OpenMP programming: 2023/2024

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1 About this tutorial

- This is an explanation. **Read it** before moving on to the ...
 - * ... exercice

If you have compilation/execution problems:

- If you get any error message(s): **Read** and try to understand them.
- This tutorial has been carefully tested and hopefully most errors have been removed: please tell me if you think you've found one!

2 Compiling and executing OpenMP programs

OpenMP is a shared memory API providing an interface to write software that can use multiple cores of a computer. This is a well-established standard (v5.0 currently) which is provided by default in nearly all compilers in nearly all operating systems for C, C++ and Fortran. It consists of a set of *compiler directives*, *library routines* and *environment variables* that influence run-time behavior.

* For the complete OpenMP 5.0 Specifications, Examples and Summary Cards visit:

```
https://www.openmp.org/specifications/
```

Don't panic! There's no need to read the whole documentation. In this tutorial you will learn about the basic OpenMP functionalities that will allow you to write your first parallel programs for shared-memory multi-core systems. If you need more detailed information and advanced features, this is the place to go.

2.1 First OpenMP program:

* Open a file called hello_world.f90 and copy in the following code:

```
PROGRAM hello_world

IMPLICIT NONE

!$omp parallel
PRINT*,'hello'
!$omp end parallel
END PROGRAM hello_world
```

Listing 1 – hello_world.f90

A block preceded by a !\somp parallel directive is called a parallel OpenMP region. It is executed by a newly created team of threads. This is an instance of the SPMD model : all threads execute the same segment of code. Note that lines starting with ! indicate a commentary in Fortran, so they will be ignored if you compile this code without specifying the -fopenmp option to the compiler.

* Compile this code with the following command :

```
$ gfortran -fopenmp hello_world.f90 -o hello_world
```

and run the executable hello world.

- * In the command line type 1scpu to find out how many cores/threads your computer has.
- * Execute the hello_world program with a varying number of threads. This can be done by setting the OMP_NUM_THREADS environment variable. For example, this

```
$ OMP_NUM_THREADS=16 ./hello_world
```

will execute hello_world with 16 OpenMP threads, while

```
$ export OMP_NUM_THREADS=5
```

sets the environment variable to 5 for all subsequently executed commands (in that shell).

 \star Compile and run the program again without the -fopenmp flag.

2.2 Threads information

We can get some information about the OpenMP threads by using *library routines* defined in the OpenMP standard, mainly the number of threads and an unique number for each thread.

* Extend the previous hello_world.f90 as follows (copy to a new file hello_threads.f90)

```
PROGRAM hello_threads

USE omp_lib
IMPLICIT NONE

INTEGER :: thread_id,threads_num
PRINT*,'Only one thread here...'

!$omp parallel private(thread_id,threads_num)
thread_id = omp_get_thread_num()
threads_num = omp_get_num_threads()
PRINT*,'hello! my thread id is',thread_id,' out of ',threads_num
!$omp end parallel

PRINT*,'back to one thread'
END PROGRAM hello_threads
```

Listing 2 – hello_threads.f90

Note that the two variables thread_id, threads_num are declared with the private data-sharing clause so each thread has his own memory copy of these variables. By default, all the variables are shared in memory and can be written by all the threads. Note also the declaration USE omp_lib, which is necessary for using OpenMP functions.

- ★ Compile and execute. Vary the number of threads and observe the output.
- * How does the output change if you remove the private clause?
- * Use the default(none) clause (instead of private(...)) and re-compile. Notice the compiler messages.

3 Parallel loop

Loops are the main way for parallelizing in OpenMP but the following rules need to be followed:

- The !\sum do work-sharing directive specifies that iterations of the loop (the one following the directive!) are distributed among the existing threads in a parallel region.
 - · NB : all iterations should be independent of each other!
- The following restrictions apply to loops associated with a !\$omp do directive :
 - · DO WHILE or infinite loops are not parallelizable with OpenMP
 - · loops should not contain statements that terminate the loop prematurely (EXIT, GOTO)

(however, recent versions of OpenMP allow premature loop cancellation using !\$omp cancel directives)

- If a parallel region contains only one parallel loop the !\$\text{omp parallel and !}\text{somp do directives can be merged into a single equivalent !}\text{somp parallel do.}
- By default, the loop iteration variables are made private.
- The parallelized loop is (only!) the one which comes immediately after the !\$omp do directive

3.1 A simple loop

* Open a file called loops.f90 and copy in the following code:

```
PROGRAM loops
USE omp_lib
IMPLICIT NONE

INTEGER, PARAMETER :: n=1002
INTEGER :: i,num_loops=0

!$omp parallel
!$omp do

DO i=1,n
    num_loops = num_loops + 1

ENDDO
```

```
!$omp end do
PRINT*,'thread-id ',omp_get_thread_num(),' : number of loops : ',num_loops
!$omp end parallel
PRINT*,'------'
PRINT*,'total number of loops',num_loops

END PROGRAM loops
```

Listing 3 – loops.f90 (WRONG!)

