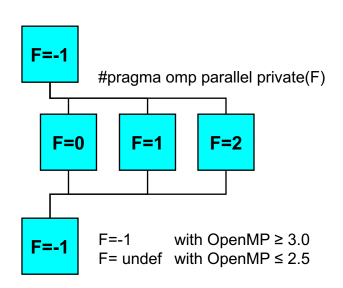
• Include file for library routines:

```
#ifdef _OPENMP
#include <omp.h>
#endif
```

OpenMP Data Scope Clauses

- private (list) Declares the variables in *list* to be private to each thread in a team
- shared (list) Makes variables that appear in *list* shared among all the threads in a team
- If not specified: default shared, but
 - stack (local) variables in called subprograms are PRIVATE
 - Loop control variable of parallel OMP • for (C)

is PRIVATE



OpenMP Environment Variables

OMP NUM THREADS

- sets the number of threads to use during execution
- when dynamic adjustment of the number of threads is enabled, the value of this environment variable is the maximum number of threads to use

```
• setenv OMP NUM THREADS 16 [csh, tcsh]
```

export OMP_NUM_THREADS=16 [sh, ksh, bash]

OMP_SCHEDULE

- applies only to do/for and parallel do/for directives that have the schedule type RUNTIME
- sets schedule type and chunk size for all such loops
- setenv OMP_SCHEDULE "GUIDED, 4" [csh, tcsh]
- export OMP_SCHEDULE="GUIDED, 4" [sh, ksh, bash]

Parallel Regions (2) - Runtime Library

```
#include "omp.h"
void main()
                                                  Sequent. region
printf("inside parallel region? %d\n", omp in parallel());
printf("number of available processors? %d\n",omp get num procs());
printf("maximum number of threads? %d\n", omp get max threads());
 omp set num threads (omp get max threads() );
 #pragma omp parallel
                                                   Parallel region
  printf("inside parallel region? %d\n", omp in parallel());
  printf("number of threads in the team %d\n", omp get num threads());
  printf("my thread id %d\n", omp get thread num() );
```

```
mint [oschenk] export OMP_NUM_THREADS=3
mint [oschenk] ./a.out
inside parallel region? 0
number of available processors? 2
maximum number of threads? 3
```

```
inside parallel region? 1
number of threads in the team 3
my thread id 0
inside parallel region? 1
number of threads in the team 3
my thread id 2
inside parallel region? 1
number of threads in the team 3
my thread id 1
```

Parallel Regions (3) - Runtime Library

	Serial region	Parallel region
<pre>void omp_set_num_threads (int)</pre>	Set # threads to use in a team	don 't
<pre>int omp_get_num_threads (void)</pre>	1	Return # threads
<pre>int omp_get_max_threads (void)</pre>	Return max # threads (OMP_NUM_THREADS)	
<pre>int omp_get_thread_num (void)</pre>	0	Return thread id
		0 #threads-1
<pre>int omp_get_num_procs (void)</pre>	Return # CPUs	
<pre>void omp_set_dynamic (int)</pre>	Control dynamic adjustment of # threads	don 't
<pre>int omp_get_dynamic(void)</pre>	.TRUE:	
	if dynamic thread adjustment enabled .FALSE. Otherwise	
<pre>int omp_in_parallel (void)</pre>	.FALSE.	.TRUE.

Parallel Regions (4)

