OpenMP Shared-Memory Parallel Programming

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Motivations for Parallel Computing



Year

Motivations for Parallel Computing

- In the years 2000's the CPU manufacturers have run out of room for boosting CPU performance.
- Instead of driving clock speeds and straight-line instruction throughput higher, they turn to hyperthreading and multicore architectures.

The parallel programming model became necessary.

- The MPI standard was introduced by the MPI Forum in May 1994 and updated in June 1995.
- The OpenMP standard was introduced in 1997 (Fortran) and 1998 (C/C++).

Distributed-Memory



- Multiple nodes
- Interconnected by a high-speed network
- Nodes consist of (a) processor(s) and local memory
- Communication is done via message passing

Process



- A process is an instance of an application
- A process is executed by at least one thread
- A process is a container describing the state of an application: code, memory mapping, shared libraries,

In scientific computing, the dominant paradigm for process parallelism is the single program multiple data model with MPI.

. . .

Shared-Memory



- At least one multi-core CPU
- All CPUs can access a single memory address space
- Systems memory may be physically distributed, but logically shared

Threads



- A thread is an independent stream of instructions that can be scheduled to run by the operating system.
- Multiple threads can exist within one process, and they share the memory
- A thread only the owns the bare essential resources to exist as an executable code: execution counter, stack pointer, registers and thread-specific data

In scientific computing the dominant paradigm for thread parallelism is OpenMP

What is OpenMP?

OpenMP is a shared-memory application programming interface which by adding directives to a sequential program describes how the work is shared among threads and order accesses to shared data.

OpenMP hides the low-level details and allows the programmer to describe the parallel code with high-level constructs.

Why use threads?

The operating system does not need to create a new memory map for a new thread. It increases efficiency on multiprocessor systems.

Shared memory makes it trivial to share data among threads (with potential drawbacks though).

Fork-Join

OpenMP use the fork-join model



- The master thread continues after the fork operation
- The children threads begin operation separate from the master thread
- Parallel execution begins

Fork-Join

OpenMP use the fork join model



- Children join after they finish
- The master thread waits until all the children join
- Join ends parallel execution. Sequential execution of the master thread continues



In practice, threads are not created or destroyed for every parallel region.

OpenMP implementations use a thread pool to avoid the cost of thread creation and destruction at each fork and join.

After the join, the children thread go idle.

OpenMP is using directives

Directives are programming language constructs that specifies how a compiler should process its input

An OpenMP program is the combination of

- a base language (C, C++ or Fortran)
- annotations with OpenMP directives

OpenMP directive in C/C++

#pragma omp construct [clauses]

OpenMP directive in Fortran



OpenMP directive in C/C++



OpenMP directive in Fortran



OpenMP directive in C/C++

#pragma omp construct [clauses]

Give instruction on what to do

OpenMP directive in Fortran

!\$omp construct [clauses]

OpenMP directive in C/C++

Additional options (optional) #pragma omp construct [clauses]

OpenMP directive in Fortran



What the directives do

An OpenMP construct can specify

- the creation of a parallel region
- how to parallelize loops
- whether the variables in the parallel region are private or shared
- how/if the threads are synchronized
- how the work is divided between threads

The Advantages of Using directive

- Does not modify the serial implementation
- You can still compile and run the program as a serial code.
- Can be added incrementally allowing a gradual parallelization
- Easier to maintain

Hard work is hidden

Directives hide the actual parallelization work from the programmer

The compiler replaces the directives by the appropriate calls to the OpenMP runtime and library

Going Parallel

The Parallel Construct

Parallel construct in C/C++

Creates a parallel region by spawning a team of threads ↓ #pragma omp parallel [clauses] structured-block

Parallel construct in Fortran

!\$omp parallel [clauses]
 structured-block
!\$omp end parallel

The Parallel Construct



Parallel construct in Fortran

!\$omp parallel [clauses]
 structured-block
!\$omp end parallel

The Parallel Construct

Parallel construct in C/C++



OpenMP Hello World

```
#include <stdio.h>
#include <omp.h>
                                                                     program main
int main(int argc, char* argv[]) {
                                                                       use omp lib
  #pragma omp parallel
                                                                       !$omp parallel
                                                                         print 100, omp get thread num(), &
    printf("Hello, I'm thread %d of %d.\n",
                                                                 &
                                                                                     omp get num threads()
            omp get thread num(),
                                                                 100
                                                                         format('Hello, I am thread ', i0, ' of ', i0)
            omp get num threads());
  }
                                                                       !somp end parallel
                                                                     end program
  return 0;
}
```

- Include omp.h or use the omp_lib module to get access to the OpenMP runtime library
- Use the omp_get_thread_num to get the ID of the thread in the team and omp_get_num_threads functions to get the number of threads in the team

Compiling the OpenMP Hello World

To compile an OpenMP program, you need to pass a specific flag to the compiler

GCC:	gcc	-fopenmp
Clang:	clang	-fopenmp
Intel:	icc	-qopenmp

This flag instructs the compiler to consider OpenMP directives

Compiling the OpenMP Hello World

Compilers with OpenMP are available for all CÉCI clusters. For example, the following modules are available:

Lemaitre3	GCC/8.3.0	intel/2019b
Hercules2	GCC/7.1.0-2.28	intel/2016b
Dragon2	GCC/8.2.0-2.31.1	intel/2018b
Vega	GCC/9.3.0	intel/2019b

\$ module load <module_name>

Executing the OpenMP Hello World

\$ gcc -fopenmp -o example omp_helloworld.c
\$ OMP_NUM_THREADS=4 ./example
Hello, I'm thread 1 of 4.
Hello, I'm thread 2 of 4.
Hello, I'm thread 3 of 4.
Hello, I'm thread 0 of 4.

