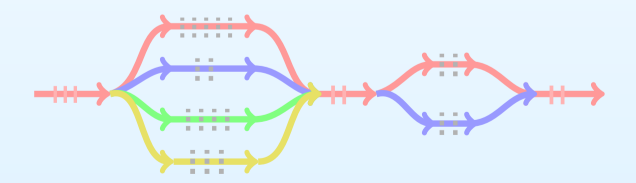


Parallel Programming in OpenMP

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Overview



- Parallel programming
 - Prerequisite for parallel computing:
 - Constructs for parallel execution
 - Data communications
 - Synchronization
- OpenMP programming: directives/pragmas, environment variables, and run-time libraries
 - Variables peculiar to OpenMP programming;
 - Loop level parallelism;
 - **Nested** thread parallelism;
 - Non-loop level parallelism;
 - Data race and false sharing;
- Summary





Parallel programming



- Parallel programming environment;
 - Essential language extensions to the existing language (Fortran 95);
 - New constructs for directives/pragmas to existing serial programs (**OpenMP** and HPF);
 - Run-time libraries that support data communication and synchronization (MPI and Pthreads);
- OpenMP stands for Open Multi-Processing (API);
- OpenMP is one of the directives/pragmas approaches that support parallelism on shared memory systems;
- **OpenMP** is supported by Fortran, and C/C++;
- **OpenMP** allows us to start from a serial code and provides an incremental approach to express parallelism;





Shared-memory parallel programming



- System level and user's application level;
- Pthreads specification is from the IEEE POSIX standard;
 - Many control knobs at low level;
 - Difficult to use and relatively heavyweight threads;

Intel Cilk Plus

- C/C++ language extensions;
- Supported by GCC and Intel C/C++ compilers;
- Fork-join mechanism and efficient load-balancing via work-stealing;
- Intel TBB (Threading Building Blocks)
 - C++ libraries instead of language extension;
 - Supports task and loop-level parallelism;
- OpenCL
 - Offload work to devices, C/C++, Python, Java APIs;
 - Low-level model;



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The "Three Knights" in OpenMP



- (1) Directives/pragmas need to express parallelism;
- (2) Run-time libraries can dynamically control or change code execution at run-time;
- (3) Environment variables specify the run-time options;
- How does OpenMP achieve parallel computing?
 - Specify parallel execution parallel constructs allowing parallel execution;
 - Data communication data constructs for communication among threads;
 - Synchronization synchronization constructs;
- OpenMP directives/pragmas:

Fortran: !\$omp, c\$omp, Or *\$omp [clauses]

C/C++: #pragma [clauses]

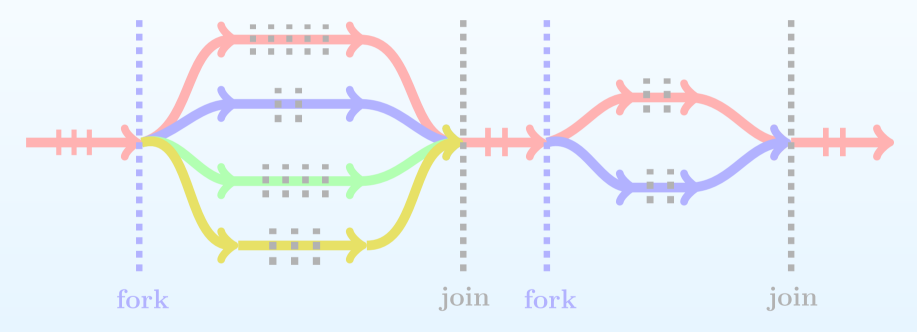




Parallel execution



 Constructs for parallel execution: OpenMP starts with a single thread, but it supports the directives/pragmas to spawn multiple threads in a fork-join model;



- OpenMP do and parallel directives;
- OpenMP also allows you to change the number of threads at run-time;





Data communication

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- Each thread was assigned to a unique thread ID from 0 to N-1. Here N is the total number of threads;
- The key point is that there are three types of variables: private, shared, and reduction variables;
- At run-time, there is always a common region in global memory that allows all threads to access it, and this memory region is used to store all shared variables;
- Each thread was also assigned a private memory region to store all private variables. Thread a cannot access the private variables stored in the private memory space for thread b;
- **Data communications** are achieved through read and write operations on **shared** variables among the threads;





Synchronization



- In OpenMP, synchronization is used to (1) control the access to shared variables and (2) coordinate the workflow;
- Event and mutual exclusion synchronization;
- Event synchronization includes barrier directives, which are either explicit or implicit; a thread has to wait until all threads reach the same point;
- Mutual exclusion is supported through critical, atomic, single, and master directives. All these are used to control how many threads, which thread, or when a thread can execute a specified code block or modify shared variables;
- Be careful with synchronization!





Compile OpenMP code



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• Compiler options that enable OpenMP directives/pragmas:

Compiler	Fortran	С	C++
Intel	ifort - openmp	icc -openmp	icpc -openmp
PGI	pgf90 -mp	pgcc -mp	pgCC -mp
GCC	gfortran -fopenmp	gcc -fopenmp	g++ -fopenmp

- Compilers support conditional compilation in disabling OpenMP. Intel compiler also provides the flag -openmp-stubs at the compiler level;
- Load modules on the HPC or LONI machines:
- \$ module load [package name]
- \$ soft add [+package name] (resoft) # intel, pgi, or gcc.
- Set up an environment variable:
- \$ export OMP_NUM_THREADS=[number of threads]