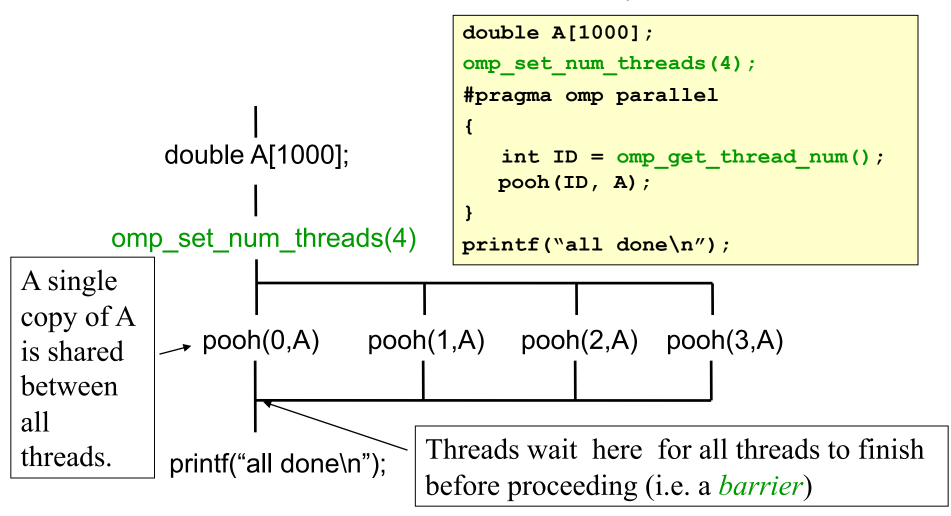
- You create threads in OpenMP with the "omp parallel" pragma.
- For example, to create a 4 thread parallel region:

Each thread executes a copy of the the code within the structured block

• Each thread calls pooh(ID,A) for ID = 0 to 3

Parallel Regions (5)

Each thread executes the same code redundantly.



OpenMP Runtime Library (3): Wall clock timers OpenMP 2.0

- Portable wall clock timers
- DOUBLE PRECISION FUNCTION OMP_GET_WTIME() provides elapsed time

```
START= OMP_GET_WTIME()
! Work to be measured
END = OMP_GET_WTIME()
printf("Work took %e seconds\n", END-START);
```

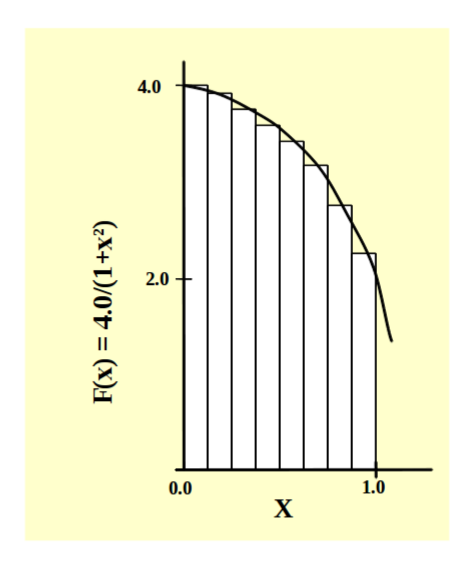
• DOUBLE PRECISION FUNCTION OMP_GET_WTICK()
returns the number of seconds between two successive clock ticks

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In-class exercise: A multi-threaded "pi" program

- On the following slide, you'll see a sequential program that uses numerical integration to compute an estimate of PI.
- Parallelize this program using OpenMP.
- Remember, you'll need to make sure multiple threads don't overwrite each other's private variables.



Mathematically, we know that:

$$\int_{0}^{1} \frac{4.0}{(1+x^{2})} dx = \pi$$

We can approximate the integral as a sum of rectangles:

$$\sum_{i=0}^{N} F(x_i) \Delta x \approx \pi$$

Where each rectangle has width Δx and height $F(x_i)$ at the middle of interval i.

```
static long num steps = 100000;
double step;
void main ()
int i;
double x, pi, sum = 0.0;
step = 1.0/(double) num steps;
for (i=0; i<num steps; i++)</pre>
{
       x = (i+0.5) *step;
       sum = sum + 4.0/(1.0+x*x);
pi = step * sum;
```