Executing the OpenMP Hello World

The OMP_NUM_THREADS environment variable allows you to specify the number of threads

- \$ OMP_NUM_THREADS=4 ./example ← ^{4 threads for this} execution of the program

Submitting an OpenMP Job

When submitting your OpenMP job to one of the CÉCI clusters set **cpus-per-task** to specify the number of threads.

```
#!/bin/bash
# Basic submission script for an openmp job
#SBATCH --time=01:00:00
#SBATCH --ntasks=1
#SBATCH --cpus-per-task=4
#SBATCH --mem-per-cpu=1024
```

export OMP_NUM_THREADS=\$SLURM_CPUS_PER_TASK

```
module load GCCcore
```

```
./your_omp_app
```

Making Things Go Parallel

Creating a parallel region does not means that that the work will be shared among the threads. For example, if we consider this piece of code:

```
max threads = omp_get_max_threads()
      allocate(iterations(0:max threads-1))
      iterations = 0
      !somp parallel private(tid)
        tid = omp get thread num()
        do i = 1.1000
          iterations(tid) = iterations(tid) + 1
        end do
      !somp end parallel
      do i = 0, max threads-1
        print 100, i, iterations(i)
100
        format('Number of iteration for thread ', i0, &
               ': ', i0)
&
      end do
```

Making Things Go Parallel

\$ gcc -fopenmp -o example omp_iterations.c
\$ OMP_NUM_THREADS=4 ./example
Number of iteration for thread 0: 1000
Number of iteration for thread 1: 1000
Number of iteration for thread 2: 1000
Number of iteration for thread 3: 1000

$\textbf{Parallel} \neq \textbf{Worksharing}$

In the last example there is no worksharing. This means that all the threads execute all the iterations of the loop.

The parallel construct means that

- a team of threads is created, i.e. there is a fork
- the code is executed redundantly by each thread
- the threads in the team join at the end of the region

Distributing iterations

One of the options for sharing the work between the threads is to define lower and higher bounds of the loop depending on the thread ID.

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```
int max threads = omp get max threads():
int* iterations = malloc(sizeof(int)*max threads);
for(int i = 0: i < max threads: ++i)</pre>
 iterations[i] = 0;
#pragma omp parallel
  int
           tid = omp get thread num();
  int nthreads = omp get num threads():
  int low = n * tid / nthreads;
  int high = n * (tid + 1) / nthreads:
  for(int i = low; i < high; ++i)</pre>
    iterations[tid]++;
}
for(int i = 0; i < max threads; ++i)</pre>
 printf("Number of iteration for thread %d: %d\n".
          i. iterations[i]):
```

```
max threads = omp get max threads()
     allocate(iterations(0:max threads-1))
      iterations = 0
     !somp parallel private(tid)
       tid = omp get thread num()
       nthreads = omp get num threads()
        low = n * tid / nthreads + 1
       high = n * (tid + 1) / nthreads
       do i = low, high
         iterations(tid) = iterations(tid) + 1
        end do
      !somp end parallel
     do i = 0, max threads-1
       print 100, i, iterations(i)
100
       format('Number of iteration for thread ', i0, &
              ': ', i0)
     end do
```

Distributing iterations

\$ gcc -fopenmp -o example omp_iterations.c
\$ OMP_NUM_THREADS=4 ./example
Number of iteration for thread 0: 250
Number of iteration for thread 1: 250
Number of iteration for thread 2: 250
Number of iteration for thread 3: 250

Distributing iterations with a directive

Instead of computing the bounds, we can use the **for** (or **do**) construct.

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```
int max threads = omp get max threads():
int* iterations = malloc(sizeof(int)*max threads);
for(int i = 0: i < max threads: ++i)</pre>
 iterations[i] = 0;
#pragma omp parallel
 #pragma omp for
 for(int i = 0; i < n; ++i)
    iterations[tid]++;
}
for(int i = 0; i < max_threads; ++i)</pre>
 printf("Number of iteration for thread %d: %d\n",
          i, iterations[i]);
```

```
max threads = omp get max threads()
allocate(iterations(0:max threads-1))
```

```
iterations = 0
      !somp parallel private(tid)
        !$omp do
         do i = 1, n
            iterations(tid) = iterations(tid) + 1
          end do
        !somp end do
      !somp end parallel
      do i = 0, max threads-1
        print 100, i, iterations(i)
100
        format('Number of iteration for thread ', i0, &
               ': ', i0)
      end do
```
Distributing iterations with a directive

\$ gcc -fopenmp -o example omp_for_iters.c
\$ OMP_NUM_THREADS=4 ./example
Number of iteration for thread 0: 250
Number of iteration for thread 1: 250
Number of iteration for thread 2: 250
Number of iteration for thread 3: 250

The Canonical for-loop

The for-loop needs to be in canonical form to be used with the for directive

var, can not be modified in the loop body. It must be an integer (signed or unsigned), a pointer or random access iterator type

start, end and incr must be loop invariant expressions, the number of iterations must be computable when the loop begins

Parallel Region Binding

In order for the iterations of a loop to be shared among the threads by a **for** /do , the construct needs a parallel region to bind to. If we take the previous example and remove the parallel region:

```
int max_threads = omp_get_max_threads();
int* iterations = malloc(sizeof(int)*max_threads);
for(int i = 0; i < max_threads; ++i)
iterations[i] = 0;
#pragma omp for
for(int i = 0; i < n; ++i)
iterations[tid]++;
for(int i = 0; i < max_threads; ++i)
printf("Number of iteration for thread %d: %d\n",
i, iterations[i]);
```

Parallel Region Binding

```
$ OMP_NUM_THREADS=4 ./example
Number of iteration for thread 0: 1000
Number of iteration for thread 1: 0
Number of iteration for thread 2: 0
Number of iteration for thread 3: 0
```

As there was no parallel region to bind to, the **for** /do construct binds to the master thread.

Combined Directive

The following code snippet,

```
#pragma omp parallel
{
    #pragma omp for
    for(int i = 0; i < n; ++i)
        do_something()
}</pre>
```

```
!$omp parallel
    !$omp do
        do i = 1,n
            call do_something()
        end do
    !$omp end do
!$omp end parallel
```

may also be written as combined **parallel** and **for** directives

```
#pragma omp parallel for
for(int i = 0; i < n; ++i)
    do_something();
```

```
!$omp parallel do
   do i = 1,n
      call do_something()
   end do
!$omp end parallel do
```



- Directives are active in the dynamic scope of a parallel region, not just its lexical scope. This allows for orphaned directives.
- Orphaning is a situation when directives related to a parallel region are outside the lexical extent of the parallel region.
- Typical situation is calling a function containing a worksharing directive from a parallel region.