Loop level parallelism





First OpenMP "Hello World!" in Fortran and C LSU INFORMAT

```
Fortran (hello f.f90)
 1
       program hello world
2
       implicit none
3
       integer :: id, omp_get_thread_num
4
5
6
    !$omp parallel
7
       id = omp get thread num()
8
       write(*,'(1x,a,i3)') "Hello World! from", id
9
    !$omp end parallel
10
       end program hello world
11
$
  export OMP_NUM_THREADS=16
  # on Mike-II in bash shell, or inline setting
 ifort -o hello hello.f90 -openmp
$
```







```
C (hello c.c)
       #include <stdio.h>
 1
2
       #include <stdlib.h>
3
       #include <omp.h>
4
5
       int main() {
6
       int id;
7
8
       #pragma omp parallel {
9
       id = omp get thread num();
       printf("Hello World! from %3d\n", id);
10
11
12
```

- \$ export OMP_NUM_THREADS=16
 # on Mike-II in bash shell, or inline setting
- \$ icc -o hello hello.c -openmp





Loop-level parallelism

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- Loop-level parallelism is one of the fine-grained approaches supported by OpenMP;
- parallel do directive in Fortran and parallel for pragma in C/C++;

```
Fortran
  !$omp parallel do [clauses]
2
      do i = imin, imax, istep
3
         loop body ...
4
      end do
5
  [!$omp end parallel do]
                                              C/C++
  #pragma omp parallel for [clauses]
1
2
      for (i = imin; i <= imax; increment expr)</pre>
3
4
        loop body ...;
      }
5
```



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Loop-level parallelism

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• Other form of parallel loops:

```
Fortran
  !$omp parallel [clauses]
 !$omp do [clauses]
2
3
      do i = imin, imax, istep
4
         loop body ...
5
      end do
6
  !$omp end do
7
  !$omp end parallel
                                              C/C++
 #pragma omp parallel [clauses]
2 #pragma omp for [clauses]
3
      for (i = imin; i <= imax; increment expr)</pre>
4
5
        loop body ...;
6
```



Loop-level parallelism

• How about nested multiple loops? Where do we add parallel for, right above outer loop or inner loop?

```
1 for (i = imin; i < imax; increment_i) C/C++
2 {
3 #pragma omp parallel for
4 for (j = jmin; j <= jmax; increment_j)
5 { loop body ...; }
6 }</pre>
```

```
1 #pragma omp parallel for C/C++ (outer loop)
2 for (i = imin; i <= imax; increment_i)
3 {
4 for (j = jmin; j <= jmax; increment_j)
5 { loop body ...; }
6 }</pre>
```

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More words on parallel loops

- OpenMP only supports Fortran do loops and C/C++ for loops that the number of loop iterations is known for at run-time;
- However, it doesn't support other loops, including do-while and repeat-until loops in Fortran and while loops and do-while loops in C/C++. In these cases, the trip count of loop is unknown before entering the loop;
- Loop body has to follow parallel do Or parallel for immediately, and **nothing** in between them!
- There is an **implicit** barrier at the end of parallel do or for loops;
- All loops must have a single entry point and single exit point.
 We are **not** allowed to jump into a loop or branch out of a loop (but ...);





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- Once we have entered the parallel region, for some variables, multiple threads need to use the same named variables, but they store different values at different memory locations; these variables are called **private** variables;
- All private variables are undefined or uninitialized before entry and after exit from parallel regions;
- The **shared** variables are also necessary to allow data communication between threads;
- **Default** scope for variables: by default all the variables are considered to be **shared** in parallel regions, unless they are explicitly declared as **private**, **reduction**, or **other** types;
- Remember, Fortran and C/C++ may have different settings regarding default rules;





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- Let's see how we can do it, for instance, in parallel loops;
- OpenMP provides a means to change the default rules;
- Clauses default(none), default(private), and default(shared) in Fortran;
- But only default(none) and default(shared) in C/C++;

```
Fortran
  ALLOCATE( da(1:nsize), db(1:nsize) )
  !$omp parallel do default(none),
2
                                           X.
  !$omp private(i,temp),
3
                                            X.
  !$omp shared(imin,imax,istep,scale,da,db)
4
5
      do i = imin, imax, istep
6
         temp = scale * da(i)
7
         da(i) = temp + db(i)
8
      end do
9
  !$omp end parallel do
```





- OpenMP reduction operations;
- The reduction variable is very special that it has both characters of private and shared variables;
- Compiler needs to know what type of operation is associated with the reduction variable; operation = +, *, max, min, etc;
- reduction(operation: variables_list)

```
Fortran
  ALLOCATE( da(1:nsize) )
2
         prod = 1.0d0
3
  !$omp parallel do default(none), private(i),
                                                    &
4
  !$omp reduction(*: prod)
5
      do i = imin, imax, istep
         prod = prod * da(i)
6
                          What happens if we compile it?
7
      end do
8
  !$omp end parallel do
```



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- Two special "private" variables: firstprivate and lastprivate; they are used to initialize and finalize some private variables;
- firstprivate: upon entering a parallel do/for, the private variable for each slave thread has a copy of the master thread's value;
- lastprivate: upon exiting a parallel do/for, no matter which thread executed the **last** iteration (sequential), the private variable was copied back to the **master** thread;
- Why do we need them? (1) all private variables are undefined outside of a parallel region, (2) they provide a simply way to exchange data to some extent through these special private variables;





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- In a parallel region, a given variable can only be one of private, shared, Or reduction, but it can be both of firstprivate and lastprivate;

```
C/C++
   double ashift = shift
   #pragma omp parallel for default(none),
2
3
                firstprivate(ashift), shared(a),
4
                private(i)
5
6
       for (i = imin; i <= imax; ++i)</pre>
7
8
          ashift = ashift + (double) i :
9
          a[i] = a[i] + ashift ;
10
11
```

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- Exception of the default rules: Fortran and C/C++ behave differently;
- The index in a parallel loop is always private. The index in a sequential loop is also private in Fortran, but is shared in C by default!
- Is the following code correct?
- Has the loop j been parallelized?

```
1 #pragma omp parallel for
2 for (i = imin; i <= imax; ++i)
3 {
4 for (j = jmin; j <= jmax; ++j)
5 a[i][j] = (double) (i + j);
6 }
```

C/C++

Do we have the same issues in the Fortran version?