The include and timing blocks are removed on the next slides

```
#include <stdio.h>
|#include <time.h>
#include <sys/time.h>
#ifdef OPENMP
   include <omp.h>
#endif
                    include block
\#define f(A) (4.0/(1.0+A*A))
const int n = 10000000;
int main(int argc, char** argv)
  int i;
  double w,x,sum,pi;
  clock t t1, t2;
  struc\overline{t} timeval tv1, tv2;
  struct timezone tz;
# ifdef OPENMP
    double wt1, wt2;
# endif
                   timing block A
  gettimeofday(&tv1, &tz);
# ifdef OPENMP
    wt1=\overline{o}mp get wtime();
```

timing block B

endif

t1=clock();

```
/* pi = integral [0..1] 4/(1+x**2) dx */
  w=1.0/n;
  sum=0.0;
  for (i=1;i<=n;i++)
  {
     x=w*((double)i-0.5);
     sum=sum+f(x);
  }
  pi=w*sum;
     the calculation</pre>
```

```
timing block C
  t2=clock();
# ifdef OPENMP
    wt2=omp get wtime();
# endif
  gettimeofday(&tv2, &tz);
 printf( "computed pi = %24.16q\n", pi);
 printf( "CPU time (clock)
   = %12.4g sec\n'', (t2-t1)/1000000.0);
# ifdef OPENMP
    printf( "wall clock time
    (omp get wtime) = %12.4g sec\n",
     wt2=wt1);
# endif
 printf( "wall clock time (gettimeofday)
   = %12.4q sec\n",
   (tv2.tv sec-tv1.tv sec) +
   (tv2.tv^-usec-tv1.tv^-usec)*1e-6);
  return 0;
```

In-class exercise: Parallel region (1)

- Goal: usage of
 - runtime library calls
 - conditional compilation, environment variables
 - parallel regions, private and shared clauses
- Serial programs: **pi0.c** on https://www2.icorsi.ch/
- Build a team of 2 students, compile **serial** program **pi0.c** on icsmaster cluster
 - ssh icsmaster
 - git pull
 - salloc -t 00:30:00 (to request one node in interactive mode for 30 min)
- Compile as OpenMP program and run on 4 core
 - gcc -O3 -fopenmp pi0.c -o pi0
 - export OMP NUM THREADS=4
 - ./pi0

expected result: program is not parallelized,

therefore same pi-value and timing,

additional output from omp_get_wtime()

In-class exercise: pi1.c

- Modify pi0.c -> pi1.c
- Directly after the declaration part, add in a parallel region that prints on each thread
 - its rank (with omp_get_thread_num()) and
 - the number of threads (with omp get num threads())
- compile gcc –O3 –fopenmp pi1.c –o pi1 and run on 4 cores
- Expected results: numerical calculation is still not parallelized, therefore still same pi-value and timing, additionally output:

```
bash$ gcc -03 -fopenmp -o pi1 pi1.c
bash$ export OMP_NUM_THREADS=4; ./pi1
I am thread 0 of 4 threads
I am thread 2 of 4 threads
I am thread 3 of 4 threads
Computed pi = 3.141592653589731
CPU time (clock) = 0.16 sec
wall clock time (omp_get_wtime) = 0.1681 sec
wall clock time (gettimeofday) = 0.1681 sec
```

OpenMP Advanced Exercise: pi1.c

- Modify pil.c
- Use a private variable for the rank of the threads
- Check, whether you can get a race-condition if you forget the private clause on the **omp parallel** directive, e.g.

```
I am thread 2 of 4 threads
```

• Don't wonder if you get always correct output because the compiler may use on each thread a private register instead of writing into the shared memory

OpenMP Advanced Exercise: pil.c

- Modify pil.c
- Guarantee with conditional compilation, that source code still works with non-OpenMP compilers (i.e., without OpenMP compile-option).
- Add an "else clause", printing a text if OpenMP is not used.
- Expected output:

If compiled with OpenMP, see previous slide.