- * Compile the program and run the executable several times with 4 threads. Is the final value of num_loops coherent? Before moving on to the next step, you should understand why the value is (in general) different from 1002.
- * Add the reduction(+:num_loops) clause to the !\$omp parallel directive in order to obtain the sum of num loops at the end of the parallel region.

3.2 Reduction : π computation

A reduction is an associative operation applied to a shared variable. The operation can be:

- arithmetic: +, -, x
- logical: .AND., .OR., .EQV., .NEQV.
- an intrinsic function: MAX, MIN, IAND, IOR, IEOR

Each thread computes a partial result independently of the others. They are then synchronized in order to update the final result.

 \star Create a serial program pi_serial.f90 that computes the value of π with the following formula. Increasing the number of iterations nsteps will improve the precision.

$$\pi/4 = \arctan(1) = \sum_{k=0}^{k=\mathsf{nsteps}} \frac{(-1)^k}{2k+1}$$

(Optional!) You can use the following piece of code to read nsteps as an argument from the command line.

```
CHARACTER(len=32) :: arg

CALL getarg(1, arg)
read(arg, *)nsteps
```

Listing 4 – read arguments

- * Copy your serial program in a file named pi_omp.c and parallelize the main loop with a reduction.
- * Use the function system_clock to measure the elapsed time taken to compute π .

```
INTEGER(kind=8) :: t1, t2, clock_rate

CALL SYSTEM_CLOCK(t1)
!pi computation here...

CALL SYSTEM_CLOCK(t2, clock_rate)
PRINT*, 'time :', REAL(t2-t1)/REAL(clock_rate)
```

Listing 5 – timing with system_clock

 \star Increase the number of iterations until the serial execution takes about 10 seconds and compare the time when increasing the number of threads. For example, you can execute your code like this :

```
1 for i in 1 2 4 6 8;do echo ${i} `OMP_NUM_THREADS=${i} ./pi_omp`;done
```

4 Load balancing: the schedule instruction

- Work in loop iterations can be imbalanced
- The distribution mode of the iterations can be specified with the schedule clause:

```
!$omp do schedule(type[,chunk])
```

where type is one of the following and [,chunk] an optional positive integer

- · static[,K]: divide loop iterations into chunks of size K, statically assigned to threads
- · dynamic[,K] : divide loop iter into chunks size K. When a thread finishes one chunk, it is dynamically assigned another. Default is K=1.
- guided[,K]: similar to dynamic, with decreasing chunk size ($\geq K$)
- · runtime: scheduling policy defined by environment variable OMP_SCHEDULE
- The choice of the scheduling policy allows a better control of the load-balancing between threads.

4.1 Prime numbers count

The following algorithm computes the number of primes contained in the interval [2, n] (in a very naive way). The j loop number of iterations depends on i so the amount of work will be different on each i loop.

```
PROGRAM main

USE omp_lib
IMPLICIT NONE

INTEGER :: i,j,n=200000,prime=0,total=0
DOUBLE PRECISION :: tstart
```

```
!$omp parallel default(none)
    tstart=omp_get_wtime()
    !$omp do reduction(+ :total)
10
    D0 i=2,n
       prime=1
12
       D0 j=2,i-1
           IF(MOD(i,j)==0)THEN
14
              prime=0
              EXIT
16
           ENDIF
18
       ENDDO
        total = total + prime
    ENDDO
    !$omp end do
    PRINT*, 'thread ', omp_get_thread_num(), ' time : ', (omp_get_wtime()-tstart)
    !$omp end parallel
24
    PRINT*, 'nb primes ', total
26 END PROGRAM main
```

Listing 6 – primes_omp.f90

- * The program won't compile because of the default(none) clause. Add appropriate shared and private data sharing clauses.
- * The program uses the omp_get_wtime() function to measure the time each thread spends in the i loop. Does this work as desired? Add the nowait clause to the omp_end_do directive and observe how this changes the (per thread) timing.
- \star Add a schedule clause to the omp do directive. Run the code with an increasing number of cores and different scheduling policies (try at least static and dynamic with chunk sizes 1, 10, 1000, 100000, 200000). How to explain the differences?