

Introduction to OpenMP

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Outline

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1 Introduction

- 2 parallel and worksharing constructs
- 3 Synchronization constructs
- 4 Data environment
- 5 Runtime library routines and environment variables
- 6 OpenMP Compilers



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Disadvantages of MPI

- Each MPI process can only access its local memory
 - The data to be shared must be exchanged with explicit inter-process communications (messages)
 - It is the responsibility of the programmer to design and implement the exchange of data between processes
- You can not adopt a strategy of incremental parallelization
 - The communication structure of the entire program has to be implemented
- The communications have a cost
- It is difficult to have a single version of the code for the serial and MPI program
 - Additional variables are needed
 - You need to manage the correspondence between local variables and global data structure

What is OpenMP?

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- De-facto **standard** Application Program Interface (API) to write shared memory parallel applications in C, C++ and Fortran
- Consists of compilers directives, run-time routines and environment variables
- "Open specifications for Multi Processing" maintained by the OpenMP Architecture Review Board (http://www.openmp.org)
- The "workers" who do the work in parallel (thread) "cooperate" through shared memory
- Memory accesses instead of explicit messages
- "local" model parallelization of the serial code
- It allows an incremental parallelization

History

- Born to satisfy the need of unification of proprietary solutions
- The past
 - October 1997 Fortran version 1
 - October 1998 C/C++ version 1
 - November 1999 Fortran version 1.1 (interpretations)
 - November 2000 Fortran version 2
 - March 2002 C/C++ version 2
 - May 2005 combined C/C++ and Fortran version 2
 - May 2008 version 3.0
- The present
 - July 2011 version 3.1
 - July 2013 version 4.0
 - November 2015 version 4.5
- The future

version 5.0

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Distributed and shared memory







UMA and NUMA systems

CPU CPU Memory CPU CPU **Bus** Router Bus Memory Bus CPU CPU CPU CPU Memory

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Process and thread

- A process is an instance of a computer program
- Some information included in a process are:
 - Text
 - Machine code
 - Data
 - Global variables
 - Stack

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- Local variables
- Program counter (PC)
 - A pointer to the istruction to be executed





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Multi-threaded processes

- The process contains several concurrent execution flows (threads)
 - Each thread has its own program counter (PC)
 - Each thread has its own private stack (variables local to the thread)
 - The instructions executed by a thread can access:
 - the process global memory (data)

the thread local stack



Execution model



Why should I use OpenMP?

Standardized

• enhance portability

2 Ease of use

- limited set of directives
- fast code parallelization
- parallelization is incremental
- coarse/fine parallelism

Ortability

- C, C++ and Fortran API
- · part of many compilers

OpenMP (possible) issues

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Performance

- may be non-portable
- increase memory traffic

2 Limitations

- shared memory systems
- mainly used for loops



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Structure of an OpenMP program

Execution model

- the program starts with an initial thread
- when a parallel construct is encountered a team is created
- parallel regions may be nested arbitrarily
- worksharing constructs permit to divide work among threads

Shared-memory model

- all threads have access to the memory
- each thread is allowed to have a temporary view of the memory
- · each thread has access to a thread-private memory
- two kinds of data-sharing attributes: private and shared
- data-races trigger undefined behavior

Programming model

• compiler directives + environment variables + run-time library



OpenMP core elements



Conditional compilation

C/C++

```
#ifdef _OPENMP
printf("OpenMP support:%d",_OPENMP);
#else
printf("Serial execution.");
#endif
```

Fortran

- !\$ print *, "OpenMP support"
- **1** The macro _OPENMP has the value yyyymm
- Portran 77 supports !\$, *\$ and c\$ as sentinels
- **3** Fortran 90 supports !\$ only

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parallel construct



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C/C++

```
int main () {
```

```
printf("Hello world\n");
```

```
return 0;
```



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C/C++

```
int main () {
    /* Serial part */
```

```
#pragma omp parallel
{
    printf("Hello world\n");
}
/* Serial part */
return 0;
}
```

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Fortran

PROGRAM HELLO

Print *, "Hello World!!!"