If compiled without OpenMP

```
bash$ gcc -o pil pil.c
bash$ export OMP_NUM_THREADS=4; ./pil

This program is not compiled with OpenMP
computed pi = 3.1415926535897931
CPU time (clock) = 0.16 sec
wall clock time (gettimeofday) = 0.1681 sec
```

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Worksharing directives

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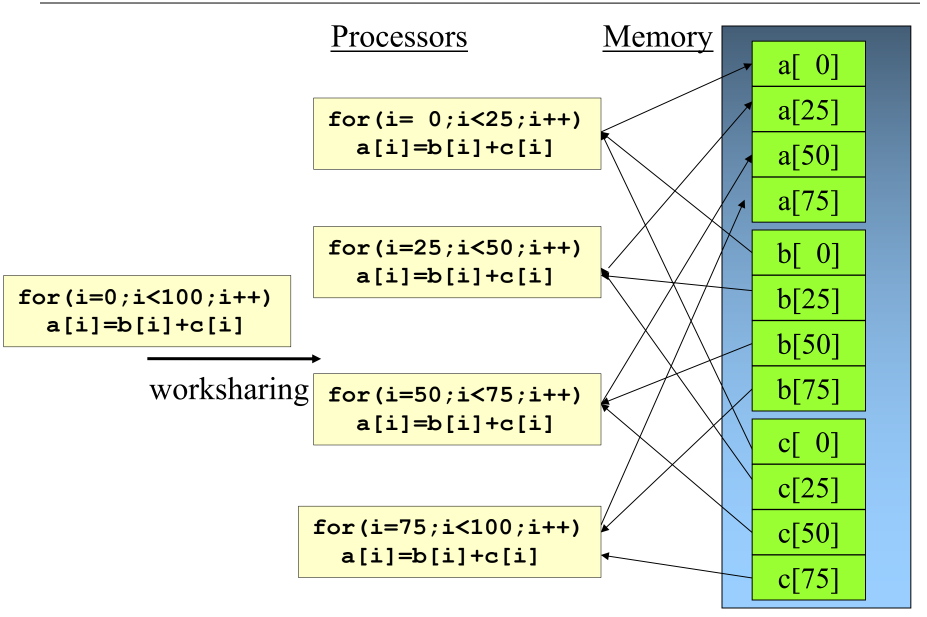
Worksharing and Synchronization

- Which thread executes which statement or operation?
- and when?
 - Worksharing constructs
 - Master and synchronization constructs
- i.e., organization of the parallel work!!!

Worksharing and Synchronization

- Divide the execution of the enclosed code region among the members of the team
- Must be enclosed dynamically within a parallel region
- They do not launch new threads
- No implied barrier on entry
 - for directive
 - sections directive
 - task directive
 - single directive

Work Sharing (1) - Principle



Sequential code

```
for(i=0;I<N;i++) { a[i] = a[i] + b[i];}
```

OpenMP parallel region

```
#pragma omp parallel
{
   int id, i, Nthrds, istart, iend;
   id = omp_get_thread_num();
   Nthrds = omp_get_num_threads();
   istart = id * N / Nthrds;
   iend = (id+1) * N / Nthrds;
   for(i=istart;I<iend;i++)
   { a[i] = a[i] + b[i];}
}</pre>
```

OpenMP

<u>parallel region</u> and a <u>work-sharing</u>

for-construct

```
#pragma omp parallel
#pragma omp for schedule(static)
for(i=0;I<N;i++) { a[i] = a[i] + b[i];}</pre>
```

Work Sharing (2) - Sharing Constructs

• The "for" work-sharing construct splits up loop iterations among the threads in a team

```
#pragma omp parallel
for (I=0;I<N;I++)
{
   NEAD_STUFF(I);
}</pre>
```

All threads select all loop indices "I"

```
#pragma omp parallel
#pragma omp for
for (I=0;I<N;I++)
{
    NEAD_STUFF(I);
}</pre>
```

The loop indices "I" are distributed among threads

By default, there is a **barrier** at the end of the "omp for". Use the **nowait** clause to turn off the barrier.

