

# Introduction to OpenMP

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# **History**

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- De-facto standard for Shared-Memory Parallelization.
- 1997: OpenMP 1.0 for FORTRAN
- 1998: OpenMP 1.0 for C and C++
- 1999: OpenMP 1.1 for FORTRAN (errata)
- 2000: OpenMP 2.0 for FORTRAN
- 2002: OpenMP 2.0 for C and C++
- 2005: OpenMP 2.5 now includes both programming languages.
- 05/2008: OpenMP 3.0 release
   07/2011: OpenMP 3.1 release

### 07/2013: OpenMP 4.0 release

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RWTH Aachen University is a member of the OpenMP Architecture Review Board (ARB) since 2006.



# OpenMP Overview & Parallel Region

### **OpenMP's machine model**



OpenMP: Shared-Memory Parallel Programming Model.



All processors/cores access a shared main memory.

Real architectures are more complex, as we will see later / as we have seen.

# Parallelization in OpenMP employs multiple threads.

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# **OpenMP Execution Model**

- OpenMP programs start with just one thread: The *Master*.
  - Worker threads are spawned at Parallel Regions, together with the Master they form the Team of threads.
  - In between Parallel Regions the Worker threads are put to sleep. The OpenMP *Runtime* takes care of all thread management work.

#### Concept: Fork-Join.

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Allows for an incremental parallelization!



### Parallel Region and Structured Blocks



### The parallelism has to be expressed explicitly.



#### Structured Block

- $\rightarrow$  Exactly one entry point at the top
- $\rightarrow$  Exactly one exit point at the bottom
- $\rightarrow$  Branching in or out is not allowed
- Terminating the program is allowed (abort / exit)

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### Specification of number of threads:

Environment variable:

OMP\_NUM\_THREADS=...

 Or: Via num\_threads clause:
 add num\_threads (num) to the parallel construct



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# Hello OpenMP World



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# Hello orphaned OpenMP World

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### **Starting OpenMP Programs on Linux**



From within a shell, global setting of the number of threads:

```
export OMP_NUM_THREADS=4
```

./program

### From within a shell, one-time setting of the number of threads:

OMP\_NUM\_THREADS=4 ./program



# **For Worksharing Construct**

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# **For Worksharing**



- If only the *parallel* construct is used, each thread executes the Structured Block.
- Program Speedup: Worksharing
- OpenMP's most common Worksharing construct: for

C/C++	Fortran
<pre>int i; #pragma omp for for (i = 0; i &lt; 100; i++) {</pre>	<pre>INTEGER :: i !\$omp do DO i = 0, 99     a[i] = b[i] + c[i]; END DO</pre>
}	

 $\rightarrow$  Distribution of loop iterations over all threads in a Team.

 $\rightarrow$  Scheduling of the distribution can be influenced.

### Loops often account for most of a program's runtime!

# **Worksharing illustrated**





Demo



# **Summing up Vector Elements**

### **Synchronization Overview**



### Can all loops be parallelized with for-constructs? No!

→ Simple test: If the results differ when the code is executed backwards, the loop iterations are not independent. BUT: This test alone is not sufficient:

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

*Data Race*: If between two synchronization points at least one thread writes to a memory location from which at least one other thread reads, the result is not deterministic (race condition).

# **Synchronization: Critical Region**



A *Critical Region* is executed by all threads, but by only one thread simultaneously (*Mutual Exclusion*).

C/C++	
<pre>#pragma omp critical (name)</pre>	
<pre>{     structured block }</pre>	

#### Do you think this solution scales well?

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

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# **Data Scoping**

# **Scoping Rules**



• Managing the Data Environment is the challenge of OpenMP.