- Exception of the default rules. Fortran and C/C++ behave differently;
- The index in a parallel loop is always private. The index in a sequential loop is also private in Fortran, but is shared in C by default!
- Is the following code correct?
- Has the loop j been parallelized?

```
1 #pragma omp parallel for private(i,j) C/C++
2 for (i = imin; i <= imax; ++i)
3 {
4 for (j = jmin; j <= jmax; ++j)
5 a[i][j] = (double) (i + j);
6 }</pre>
```

Do we have the same issues in the Fortran version?





How to control loops?

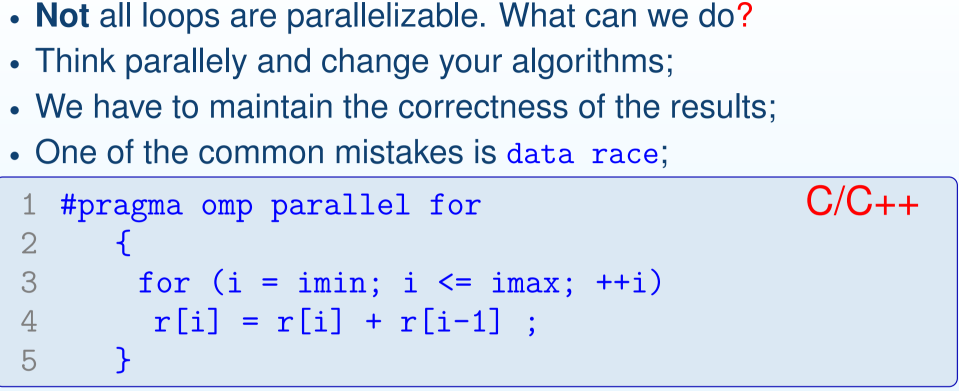
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- Parallelize multiple nested loops;
- The collapse(n) for nested parallel loops (n \geqslant 1);
- Each thread takes a chunk of the i loop and a chunk of the j loop at the same time;
- No statements in between;

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Restrictions on parallel loops

- Data race means that in a parallel region, the same memory location is referred by two or more statements, and at least one of them is a write operation;
- Data race requires more attention and might lead to incorrect results!







Restrictions on parallel loops

- A closer look at at the data race: let's run it on 2 threads and assume that r[0]=a; r[1]=b; r[2]=c; and imin=1; imax=2;
- Note, r[1] is referred **twice**, and thus we have two scenarios:

if thread () finished first	if thread	1 finished first
thread O	thread 1	thread 1	thread O
i = 1	i = 2	i = 2	i = 1
r[0]=a		r[1]=b	r[0]=a
r[1]=a+b	r[2]=a+b+c	r[2]=b+c	r[1]=b+a
_			
time		time	

OpenMP standard does not guarantee which thread finishes first or later;





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- How to identify data dependence and possibly remove it;
- A rule of thumb: (1) focus on those variables that are accessed twice or more. If the same memory location is only accessed (r/w) once, we don't have data dependence for that variable. (2) analyze all variables (mostly array elements) in a loop body. (3) pay attention to the global variables.

1 2	<pre>for (i = imin; i <= imax; ++i) { b[i] = b[i] + b[i-1] ; }</pre>	C/C++
1 2	for (i = imin; i <= imax; ++i) { b[i] = b[i] + a[i-1] ; }	
1 2	for (i = 2; i <= imax; i+=2) { a[i] = a[i] + a[i-1] ; }	



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- How to identify data dependence and possibly remove it;
- A rule of thumb: (1) focus on those variables that are accessed twice or more. If the same memory location is only accessed (r/w) once, we don't have data dependence for that variable. (2) analyze all variables (mostly array elements) in a loop body. (3) pay attention to the global variables.

1 2	<pre>for (i = imin; i <= imax; ++i) { b[i] = b[i] + b[i-1] ; }</pre>	C/C++
1 2	for (i = imin; i <= imax; ++i) { b[i] = b[i] + a[i-1] ; }	
1 2	<pre>for (i = 1; i <= imax/2; ++i) { b[i] = b[i] + b[i+imax/2] ; }</pre>	



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- **Dataflow** analysis for potential data dependence;
- Access sequence in two or more statements is critical;
- Three types of dataflows:

(1) flow dep. (2) anti-flow dep. (3) output dep.

3

- 1 for (i=0;i<10;++i)
- 2 { tmpi=sin(i);
- 3 b[i]=a[i]+tmpi; }





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- **Dataflow** analysis for potential data dependence;
- Access sequence in two or more statements is critical;
- Three types of dataflows:
 - (1) flow dep. (2) anti-flow dep. (3) output dep.

 S_1 writes the var. S_2 reads the same var.

(1) Flow dependence

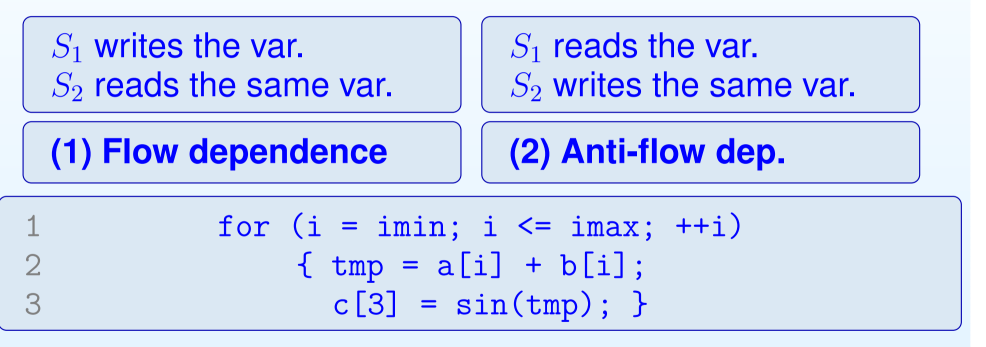
 S_1 reads the var. S_2 writes the same var.

