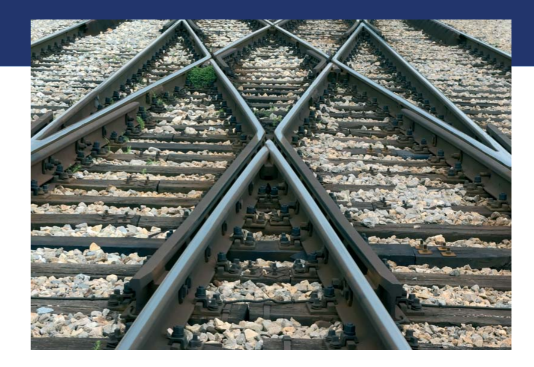
Parallel programming Programming with OpenMP Part 1







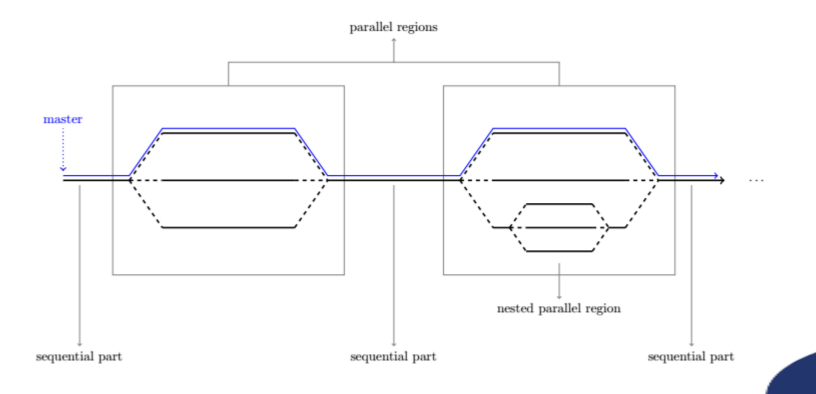
Introduction to OpenMP

- OpenMP (Open Multi-Processing) provides constructs for parallel programming in C++, C, and Fortran on Linux, MacOS, and Windows.
- A sequential code is transformed to a parallel one by adding compiler pragmas, so if a compiler does not support OpenMP, the pragmas are skipped and the output is a sequential program.
 - OpenMP manual: 1.3 Execution model: The OpenMP API is intended to support programs that will execute correctly both as parallel programs (multiple threads of execution and a full OpenMP support library) and as sequential programs (directives ignored and a simple OpenMP stubs library). However, it is possible and permitted to develop a program that executes correctly as a parallel program but not as a sequential program, or that produces different results when executed as a parallel program compared to when it is executed as a sequential program. Furthermore, using different numbers of threads may result in different numeric results because of changes in the association of numeric operations.
- OpenMP is widely used in software like Blender, fftw, OpenBLAS, and eigen to accelerate computations.
- It is easy to use!



Execution model

- OpenMP program starts as a single thread only (master thread).
- It is executed sequentially until it reaches a **parallel region** defined by OpenMP pragma.
- At the entry of parallel region, new *team of threads* **is** created. Each thread executes concurrently with the others in order to share the work to be done.
- An OpenMP program alters between sequential regions and parallel regions.





Using OpenMP

- Include header file #include <omp.h>
- Cmake (multi-platform)

```
find_package(OpenMP)
if (OPENMP_FOUND)
   set(CMAKE_C_FLAGS "${CMAKE_C_FLAGS} ${OpenMP_C_FLAGS}")
   set(CMAKE_CXX_FLAGS "${CMAKE_CXX_FLAGS} ${OpenMP_CXX_FLAGS}")
   set(CMAKE_EXE_LINKER_FLAGS "${CMAKE_EXE_LINKER_FLAGS} ${OpenMP_EXE_LINKER_FLAGS}")
endif()
```

• gcc

g++ -fopenmp ...