#### **Scoping** in OpenMP: Dividing variables in *shared* and *private*:

- → private-list and shared-list on Parallel Region
- → private-list and shared-list on Worksharing constructs
- → General default is *shared* for Parallel Region, *firstprivate* for Tasks.
- → Loop control variables on for-constructs are private
- $\rightarrow$  Non-static variables local to Parallel Regions are *private*
- → private: A new uninitialized instance is created for each thread
  - $\rightarrow$  *firstprivate*: Initialization with Master's value
  - $\rightarrow$  *lastprivate*: Value of last loop iteration is written back to Master

→ Static variables are shared Introduction to OpenMP Dirk Schmidl | IT Center der RWTH Aachen University

# Privatization of Global/Static Variables



- Global / static variables can be privatized with the *threadprivate* ... or eated for each thread
  > Before the first parallel region is encountered read private additional of the program ends of th directive
  - $\rightarrow$  One instance is created for each thread

- $\rightarrow$  Does not work (well) with nested
- → Based on thread-local ste
  - →TIsAlloc (Win32 ead\_key\_create (Posix-Threads), keyword

threadprivate(i)

Fortran

```
INTEGER ::
SAVE
!$omp threadprivate(i)
```

C/C++



# **The Barrier Construct**

### **The Barrier Construct**



### OpenMP barrier (implicit or explicit)

→ Threads wait until all threads of the current *Team* have reached the barrier

C/C++

#pragma omp barrier

All worksharing constructs contain an implicit barrier at the end



# Back to our bad scaling example

C/C++







} // end parallel

### **The Reduction Clause**



In a *reduction*-operation the operator is applied to all variables in the list. The variables have to be *shared*.

```
→ reduction (operator:list)
```

 $\rightarrow$  The result is provided in the associated reduction variable

```
C/C++
#pragma omp parallel for reduction(+:s)
for(i = 0; i < 99; i++)
{
    s = s + a[i];
}</pre>
```

 $\rightarrow$  Possible reduction operators with initialization value:





Ρ

# Example: Pi (1/2)

```
double f(double x)
{
    return (4.0 / (1.0 + x*x));
}
```

```
double CalcPi (int n)
```

{

```
const double fH = 1.0 / (double) n;
double fSum = 0.0;
double fX;
int i;
```

```
#pragma omp parallel for
for (i = 0; i < n; i++)
{
    fX = fH * ((double)i + 0.5);
    fSum += f(fX);
}
return fH * fSum;
}
```







# Example: Pi (1/2)

```
double f(double x)
{
    return (4.0 / (1.0 + x*x));
}
```

```
double CalcPi (int n)
```

{

```
const double fH = 1.0 / (double) n;
double fSum = 0.0;
double fX;
int i;
```

```
#pragma omp parallel for private(fX,i) reduction(+:fSum)
    for (i = 0; i < n; i++)
    {
        fX = fH * ((double)i + 0.5);
        fSum += f(fX);
    }
    return fH * fSum;
}</pre>
```







# Example: Pi (2/2)



### **Results:**

# Threads	Runtime [sec.]	Speedup
1	1.11	1.00
2		
4		
8	0.14	7.93

### Scalability is pretty good:

- $\rightarrow$  About 100% of the runtime has been parallelized.
- → As there is just one parallel region, there is virtually no overhead introduced by the parallelization.
- $\rightarrow$  Problem is parallelizable in a trivial fashion ...



# **Correctness Checking Tools**

### **Race Condition**



#### **Data Race: the typical OpenMP programming error, when:**

- $\rightarrow$  two or more threads access the same memory location, and
- $\rightarrow$  at least one of these accesses is a write, and
- $\rightarrow$  the accesses are not protected by locks or critical regions, and
- $\rightarrow$  the accesses are not synchronized, e.g. by a barrier.
- Non-deterministic occurrence: e.g. the sequence of the execution of parallel loop iterations is non-deterministic and may change from run to run
- In many cases private clauses, barriers or critical regions are missing
- Data races are hard to find using a traditional debugger
  - → Use the Intel Inspector XE

### **Intel Inspector XE**



#### Detection of

→ Memory Errors

#### → Dead Locks

→ Data Races

### Support for

- → Linux (32bit and 64bit) and Windows (32bit and 64bit)
- → WIN32-Threads, Posix-Threads, Intel Threading Building Blocks and OpenMP

# **PI Example Code**

```
double f(double x)
{
    return (4.0 / (1.0 + x*x));
}
```

```
double CalcPi (int n)
```

{

```
const double fH = 1.0 / (double) n;
double fSum = 0.0;
double fX;
int i;
```

```
#pragma omp parallel for private(fX,i) reduction(+:fSum)
    for (i = 0; i < n; i++)
    {
        fX = fH * ((double)i + 0.5);
        fSum += f(fX);
    }
    return fH * fSum;
}</pre>
```







# **PI Example Code**

```
double f(double x)
{
    return (4.0 / (1.0 + x*x));
}
double CalcPi (int n)
{
    const double fH = 1.0 / (double) n;
    double fSum = 0.0;
```

```
double fX;
int i;
```

```
#pragma omp parallel for private(fX,i) reduction(+:fSum)
    for (i = 0; i < n; i++)
    {
        fX = fH * ((double)i + 0.5);
        fSum += f(fX);
    }
    return fH * fSum;
}</pre>
```



What if we would have forgotten this?

