

# Introduction to OpenMP

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# Outline

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

## 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

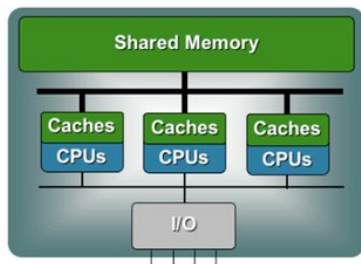
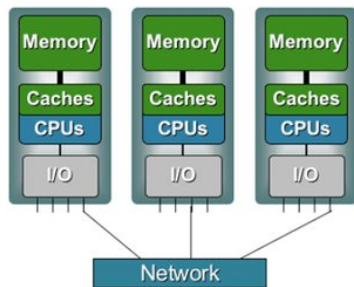
## What is OpenMP?

- 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

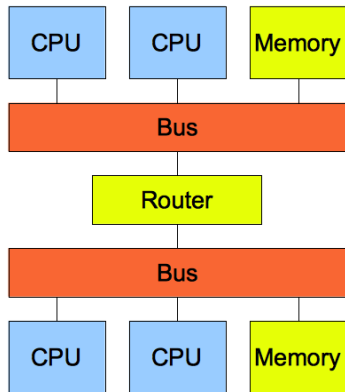
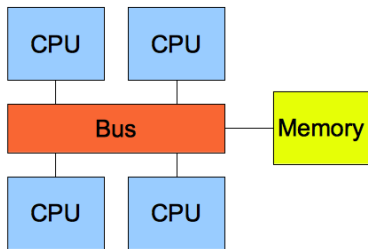
# History

- 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 (task)
- **The present**
  - July 2011 - version 3
  - July 2013 - version 4.0 (Accelerator, SIMD extensions, Affinity, Error handling, User-defined reduction)
- **The future**
  - version 4.1/4.5/5.0

## Distributed and shared memory

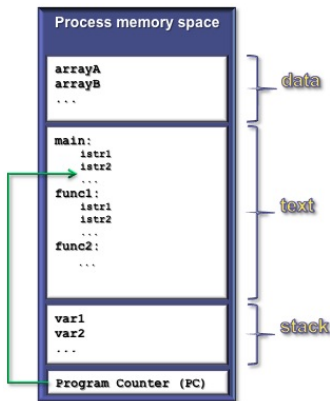


## UMA and NUMA systems



## Process and thread

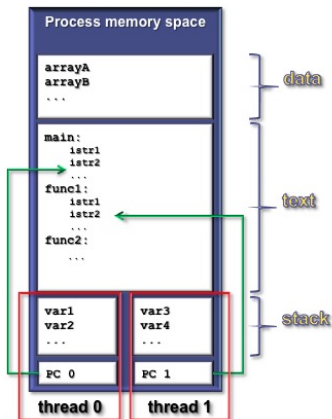
- A process is an instance of a computer program
- Some information included in a process are:
  - Text
    - Machine code
  - Data
    - Global variables
  - Stack
    - Local variables
  - Program counter (PC)
    - A pointer to the instruction to be executed



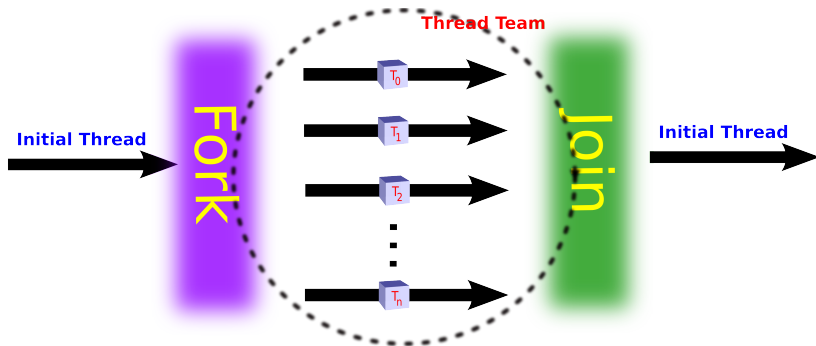


# Multi-threaded processes

- 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?