Orphaning Example

```
void ax(int n, double alpha, double* x) {
  int nthreads = omp_get_num_threads();
  int tid = omp get thread num();
  printf("Executing ax by thread %d of %d threads.\n", tid.
         nthreads);
  int niters = 0:
  #pragma omp for
  for (int i = 0: i < n: ++i) {</pre>
    x[i] = alpha * x[i];
    niters++;
  printf("Thread with id %d did %d iterations.\n", tid,
        niters);
}
int main ( int argc, char *argv[] ) {
  #pragma omp parallel
    ax(n, 3.0, x):
  ax(n, 5.0, y);
```

```
!$omp parallel
 call ax(n, 3.0d0, x)
!somp end parallel
```

call ax(n, 5.0d0, y)

100

&

```
contains
 subroutine ax(n, alpha, x)
   print 100, tid, nthreads
    format('Executing ax by thread ', i0,
           ' of ', i0, ' threads.')
```

```
!$omp do
 do i = 1.n
    x(i) = alpha * x(i)
    niters = niters + 1
  end do
!$omp end do
```

	print 200, tid, niters
200	format('Thread with id ', i0, &
&	' did ', i0, ' iterations.')

Orphaning Example

\$ gcc -fopenmp -o example omp orphaned.c \$ OMP NUM THREADS=4 ./example Executing ax by thread 0 of 4 threads. Executing ax by thread 2 of 4 threads. Executing ax by thread 1 of 4 threads. Executing ax by thread 3 of 4 threads. Thread with id 0 did 250 iterations. Thread with id 1 did 250 iterations. Thread with id 2 did 250 iterations. Thread with id 3 did 250 iterations. Executing ax by thread 0 of 1 threads. Thread with id 0 did 1000 iterations.

Nested Parallelism

This code snippet is not a valid OpenMP code: work-sharing region may not be closely nested inside of an other work-sharing region

```
#pragma omp parallel for
for (int i = 0; i < n; ++i)
#pragma omp for
for (int j = 0; j < n; ++j)
do_something();
}
```

```
!$omp parallel do
do i = 1,n
  !$omp do
  do j = 1,n
      call do_something()
      end do
  !$omp end do
end do
!$omp end parallel do
```

However, inclosing this nested work-sharing contruct inside a nested parallel region is valid.

```
#pragma omp parallel for
for (int i = 0; i < n; ++i)
#pragma omp parallel for
for (int j = 0; j < n; ++j)
do_something();
}
```

```
!$omp parallel do
 do i = 1,n
  !$omp parallel do
    do j = 1,n
        call do_something()
        end do
  !$omp end parallel do
!$omp end parallel do
```

Nested Parallelism

OpenMP parallel regions can be nested inside each other but it is disabled by default, meaning that

- If nested parallelism is disabled, then the new team created by a thread encountering a parallel construct inside a parallel region consists only of the encountering thread. This is the default.
- If nested parallelism is enabled, then the new team may consist of more than one thread (OMP_NESTED=TRUE).
- The maximum level of nested parallelism can be set by the OMP_MAX_ACTIVE_LEVELS environment variable.

Nested Parallelism

```
void report_num_threads(int level) {
    #pragma omp single
    printf("Level %d - number of threads: %d\n",
        level, omp_get_num_threads());
}
```

```
#pragma omp parallel num_threads(2)
{
    report_num_threads(1);
    #pragma omp parallel num_threads(2)
    {
        report_num_threads(2);
        #pragma omp parallel num_threads(2)
        {
            report_num_threads(3);
        }
    }
}
```

```
!$omp parallel num_threads(2)
call report_num_threads(1)
!$omp parallel num_threads(2)
call report_num_threads(2)
!$omp parallel num_threads(2)
call report_num_threads(3)
!$omp end parallel
!$omp end parallel
!$omp end parallel
```

```
contains
subroutine report_num_threads(level)
integer, intent(in) :: level
```

integer :: nthreads

```
nthreads = omp_get_num_threads()
```

```
!$omp single
    print 100, level, nthreads
100 format('Level ', i0, ': the number of threads
in the team is ', i0)
    !$omp end single
end subroutine
```

Nested Parallelism : Disabled

\$ gcc -fopenmp -o example omp_nested.c
\$ OMP_NUM_THREADS=4 ./example
Level 1: number of threads in the team - 2
Level 2: number of threads in the team - 1
Level 3: number of threads in the team - 1
Level 2: number of threads in the team - 1
Level 3: number of threads in the team - 1

Nested Parallelism: Enabled

- \$ gcc -fopenmp -o example example omp_nested.c
- \$ OMP_NUM_THREADS=4 OMP_NESTED=TRUE \

./example

Level 1: number of threads in the team - 2

- Level 2: number of threads in the team 2
- Level 2: number of threads in the team 2
- Level 3: number of threads in the team 2
- Level 3: number of threads in the team 2
- Level 3: number of threads in the team 2
- Level 3: number of threads in the team 2

Closely Nested Loops

Even if, nested parallelism is possible, try to avoid it: nesting parallel regions can easily create too many threads and oversubscribe the system.

Most of the time, parallelizing the outer loop is enough. Then, you have to be careful to have the inner loop to access consecutive elements in memory in order to maximize cache use.

Loop collapsing

In some cases, you can collaspe the loops into one in order to increase the run trip of the loop.

= 1.n

j) = do something()

```
parallel do collapse(2)
#pragma omp parallel for collapse(2)
                                                                   do i
                                                                       = 1.3
for (int i = 0; i < 3; ++i) {
  for (int j = 0; j < n; ++j) {
    a[i][j] = do something();
                                                                     end do
}
                                                                       end parallel do
```

This is particularly useful when one of the loops is not of sufficient length to have efficient parallelization.

Loop collapsing

The **collapse** clause, collapse the iterations of the n-associated loops to which the clause applies into one larger iteration space. This clause can only apply on tightly nested loops, meaning that there is no code between the loops.

#pragma omp for collapse(n)
 nested-for-loops

!\$omp do collapse(n)
 nested-do-loops

Data Sharing in a Parallel World



Let's go back to the hello world code:

```
#include <stdio.h>
                                                                    program main
#include <omp.h>
                                                                       use omp lib
int main(int argc, char* argv[]) {
                                                                       implicit none
  int nthreads, tid;
                                                                       integer :: nthreads, tid
  #pragma omp parallel
                                                                       !$omp parallel
    tid = omp get thread num();
                                                                        tid = omp_get_thread_num()
    nthreads = omp_get_num_threads();
    printf("Hello, I'm thread %d of %d.\n", tid, nthreads);
                                                                        print 100, tid, nthreads
                                                                       !somp end parallel
  return 0;
                                                                    end program
```

```
implicit none
integer :: nthreads, tid
!$omp parallel
  tid = omp_get_thread_num()
    nthreads = omp_get_num_threads()
    print 100, tid, nthreads
100 format('Hello, I am thread ', i0,' of ', i0, '.')
!$omp end parallel
end program
```

Hello Again

Most of the time, the program output is what is expected but, ...

```
$ OMP_NUM_THREADS=4 ./example
Hello, I'm thread 2 of 4.
Hello, I'm thread 2 of 4.
Hello, I'm thread 2 of 4.
Hello, I'm thread 3 of 4.
```

... occasionally, we have imposters pretending to be thread 2.