Hello world! In OpenMP...

lab_codes/src/HelloWorld.cpp



Runtime Library Routines

omp_get_num_procs()

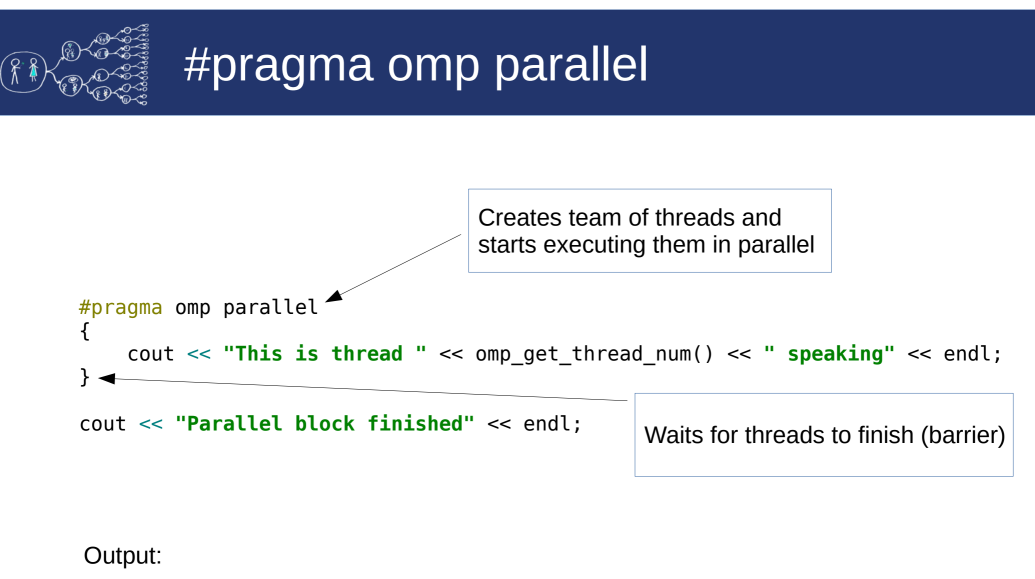
Returns the number of processors that are available to the program

omp_get_num_threads()

Returns the number of threads that are currently in the team executing the parallel region from which it is called

omp_get_thread_num()

Returns the calling thread index within the current team



This is thread 0 speaking This is thread 3 speaking This is thread 2 speaking This is thread 1 speaking Parallel block finished

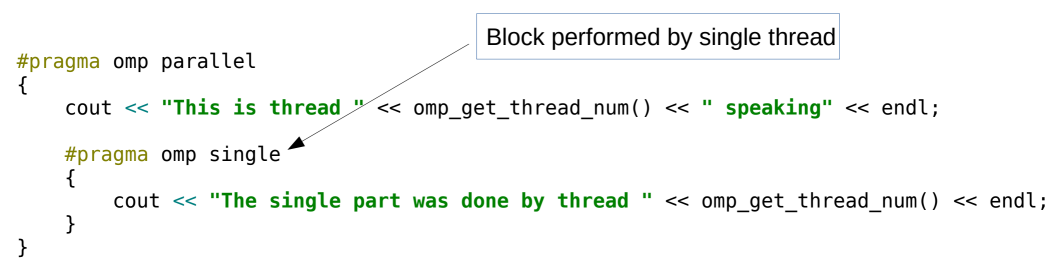


Output:

This is thread 0 speaking This is thread 3 speaking This is thread 6 speaking This is thread 1 speaking This is thread 2 speaking This is thread 7 speaking This is thread 4 speaking This is thread 5 speaking