## ① Standardized

- enhance **portability**

## ② Lean and mean

- **limited set** of directives
- **fast** code parallelization

## ③ Ease of use

- parallelization is **incremental**
- coarse / fine parallelism

## ④ Portability

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

## ① Performance

- may be non-portable
- increase memory traffic

## ② Limitations

- shared memory systems
- mainly used for **loops**

# Structure of an OpenMP program

## ① 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

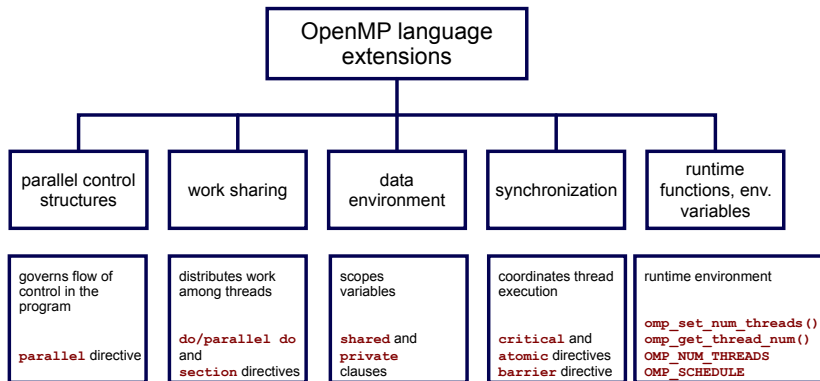
## ② Shared-memory model

- 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

## ③ 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 `yyyyymm`
- 2 Fortran 77 supports `!$, *$` and `c$` as sentinels
- 3 Fortran 90 supports `!$` only

## Directive format

### C/C++

```
#pragma omp directive-name [clause...]
```

### Fortran

```
sentinel directive-name [clause...]
```

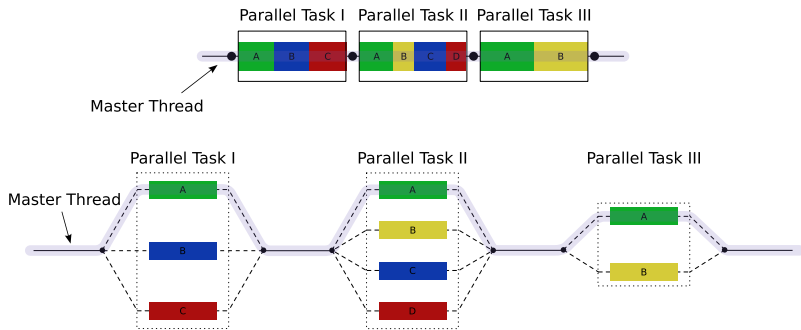
- 1 Follows conventions of C and C++ compiler directives
- 2 From here on free-form directives will be considered

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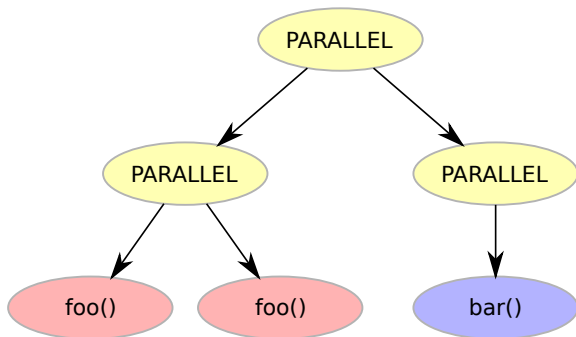


# parallel construct



- 1 The encountering thread becomes the **master** of the new team
- 2 All threads execute the parallel region
- 3 There is an **implied barrier** at the end of the parallel region

## Nested parallelism



- 1 Nested parallelism is allowed in OpenMP 3.1
- 2 Most constructs bind to the **innermost parallel region**

# OpenMP: Hello world

C/C++

```
int main () {  
  
    printf("Hello world\n");  
  
    return 0;  
}
```

# OpenMP: Hello world

C/C++

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

# OpenMP: Hello world

## Fortran

```
PROGRAM HELLO
```

```
Print *, "Hello World!!!"
```

```
END PROGRAM HELLO
```

# OpenMP: Hello world

## Fortran

```
PROGRAM HELLO
! Serial code

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

! Resume serial code

END PROGRAM HELLO
```