All variables declared outside of the scope of an OpenMP parallel construct is shared by all threads by default.

We created a data race



A data race is when one or more threads concurrently access a location in memory or a variable, at least one of which is a write.

As an example, we will consider this simple construct

```
int x = 0;
#pragma omp parallel
{
    x = x + 1;
}
```



Thread 1 fetch × from memory and store its value in a register



 Thread 1 add one to the value stored in the register

Thread 2 fetch × from memory and store its value in a register



Thread 1 store the result of the addition back in the shared memory

 Thread 2 adds one to the value stored in the register



Thread 2 stores the result of the addition back in the shared memory



Because of the potential data races in shared-memory parallel programs, extra care is needed as this is not always easy to spot

- with floating-point data, it may be difficult to distinguish from a numerical side effect
- changing the number of threads can cause the problem to seemingly (dis)appear
- may depend on the load on the system
- may only show up using many threads



In the previous example, we executed x+1 twice and get 1 as a result (while 2 was expected)

We need a way to prevent data race from happening: only share data that are not modified by other threads.

Hello Again (Data Race Free)

The solution to avoid a data race is to declare the **nthreads** and **tid** variables as private to the threads.

```
#include <stdio.h>
#include <omp.h>
int main(int argc, char* argv[]) {
    int nthreads, tid;
    #pragma omp parallel private(nthreads, tid)
    {
        tid = omp_get_thread_num();
        nthreads = omp_get_num_threads();
        printf("Hello, I'm thread %d of %d.\n", tid, nthreads);
    }
    return 0;
}
```

```
program main
    use omp_lib
implicit none
integer :: nthreads, tid
!$omp parallel private(nthreads, tid)
    tid = omp_get_thread_num()
    nthreads = omp_get_num_threads()
    print 100, tid, nthreads
100 format('Hello, I am thread ', i0,' of ', i0, '.')
!$omp end parallel
end program
```

private(list)

The parallel construct can take one or more data-sharing clause. The first one is private, which instruct that each thread should have its own instance of the listed variables. The initial value when we enter the parallel region is undefined.

firstprivate(list)

If the value of the variable before entering the parallel region matters, we can use firstprivate which is the same as private but, the variable should be initialized with its value before the parallel construct.

shared(list)

The third option is to declare a variable as **shared** which indicates that the variables listed should be shared among all threads. This is the default.

default(shared | none)

You can change the default data-sharing attribute with the default clause. Setting the value to none will force you to specify the data-sharing attribute for all your .
```
int x = 1, y = 2:
int z = 3, a = 4;
#pragma omp parallel private(x) firstprivate(v) shared(z)
{
  x = x + z;
 y = y + z;
  a = a + 1:
  printf("Thread %d: x = %d, y = %d, z = %d n",
          tid, x, y, z);
}
printf("Final: x = %d, y = %d, z = %d, a = %d n".
        x, y, z, a);
```

```
integer :: x = 1, y = 2
    integer :: z = 3, a = 4
    !$omp parallel private(x) firstprivate(y) shared(z)
      x = x + z
      y = y + z
      a = a + 1
      print 100, tid, x, y, z
100 format('Thread ', i0, ': x = ', i0, &
             ', y = ', i0, ', z = ', i0)
    !$omp end parallel
   print 200, x, y, z, a
200 format('Final: x = ', i0, ', y = ', i0, &
          ', z = ', i0, ', a = ', i0)
```

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&

\$ OMP_NUM_THREADS=4 ./example Thread 0: x = 3, y = 5, z = 3 Thread 1: x = 32674, y = 5, z = 3 Thread 3: x = 32674, y = 5, z = 3 Thread 2: x = 32674, y = 5, z = 3 Final: x = 1, y = 2, z = 3, a = 7

\$ OMP_NUM_THREADS=4 ./example Thread 0: x = 4, y = 5, z = 3Thread 1: x = 32674, y = 5, z = 3Thread 3: x = 32674, y = 5, z = 3Thread 2: x = 32674, y = 5, z = 3Final: x = 1, y = 2, z = 3, a = 7

The value of x is wrong because using the private clause the value of the variable is undefined when entering the parallel construct.

\$ OMP_NUM_THREADS=4 ./example Thread 0: x = 4, y = 5, z = 3Thread 1: x = 32674, y = 5, z = 3Thread 3: x = 32674, y = 5, z = 3Thread 2: x = 32674, y = 5, z = 3Final: x = 1, y = 2, z = 3, a = 7

The value of y, on the other hand, is correct as the **firstprivate** clause sets the initial value in the **parallel** construct to be the value before entering the construct.

\$ OMP_NUM_THREADS=4 ./example Thread 0: x = 4, y = 5, z = 3Thread 1: x = 32674, y = 5, z = 3Thread 3: x = 32674, y = 5, z = 3Thread 2: x = 32674, y = 5, z = 3Final: x = 1, y = 2, z = 3, a = 7

After the **parallel** region, the values of the private variables are the same as before the region.

\$ OMP_NUM_THREADS=4 ./example Thread 0: x = 4, y = 5, z = 3Thread 1: x = 32674, y = 5, z = 3Thread 3: x = 32674, y = 5, z = 3Thread 2: x = 32674, y = 5, z = 3Final: x = 1, y = 2, z = 3, a = 7

The value of z never changes. This variable is shared but never modified.

\$ OMP_NUM_THREADS=4 ./example Thread 0: x = 4, y = 5, z = 3Thread 1: x = 32674, y = 5, z = 3Thread 3: x = 32674, y = 5, z = 3Thread 2: x = 32674, y = 5, z = 3Final: x = 1, y = 2, z = 3, a = 7

We did not specify any data-sharing attribute for a. Thus, this variable has the default attribute and is shared. We see that there is a data race problem as we expected the final value to be 8 with 4 threads.