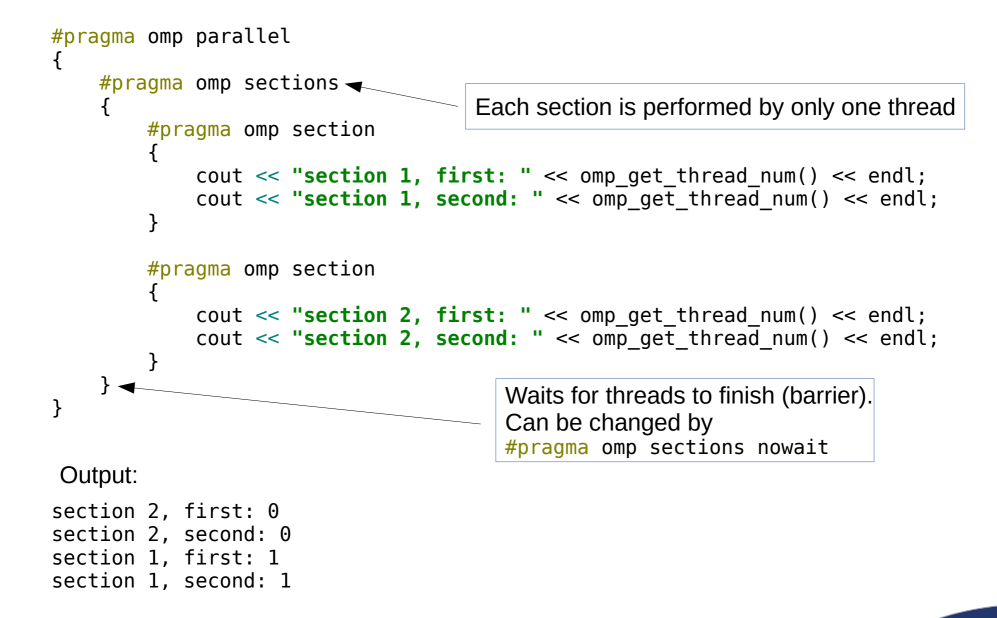
#pragma omp single



Output: This is thread 3 speaking The single part was done by thread 3 This is thread 1 speaking This is thread 2 speaking This is thread 0 speaking



#pragma omp sections





#pragma omp critical

int sum;

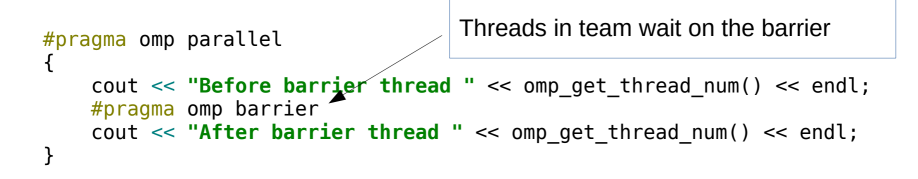
```
#pragma omp parallel
{
    #pragma omp critical 
    cout << "Thread " << omp_get_thread_num() << " in critical region" << endl;
    sum += omp_get_thread_num();
    }
}
cout << sum << endl;</pre>
```

Output:

Thread 1 in critical region Thread 0 in critical region Thread 3 in critical region Thread 2 in critical region 6



#pragma omp barrier



Output:

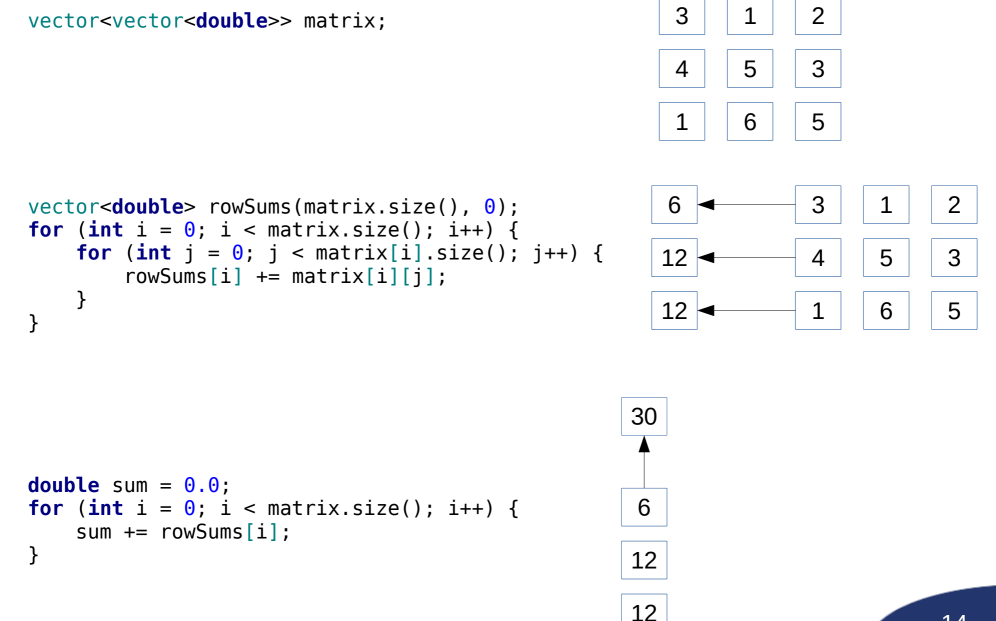
Before barrier thread 0 Before barrier thread 3 Before barrier thread 1 Before barrier thread 2 After barrier thread 1 After barrier thread 2 After barrier thread 0 After barrier thread 3