# **Inspector XE – Configure Analysis**

**Threading Error Analysis Modes** 

- 1. Detect Deadlocks
- 2. Detect Deadlocks and Data Races
- 3. Locate Deadlocks and Data Races



more details, more overhead

Configure Analys	sis Type		Intel Inspector XE 20
<ul> <li>Memory Error Analysis</li> <li>Å Detect Leaks</li> <li>Å Detect Memory Proble</li> </ul>	<b>Locate Deadlocks and Data Races</b> Widest scope threading error analysis type. Maximizes the load on the system. Maximizes the time required to perform the analysis. Maximizes the chances the analysis will fail because the system may run out of resources. Press F1 for more	$\langle$	Start Stop
✓ ☐ Threading Error Analysis Å Detect Deadlocks	details.	<u>_</u>	Close
A Locate Deadlocks and Custom Analysis Types	Stack frame depth: 16 Scope: Normal		

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### **Inspector XE – Results**

detected problems

1

2

filters

3 code location



The missing reduction is detected.

r001ti3 🕱		
Locate Deadlocks and Data Races	Intel Ins	spector XE 2011
🔮 🔮 Target 🛱 🏧 Analysis Type 🛛 🛃 Collection Log 📄 🧶 Summary		
Problems 2	Filters	Sort 🗸 💥 😨
ID 🔺 🎕 Problem Sources Modules State	Severity	
P1 🥝 Data race pi.c pi.exe New	Error	1 item(s)
	Problem	
	Data race	1 item(s)
	Source	
	pi.c	1 item(s)
	Module	
	pi.exe	1 item(s)
	State	
	New	1 item(s)
	Suppressed	
Code Locations Code Locations / Timeline @	Not suppressed	1 item(s)
ID Description  Source Function Module	Investigated	
▼X1 Read   pi.c:71 CalcPi pi.exe	Not investigated	1 item(s)
69 {		
70 $fX = fH * ((double)i + 0.5);$ 71 $fSum += f(fX):$		
72 }		
73 return fH * fSum;		
▼X2 Write   pi.c:71 CalcPi pi.exe		
69 {		
70 $TX = TH * ((double)1 + 0.5);$ 71 $fSum += f(fX);$		
72 }		
73 return fH * fSum;	1	
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# **PI Example Code**

```
double f(double x)
{
    return (4.0 / (1.0 + x*x));
}
double CalcPi (int n)
```

{

```
const double fH = 1.0 / (double) n;
double fSum = 0.0;
double fX;
int i;
```

```
#pragma omp parallel for private(fX,i,fSum)
    for (i = 0; i < n; i++)
    {
        fX = fH * ((double)i + 0.5);
        fSum += f(fX);
    }
    return fH * fSum;
}</pre>
```

What if we just made the variable private?



# **Inspector XE – Static Security Analysis**



At runtime no Error is detected!

Compiling with the argument "-diag-enable sc-full" delivers:

Problems						8	Filters	Sort 🗸 😚 💡		
ID	۹	Туре	Sources	ď	State	Weight 🔻	Category		Severity	
P1	8	Misuse of PRIVATE	pi.c		隆 New	100	Threading		Error	2 item(s)
5	pi.c(73):	error #12358: variabl	e "fSum" us	ed here	was last as	signed at (file	pi.c line:71) in a parallel region where it		Туре	
	was mark consider	ced PRIVATE at (file:pi.o	c line:67). PF	NATE value o	ariables ha	ave indetermin	nate value after leaving a parallel region;		Misuse of PRIVATE	1 item(s)
Ľ			copy out ius	t value o	II CAIC				Uninitialized PRIVATE	1 item(s)
P2	<b>(3)</b>	Uninitialized PRIVATE	pi.c		Rew New	100	Initialization	_	Source	
Ì	pi.c(71):	error #12361: PRIVAT	E variable "f	Sum" is	uninitializ	ed in region at	t (file:pi.c line:67).		pi.c	2 item(s)
									State	
									New	2 item(s)
Cod	e Locati	ons: Misuse of PRIV	/ATE					2	Suppressed	
Dece	ription	Fourse	Function	Variabl					Not suppressed	2 item(s)
Description Source Function Variable							-	Investigated		
	69	ywnite pi.c:71	CalCPI			Cal	lcPi - pi c:71	1 =	Not investigated	2 item(s)
	70	t fX = fH * ((do	ouble)i + 0	.5);		Cat	teri - pitet/i		Category	
	71	fSum += f(fX);							Initialization	1 item(s)
	72	}							Threading	1 item(s)
L	7.5 return TH * TSUM;									
		int is	CalcPl			(c)	lcDi pi c.67			
	66	100 1;				Cat	(CF1 - p1/C:0/			
	67 #pr	ragma omp parallel f	for private	(fX,i,f	Sum)					
	68	for (i = 0; i < n;	i++)					~		
	69	1						C)		

#### At compile-time this error can be found!

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# **Single and Master Construct**

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# **The Single Construct**





The single construct specifies that the enclosed structured block is executed by only on thread of the team.

 $\rightarrow$  It is up to the runtime which thread that is.

### **Useful for:**

 $\rightarrow$  I/O

- → Memory allocation and deallocation, etc. (in general: setup work)
- Implementation of the single-creator parallel-executor pattern as we will see now...

# **The Master Construct**



C/C++	Fortran
<pre>#pragma omp master[clause] structured block</pre>	<pre>!\$omp master[clause] structured block</pre>
	!\$omp end master

The master construct specifies that the enclosed structured block is executed only by the master thread of a team.

Note: The master construct is no worksharing construct and does not contain an implicit barrier at the end.



# **Section and Ordered Construct**

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### How to parallelize a Tree Traversal?



```
How would you parallelize this code?
void traverse (Tree *tree)
{
    if (tree->left) traverse(tree->left);
    if (tree->right) traverse(tree->right);
    process(tree);
}
```

One option: Use OpenMP's parallel sections.

# **The Sections Construct**





The sections construct contains a set of structured blocks that are to be distributed among and executed by the team of threads.

### How to parallelize a Tree Traversal?!

```
How would you parallelize this code?
  void traverse (Tree *tree)
   Ł
                                                  Nested Parallel Regions
#pragma omp parallel sections
#pragma omp section
       if (tree->left)
                          traverse(tree->left);
#pragma omp section
       if (tree->right) traverse(tree->right);
                                                          Barrier here!
} // end omp parallel
       process(tree);
```

# We will later see how this can be done with tasks in a better way.

→ Not always well supported (how many threads to be used?)

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### **The ordered Construct**



Allows to execute a structured block within a parallel loop in sequential order

→ In addition, an ordered clause has to be added to the for construct which any

ordered construct may occur

Use Cases:

- $\rightarrow$  Can be used e.g. to enforce ordering on printing of data
- $\rightarrow$  May help to determine whether there is a data race



# **Runtime Library**

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### **Runtime Library**



### **C** and C++:

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- → If OpenMP is enabled during compilation, the preprocessor symbol \_OPENMP is defined. To use the OpenMP runtime library, the header omp.h has to be included.
- > omp\_set\_num\_threads(int): The specified number of threads will be used for the parallel region encountered next.
- > int omp\_get\_num\_threads: Returns the number of threads in the
  current team.
- > int omp\_get\_thread\_num(): Returns the number of the calling thread
  in the team, the Master has always the id 0.

# Additional functions are available, e.g. to provide locking functionality.



# **Tasking**

### Recursive approach to compute Fibonacci



```
int fib(int n) {
    if (n < 2) return n;
    int x = fib(n - 1);
    int y = fib(n - 2);
    return x+y;
}</pre>
```

On the following slides we will discuss three approaches to parallelize this recursive code with Tasking.