Variables with automatic storage duration that are declared in a scope inside the construct are private.

```
#pragma omp parallel
{
    int a = 3; // private
}
```

Variables with static storage duration that are declared in a scope inside the construct are shared.

```
#pragma omp parallel
{
   static int a; // shared
}
```

The loop iteration variable(s) in the associated for-loop(s) of a for construct is (are) private.

```
int i;
#pragma omp parallel
{
  #pragma omp for
  for(i = 0; i < n; ++i) // i is private</pre>
  ł
   // ...
  }
}
```

Objects with dynamic storage duration are shared (allocated by malloc).

```
int* a = (int*)malloc(n * sizeof(int));
#pragma omp parallel
{
  // the array a can not be privatized
  // all threads can read and write the
  // whole array
  a[0] = 3;
}
```



Set the default data attribute to none with default (none).

- You will get a compiler error if you do not explicitly specify the data attribute of your variables
- It forces you to think about the data attribute of your variables

Loop Carried Data Dependency

The fact that dynamically allocated objects cannot be private implies that particular care must be taken when handling arrays.

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

```
!$omp parallel
 !$omp do
   do i = 1,n-1
      a(i) = a(i+1) + b(i)
   end do
 !$omp end do
!$omp end parallel
```

Loop Carried Data Dependency

A **loop carried data dependency** occurs when a value written in one loop iteration is read or written by another iteration.

Thread 1	Thread 2
a[0] = a[1] + b[0]	a[4] = a[5] + b[4]
a[1] = a[2] + b[1]	a[5] = a[2] + b[5]
a[2] = a[3] + b[2]	a[6] = a[3] + b[6]
a[3] = a[4] + b[3]	a[7] = a[4] + b[7]



Synchronization ensures that two or more threads do not simultaneously execute some part of the program.

Synchronization may be needed for various reasons:

- makes sure that a particular operation is only executed once
- to avoid conflicts when accessing shared data
- ensure the order in which tasks are executed



A **barrier** directive is a synchronization point at which the threads in a parallel region will wait until all other threads in that section reach the same point.

- When a first thread reaches the barrier, it waits
- When a second thread reaches the barrier, it does the same thing and so on
- When the last thread reaches the barrier, all the threads resume execution



Most common usage of a barrier is to make sure that the value set by a thread is correctly defined before reading it from another thread.

```
#pragma omp parallel private(tid, neighb)
{
   tid = omp_get_thread_num();
   neighb = tid - 1;
   if (tid == 0)
        neighb = omp_get_num_threads() - 1;
   a[tid] = a[tid] * 3.5;
   #pragma omp barrier
   b[tid] = a[neighb] + c;
}
```

```
!$omp parallel private(tid, neighb, nthreads)
    tid = omp_get_thread_num()
    nthreads = omp_get_num_threads()
    neighb = tid - 1
    if (tid .eq. 0) neighb = nthreads - 1
    a(tid) = a(tid) * 3.5
    !$omp barrier
    b(tid) = a(neighb) + c
!$omp end parallel
```



Some constructs in OpenMP have an implicit barrier. This is the case for the parallel and for /do constructs.

```
#pragma omp parallel
{
    #pragma omp for
    for (int i = 0; i < n; ++i) {
        // ...
    }
    Implicit barrier, wait for all the threads to finish their iterations
    // ...
}    Implicit barrier, wait for all the threads to join</pre>
```

Master Directive

A master construct specifies a block of code that should be executed only by the master thread of the team.





Hello World, Master Edition

Let's revisit the hello world program but, this time, only the master thread print the number of threads in the team.

Hello World, Master Edition

```
$ gcc -fopenmp -o example omp_helloworld_master.c
$ OMP_NUM_THREADS=4 ./example
Hello, I'm thread 3
Hello, I'm thread 0
There is 4 threads in the team
Hello, I'm thread 2
Hello, I'm thread 1
```

Single Directive

A **single** directive is executed by only one of the threads in the team (not necessarily the master thread). There is an implicit barrier at the end.

```
#pragma omp single
structured-block
!$omp single
structured-block
!$omp end single
```



Hello World, Single Edition

Let's revisit the hello world program using the single construct. This time we illustrate the most common usage of the single construct, that is, assign a value to a shared variable.

```
!$omp parallel private(tid)
!$omp single
    nthreads = omp_get_num_threads()
    print 100, tid, nthreads
100    format('Hello, I am thread ', i0, ' of ', i0, &
        ' in the single construct.')
!$omp end single
    tid = omp_get_thread_num()
    print 200, tid, nthreads
200    format('Hello, I am thread ', i0, ' of ', i0, '.')
!$omp end parallel
```

Hello World, Single Edition

\$ gcc -fopenmp -o example omp_helloworld_single.c \$ OMP_NUM_THREADS=4 ./example Hello, I'm thread 3 of 4 in the single construct. Hello, I'm thread 3 of 4. Hello, I'm thread 2 of 4. Hello, I'm thread 0 of 4. Hello, I'm thread 1 of 4.

A **critical** section restricts execution of the associated structured block to one thread at a time.





Critical section is mostly used to update shared variables avoiding a data race.

```
#pragma omp parallel private(tid, local_sum)
{
   tid = omp_get_thread_num();
   local_sum = 0;
   #pragma omp for
   for (int i = 0; i < n; ++i)
        local_sum += a[i];
   #pragma omp critical
   {
      sum += local_sum;
      printf("Thread %d: local sum = %d, sum = %d.\n",
            tid, local_sum, sum);
   }
printf("Sum after parallel region: %d.\n", sum);
</pre>
```

```
!somp parallel private(tid, local_sum)
      tid = omp get thread num()
      local sum = 0
      !$omp do
        do i = 1, n
          local sum = local sum + a(i)
        end do
      !$omp end do
      !somp critical
        qlobal sum = qlobal sum + local sum
        print 100, tid, local_sum, global_sum
        format('Thread ', i0, ': local sum = ', i0, &
100
               ', sum = ', i0, '.')
&
      !somp end critical
    !$omp end parallel
    print*, 'Sum after parallel region:', global_sum
```

Critical section is mostly used to update shared variables avoiding a data race.

```
#pragma omp parallel shared(sum) private(tid, local_sum)
ł
  tid = omp_get_thread_num();
  local_sum = 0;
  #pragma omp for
  for (int i = 0; i < n; ++i)
      local sum += a[i];
                                 Critical section to update the global sum. Without the critical section,
  #pragma omp critical
                                 there is a potential data race here
    sum += local_sum;
    printf("Thread %d: local sum = %d, sum = %d.\n",
            tid, local_sum, sum);
  }
}
printf("Sum after parallel region: %d.\n", sum);
```

\$ gcc -fopenmp -o example omp_critical.c
\$ OMP_NUM_THREADS=4 ./example
Thread 0: local sum = 300, sum = 300.
Thread 3: local sum = 2175, sum = 2475.
Thread 1: local sum = 925, sum = 3400.
Thread 2: local sum = 1550, sum = 4950.
Sum after parallel region: 4950.

