## Parallel Programming with OpenMP

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## A Programmer's View of OpenMP

- What is OpenMP?
  - Open specification for Multi-Processing
  - "Standard" API for defining multi-threaded shared-memory programs
  - <u>openmp.org</u> Talks, examples, forums, etc.
- OpenMP is a portable, threaded, shared-memory programming *specification* with "light" syntax
  - Exact behavior depends on OpenMP implementation!
  - Requires compiler support (<u>C</u> or Fortran)
- OpenMP will:
  - Allow a programmer to separate a program into *serial regions* and *parallel regions*, rather than T concurrently-executing threads.
  - Hide stack management
  - Provide synchronization constructs
- OpenMP will not:
  - Parallelize automatically
  - Guarantee speedup
  - Provide freedom from data races

```
int main() {
```

// Do this part in parallel

printf( "Hello, World!\n" );

```
return 0;
}
```

#### Motivation – OpenMP

omp set num threads(4);

#pragma omp parallel

// Do this part in parallel



#### All OpenMP directives begin: #pragma

```
printf( "Hello, World!\n" );
}
return 0;
}
```

int main() {

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#### **OpenMP parallel region construct**

- Block of code to be executed by multiple threads in parallel
- Each thread executes the same code redundantly (SPMD)
  - Work within work-sharing constructs is distributed among the threads in a team
- Example with C/C++ syntax

#pragma omp parallel [clause[clause]...]new-line
 structured-block

• clause can include the following:

private (list)

shared (list)

Example: OpenMP default is *shared* variables
 To make private, need to declare with pragma:

 #pragma omp parallel private (x)

## **OpenMP Programming Model - Review**

• Fork - Join Model:



- OpenMP programs begin as single process (*master thread*) and executes sequentially until the first parallel region construct is encountered
  - FORK: Master thread then creates a team of parallel threads
  - Statements in program that are enclosed by the parallel region construct are executed in parallel among the various threads
  - JOIN: When the team threads complete the statements in the parallel region construct, they synchronize and terminate, leaving only the master thread

#### parallel Pragma and Scope – More Examples

#pragma omp parallel num threads(2) x=1; y=1+x;x=1;
y=1+x; X=1; v=1+x;

#### X and y are shared variables. There is a risk of data race

## parallel Pragma and Scope - Review

#pragma omp parallel

{
 x=1;
 y=1+x;
}



Thread 0 X=1; y=1+x;



X and y are shared variables. There is a risk of data race

## parallel Pragma and Scope - Review

```
#pragma omp parallel num_threads(2)
{
    x=1; y=1+x;
}
```



#### X and y are shared variables. There is a risk of data race

#### **Divide for-loop for parallel sections**





#### for (int i=0; i<8; i++) x[i]=0;</pre>



#pragma omp parallel for
{
 for (int i=0; i<8; i++)
 x[i]=0;</pre>

}

System divides loop iterations to threads

Id=0;	Id=1;	Id=2;	Id=3;
<b>x</b> [0]=0;	<b>x</b> [1]=0;	<b>x</b> [2]=0;	<b>x</b> [3]=0;
X[4]=0;	X[5]=0;	X[6]=0;	X[7]=0;

## **OpenMP Data Parallel Construct: Parallel Loop**

- Compiler calculates loop bounds for each thread directly from *serial* source (computation decomposition)
- Compiler also manages data partitioning
- Synchronization also automatic (barrier)

Serial Program:	Parallel Program:
void main()	void main()
{	{
double Res[1000];	double Res[1000];
	#pragma omp parallel for
for(int i=0;i<1000;i++) {	for(int i=0;i<1000;i++) {
do_huge_comp(Res[i]);	do_huge_comp(Res[i]);
}	}
}	}

#### **Programming Model – Parallel Loops**

- Requirement for parallel loops
  - No data dependencies (reads/write or write/write pairs) between iterations!
- Preprocessor calculates loop bounds and divide iterations among parallel threads

```
#pragma omp parallel for
for( i=0; i < 25; i++ )
{
    printf("Foo");
}</pre>
```



#### for (i=0; i<max; i++) zero[i] = 0;</pre>

- Breaks *for loop* into chunks, and allocate each to a separate thread
  - e.g. if max = 100 with 2 threads: assign 0-49 to thread 0, and 50-99 to thread 1
- Must have relatively simple "shape" for an OpenMP-aware compiler to be able to parallelize it
  - Necessary for the run-time system to be able to determine how many of the loop iterations to assign to each thread
- No premature exits from the loop allowed
  - i.e. No break, return, exit, goto statements outside of
    - outside of any pragma block

In general,

## **Parallel** Statement Shorthand



• Also works for sections

## Numerical Integration



Mathematically, we know that:

$$\int_{0}^{1} \frac{4.0}{(1+x^2)} \, dx = \pi$$

We can approximate the integral as a sum of rectangles:

 $\sum_{i=0}^{N} F(\mathbf{x}_{i}) \Delta \mathbf{x} \approx \pi$ 

Where each rectangle has width  $\Delta x$  and height  $F(x_i)$  at the middle of interval i.