# The Task Construct



C/C++	Fortran
<pre>#pragma omp task [clause]</pre>	!\$omp task [clause]
structured block	structured block
	] !\$omp end task

#### Each encountering thread/task creates a new Task

- → Code and data is being packaged up
- $\rightarrow$  Tasks can be nested
  - →Into another Task directive
  - →Into a Worksharing construct

#### Data scoping clauses:

- → shared(*list*)
- > private(list) firstprivate(list)
- → default(shared | none)

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# Tasks in OpenMP: Data Scoping



#### Some rules from *Parallel Regions* apply:

- → Static and Global variables are shared
- → Automatic Storage (local) variables are private

### If shared scoping is not derived by default:

- Orphaned Task variables are firstprivate by default!
- Non-Orphaned Task variables inherit the shared attribute!
- > Variables are firstprivate unless shared in the enclosing context

# First version parallelized with Tasking (omp-v1)



```
int fib(int n) {
int main(int argc,
                                        if (n < 2) return n;
         char* argv[])
                                        int x, y;
{
                                        #pragma omp task shared(x)
   [...]
                                        Ł
  #pragma omp parallel
                                             x = fib(n - 1);
   {
       #pragma omp single
                                        #pragma omp task shared(y)
        {
                fib(input);
                                             v = fib(n - 2);
        }
   }
                                        #pragma omp taskwait
   [...]
                                             return x+y;
}
```

- Only one Task / Thread enters fib() from main(), it is responsable for creating the two initial work tasks
- $\circ$  Taskwait is required, as otherwise  ${\bf x}$  and  ${\bf y}$  would be lost

# **Fibonacci Illustration**

- T1 enters fib(4)
- T1 creates tasks for fib(3) and fib(2)
- T1 and T2 execute tasks from the queue
- T1 and T2 create 4 new tasks
- T1 T4 execute tasks





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# **Fibonacci Illustration**

- T1 enters fib(4)
- T1 creates tasks for fib(3) and fib(2)
- T1 and T2 execute tasks from the queue
- T1 and T2 create 4 new tasks
- T1 T4 execute tasks



### **Scalability measurements (1/3)**



#### Overhead of task creation prevents better scalability!



Speedup of Fibonacci with Tasks

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# Improved parallelization with Tasking (omp-v2)



# Improvement: Don't create yet another task once a certain (small enough) n is reached

```
int main(int argc,
         char* argv[])
{
   [...]
#pragma omp parallel
Ł
#pragma omp single
{
   fib(input);
}
}
   [...]
}
```

```
int fib(int n) {
   if (n < 2) return n;
int x, y;
\#pragma omp task shared(x) \setminus
  if(n > 30)
{
   x = fib(n - 1);
}
#pragma omp task shared(y) \
  if(n > 30)
ł
   y = fib(n - 2);
#pragma omp taskwait
   return x+y;
```

### Scalability measurements (2/3)



Speedup is ok, but we still have some overhead when running with 4 or 8 threads



Speedup of Fibonacci with Tasks

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# Improved parallelization with Tasking (omp-v3)



Improvement: Skip the OpenMP overhead once a certain n is reached (no issue w/ production compilers)

```
int main(int argc,
          char* argv[])
{
   [...]
#pragma omp parallel
#pragma omp single
{
   fib(input);
}
}
   [...]
```

```
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```

```
int fib(int n) {
   if (n < 2) return n;
   if (n <= 30)
      return serfib(n);
int x, y;
#pragma omp task shared(x)
{
  x = fib(n - 1);
}
#pragma omp task shared(y)
ł
  v = fib(n - 2);
#pragma omp taskwait
   return x+y;
```

**Scalability measurements (3/3)** 



#### Everything ok now ③



#### Speedup of Fibonacci with Tasks

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# Data Scoping Example (1/7)



```
int a = 1;
void foo()
{
   int b = 2, c = 3;
   #pragma omp parallel shared(b)
   #pragma omp parallel private(b)
   {
        int d = 4;
       #pragma omp task
        {
                int e = 5;
                // Scope of a:
                // Scope of b:
                // Scope of c:
                // Scope of d:
                // Scope of e:
```

# Data Scoping Example (2/7)