- A thread waits at the beginning of a critical section until no other thread is executing a critical section with the same name
- All unnamed critical directives map to the same name
- Critical section names are global to the program

```
#pragma omp critical (sum)
ł
 sum += local_sum;
 printf("Thread %d: local sum = %d,"
          " sum = %d. n",
          tid. local sum. sum):
}
#pragma omp critical (max)
 max = MAX(max, local_max);
 printf("Thread %d: local max = %d,"
         " max = %d. n",
          tid. local max. max):
}
```

```
!$omp critical (sum)
global_sum = global_sum + local_sum;
print 100, tid, 'sum', local_sum, &
& 'sum', global_sum
!$omp end critical (sum)
!$omp critical (max)
global_max = max(global_max, local_max)
print 100, tid, 'max', local_max, &
```

```
& 'max', global_max
!$omp end critical (max)
```

```
ł
 sum += local sum;
 printf("Thread %d: local sum = %d, sum = %d.\n",
        tid, local sum, sum);
}
#pragma omp critical (max)
ł
 max = MAX(max, local max);
 printf("Thread %d: local max = %d, max = %d.\n",
        tid, local_max, max);
}
```

```
ł
  sum += local sum;
  printf("Thread %d: local sum = %d, sum = %d.\n",
          tid, local sum, sum);
}
                                 Second critical section for the global maximum.
#pragma omp critical (max) 	 A thread can be in the first section while an other
ł
                                 is in the second one
  max = MAX(max, local max);
  printf("Thread %d: local max = %d, max = %d.\n",
          tid, local_max, max);
}
```

\$ gcc -fopenmp -o example omp critical named.c \$ OMP NUM THREADS=4 ./example Thread 3: local sum = 2175, sum = 2175. Thread 3: local max = 99, max = 99. Thread 1: local sum = 925, sum = 3100. Thread 1: local max = 49, max = 99. Thread 2: local sum = 1550, sum = 4650. Thread 0: local sum = 300, sum = 4950. Thread 2: local max = 74, max = 99. Thread 0: local max = 24, max = 99. Sum after parallel region: 4950. Max after parallel region: 99.

The nowait Clause

//...

}

```
#pragma omp for
for (int i = 0; i < n; ++i) {</pre>
    local_sum += a[i];

    There is an implicit barrier here

#pragma omp critical
ł
  sum += local_sum;
  printf("Thread %d: local sum = %d, sum = %d.\n",
           tid, local_sum, sum);
}
```

The nowait Clause

//...

```
#pragma omp for
for (int i = 0; i < n; ++i) {</pre>
     local_sum += a[i];
}
                                 There is no need to wait for the other threads to finish
#pragma omp critical +
                                 the iterations to execute the critical section
ł
  sum += local_sum;
  printf("Thread %d: local sum = %d, sum = %d.\n",
            tid, local_sum, sum);
}
```
```
//...
```

The **nowait** clause applied to a **for** construct remove the implicit barrier at the end of the construct.

```
#pragma omp for nowait
    structured-block
```

```
!$omp do
   structured-block
!$omp end do nowait
```

The **nowait** clause can also be applied to a **single** directive.

#pragma omp single nowait
 structured-block

!\$omp do
 structured-block
!\$omp end single nowait

The **nowait** clause can also be convenient when the work in two different loops are independent from each other.

```
#pragma omp parallel
{
    #pragma omp for
    for (int i = 0; i < n; ++i) {
        d[i] = a[i] + b[i];
    }
    #pragma omp for nowait
    for (int i = 0; i < n; ++i) {
        e[i] = a[i] + c[i];
    }
}</pre>
```

```
!$omp parallel
{
    !$omp do
        do i = 1,n
        d(i) = a(i) + b(i)
        end do
    !$omp end do nowait
    !$omp do
        do i = 1,n
            e(i) = a(i) + c(i)
        end do
    !$omp end do nowait
!$omp end parallel
```

The **nowait** clause can also be convenient when the work in two different loops are independent from each other.

```
#pragma omp parallel
ł
  #pragma omp for nowait
  for (int i = 0; i < n; ++i) {</pre>
       d[i] = a[i] + b[i];
    No barrier at the end of the loop
  #pragma omp for nowait
                                            The threads start the iterations of this loop as
  for (int i = 0; i < n; ++i) { ←
                                            soon as they finish their work in the first loop
       e[i] = a[i] + c[i]:
  }
```

The **nowait** clause can also be convenient when the work in two different loops are independent from each other.

```
#pragma omp parallel
ł
  #pragma omp for nowait
  for (int i = 0; i < n; ++i) {</pre>
       d[i] = a[i] + b[i]:
  }
  #pragma omp for nowait
  for (int i = 0; i < n; ++i) {</pre>
       e[i] = a[i] + c[i]:
  }
        Implicit barrier at the end of the parallel region
```



The **reduction** clause avoid data races when summing or combining values. This clause can be applied to the **parallel** and **for** constructs

reduction(op:list)

op is an operator:

- Arithmetic reductions: + * max min
- Logical operator reductions: & && | ||

Reduction

The sum and maximum example using critical region can be rewritten with **reduction** clauses instead

```
!$omp parallel for reduction(+:sum) &
!$omp& reduction(max:imax)
   do i = 1,n
      sum += sum + a(i)
      imax = max(imax, a(i))
   end do
!$omp end parallel for
print*, 'Sum after parallel region: ', sum
print*, 'Max after parallel region: ', imax
```

Atomic operation

An atomic operation is an operation that will always be executed without any other thread being able to read or change state that is read or changed during the operation.

```
#pragma omp atomic [atomic-clause]
    expression-statement
```

Atomic operation

#pragma omp atomic atomic-clause expression-statement

The value of atomic-clause can be one of the following: read, write, update and capture. If no atomic-clause is specified, the default value is update.