(2) Anti-flow dep.





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- **Dataflow** analysis for potential data dependence;
- Access sequence in two or more statements is critical;
- Three types of dataflows:
 - (1) flow dep. (2) anti-flow dep. (3) output dep.







- LSU INFORMATIO TECHNOLOG SERVICES
- **Dataflow** analysis for potential data dependence;
- Access sequence in two or more statements is critical;
- Three types of dataflows:
 - (1) flow dep. (2) anti-flow dep. (3) output dep.

 S_1 writes the var.
 S_2 reads the same var. S_1 reads the var.
 S_2 writes the same var.(1) Flow dependence(2) Anti-flow dep.1for (i = imin; i <= imax; ++i)
{ tmp = a[i] + b[i];
c[3] = sin(tmp); }

Multiple statements write to the same memory location (3) Output dependence





- Is it possible to remove anti-flow and output dependences?
- The answer is yes in most cases: change the data structure;
- Can we parallelize the following serial code?

1	<pre>// array a[] and b[] are ready to use.</pre>	v0 C/C++
2	<pre>for (i=0; i<nsize, i++)<="" pre=""></nsize,></pre>	
3	a[i] = a[i+1] + b[i];	

- This is a typical example of **anti-flow** dependence;
 - (1) For the given i-th iteration, read a[i+1] (RHS, S_1); (2) In the next i+1-th iteration, write a[i+1] (LFS, S_2);





- Is it possible to remove anti-flow and output dependences?
- The answer is yes in most cases: change the data structure;
- Data dependence follows a certain pattern;
- Define a new array with **shifted** indices;

```
1 // make a new array by shifting index. v1 C/C++
2 #pragma omp parallel for
3 for (i=0; i<nsize, i++) a_new[i] = a[i+1];
4
5 // array a[] and b[] are ready to use.
6 #pragma omp parallel for
7 for (i=0; i<nsize, i++)
8 a[i] = a_new[i] + b[i];</pre>
```

• Now the data dependences were removed;





How to control loops again?



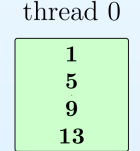
- OpenMP supports three loop schedulings as clauses: static, dynamic, and guided in the code, plus run-time scheduling;
- schedule(type[, chunk_size])

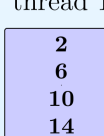
For static, if chunk size is given, loop iterations are divided into multiple blocks and each block contains chun size iterations. The iterations will be assigned to threads in a round-robin fashion. If chunk size is not present, the loop iterations will be (nearly) evenly divided and assigned to each thread.

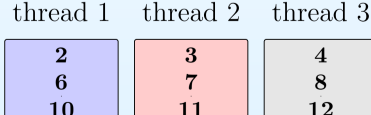
4

8

12







14 iterations on 4 threads in round-robin fashion





How to control loops again?

- For dynamic, if chunk_size is given, the partition is almost the same as those of static. The difference is that with static, the mapping between loop iterations and threads are done during **compilation**, while for dynamic, it will be done at **run-time** (therefore, more potentially overhead); if chunk_size is not present, then it was set to 1.
- The guided scheduling means the chunk_size assigned to threads decreases exponentially;
- Run-time scheduling: set the environment variable OMP_SCHEDULE;
- \$ export OMP_SCHEDULE=10, for instance;
- Each scheduling has its own pros and cons, so be careful with chunk_size and potential overhead;







- Let's consider the following question: For a given integer matrix, how do we count the total number of even matrix elements?
- This is how the serial code looks like:

```
v0 C/C++
  // count the number of even integers.
2
      counter even = 0;
3
     for (i=0; i<nosize; i++)</pre>
4
     for (j=0; j<nosize; j++)</pre>
5
      { itmp = matrix[i][j]%2;
        if (itmp == 0) ++counter even;
6
7
      }
8
      printf("Number of even integers is d\n", \
9
           counter even);
```

How do we parallelize these two loops?







- Generally, loop-level parallelism is considered fine-grained parallelism;
- Let's try something different: manually decompose the data;
- This is called **coarse-grained** parallelism:
 - We decompose the matrix in a row-wise and each thread takes care of one block (for the outer loop);
 - Then we define a counter array: each thread has its own counter;
 - Each thread counts the number of even matrix elements in its own block;
 - Finally, the **master** thread makes a summation;
- Reduce the overhead in the loop scheduling and maintain good scalability and performance;