Work Sharing (7) - Combined parallel/work-share

- OpenMP shortcut:
 - Put the "parallel" and the work-share on the same line

```
double res[MAX];
int i;
#pragma omp parallel
{
    #pragma omp for
    for (i=0;i< MAX; i++)
    {
       res[i] = huge();
    }
}</pre>
```

```
double res[MAX];
int i;
#pragma omp parallel for
for (i=0;i< MAX; i++)
{
   res[i] = huge();
}</pre>
```

These are equivalent

OpenMP for Directives – Syntax

- Immediately following loop executed in parallel
 #pragma omp for [clause[[,]clause] ...] new-line
 for-loop
- The corresponding **for** loop must have *canonical shape*:

```
var: must not be modified in the loop body;
integer (signed or unsigned),
or pointer type (C only), (OpenMP ≥ 3.0)
or random access iterator type (C++ only)
```

Work Sharing (3) - The schedule clause

- The schedule clause effects how loop iterations are mapped onto threads
 - schedule(static [,chunk])
 - Deal-out blocks of iterations of size "chunk" to each thread.
 - In our example: thread #0: i=0 to24; thread #1: i=25 to 49; ...

- schedule(dynamic[,chunk])

- Each thread grabs "chunk" iterations off a queue until all iterations have been handled.
- In our example: thread #0: i=0, 3, 8, ..; thread #1: i=1, 2, 5, ..;

- schedule(guided[,chunk])

• Threads dynamically grab blocks of iterations. The size of the block starts large and shrinks down to size "chunk" as the calculation proceeds.

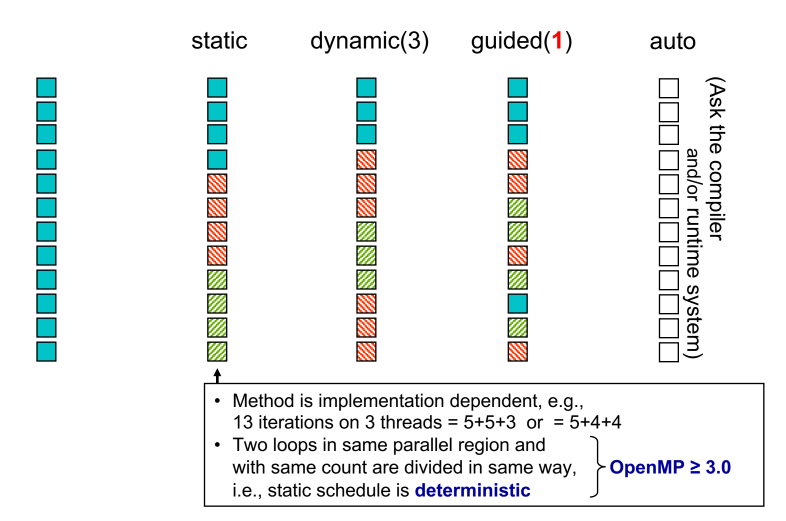
schedule(auto)

Scheduling is delegated to the compiler and/or runtime system
 (OpenMP ≥ 3.0)

schedule(runtime)

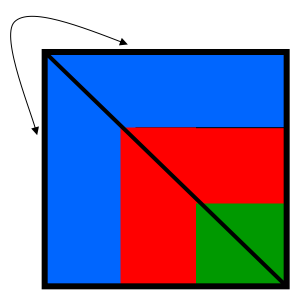
• Schedule and chunk size taken from the OMP_SCHEDULE environment variable.

Loop scheduling



OpenMP

```
#pragma omp parallel for private(h,i,j)
schedule static numthreads(3)
for (i=0; i<n; i++) {
   for (j=i+1; j<n; j++) {
     h = A[i][j];
     A[i][j] = A[j][i];
     A[j][i] = h;
}
} // end parallel for</pre>
```



Thread 0 would have much more work than thread 2!