Atomic operation: Read and Write

The **read** clause allows for the atomic read of **x**.

#pragma omp atomic read
v = x;

The write clause allows for the atomic write of x. Here, expr is an expression with scalar type, i.e. the result of the expression is a scalar.

#pragma omp atomic write
x = expr;

Atomic operation: Update

The update clause allows for the atomic update of x.

#pragma omp atomic update
 expression-statement

Expression statement

x++; x--; ++x; --x; x op= expr; x = x op expr; x = expr op x;

Atomic operation: Capture

The capture clause allows for atomic update of the location designated by \times while also capturing the original or final value of the location designated by \times .

#pragma omp atomic update
 expression-statement

Expression statement

v = x++; v = x--; v = ++x; v = --x; v = x op= expr; v = x = x op expr; v = x = expr op x;

Atomic operation: Capture

The capture clause allows for atomic update of the location designated by x while also capturing the original or final value of the location designated by x.

#pragma omp atomic update
 structured-block

Structured block (part. 1)

{ v = x; x op= expr; } { x op= expr; v = x; }
{ v = x; x = x op expr; } { v = x; x = expr op x; }
{ x = x op expr; v = x; } { x = expr op x; v = x; }

Atomic operation: Capture

The capture clause allows for atomic update of the location designated by x while also capturing the original or final value of the location designated by x.

#pragma omp atomic update
 structured-block

Structured block (part. 2)

{ v = x; x++; } { v = x; ++x; } { ++x; v = x; } { x++; v = x; } { v = x; x--; } { v = x; --x; } { --x; v = x; } { x--; v = x; }

Atomic example

The previous example of the summation of the elements of an array using a **critical** construct can be rewritten using an **atomic** update.

```
#pragma omp for
for (int i = 0; i < n; ++i) {
    local_sum += a[i];
}</pre>
```

```
#pragma omp atomic
    sum += local_sum;
```

```
!$omp do
do i = 1,n
    local_sum += local_sum + a(i)
end do
!$omp end do
!$omp atomic
    sum = sum + local_sum
!$omp end atomic
```

Atomic example

```
for (i = 0; i < 10000; ++i) {</pre>
  index[i] = i % 1000;
  y[i] = 0.0;
}
for (i = 0; i < 1000; ++i)</pre>
  x[i] = 0.0;
#pragma omp parallel for
for (i = 0; i < 10000; ++i) {</pre>
  #pragma omp atomic update
  x[index[i]] += 1.0 * i;
 y[i] += 2.0 * i;
}
```

```
do i = 1,10000
 inds(i) = mod(i, 1000)
  y(i) = 0.0
end do
do i = 1,1000
  x(i) = 0.0
end do
!$omp parallel do
  do i = 1,10000
    !$omp atomic update
    x(inds(i)) = x(inds(i)) + 1.0 * i
    y(i) = y(i) + 2.0 * i
  end do
!$omp end parallel do
```

Atomic example

```
for (i = 0; i < 10000; ++i) {
    index[i] = i % 1000;
    y[i] = 0.0;
}
for (i = 0; i < 1000; ++i)
    x[i] = 0.0;</pre>
```

```
#pragma omp parallel for In
for (i = 0; i < 10000; ++i) {
    #pragma omp atomic update 
    x[index[i]] += 1.0 * i;
    y[i] += 2.0 * i;
}</pre>
```

The advantage of using atomic in this example is that it allows updates of two different elements of x in parallel. If a critical construct were used, all updates to elements of x would be executed serially

Atomic vs. Critical

Safely increasing the value of **count** in parallel can be done either by using an **atomic** Or a **critical** directive

```
#pragma omp atomic
    count++;
```

- An atomic operation has much lower overhead but the set of possible operations is restricted
- It can take advantage of hardware support for atomic operations

```
#pragma omp critical
    count++;
```

- A critical section can surround any arbitrary block of code
- There is a significant overhead when a thread enters and exits the critical section

Atomic vs. Reduction

Don't use **atomic** operation this way:

```
#pragma omp parallel for
for (int i = 0; i < n; ++i) {
    #pragma omp atomic
    sum += a[i];
}
```

```
!$omp parallel do
   do i = 1,n
      !$omp atomic
      sum = sum + a(i)
   end do
!$omp end parallel do
```

It is better to use a **reduction** clause:

```
#pragma omp parallel for reduction(+sum)
for (int i = 0; i < n; ++i) {
    sum += a[i];
}</pre>
```

```
!$omp parallel do reduction(+sum)
  do i = 1,n
    sum = sum + a(i)
    end do
!$omp end parallel do
```

Performance Considerations

- Avoid or minimize the use of barrier and critical sections.
- Use the nowait clause where possible to eliminate unnecessary barriers
- Favour the use of master instead of single

Loop Scheduling

Loop Scheduling

Loop scheduling, specify how iterations of a loop are divided into contiguous non-empty subsets (chunks), and how these chunks are distributed to the threads. Changing the loop scheduling is possible to use the **schedule** clause.

#pragma omp for schedule(kind, chunk)
 for-loop

```
!$omp do schedule(kind, chunk)
    do-loop
!$omp end do
```

Where the value of kind can be static, dynamic, guided or runtime. The default scheduling is static. The optional chunk may have different behaviour depending on the scheduling.

Static Loop Scheduling

With **static** loop scheduling, iterations are divided into chunks and the chunks are assigned to the threads. Each chunk contains the same number of iterations, except for the chunk that contains the last iteration, which may have fewer iterations.

#pragma omp for schedule(static)
 for-loop

!\$omp do schedule(static)
 do-loop
!\$omp end do



Dynamic Loop Scheduling

With dynamic loop scheduling, the iterations are distributed to threads in chunks. Each thread executes a chunk of iterations, then requests another chunk, until no chunks remain to be distributed.

#pragma omp for schedule(dynamic)
 for-loop

!\$omp do schedule(dynamic)
 do-loop
!\$omp end do



Guided Loop Scheduling

The **guided** loop scheduling is similar to the **dynamic** scheduling except that the size of each chunk is proportional to the number of unassigned iterations, decreasing to one.

#pragma omp for schedule(guided)
 for-loop

!\$omp do schedule(guided)
 do-loop
!\$omp end do



Why Using the Scheduling Clause?

- The default scheduling, static with a chunk equals to niter/nthreads is not ideal for all workload.
- It may be the case that iterations of high index represent more work. In that case, some of the threads will finish early and have nothing to do. We have a load imbalance.
- Changing the scheduling may help balance the amount of work between the threads.