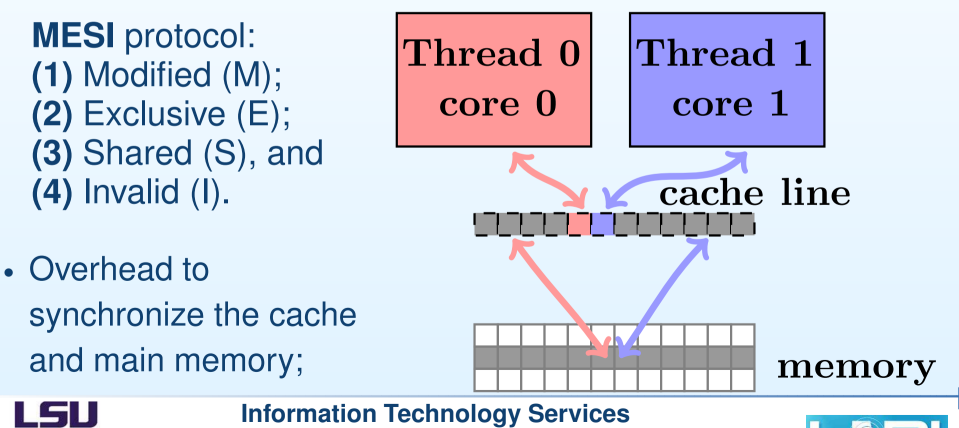






```
// define the arrays.
 2 counter = (int*)malloc(nothread*sizeof(int));
 3
    start idx = (int*)malloc(nothread*sizeof(int));
 4 end_idx = (int*)malloc(nothread*sizeof(int));
   chunk size = nosize / nothread + 1;
 5
                                               v1 C/C++
  for (id=0; id<nothread; id++)</pre>
 6
 7
   { start idx[id] = id*chunk size;
      end idx[id] = MIN(start idx[id]+chunk_size-1,\
 8
 9
     nosize-1);i }
10
    for(id=0;id<nothread;id++) counter[id] = 0;</pre>
   #pragma omp parallel private(id,i,j,itmp)
11
12
      { id = omp get thread num();
      for (i=start idx[id]; i<=end idx[id]; i++)</pre>
13
      for (j=0; j<nosize; j++)</pre>
14
      { itmp = matrix[i][j]%2;
15
      if (itmp == 0) ++counter[id]; } }
16
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```

- False sharing affects performance: the array counter [];
- Thread only accesses its array element, but adjacent;
- What is false sharing? If two or more threads that access the same cache line, at least one of them is to modify (write) that cache line.

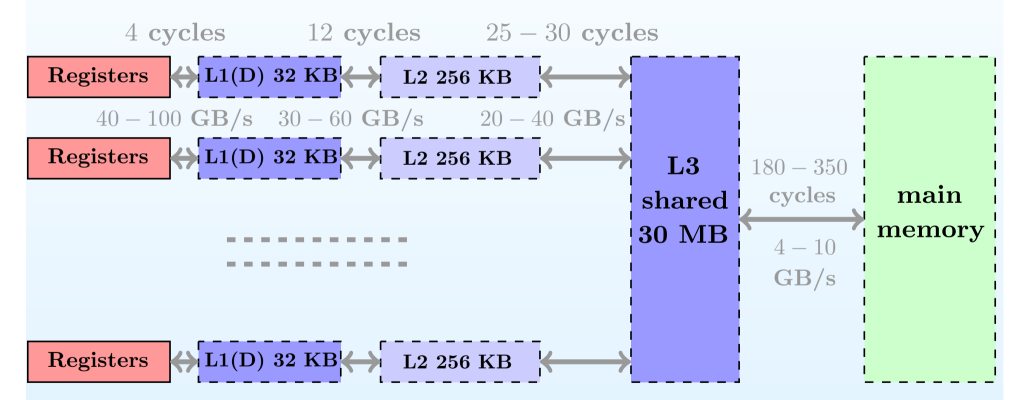


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- Understand memory and cache hierarchy:
- Lower-level cache (smaller, faster), but higher-level cache (larger, slower);



Memory Hierarchy of Xeon Sandy-Bridge Processor





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- Can we **remove** the false sharing?
- The shared array counter[] is the root of the problems;

```
v2 C/C++
   #pragma omp parallel \
2
      private(id,i,j,itmp,counter own)
3 { counter own = 0;
   id = omp get thread num();
4
5 for (i=start_idx[id]; i<=end_idx[id]; i++)</pre>
6 for (j=0; j<nosize; j++)
7 { itmp = matrix[i][j]%2;
8 if (itmp == 0) ++counter own; }
9
   counter[id] = counter own; }
10 counter even = 0;
   for(id=0; id<nothread; id++) \</pre>
11
12
      counter even += counter[id];
```



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- Any improvement?
- Has the false sharing been removed **completely**?
- What happens if we simple do parallel for?