Work Sharing (5) - The schedule clause

Schedule Clause	When To Use
STATIC	Predictable and similar work per iteration
	thread #0: i=0 to24; thread #1: i=25 to 49 thread #2: i=50 to74; thread #0: i=75 to 99
DYNAMIC	Unpredictable, highly variable work per iteration
	thread #0: i=0,2, 8,, thread #1: 4,6,7, thread #2: i=2,3,9,; thread #0: i=15,18,91,
GUIDED	Special case of dynamic to reduce scheduling overhead

Work Sharing (6) - The section clause

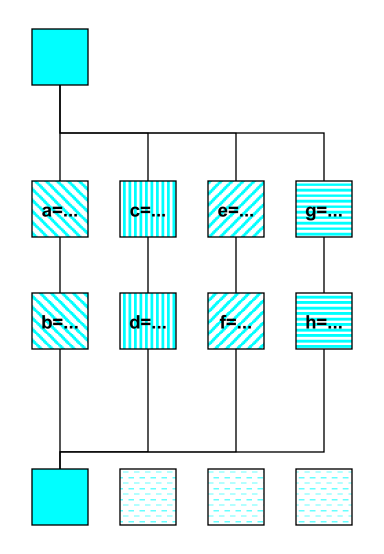
• The **section work-sharing** construct gives a different structured block to each thread.

```
#pragma omp parallel
#pragma omp sections
{
    #pragma omp section
    X_calculation(); // only one thread
    #pragma omp section
    y_calculation(); // only one thread
    #pragma omp section
    z_calculation(); // only one thread
}
```

By default, there is a **barrier** at the end of the "omp sections". Use the "**nowait**" clause to turn off the barrier.

Work Sharing (6) - The section clause

```
C / C++: #pragma omp parallel
          #pragma omp sections
          #pragma omp section
             { a=...;
               b=...; }
          #pragma omp section
             { c=...;
               d=...; }
          #pragma omp section
             { e=...;
               f=...; }
          #pragma omp section
             { g=...;
               h=...; }
           } /*omp end sections*/
          } /*omp end parallel*/
```



OpenMP task Directive – Parallelized traversing of a tree

```
C/C++
```

```
struct node {
                                                                 Starting the parallel
                                                                 team of threads
   struct node *left;
   struct node *right;
                                                                 Using only one thread
};
                                                                 for starting the
extern void process(struct node *);
                                                                 traversal
void traverse( struct node *p ) {
                                                                 First execution with
    if (p->left)
                                                                 single thread
#pragma omp task // p is firstprivate by default
                                                                 (= 1<sup>st</sup> task)
                   traverse(p->left);
                                                                A new task is started
    if (p->right)
                                                                 (on a new thread)
#pragma omp task // p is firstprivate by default

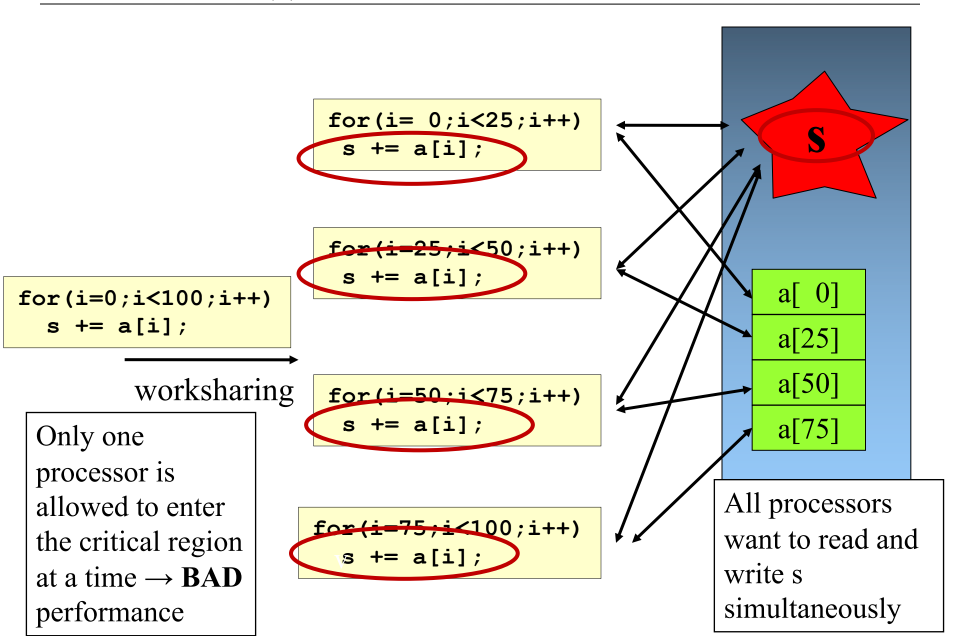
    A recursive call to

                   traverse(p->right);
    process(p); // significant work with p
                                                                 traverse() in this
                                                                 2<sup>nd</sup> task
int main(int argc, char **argv)
                                                                 3<sup>rd</sup> task is started
{ struct node tree;
                                                                 Work is done
  ... // producing the tree
                                                                 in 1<sup>st</sup> task
#pragma omp parallel ←
                                                                 Recursive calls start-
                                                                 ing 4<sup>th</sup>, 5<sup>th</sup>, ... tasks
#pragma omp single
        traverse(&tree);//traversing the existing tree
      } // end of omp single
  } // end of omp parallel
```