- Write function for computing vector normalization. Split the vector into two halves, each is processed by one section.
 - You may use skeleton lab_codes/src/VectorNormalization.cpp

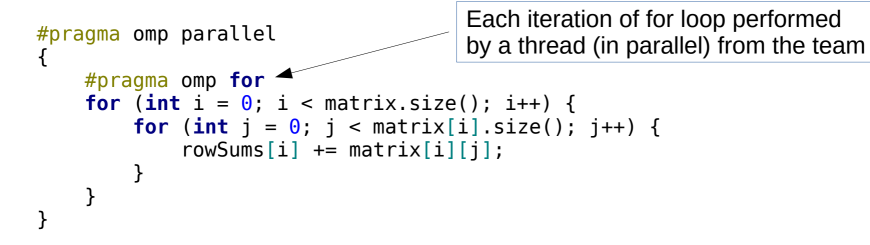


Sequential summing of matrix rows





Parallel summing of matrix rows



A shorter code...

Question: what happens if you write the pragma on the inner loop?





```
#pragma omp parallel for if(matrix.size() >= 10)
for (int i = 0; i < matrix.size(); i++) {
    for (int j = 0; j < matrix[i].size(); j++) {
        rowSums[i] += matrix[i][j];
    }
}</pre>
```

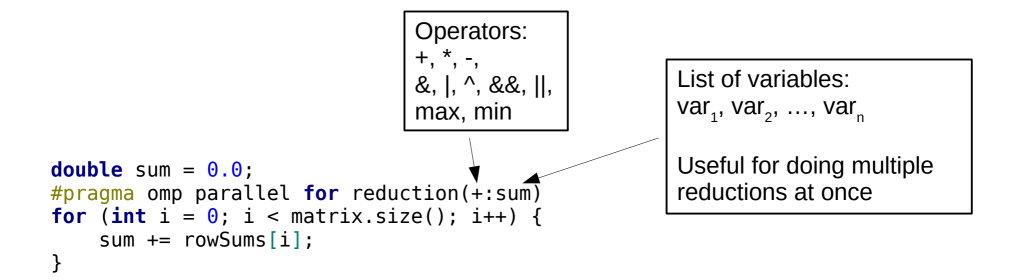
Threads are only created for large matrices. Small matrices are summed sequentially since it does not pays off to create threads.



Reductions

• Parallel aggregation of an expression, e.g., a sum

sum = rowSums[0] + rowSums[1] + rowSums[2] + ... + rowSums[matrix.size() - 1];





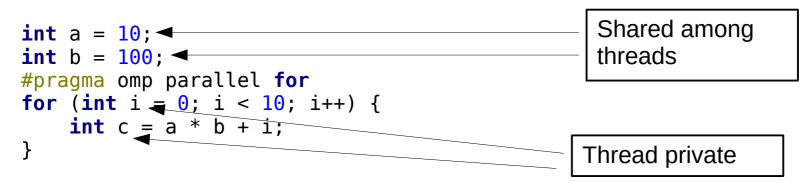
Collapse

Collapse for loops into one for distribution of the work among threads

```
int numRows = matrix.size();
int numCols = matrix[0].size();
double sum = 0.0;
#pragma omp parallel for collapse(2) reduction(+:sum)
for (int i = 0; i < numRows; i++) {
   for (int j = 0; j < numCols; j++) {
      sum += matrix[i][j];
   }
}
```



Data sharing



- The sharing can be stated explicitly as a clause
 - #pragma omp parallel for private(a, b)
 - Variables a and b are private to each thread (without global initialization)
 - #pragma omp parallel for firstprivate(a, b)
 - Variables a and b are private to each thread (with global initialization)
 - #pragma omp parallel for shared(a, b)
 - Variables a and b are shared among threads
- The default policy can be set to
 - #pragma omp parallel for default(shared)
 - By default, all the variables outside of the parallel section are shared
 - #pragma omp parallel for default(none)
 - The programmer must explicitly state the sharing policy of the variables



- Vector normalization using parallel for (reduction, critical section ...)
- Computation of pi using Monte Carlo