```
v3 C/C++
  // count the number of even integers.
2
     counter even = 0;
3
  #pragma omp parallel for private(i,j,itmp)
4
     for (i=0; i<nosize; i++)</pre>
     for (j=0; j<nosize; j++)</pre>
5
6
     { itmp = matrix[i][j]%2;
7
  #pragma omp critical
     if (itmp == 0) ++counter even;
8
9
     }
```

- Using critical to protect the access to the shared variable;
- Any room to improve it?







• Using reduction instead of critical:

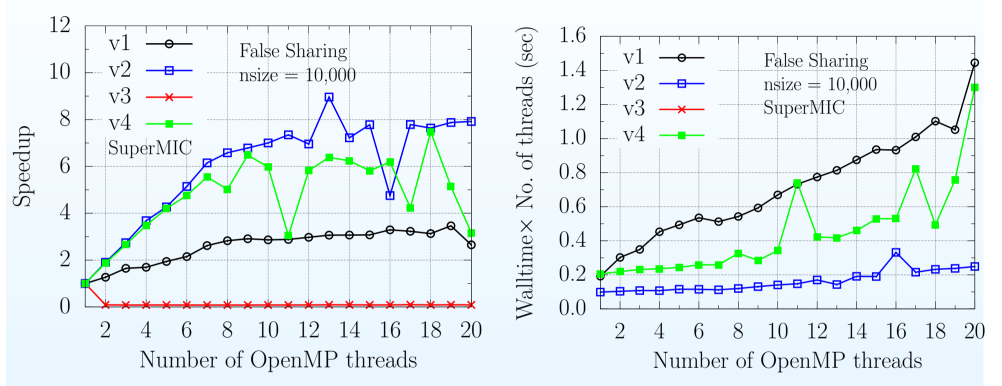
```
v4 C/C++
  // start a timer.
2
     time start = omp get wtime();
3
  // count the number of even integers.
  counter even = 0;
4
5
  #pragma omp parallel for private(i,j,itmp) \
     reduction(+:counter even)
6
     for (i=0; i<nosize; i++)</pre>
     for (j=0; j<nosize; j++)</pre>
8
     { itmp = matrix[i][j]%2;
9
   if (itmp == 0) ++counter even;
10
11
      }
      time_end = omp_get_wtime();
12
   printf("Elapsed time (sec) is \
13
      %.5f\n",time end-time start);
14
```



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• Compare the performance of the different versions;



- Which version is the best?
- In the ideal case, the walltime \times No. of threads is **flat**;
- The coarse-grained data decomposition(v2) is the best in terms of both speedup and walltime;

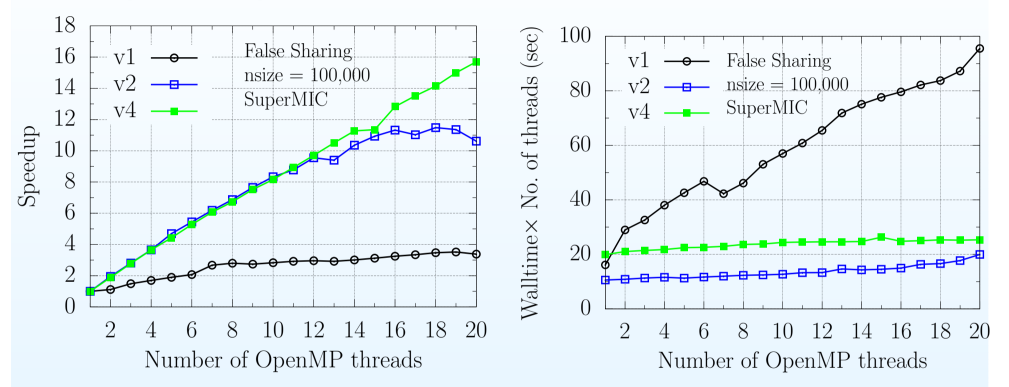


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• Compare the performance of the different versions;



- Which version is the best?
- Performance also depends on the **problem size**;
- The coarse-grained data decomposition(v2) is the best in terms of the speedup (v4) and walltime (v2);







v1: manually decomposition with false sharing;

- v2: manually decomposition with false sharing removed
 (almost);
- **v3**: using parallel for with critical;
- v4: using parallel for with reduction;
- Don't confuse **false sharing** with **data race**!
- Data race affects the correctness of your results;
- False sharing affects the code performance, but has nothing to do with the correctness of data;
- Both are caused by shared variables;

"Sharing is the root of all contention!" (H. Sutter 2009)











- ILS U INFORMATION TECHNOLOGY SERVICES
- The Intel MKL contains BLAS, LAPACK, Sparse BLAS and solvers, FFT routines, ...
- All BLAS/LAPACK routines were written in Fortran, and Intel compiler provides the flag -mkl to link to it. It also supports a CBLAS wrapper for C/C++ code;
- In some cases we need to call the MKL routines in a parallel region. Note the problems of oversubscribing resources;
- -mkl=parallel Uses the threaded MKL routines. It is the same as -mkl (default);
- -mkl=sequential Uses the non-threaded MKL routine.
- -mkl=cluster Uses the cluster part and the sequential part of the MKL routines on multiple nodes. It is possible to link to the cluster part and parallel routines at the same time;





- Consider matrix-matrix products that need to be repeated multiple times;
- Link it to the MKL routine dgemm in Fortran or cblas_dgemm routine in C;
- 1 // repeat "iteration" times: C = A×B. C/C++
 2 #pragma omp parallel for
 3 for (k=0; k<iteration; k++)
 5 { cblas_dgemm(CblasRowMajor, CblasNoTrans, \
 6 CblasNoTrans, nsize, nsize, nsize, \
 7 alpha, matrix_a, nsize, matrix_b, nsize, \
 8 beta, matrix_c, nsize); }</pre>
 - The MKL routine was embedded in the parallel region;
 - Explicit OpenMP threading and implicit MKL threading;







- Compile it with -openmp -mkl flags;
- On a **compute node**, run the following commands and monitor the **load** average:
- (1) Without setting anything but run \$./threaded_dgemm How many threads from **OpenMP parallel for** and how many threads from the implicit **MKL** routine?
- (2) How do we **explicitly** control the no. of the **OpenMP** threads and the no. of the **MKL** threads?
 - \$ OMP_NESTED=true MKL_DYNAMIC=false \
 - OMP_NUM_THREADS=2 MKL_NUM_THREADS=4
 - ./threaded_dgemm

In this case, we ask 2 OpenMP threads and each of them spawns 4 MKL threads;







- OMP_NUM_THREADS controls the number of OpenMP threads;
- MKL_NUM_THREADS controls the number of MKL threads;
- Setting MKL_NUM_THREADS=1 at run-time is kind of equivalent to -mkl=sequential during compilation;
- What if we **don't** explicitly specify both of the thread counts? \$ OMP NESTED=true MKL DYNMAIC=false ./threaded dgemm
- How many threads spawned in total?
- OMP_NESTED enable/disable **nested parallel regions**. If not defined, nested parallel regions are **disabled** by default;
- MKL_DYNAMIC=true allows the run-time system to detect and decide how many threads need to spawn. This is good to avoid oversubscription. If it is false, each **MKL** thread has a chance to spawn the max of thread counts, which is bad;







- The **default** value of MKL_DYNAMIC is true.
- By default the MKL routines will use only 1 thread if they are called in an OpenMP parallel region;
- "I'm not a programmer, why should I be concerned about it?"
- Some third-party applications or packages that were built on the top of the Intel MKL may **overwrite** the default behavior at run-time (call omp_set_nested() and mkl_set_dynamic());
- (1) Modify the threaded_dgemm.c code by adding omp_set_nested(1); mkl_set_dynamic(0);
- (2) Compile it and run it with ./thread_dgemm
- (3) Are there any problems and how can we fix it?
- (4) Run it with \$ OMP_NUM_THREADS=4 ./threaded_dgemm How many threads in total?
- Use the env OMP_NUM_THREADS to control it;







Non-loop-level parallelism





Fortran

C/C++

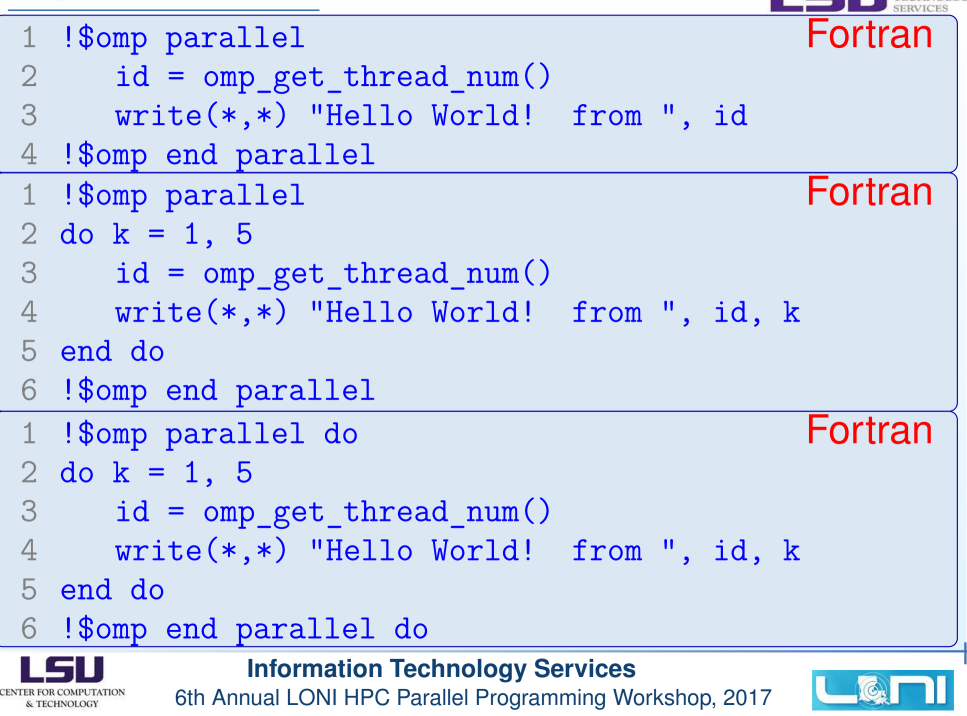
Parallel regions

- In addition to parallel do or for, most importantly, OpenMP supports the parallelism beyond loop levels;
- 1 !\$omp parallel [clauses]
- 2 code block
- 3 !\$omp end parallel
- 1 #pragma omp parallel [clauses]
 2 { code block ; }
 - { code block ; }
- Each thread in the parallel team executes the same block of code, but with different data;
- In parallel directives, clauses include: private(list), shared(list), reduction(operation : list), default(none | private | shared), if(logical operation), copyin(list);





Any differences?



Global variables in OpenMP



- In addition to automatic or static variables in Fortran and C/C++, we also need global variables;
- common blocks or modules in Fortran, while globals in C/C++, and we might have issues with private variables;
- Global/local variables between different code units for a given thread;
- Private/shared variables between multiple threads in a given code unit;
- The default data scoping rule is only apply to its lexical region, and all rest are shared; How can we make private variables "propagate" to other code units?
- OpenMP introduced the threadprivate directive to solve data scoping issues;





Global variables in OpenMP

- !\$omp threadprivate (list_common_variables) in Fortran;
- #pragma omp threadprivate (list_variables) in C/C++;
- We have global but private variables;
- The threadprivate variables are special private variables; thus thread a cannot access the threaprivate variables stored on thread b;
- The threadprivate variables persist from one parallel region to another, because they are globals;
- Furthermore, OpenMP supports the copyin (list) clause to initialize global variables on slave threads to be the values on the master thread;
- #pragma omp parallel copyin (a,b,c) { code block; }
- Sounds similar to the Intel Xeon Phi programming?





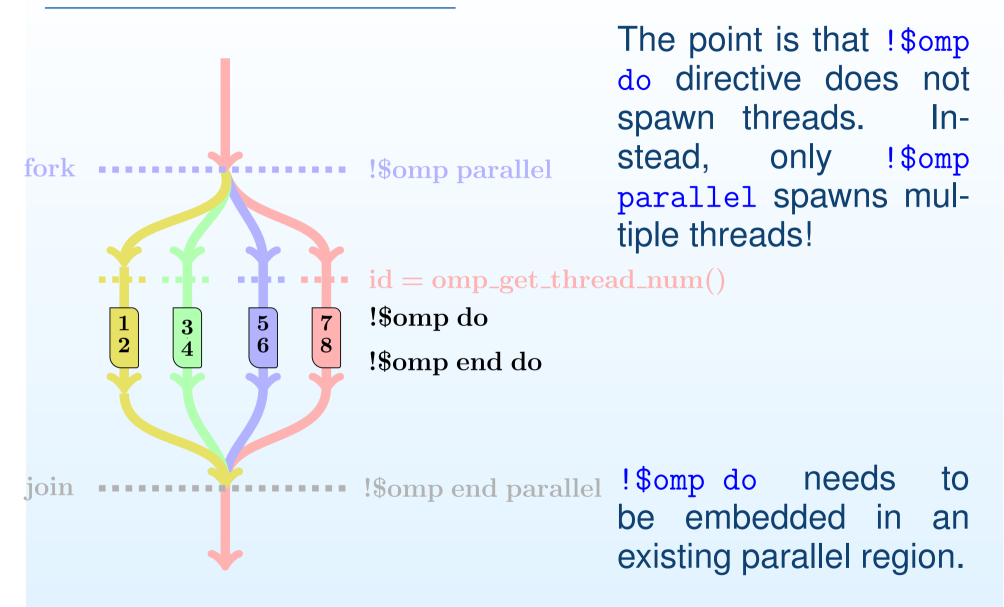