Critical sections (1)

Processors **Memory** for(i= 0;i<25;i++) s += a[i]; for(i=25;i<50;i++) s += a[i]; a[0] for(i=0;i<100;i++) s += a[i];a[25] a[50] worksharing for(i=50;i<75;i++) s += a[i]; a[75] All processors for(i=75;i<100;i++) want to read and s += a[i]; write s simultaneously

Critical sections (2)



Critical sections (3)

• Only one thread at a time can enter a critical section.

Threads wait their turn – only one at a time calls consum()

```
float res;
#pragma omp parallel
   float B;
   int i;
   #pragma omp for
   for(i=0;i<niters;i++)</pre>
      B = big_job(i);
      #pragma omp critical
      consum (B, RES);
```

Critical sections (4) – Critical / end critical

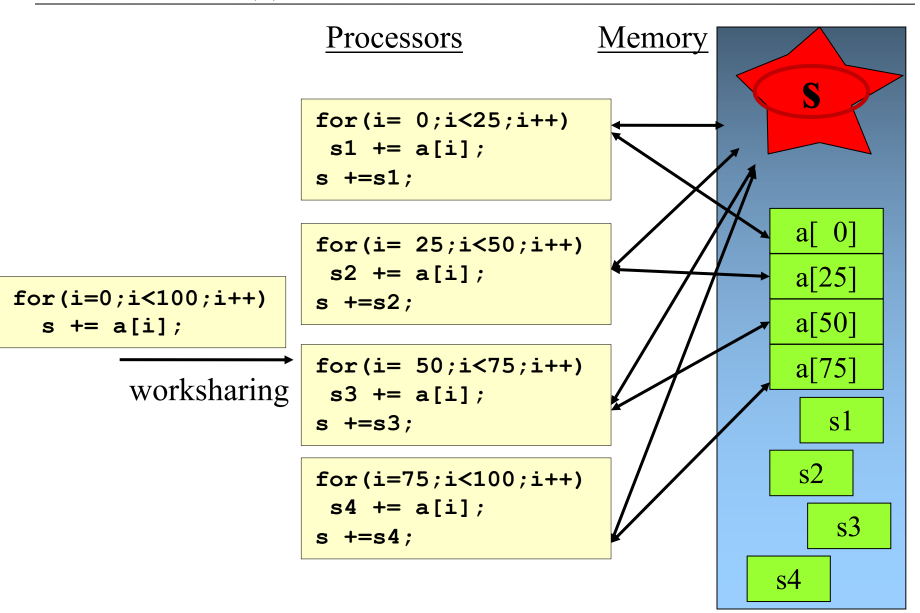
```
for(i=0;i<100;i++)
s= s + a[i];
```

```
#pragma omp parallel for private (i)
for (i=0; i<100; i++) {
    #pragma omp critical
    { s += a[i]; }
}</pre>
```