Example: Number of Prime Numbers

```
int prime, sum = 0;
#pragma omp parallel shared(n) private(prime)
{
  #pragma omp for reduction(+:sum)
  for (int i = 2; i <= n; i++) {</pre>
    prime = 1;
                                              Trip count of this loop may be very low or
    for (int j = 2; j < i; j++) { \leftarrow very high depending if the number is prime
       if ( i % j == 0 ) {
                                              or not
         prime = 0;
         break;
     }
    sum += prime;
  }
```

Example: Number of Prime Numbers

```
int prime, sum = 0;
#pragma omp parallel shared(n) private(prime)
{
 #pragma omp for reduction(+:sum)
 for (int i = 2; i <= n; i++) {</pre>
   prime = 1;
   for (int j = 2; j < i; j++) {
     if ( i % j == 0 ) {
      prime = 0;
      }
   }
   sum += prime;
 }
}
```

Example: Number of Prime Numbers

\$ gcc -fopenmp -o example omp_schedule_prime.c
\$ OMP NUM THREADS=4 ./example

		Default	Static	Dynamic	Guided
Ν	Pi(N)	Time	Time	Time	Time
1024	172	0.000182	0.000120	0.000104	0.000121
2048	309	0.000561	0.000359	0.000425	0.000393
4096	564	0.001987	0.001309	0.001216	0.001239
8192	1028	0.007116	0.004474	0.004375	0.005114
16384	1900	0.029730	0.015594	0.015902	0.015161
32768	3512	0.099248	0.058475	0.056940	0.057160
65536	6542	0.358250	0.218291	0.244626	0.254815
131072	12251	1.416871	0.848736	0.788619	0.819390
262144	23000	5.207946	3.193940	3.062080	3.064527
524288	43390	20.565462	12.638959	12.086839	12.102800

Example: Triangular Loop

```
#pragma omp parallel shared(a, n)
{
    #pragma omp for
    for (int i = 0; i < n; ++i) {
        a[i] = 0.0;
        for (int j = 0; j < i; ++j) {
            a[i] += cos( -3.1 * sin( 2.3 * cos ( (double) j ))) ;
        }
    }
}</pre>
```

Example: Triangular Loop

\$ gcc -fopenmp -o example omp_schedule_triangular.c
\$ OMP_NUM_THREADS=4 ./example

	Default	Static	Dynamic	Guided
Ν	Time	Time	Time	Time
1024	0.025865	0.016811	0.018739	0.018241
2048	0.100062	0.070023	0.082587	0.091206
4096	0.383107	0.238520	0.232556	0.226914
8192	1.515341	0.905186	0.895541	0.880046
16384	6.064787	3.540685	3.526388	3.590453
32768	24.041088	15.465762	14.088137	14.539937
65536	97.495829	59.291353	59.403173	60.252156

OpenMP and Cache

Your Typical Compute Node



OpenMP and cc-NUMA

You also have two option for the placement of your threads. The first is put the threads far apart, i.e. on different sockets.

- may improve the aggregated memory bandwidth available to your application
- may improve the combined cache size available to your application
- may decrease performance of synchronization constructs

The second option is to put the threads close together, i.e. on two adjacent cores.

- may improve performance of synchronization constructs
- may decrease the available memory bandwidth and cache size

OpenMP and cc-NUMA

For the placement, you can use the **OMP_PROC_BIND** environment variable with the values:

- **close** : successively through the available places
- spread : which spreads the threads over the places

The second option is the **OMP_PLACES** environment variable with the values:

- **core** : places correspond to the cores
- socket : places correspond to the sockets

OpenMP and cc-NUMA

```
double* A = (double*)malloc(N * sizeof(double));
```

```
#pragma omp parallel for
for (int i = 0; i < N; i++) {
    A[i] = 0.0;
}</pre>
```

- For a serial serial code: all array elements are allocated in the memory of the NUMA node containing the core executing the thread
- For a parallel code on an OS with a first touch policy the array elements are allocated in the memory of the NUMA node containing the core executing the thread initializing
False Sharing in Action

Another thing you need to consider if you want to get the best out of OpenMP is false sharing. To discuss this we will start with these piece of code:

```
double local sum[omp get max threads()];
double sum = 0.0;
#pragma omp parallel shared(sum)
ł
  int tid = omp get thread num();
  local sum[tid] = 0.0;
  #pragma omp omp for
  for (int i = 0; i < n; ++i)</pre>
      local sum[tid] += 0.5 * x[i] + y[i];
  #pragma omp atomic
  sum += local_sum[tid];
}
```

False Sharing in Action

Let's measure the time spend in the parallel region (using the **omp_get_wtime(**) function).

Threads	Time (s)	
1	0.535418	
2	0.421140	
4	0.554419	
8	0.597622	

- The speedup from 1 thread to 2 theads is bad
- When going to 4 and 8 threads the time spend in the parallel region is worst than with 1 thread

False Sharing

False sharing is when threads impact the performance of each other while modifying independent variables sharing the same cache line



- If one core writes, the cache line holding the memory line is invalidated on other cores.
- Even though another core is not using that data, the second core will need to reload the line before it can access its own data again.

False Sharing: Solution

Solution: introduce a padding.

}

```
double local sum[LINESIZE*omp get max threads()];
double sum = 0.0;
#pragma omp parallel shared(sum)
{
  int tid = omp_get_thread_num();
  local sum[LINESIZE*tid] = 0.0;
  #pragma omp omp for
  for (int i = 0; i < n; ++i)</pre>
      local sum[LINESIZE*tid] += 0.5 * x[i] + y[i];
  #pragma omp atomic
  sum += local sum[LINESIZE*tid];
```

False Sharing: Solution

Timing for different paddings on a CPU with a cache line size of 64 bytes.

Threads	Time (s)	Time (s)	Time (s)
	padding = 4	padding = 8	padding = 16
1	0.535418	0.535418	0.535418
2	0.601417	0.270089	0.270843
4	0.441149	0.152651	0.149363

False sharing

- When threads access global or dynamically allocated shared data structures there is a potential sources of false sharing
- False sharing may be difficult to spot. For example, when theads access completely different global variables that happen to be relatively close together in memory.
- Use thread-local copies of data when possible. The thread-local copy can be read and modified frequently and only when complete, be copied back to the global data structure



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