```
Fortran
 1 program mapping
2 implicit none
3 integer :: i, id, nothread, &
4
           omp_get_thread_num, omp_get_num_threads
5
6
   !$omp parallel private (k,id), shared(nothread)
7
      id = omp_get_thread_num()
8
      nothread = omp get num threads()
9
   !$omp do
10
   do k = 1, 40
11 write(*,'(1x,2(a,i4))') "id = ",id, " k = ",k
12 end do
13
  !$omp end do [nowait]
   !$omp end parallel
14
15 end program mapping
```













- Work-sharing constructs do not spawn multiple threads; they need to be embedded in a parallel region; if not, only one thread will run work-sharing constructs;
- There is an **implicit** barrier at the **end** of a work-sharing construct, but no implicit barrier upon the entry to it;
- Three work-sharing constructs:

!\$omp	do	#pragma	for
!\$omp	section	<pre>#pragma</pre>	section
!\$omp	single	<pre>#pragma</pre>	single

- A thread may work on zero, one, or more omp sections; but only one thread runs omp single at a given time;
- Be sure there are no data dependencies between sections;
- All threads must encounter the same workflow (though it may or may not execute the same code block at run-time);



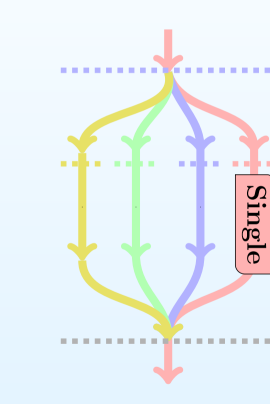
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6

!\$omp do !\$omp section #pragma for #pragma section do / 7 5 3 S_3

8



 S_2

!\$omp single

#pragma single



for

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```
C/C++
 1 #include <omp.h>
2 #define nsize 500
3 main() { int i, j, k ;
4 double a[nsize], b[nsize], c[nsize] ;
 5 for (k = 0; k \le nsize, ++k) {
 6 a[k] = (double) k; b[k] = a[k]; c[k] = 0.5*a[k];
 7
8
  #pragma omp parallel {
9
       #pragma omp sections {
10
       #pragma omp section { code block 1; }
       #pragma omp section { code block 2; }
11
12
       #pragma omp section { code block 3; }
13
14
           }
15
```