# OpenMP: Hello world

## What's wrong?

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

# Race condition

- A **race condition** (or data race) is when two or more threads access the same memory location:
  - asynchronously 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**



# Worksharing constructs

- 1 **Distribute the execution** of the associated region
- 2 A worksharing region has **no barrier** on entry
- 3 An **implied barrier** exists at the end, unless `nowait` is present
- 4 Each region **must** be encountered by all threads or none
  - Every thread must encounter the **same sequence** of worksharing regions and barrier regions
- 5 The OpenMP API defines **four worksharing** constructs:
  - **loop** construct
  - **sections** construct
  - **single** construct
  - **workshare** construct

## Loop construct: syntax

### C/C++

```
#pragma omp for [clause[[,] clause] ... ]  
for-loops
```

### Fortran

```
!$omp do [clause[[,] clause] ... ]  
do-loops  
[!$omp end do [nowait] ]
```

## Loop construct: restrictions

### C/C++

```
for (init-expr; test-expr; incr-expr)  
    structured-block
```

```
init-expr:    var = lb  
              integer-type var = lb
```

```
test-expr:    relational expr.
```

```
incr-expr:    addition or subtraction expr.
```

## Loop construct: the rules

- 1 The iterations of the loop are **distributed** over the threads that already exist in the team
- 2 The inner loops are executed sequentially by each thread
- 3 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
- 4 The `schedule` clause:
  - may be used to specify **how** iterations are divided into chunks
- 5 The `collapse` clause:
  - may be used to specify how many loops are parallelized
  - valid values are constant positive integer expressions

## Loop construct: scheduling

### C/C++

```
#pragma omp for schedule(kind[, chunk_size])  
for-loops
```

### Fortran

```
!$omp do schedule(kind[, chunk_size])  
do-loops  
[!$omp end do [nowait] ]
```

## Loop construct: schedule kind

### 1 static

- iterations are divided into chunks of size `chunk_size`
- the chunks are assigned to the threads in a **round-robin** fashion
- must be **reproducible** within the same parallel region

### 2 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

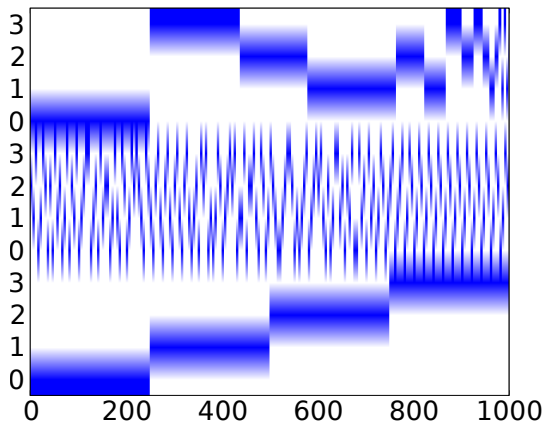
### 3 guided

- 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
- Special case of dynamic to reduce scheduling overhead

### 4 auto

- When the runtime can "learn" from previous executions of the same loop

## Loop construct: schedule kind



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

## Loop construct: nowait clause

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++)
        b[i] = (a[i] + a[i-1]) / 2.0;
#pragma omp for nowait
    for (int i=0; i<m; i++)
        y[i] = sqrt(z[i]);
}
}
```



## 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++)
        c[i] = (a[i] + b[i]) / 2.0f;
#pragma omp for schedule(static) nowait
    for (i=0; i<n; i++)
        z[i] = sqrtf(c[i]);
#pragma omp for schedule(static) nowait
    for (i=1; i<=n; i++)
        y[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;
  }
}
}
```

## 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;
    }
  }
}
```

## Loop collapse

- 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  $N \times M$ ) and then parallelizes it
- Useful if  $N < \text{no. of threads}$ , so parallelizing the outer loop makes balancing the load difficult.

## Loop dependencies

### loop carried dependencies

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

<i>i</i>	<i>j</i>
0	3+2
1	3+4
2	3+6
...	...
<i>n</i>	$3+(2*n+2)$

# Removing loop dependencies

## removing 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

### C/C++

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

## Sections construct: syntax

### Fortran