END PROGRAM HELLO



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Fortran

PROGRAM HELLO ! Serial code

!\$OMP PARALLEL
 Print *, "Hello World!!!"
!\$OMP END PARALLEL

! Resume serial code

END PROGRAM HELLO

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What's wrong?

```
int main() {
    int i;
#pragma omp parallel
    {
        for(i = 0; i < 10; ++i)
            printf("iteration %d\n", i);
    }
    return 0;
}</pre>
```

Race condition

- A race condition (or data race) is when two or more threads access the same memory location:
 - asyncronously and,
 - · without holding any common exclusive locks and,
 - at least one of the accesses is a write/store
- In this case the resulting values are undefined

```
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Defined or undefined?
                                                  Computing 2017
   What's the result?
   #include <stdio.h>
   #include <omp.h>
   void main() {
        int a;
       a = 0;
       #pragma omp parallel
            // omp_get_thread_num returns the id
            // of the thread
            a = a + omp_get_thread_num();
       printf("%d\n", a);
```



Worksharing constructs: rules

- Distribute the execution of the associated region
- A worksharing region has no barrier on entry
- 3 An implied barrier exists at the end, unless nowait is specified
- 4 Each region must be encountered by all threads or none
 - Every thread must encounter the same sequence of worksharing regions and barrier regions



Worksharing constructs: types

- The OpenMP API defines four worksharing constructs:
 - loop
 - sections
 - single
 - workshare







Loop construct: restrictions

- Only loops with canonical forms are allowed
 - The iteration count needs to be computed before executing the loops
 - incr-expr: addition or subtraction expression.

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Wrong loop construct

wrong incremental expression

```
#include <stdio.h>
#include <omp.h>
```

```
void incr(int *var) {
        *var = *var + 1;
   }
   void main() {
        int a;
   #ifdef OPENMP
        #pragma omp parallel for
   #endif
        for (a=0;a<10;incr(&a))</pre>
            printf("%d\n", a);
9911
```

Loop construct: the rules

- The iterations of the loop are **distributed** over the threads that already exist in the team
- 2 The iteration variable in the for loop
 - if shared, is implicitly made private
 - must not be modified during the execution of the loop
 - has an unspecified value after the loop
- **3** The schedule clause:
 - may be used to specify how iterations are divided into chunks
- **4** The collapse clause:
 - may be used to specify how many loops are parallelized
 - · valid values are constant positive integer expressions

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Loop construct: schedule kind

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static

- if no chunk_size is specified the iterations space is diveded in chunks of equal size and one chunk per thread
- if chunk_size is specified, chunks are assigned to the threads in a round-robin fashion
- must be reproducible within the same parallel region

Ø dynamic

- iterations are divided into chunks of size chunk_size
- the chunks are assigned to the threads as they request them
- the default chunk_size is 1

guided

- · iterations are divided into chunks of decreasing size
- · the chunks are assigned to the threads as they request them
- chunk_size controls the minimum size of the chunks



Figure: Different scheduling for a 1000 iterations loop with 4 threads: guided (top), dynamic (middle), static (bottom)


Loop construct: nowait clause

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Where are the implied barriers?

```
void nowait_example(int n, int m, float *a,
  float *b, float *y, float *z) {
#pragma omp parallel
#pragma omp for
    for (int i=1; i<n; i++)</pre>
      b[i] = (a[i] + a[i-1]) / 2.0;
#pragma omp for
    for (int i=0; i<m; i++)</pre>
      y[i] = sqrt(z[i]);
  }
```

Loop construct: nowait clause

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Where are the implied barriers?

```
void nowait_example(int n, int m, float *a,
  float *b, float *y, float *z) {
#pragma omp parallel
#pragma omp for nowait
    for (int i=1; i<n; i++)</pre>
      b[i] = (a[i] + a[i-1]) / 2.0;
#pragma omp for nowait
    for (int i=0; i<m; i++)</pre>
      y[i] = sqrt(z[i]);
  }
```

Loop construct: nowait clause

Is the following snippet semantically correct?

```
. . .
   int i;
   #pragma omp parallel
   #pragma omp for
        for (i=0; i<n; i++)</pre>
          c[i] = (a[i] + b[i]) / 2.0f;
   #pragma omp for
        for (i=0; i<n; i++)</pre>
          z[i] = sqrtf(c[i]);
   #pragma omp for
        for (i=1; i<=n; i++)</pre>
          y[i] = z[i-1] + a[i];
****
```

Loop construct: nowait clause

We can explot the static schedule. Say we have 100 iterations and 10 threads

th 0	for	i	knows
c[i]	i=0,99	0,,9	
z[i]	i=0,99	0,,9	c[0,,9]
y[i]	i=1,100	1,,10	z[0,,9]



Loop construct: nowait clause

Is the following snippet semantically correct?

```
. . .
int i;
#pragma omp parallel
#pragma omp for schedule(static) nowait
    for (i=0; i<n; i++)</pre>
      c[i] = (a[i] + b[i]) / 2.0f;
#pragma omp for schedule(static) nowait
    for (i=0; i<n; i++)</pre>
      z[i] = sqrtf(c[i]);
#pragma omp for schedule(static) nowait
    for (i=1; i<=n; i++)</pre>
      v[i] = z[i-1] + a[i];
```