#### Sequential Calculation of $\pi$ in C

```
#include <stdio.h> /* Serial Code */
static long num steps = 100000;
double step;
void main () {
    int i;
    double x, pi, sum = 0.0;
   step = 1.0/(double)num steps;
    for (i = 1; i <= num steps; i++) {</pre>
     x = (i - 0.5) * step;
      sum = sum + 4.0 / (1.0 + x*x);
   }
                                                F(x_i)\Delta x \approx \pi
   pi = sum / num steps;
   printf ("pi = %6.12f\n", pi);
```

#### **Parallel OpenMP Version (1)**

```
#include <omp.h>
#define NUM THREADS 4
static long num steps = 100000; double step;
void main () {
  int i; double x, pi, sum[NUM THREADS];
  step = 1.0/(double) num steps;
  #pragma omp parallel private ( i, x )
    int id = omp get thread num();
    for (i=id, sum[id]=0.0; i< num steps; i=i+NUM THREADS)</pre>
      x = (i+0.5) * step;
      sum[id] += 4.0/(1.0+x*x);
  for(i=1; i<NUM THREADS; i++)</pre>
    sum[0] += sum[i]; pi = sum[0] / num steps
  printf ("pi = %6.12f\n", pi);
}
```

#### **OpenMP Reduction**



- Problem is that we really want sum over all threads! Sum+=A[3]
- Reduction: specifies that 1 or more variables that are private to each thread are subject of reduction operation at end of parallel region: reduction(operation:var) where
  - *Operation*: operator to perform on the variables (var) at the end of the parallel region
  - Var: One or more variables on which to perform scalar reduction.

Sum+=A[1

#### **OpenMp: Parallel Loops with Reductions**

OpenMP supports reduce operation

```
sum = 0;
```

```
#pragma omp parallel for reduction(+:sum)
```

```
for (i=0; i < 100; i++) {
sum += array[i];
}
```

- Reduce ops and init() values (C and C++):
- + 0 bitwise & ~0 logical & 1
- 0 bitwise | 0 logical | 0
- \* 1 bitwise ^ 0

#### Calculating $\pi$ Version (1) - review

```
#include <omp.h>
#define NUM THREADS 4
static long num steps = 100000; double step;
void main () {
  int i; double x, pi, sum[NUM THREADS];
  step = 1.0/(double) num steps;
  #pragma omp parallel private ( i, x )
    int id = omp get thread num();
    for (i=id, sum[id]=0.0; i< num steps; i=i+NUM THREADS)</pre>
    Ł
      x = (i+0.5) * step;
      sum[id] += 4.0/(1.0+x*x);
  for(i=1; i<NUM THREADS; i++)</pre>
    sum[0] += sum[i]; pi = sum[0] / num steps
  printf ("pi = %6.12f\n", pi);
}
```

#### **Version 2: parallel for, reduction**

```
#include <omp.h>
#include <stdio.h>
/static long num steps = 100000;
double step;
void main ()
{
   int i; double x, pi, sum = 0.0;
   step = 1.0/(double) num steps;
#pragma omp parallel for private(x) reduction(+:sum)
   for (i=1; i<= num steps; i++) {</pre>
        x = (i-0.5) * step;
        sum = sum + 4.0/(1.0+x*x);
   }
   pi = sum / num steps;
 printf ("pi = %6.8f\n", pi);
}
```

#### Loop Scheduling in Parallel for pragma

```
#pragma omp parallel for
for (i=0; i<max; i++) zero[i] = 0;</pre>
```

- Master thread creates additional threads, each with a separate execution context
- All variables declared outside for loop are shared by default, except for loop index which is private per thread (Why?)
- Implicit "barrier" synchronization at end of for loop
- Divide index regions sequentially per thread
  - Thread 0 gets 0, 1, ..., (max/n)-1;
  - Thread 1 gets max/n, max/n+1, ..., 2\*(max/n)-1
  - Why?

#### **Impact of Scheduling Decision**

- Load balance
  - Same work in each iteration?
  - Processors working at same speed?
- Scheduling overhead
  - Static decisions are cheap because they require no run-time coordination
  - Dynamic decisions have overhead that is impacted by complexity and frequency of decisions
- Data locality
  - Particularly within cache lines for small chunk sizes
  - Also impacts data reuse on same processor

#### **OpenMP environment variables**

#### OMP\_NUM\_THREADS

- sets the number of threads to use during execution
- when dynamic adjustment of the number of threads is enabled, the value of this environment variable is the maximum number of threads to use
- For example,

```
setenv OMP_NUM_THREADS 16 [csh, tcsh]
export OMP_NUM_THREADS=16 [sh, ksh, bash]
```

#### **OMP\_SCHEDULE**

- applies only to do/for and parallel do/for directives that have the schedule type RUNTIME
- sets schedule type and chunk size for all such loops
- For example,

```
setenv OMP_SCHEDULE GUIDED,4 [csh, tcsh]
export OMP_SCHEDULE= GUIDED,4 [sh, ksh, bash]
```