Only one processor is allowed to enter the **critical section** at a time

As the loop body consists of a critical section only, the parallel program will run much slower

Critical section (5)



Critical section (6) – Critical / end critical

```
for(i=0;i<100;i++)
s= s + a[i];
```

```
#pragma omp parallel \
    private (i,s_local)
{

    s_local = 0;
    #pragma omp for
        for (i=0; i<100; i++)
            { s_local += a[i]; }

    #pragma omp critical
            { s += s_local; }
}</pre>
```

Only one processor is allowed to enter the **critical section** at a time

As the loop body consists of a critical region only, the parallel program will run much slower

Now the partial sums are calculated in parallel. The critical region is entered only once per thread.

Outline

- Introduction into OpenMP
- Programming and Execution Model
 - Parallel regions: team of threads
 - Syntax
 - Data environment (part 1)
 - Environment variables
 - Runtime library routines
 - Exercise 1: Parallel region / library calls / privat & shared variables
- Worksharing directives
 - Which thread executes which statement or operation?
 - Synchronization constructs, e.g., critical section
 - Exercise 2: Pi
- Data environment and combined constructs
 - Nesting and Binding
 - Private and shared variables, Reduction clause
 - Combined parallel worksharing directives
 - Exercise 3: Pi with reduction clause and combined constructs
 - Exercise 4: Heat
- Summary of OpenMP API
- OpenMP Pitfalls & Optimization Problems

In-class exercise 2: pi Program (1)

- Goal: usage of
 - worksharing constructs: #pragma omp for
 - #pragma omp critical directive
 - Use your result **pi1.c** from the last in-class exercise
 - Modify pi1.c -> pi2.c
 - compile serial program **pi2.c** and run
- add parallel region and pragma omp for directive and compile
- set environment variable **OMP_NUM_THREADS** to **2** and run value of pi? (should be wrong!)
- run againvalue of pi? (...wrong and unpredictable)
- set environment variable **OMP_NUM_THREADS** to **4** and run value of pi? (...and stays wrong)
- run again value of pi? (...but where is the race-condition?)

In-class exercise 2: pi Program (2)

- add private (x) clause in pi2.c and compile
- set environment variable **OMP_NUM_THREADS** to **2** and run value of pi? (should be still incorrect ...)
- run again value of pi?
- set environment variable **OMP_NUM_THREADS** to **4** and run value of pi?
- run again value of pi? (... and where is the second race-condition?)

In-class exercise 2: pi Program (3)

- add critical directive in pi2.c around the sum-statement and compile
- set environment variable **OMP_NUM_THREADS** to **2** and run value of pi? (should be now correct!, but huge CPU time!)
- run againvalue of pi? (but not reproducible in the last bit!)
- set environment variable **OMP_NUM_THREADS** to **4** and run value of pi? execution time? (Oh, does it take longer?)
- run againvalue of pi? execution time?How can you optimize your code?

In-class advanced exercise 2: pi Program (4)

- Modify the printing of the thread rank and the number of threads from Exercise 1:
 - Only one thread should print the real number of threads used in parallel regions.
 - For this, use a **single** construct
 - Expected result:

```
OpenMP-parallel with 4 threads
computed pi = 3.14159265358967

CPU time (clock) = 0.01659 sec
wall clock time (omp_get_wtime) = 0.01678 sec
wall clock time (gettimeofday) = 0.01679 sec
```