Loop construct: nested loops

Am I allowed to do the following?

```
#pragma omp parallel
{
    #pragma omp for
    for(int ii = 0; ii < n; ii++) {
        #pragma omp for
        for(int jj = 0; jj < m; jj ++) {
            A[ii][jj] = ii*m + jj;
        }
    }
}</pre>
```



Loop construct: collapse clause

The right way to collapse nested loops

```
#pragma omp parallel
{
    #pragma omp for collapse(2)
    for(int ii = 0; ii < n; ii++) {
        for(int jj = 0; jj < m; jj ++) {
            A[ii][jj] = ii*m + jj;
            }
        }
    }
}</pre>
```

Loop collapse

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- Allows parallelization of perfectly nested rectangular loops
- The collapse clause indicates how many loops should be collapsed
- Compiler forms a single loop (e.g. of length NxM) and then parallelizes it
- Useful if N < no. of threads, so parallelizing the outer loop makes balancing the load difficult.

Loop dependencies

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loop carried dependencies

```
int i, j, ARR[N];
j = 3;
for (i=0; i<N; i++) {
    j+=2;
    ARR[i] = func(j);
}</pre>
```

i	j
0	3+2
1	3+4
2	3+6
n	3+(2*n+2)





Removing loop dependencies

removig loop carried dependencies

```
int i, j, ARR[N];
for (i=0; i<N; i++) {
    j = 3+(2*i+2);
    ARR[i] = func(j);
}
```

Sections construct: syntax

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C/C++

```
#pragma omp sections [clause[[,] clause]...]
{
#pragma omp section
  structured-block
#pragma omp section
  structured-block
...
```



Sections construct: syntax

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Fortran

```
!$omp sections [clause[[,] clause]...]
```

- !\$omp section
 - structured-block
- !\$omp section
 - structured-block
- • •
- !\$omp end sections [nowait]



Sections construct: some facts

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- it contains a set of structured-blocks
- each one is executed once by one of the threads
- Scheduling of the sections is implementation defined
- O There is an implied barrier at the end of the construct







Single construct: some facts

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- The associated structured block is executed by only one thread
- 2 The other threads wait at an implicit barrier
- ③ The method of choosing a thread is implementation defined



Workshare construct: syntax

Fortran

!\$omp workshare
 structured-block
!\$omp end workshare [nowait]

Divides the following into shared units of work:

- array assignments
- 2 FORALL statements or constructs
- **3** WHERE statements or constructs

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Reduction clause: some facts

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1 The reduction clause:

- is valid on parallel, loop and work-sharing constructs
- specifies an operator and one or more list items
- 2 A list item that appears in a reduction clause must be shared
- 3 For each item in the list:
 - a local copy is created and initialized appropriately based on the reduction operation (e.g * -> 1)
 - updates occur on the local copy.
 - local copies are **reduced** into a single value and combined with the original global value.
- Items must not be const-qualified



Reduction clause: example

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Sum over many iterations

```
int a = 5;
#pragma omp parallel
{
  #pragma omp for reduction(+:a)
    for(int i = 0; i < 10; ++i)
        ++a;
}
printf("%d\n", a);
```

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Master construct: syntax

C/C++

#pragma omp master
 structured-block

Fortran

!\$omp master
 structured-block
!\$omp end master



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Master construct: some facts

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1 The master construct specifies a structured block:

- that is executed by the master thread
- with no implied barrier on entry or exit
- Oused mainly in:
 - hybrid MPI-OpenMP programs
 - progress/debug logging



Critical construct: syntax

C/C++

#pragma omp critical [name]
 structured-block

Fortran

!\$omp critical [name]
 structured-block
!\$omp end critical [name]



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Critical contruct: some facts

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1 The critical construct restricts the execution:

- to a single thread at a time (wait on entry)
- An optional name may be used to identify a region. All critical without a name share the same unspecified tag
- When a thread encounters a critical construct, it waits until no other thread is executing a critical region with the same name.
- In Fortran the names of critical constructs:
 - are global entities of the program
 - may conflict with other names (and trigger undefined behavior)



Critical construct: example

Named critical regions

```
#pragma omp parallel
{
#pragma omp critical(long_critical_name)
    doSomeCriticalWork_1();
#pragma omp critical
    doSomeCriticalWork_2();
#pragma omp critical
    doSomeCriticalWork_3();
}
```



at which the construct appears



Barrier construct: example

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Waiting for the master

```
int counter = 0;
#pragma omp parallel
{
  #pragma omp master
    counter = 1;
#pragma omp barrier
    printf("%d\n", counter);
}
```

Atomic construct: syntax

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C/C++

```
#pragma omp atomic \
  [read | write | update | capture]
  expression-stmt
```