```
int a = 1;
void foo()
{
   int b = 2, c = 3;
   #pragma omp parallel shared(b)
   #pragma omp parallel private(b)
   {
        int d = 4;
        #pragma omp task
        {
                int e = 5;
                // Scope of a: shared
                // Scope of b:
                // Scope of c:
                // Scope of d:
                // Scope of e:
```

} } } 60 Introduction to OpenMP Dirk Schmidl | IT Center der RWTH Aachen University

# Data Scoping Example (3/7)



```
int a = 1;
void foo()
{
   int b = 2, c = 3;
   #pragma omp parallel shared(b)
   #pragma omp parallel private(b)
   {
        int d = 4;
        #pragma omp task
        {
               int e = 5;
               // Scope of a: shared
               // Scope of b: firstprivate
               // Scope of c:
               // Scope of d:
               // Scope of e:
```

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# Data Scoping Example (4/7)



```
int a = 1;
void foo()
{
   int b = 2, c = 3;
   #pragma omp parallel shared(b)
   #pragma omp parallel private(b)
   {
       int d = 4;
       #pragma omp task
        {
               int e = 5;
               // Scope of a: shared
               // Scope of b: firstprivate
               // Scope of c: shared
               // Scope of d:
               // Scope of e:
```

# Data Scoping Example (5/7)



```
int a = 1;
void foo()
{
   int b = 2, c = 3;
   #pragma omp parallel shared(b)
   #pragma omp parallel private(b)
   {
       int d = 4;
       #pragma omp task
        {
               int e = 5;
               // Scope of a: shared
               // Scope of b: firstprivate
               // Scope of c: shared
               // Scope of d: firstprivate
               // Scope of e:
```

### Data Scoping Example (6/7)

```
int a = 1;
void foo()
{
  int b = 2, c = 3;
   #pragma omp parallel shared(b)
  #pragma omp parallel private(b)
   Ł
       int d = 4;
       #pragma omp task
        {
               int e = 5;
               // Scope of a: shared
               // Scope of b: firstprivate
               // Scope of c: shared
               // Scope of d: firstprivate
               // Scope of e: private
```



Hint: Use default(none) to be forced to think about every variable if you do not see clear.

# Data Scoping Example (7/7)



```
int a = 1;
void foo()
{
  int b = 2, c = 3;
  #pragma omp parallel shared(b)
  #pragma omp parallel private(b)
   {
       int d = 4;
       #pragma omp task
       {
              int e = 5;
              // Scope of a: shared,
                                           value of a: 1
              // Scope of b: firstprivate, value of b: 0 / undefined
              // Scope of c: shared,
                                         value of c: 3
              // Scope of d: firstprivate, value of d: 4
              // Scope of e: private, value of e: 5
```



### OpenMP barrier (implicit or explicit)

→ All tasks created by any thread of the current *Team* are guaranteed to be completed at barrier exit

C/C++

#pragma omp barrier

#### Task barrier: taskwait

→ Encountering Task suspends until child tasks are complete

 $\rightarrow$ Only direct childs, not descendants!

C/C++

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#pragma omp taskwait

### **Task Synchronization**



Task Synchronization explained:





# **More Environment Variables**

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- OMP\_NUM\_THREADS: Controls how many threads will be used to execute the program.
- OMP\_SCHEDULE: If the schedule-type runtime is specified in a schedule clause, the value specified in this environment variable will be used.
- OMP\_DYNAMIC: The OpenMP runtime is allowed to smartly guess how many threads might deliver the best performance. If you want full control, set this variable to *false*.
- OMP\_NESTED: Most OpenMP implementations require this to be set to *true* in order to enabled nested Parallel Regions. Remember: Nesting Worksharing constructs is not possible.



### **Define interaction with system environment:**

- → Env. Var. OMP\_MAX\_NESTED\_LEVEL + API functions
  - $\rightarrow$  Controls the maximum number of active parallel regions
- → Env. Var. OMP\_THREAD\_LIMIT + API functions

→Controls the maximum number of OpenMP threads

- → Env. Var. OMP\_STACKSIZE
  - →Controls the stack size of child threads
- → Env. Var. OMP\_WAIT\_POLICY

 $\rightarrow$  Control the thread idle policy:

 $\rightarrow$  active: Good for dedicated systems (e.g. in batch mode)

→passive: Good for shared systems



# **Questions?**