```
!$omp sections [clause[[,] clause]...]  
!$omp section  
    structured-block  
!$omp section  
    structured-block  
...  
!$omp end do [nowait]
```



## Sections construct: some facts

- 1 `sections` is a non-iterative worksharing construct
  - it contains a set of `structured-blocks`
  - each one is executed **once** by one of the threads
- 2 Scheduling of the sections is **implementation defined**
- 3 There is an implied barrier at the end of the construct

## Single construct: syntax

### C/C++

```
#pragma omp single [clause[[,] clause]...]
    structured-block
```

### Fortran

```
!$omp single [clause[[,] clause] ... ]
    structured-block
[!$omp end single [nowait] ]
```

## Single construct: some facts

- 1 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**:

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

## Reduction clause: some facts

- 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.
- 4 Items must not be `const`-qualified

## Reduction clause: example

### 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);
```

## Reduction clause: example

### Fortran features?

```
PROGRAM REDUCTION_WRONG
  MAX = HUGE(0)
  M = 0
  !$OMP PARALLEL DO REDUCTION(MAX: M)
  DO I = 1, 100
    CALL SUB(M, I)
  END DO
END PROGRAM REDUCTION_WRONG
```

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## Master construct: syntax

### C/C++

```
#pragma omp master  
    structured-block
```

### Fortran

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

## Master construct: some facts

- 1 The `master` construct specifies a structured block:
  - that is **executed by the master** thread
  - with **no implied barrier** on entry or exit
- 2 Used 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]
```

## Critical construct: some facts

- 1 The `critical` construct restricts the execution:
  - to a single thread at a time (wait on entry)
  - disregarding team information
- 2 An optional `name` may be used to identify a region
- 3 All `critical` without a name `share` the same unspecified tag
- 4 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();
}
```

## Barrier construct: syntax

### C/C++

```
#pragma omp barrier
```

### Fortran

```
!$omp barrier
```

The `barrier` construct specifies an **explicit barrier** at the point at which the construct appears

## Barrier construct: example

### Waiting for the master to come

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

## Atomic construct: syntax

### C/C++

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

```
#pragma omp atomic capture  
    structured-block
```



# Atomic construct: syntax

## Fortran

```
!$omp atomic read  
  capture-statement  
[!$omp end atomic]
```

```
!$omp atomic write  
  write-statement  
[!$omp end atomic]
```

# Atomic construct: syntax

## Fortran

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

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

## Atomic construct: some facts

### 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

### 4 Accesses to the same location must have **compatible** types

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## Data-sharing attributes: C/C++

- 1 The following are always **shared**:
  - 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
- 3 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

## Data-sharing clauses: syntax

### C/C++

```
#pragma omp directive-name [clause[[,]clause]
```

### Fortran

```
!$omp directive-name [clause[[,]clause]
```

```
...
```

```
!$omp end directive-name [clause]
```

# 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` construct
  - declares one or more list items to be shared
- 3 The clause `private(list)`:
  - is valid on `parallel`, and `worksharing` constructs
  - declares one or more list items to be private
  - allocates a new item of the same type with undefined value



## 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);
}
```

## 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

# 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];
```

- 1 valid on for, sections
- 2 the value of each new list item is the sequentially last value

# Copyprivate clause

## C/C++

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

- 1 Valid only on `single`
- 2 Broadcasts 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

## 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
```

- 1 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

- OMP\_NUM\_THREAD** sets the number of threads for parallel regions
- OMP\_STACKSIZE** specifies the size of the stack for threads
- OMP\_SCHEDULE** controls schedule type and chunk size of `runtime`
- OMP\_PROC\_BIND** controls whether threads are bound to processors
- OMP\_NESTED** enables or disables nested parallelism



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# OpenMP Compilers

## 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.

## Sun Microsystems:

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

# OpenMP Compilers

## Intel:

Compile with **-Qopenmp** on Windows, or just **-openmp** on Linux or Mac Emit useful information to stderr.

**-openmp-report2**

## Portland Group Compilers:

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

# OpenMP: THE END!!!

Good luck and enjoy OpenMP!!!