#pragma omp atomic capture
 structured-block



Atomic construct: syntax

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Fortran

- !\$omp atomic read capture-statement [!\$omp end atomic]
- !\$omp atomic write write-statement [!\$omp end atomic]



Atomic construct: syntax

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Fortran

!\$omp atomic [update]
 update-statement
[!\$omp end atomic]

!\$omp atomic capture update-statement capture-statement !\$omp end atomic



Atomic construct: some facts

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1 The atomic construct:

- ensures a specific storage location to be updated atomically
- · does not expose it to multiple, simultaneous writing threads
- 2 The binding thread set for an atomic region is all threads

3 The atomic construct with the clause:

read forces an atomic read regardless of the machine word size
write forces an atomic write regardless of the machine word size
update forces an atomic update (default)

capture same as an update, but captures original or final value

Accesses to the same location must have compatible types

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Data-sharing attributes: C/C++



- objects with dynamic storage duration
- variables with static storage duration
- file scope variables
- 2 The following are always private:
 - · loop iteration variable in the loop construct
 - variables with automatic storage duration
- O Arguments passed by reference inherit the attributes



Data-sharing attributes: Fortran

1 The following are always private:

- variables with automatic storage duration
- loop iteration variable in the loop construct
- 2 The following are always shared:
 - assumed size arrays
 - variables with save attribute
 - · variables belonging to common blocks or in modules
- 3 Arguments passed by reference inherit the attributes



Default/shared/private clauses

1 The clause default:

- is valid on parallel
- accepts only shared or none in C/C++ and Fortran
- accepts also private and firstprivate in Fortran
- default (none) requires each variable to be listed in a clause
- 2 The clause shared(list):
 - is valid only on parallel contruct
 - · declares one or more list items to be shared
- 3 The clause private(list):
 - is valid on parallel, and worksharing contructs
 - · declares one or more list items to be private
 - allocates a new item of the same type with undefined value

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Default/shared/private clauses

Example

```
int q,w;
#pragma omp parallel private(q) shared(w)
{
    q = 0;
#pragma omp single
    w = 0;
#pragma omp critical(stdout_critical)
    printf("%d %d\n", q, w);
}
```



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Firstprivate clause

Example

```
int q = 3, w;
#pragma omp parallel firstprivate(q) shared(w)
{
  #pragma omp single
  w = 0;
#pragma omp critical(stdout_critical)
  printf("%d %d\n", q, w);
}
```

Same as private, but initializes items



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Lastprivate clause

Example

```
#pragma omp parallel
{
    #pragma omp for lastprivate(i)
    for(i = 0; i < (n1); ++i)
        a[i] = b[i] + b[i + 1];
    }
    a[i] = b[i];</pre>
```

1 valid on for, sections

2 the value of each new list item is the sequentially last value

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Copyprivate clause

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C/C++

```
#pragma omp single copyprivate(tmp)
{
   tmp = (float *) malloc(sizeof(float));
} /* copies the pointer only */
```

1 Valid only on single

Proadcasts the value of a private variable



False sharing

- · say we have array elements contiguous in memory
- if independent data elements are on the same cache line threads might share the same cache line
- each update on one element will cause the cache lines of the remaining threads to be trashed
- this is called false sharing
- poor scalability
- Solution:
 - When updates to an item are frequent, work with local copies of data instead of an array indexed by the thread ID.
 - Pad arrays so elements you use are on distinct cache lines.

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Runtime library routines

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Most used functions

int omp_get_num_threads(void);// # of threads
int omp_get_thread_num(void);// thread id
double omp_get_wtime(void);// get wall-time

- Prototypes for C/C++ runtime are provided in omp.h
- 2 Interface declarations for Fortran are provided as:
 - a Fortran include file named omp_lib.h
 - a Fortran 90 module named omp_lib



Environment variables

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OMP_NUM_THREADSsets the number of threads for parallel regionsOMP_STACKSIZEspecifies the size of the stack for threadsOMP_SCHEDULEcontrols schedule type and chunk size of runtimeOMP_PROC_BINDcontrols whether threads are bound to processorsOMP_NESTEDenables or disables nested parallelism



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OpenMP Compilers

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GNU:

(Version \geq = 4.3.2) Compile with **-fopenmp** For Linux, Solaris, AIX, MacOSX, Windows.

IBM:

Compile with -qsmp=omp for Windows, AIX and Linux.

Intel:

Compile with **-Qopenmp** on Windows, or **-qopenmp** on Linux or Mac



OpenMP Compilers

Sun Microsystems:

Compile with **-xopenmp** for Solaris and Linux.

Portland Group Compilers:

Compile with -mp Emit useful information to stderr. -Minfo=mp

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OpenMP: THE END!!!

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Good luck and enjoy OpenMP!!!