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- OpenMP provides the constructs for mutual exclusion: critical, atomic, master, barrier, and run-time routines;
 !\$omp critical [name] code block
 !\$omp end critical [name] in Fortran;
 #pragma omp critical [name] {code block;} in C/C++;
- [name] is an optional; But in Fortran, name here should be unique (cannot be the same as those of do loops or if/endif blocks, etc);
- At a given time, critical only allows **one** thread to run it, and all other threads also need to go through the critical section, but have to wait to enter the critical section;
- Don't jump into or branch out of a critical section;
- It is useful in a parallel region;
- It might have a tremendous impact on code performance;





• The other way to think of reduction variable (say addition):

```
Fortran
       tsum = 0.0d0; nsize = 10000
 2
   !$omp parallel private(temp), shared(tsum,nsize)
3
       temp = 0.0d0
4
   !$omp do
 5
       do i = 1, nsize
 6
           temp = temp + array(i)
7
       end do
8
   !$omp end do
9
   !$omp critical
10
11
          tsum = tsum + temp
12
   !$omp end critical
13
  !$omp end parallel
14
```







• Using atomic to protect a shared variable: C/C++1 #include <omp.h> 2 #include <stdio.h> 3 #include <stdlib.h> 4 #define nsize 1000 5 int main () $\{$ 6 int i; double x = 0.0, answer; #pragma omp parallel for private(i) shared(x) { 8 for (i = 1; i <= nsize; ++i) { 9 #pragma omp atomic x += (double) i; } 10 11 answer = (double) 0.5*(nsize+1)*nsize; 12 printf("%f n", x); 13 printf("correct answer is %f\n", answer); 14 }

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OpenMP run-time libraries

- integer omp_get_num_threads()
 int omp_get_num_threads(void)
 # No. of threads in the current collaborating parallel region;
- integer omp_get_thread_num()
 int omp_get_thread_num(void)
 # Return the thread IDs in a parallel team;
- integer omp_get_num_procs()
 int omp_get_num_procs(void)
 # Get the number of "processors" available to the code;
- call omp_set_num_threads(num_threads)
 omp_set_num_threads(num_threads)
 # Set number of threads to be num_threads for the following
 parallel regions;
- omp_get_wtime() # Measure elapsed wall-clock time (in seconds) relative to an arbitrary reference time;



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Summary and Further Reading



- OpenMP loop-level, nested thread parallelism, non-loop level parallelism, synchronization, and run-time libraries;
- How to protect shared variables; pay attention to them and synchronization;
- Global and local variables in OpenMP programming (global private variables);
- Data race and false sharing;
- Develop a defensive programming style;

Questions?

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