# TP - Introduction to OpenMP

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To compile the program program.cpp with OpenMP and generate the executable program, type the following command in the terminal:

g++ -02 -std=c++11 -fopenmp program.cpp -o program

#### Part 1

# Hello World using OpenMP

#### Ex. 1

- a) Write a program hello-openmp.cpp (skeleton code given) having a parallel region in which each thread prints its identifier as well as the total number of threads.
- b) Next, in the same parallel region, print "Hello World from threadId=??" by a single thread with its thread id. Try using both omp single and omp master constructs and observe the difference.
- c) Try to modify the number of threads in the parallel region using three different methods: modifying the environment variable OMP\_NUM\_THREADS, calling the function omp\_set\_num\_threads(...), and adding the num\_threads(...) clause to the omp parallel construct. What is the order of precedence among these three methods?

#### Part 2 -

## Sum of an array using OpenMP

### Ex. 2

- a) Write a C/C++ program (use the given skeleton code sum-array-openmp.cpp) that initializes an array A of N floating point numbers so that A[i] = i for all  $0 \le i < N$ . Use a value of N sufficiently large (>1M) to see gains in parallel execution.
- b) Add a second for loop that computes the sum of all elements in A.
- c) Now add a parallel region enclosing both loops, and parallelize each loop using a separate #pragma omp for.
- d) Can we add a nowait clause to the first loop's #pragma omp for? Explain.
- e) Now, parallelize the second loop with a #pragma omp sections construct having 4 sections. Each section should iterate over N/4 elements of A, find the sum of the N/4 elements, and finally add it to the global sum of all N elements. Make sure to eliminate the race conditions efficiently using #pragma omp critical and #pragma omp atomic.
- f) Query the number of hardware threads supported by your processor using the 1scpu command, then execute your program with a timer using  $1, 2, \ldots, P$  threads if your CPU has P threads. Compute the speedup/acceleration and efficiency for each execution, and plot these results.

#### Part 3 -

# Parallel mergesort using OpenMP sections

The goal of this exercise is to sort an array of numbers using the mergesort algorithm in parallel using OpenMP sections. A skeleton code is already provided in the file mergesort.cpp. This code allocates and initializes an array A of N numbers as well as another temporary buffer array temp of the same size.

#### Ex. 3

- a) Create a parallel region with 4 sections (or 8, if there is at least 8 hardware threads in your machine). Each section should sort N / 4 consecutive elements of the array A. You can use either std::sort of the STL library or the mergesort function in the skeleton code.
- b) In the same parallel region, after having finished these 4 sections for sorting, create 2 sections where each section merges two sorted arrays of N/4 elements, to generate a sorted array of N/2 elements. To do this, use the provided merge function that performs this merge operation in-place on A.
- c) Finally, outside the parallel region, merge these two subarrays of size  $\mathbb{N}/2$  to obtain the final sorted array  $\mathbb{A}$  of size  $\mathbb{N}$ .
- d) Compare the sequential execution time with the parallel execution time using 4 (or 8) threads. What is the speedup/acceleration? What is the parallel efficiency?

Part 4

### Computing the $\pi$

The  $\pi$  number can be defined as the integral of  $f(x) = \frac{4}{1+x^2}$  from 0 to 1. An easy way to approximate this integral is to uniformly discretize the domain using N points with  $s = \frac{1}{N}$  distance between two consecutive points:

$$\pi \approx \int_0^1 \frac{4}{1+x^2} dx \approx \sum_{i=0}^{N-1} s \times \frac{f(i \times s) + f((i+1) \times s)}{2}$$

We will write a C++ program that computes this approximation for  $\pi$ , then parallelize it using two different methods using OpenMP. A skeleton code is provided in the file calcul-pi.cpp.

Ex. 4

- a) First, write a sequential code in the skeleton code that correctly computes the value of  $\pi$  using this formula.
- b) We can then distribute this computation among P threads available in your machine. First, simply parallelize the main loop of computation using pragma omp for. Be careful about the race condition here; you need to add an OpenMP clause to avoid it. Test the performance and acceleration using different number of threads (1, 2, ..., P).
- c) The second parallelization strategy is "by hand"; you are not allowed to use the OpenMP constructs omp for/omp sections or the clause reduction. Each thread should execute a standard for loop, but with a different domain of iteration (begin/end) of size N / P depending on its thread id, and compute a partial value of  $\pi$  for its domain. Next, each thread will merge these partial values into the final value of  $\pi$ . In doing so, make sure that you avoid race conditions, using omp atomic clause. Your code should perform correctly and efficiently for any number of threads P.