#### Programming Model – Loop Scheduling

- schedule clause determines how loop iterations are divided among the thread team
  - **static([chunk])** divides iterations statically between threads
    - Each thread receives [chunk] iterations, rounding as necessary to account for all iterations
    - Default [chunk] is ceil ( # iterations / # threads )
  - dynamic([chunk]) allocates [chunk] iterations per thread, allocating an additional [chunk] iterations when a thread finishes
    - Forms a logical work queue, consisting of all loop iterations
    - Default [chunk] is 1
  - guided ([chunk]) allocates dynamically, but [chunk] is exponentially reduced with each allocation

#### Loop scheduling options



## **Programming Model – Data Sharing**

}

- Parallel programs often employ two types of data
  - Shared data, visible to all threads, similarly named
  - Private data, visible to a single thread (often stack-allocated)
- PThreads:
  - Global-scoped variables are shared
  - Stack-allocated variables are private
- OpenMP:
  - shared variables are shared
  - private variables are private

```
// shared, globals
```

int bigdata[1024];

```
void* foo(void* bar) {
  intpridate, stack
  int tid;
  #pragma omp parallel \
  /shearedulabigdagees \
   prhezee* ( tid )
} {
    /* Calc. here */
  }
```

#### **Programming Model - Synchronization**

<ul> <li>OpenMP Synchronization</li> <li>OpenMP Critical Sections <ul> <li>Named or unnamed</li> <li>No <i>explicit</i> locks / mutexes</li> </ul> </li> </ul>	<pre>#pragma omp critical {    /* Critical code here */ }</pre>
<ul> <li>Barrier directives</li> </ul>	#pragma omp barrier
<ul> <li>Explicit Lock functions</li> <li>When all else fails – may require flush directive</li> </ul>	<pre>omp_set_lock( lock l ); /* Code goes here */ omp_unset_lock( lock l );</pre>
<ul> <li>Single-thread regions within parallel regions</li> <li>master, single directive</li> </ul>	<pre>#pragma omp single { {    S /* Only executed once */ }</pre>

## Omp critical vs. atomic

```
int sum=0
#pragma omp parallel for
 for(int j=1; j <n; j++){
       int x = j^*j;
       #pragma omp critical
           Sum=sum+x;// One thread enters the critical section at a time.
        ł
* May also use
        #pragma omp atomic
          x += exper
          Faster, but can support only limited arithmetic operation such as
        •
          ++, --, +=, -=, *=, /=, &=, |=
```

• Elapsed wall clock time:

double omp\_get\_wtime(void);

- Returns elapsed wall clock time in seconds
- Time is measured per thread, no guarantee can be made that two distinct threads measure the same time
- Time is measured from "some time in the past," so subtract results of two calls to omp\_get\_wtime to get elapsed time

# Parallel Matrix Multiply: Run Tasks Ti in parallel on multiple threads

$$\begin{pmatrix} 1 \\ 1 & 2 \\ 3 & 4 \end{pmatrix} * \begin{pmatrix} 5 & 7 \\ 6 & 8 \end{pmatrix} = \begin{pmatrix} 1*5+2*6 & 1*7+2*8 \\ 3*5+4*6 & 3*7+4*8 \end{pmatrix} = \begin{pmatrix} 17 & 23 \\ 39 & 53 \end{pmatrix}$$

for 
$$i = 1$$
 to  $n$  do  
 $T_i$ : for  $j = 1$  to  $n$  do  
 $sum = 0;$   
for  $k = 1$  to  $n$  do  
 $sum = sum + a[i, k] * b[k, j];$   
endfor  
 $c[i, j] = sum;$   
endfor  
 $T_1$   $T_2$ 

# Parallel Matrix Multiply: Run Tasks Ti in parallel on multiple threads

$$\begin{pmatrix} 2 \\ 1 & 2 \\ 3 & 4 \end{pmatrix} * \begin{pmatrix} 5 & 7 \\ 6 & 8 \end{pmatrix} = \begin{pmatrix} 1*5+2*6 & 1*7+2*8 \\ 3*5+4*6 & 3*7+4*8 \end{pmatrix} = \begin{pmatrix} 17 & 23 \\ 39 & 53 \end{pmatrix}$$

for 
$$i = 1$$
 to  $n$  do  
 $T_i$ : for  $j = 1$  to  $n$  do  
 $sum = 0$ ;  
for  $k = 1$  to  $n$  do  
 $sum = sum + a[i, k] * b[k, j]$ ;  
endfor  
 $c[i, j] = sum$ ;  
endfor  
 $T_1$   $T_2$  33

#### **Matrix Multiply in OpenMP**



## **OpenMP Summary**

- OpenMP is a compiler-based technique to create concurrent code from (mostly) serial code
- OpenMP can enable (easy) parallelization of loop-based code with fork-join parallelism
  - •pragma omp parallel
  - •pragma omp parallel for
  - pragma omp parallel private ( i, x )
  - pragma omp atomic
  - •pragma omp critical
  - #pragma omp for reduction(+ : sum)
- OpenMP performs comparably to manually-coded threading
  - Not a silver bullet for all applications