# **Supercomputing in Plain English** Part V: Shared Memory Multithreading

### **Henry Neeman, Director**

OU Supercomputing Center for Education & Research University of Oklahoma Information Technology Tuesday March 3 2009





## This is an experiment!

### It's the nature of these kinds of videoconferences that FAILURES ARE GUARANTEED TO HAPPEN! NO PROMISES!

- So, please bear with us. Hopefully everything will work out well enough.
- If you lose your connection, you can retry the same kind of connection, or try connecting another way.
- Remember, if all else fails, you always have the toll free phone bridge to fall back on.





### **Access Grid**

#### This week's Access Grid (AG) venue: Titan. If you aren't sure whether you have AG, you probably don't.

Tue March 3	Titan
Tue March 10	NO WORKSHOP
Tue March 17	NO WORKSHOP
Tue March 24	Axon
Tue March 31	Cactus
Tue Apr 7	Walkabout
Tue Apr 14	Cactus
Tue Apr 21	Verlet

Many thanks to John Chapman of U Arkansas for setting these up for us.





If you want to use H.323 videoconferencing – for example, Polycom – then dial

### 69.77.7.203##12345

any time after 2:00pm. Please connect early, at least today.

For assistance, contact Andy Fleming of <u>KanREN</u>/Kan-ed (<u>afleming@kanren.net</u> or 785-865-6434).

KanREN/Kan-ed's H.323 system can handle up to 40 simultaneous H.323 connections. If you cannot connect, it may be that all 40 are already in use.

Many thanks to Andy and KanREN/Kan-ed for providing H.323 access.





We have unlimited simultaneous iLinc connections available.

- If you're already on the SiPE e-mail list, then you should receive an e-mail about iLinc before each session begins.
- If you want to use iLinc, please follow the directions in the iLinc e-mail.
- For iLinc, you <u>MUST</u> use either Windows (XP strongly preferred) or MacOS X with Internet Explorer.
- To use iLinc, you'll need to download a client program to your PC. It's free, and setup should take only a few minutes.
- Many thanks to Katherine Kantardjieff of California State U Fullerton for providing the iLinc licenses.





### **QuickTime Broadcaster**

If you cannot connect via the Access Grid, H.323 or iLinc, then you can connect via QuickTime:

#### rtsp://129.15.254.141/test\_hpc09.sdp

We recommend using QuickTime Player for this, because we've tested it successfully.

We recommend upgrading to the latest version at:

http://www.apple.com/quicktime/

When you run QuickTime Player, traverse the menus

File -> Open URL

Then paste in the rstp URL into the textbox, and click OK.

Many thanks to Kevin Blake of OU for setting up QuickTime Broadcaster for us.





### **Phone Bridge**

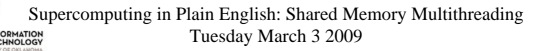
If all else fails, you can call into our toll free phone bridge:

1-866-285-7778, access code 6483137#

Please mute yourself and use the phone to listen.

Don't worry, we'll call out slide numbers as we go.

- Please use the phone bridge <u>ONLY</u> if you cannot connect any other way: the phone bridge is charged per connection per minute, so our preference is to minimize the number of connections.
- Many thanks to Amy Apon and U Arkansas for providing the toll free phone bridge.





No matter how you connect, please mute yourself, so that we cannot hear you.

- At OU, we will turn off the sound on all conferencing technologies.
- That way, we won't have problems with echo cancellation.
- Of course, that means we cannot hear questions.
- So for questions, you'll need to send some kind of text.

### Also, if you're on iLinc: **SIT ON YOUR HANDS! Please DON'T touch ANYTHING!**





## **Questions via Text: iLinc or E-mail**

Ask questions via text, using one of the following:

- iLinc's text messaging facility;
- e-mail to <u>sipe2009@gmail.com</u>.

All questions will be read out loud and then answered out loud.





## **Thanks for helping!**

- OSCER operations staff (Brandon George, Dave Akin, Brett Zimmerman, Josh Alexander)
- OU Research Campus staff (Patrick Calhoun, Josh Maxey, Gabe Wingfield)
- Kevin Blake, OU IT (videographer)
- Katherine Kantardjieff, CSU Fullerton
- John Chapman and Amy Apon, U Arkansas
- Andy Fleming, KanREN/Kan-ed
- This material is based upon work supported by the National Science Foundation under Grant No. OCI-0636427, "CI-TEAM Demonstration: Cyberinfrastructure Education for Bioinformatics and Beyond."





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### **Supercomputing Exercises**

Want to do the "Supercomputing in Plain English" exercises?

- The first several exercises are already posted at: <u>http://www.oscer.ou.edu/education.php</u>
- If you don't yet have a supercomputer account, you can get a temporary account, just for the "Supercomputing in Plain English" exercises, by sending e-mail to:

hneeman@ou.edu

Please note that this account is for doing the <u>exercises only</u>, and will be shut down at the end of the series.

• This week's OpenMP exercise will give you experience coding for, and benchmarking, OpenMP shared memory parallel code.



# OK Supercomputing Symposium 2009



2003 Keynote: Peter Freeman NSF Computer & Information Science & Engineering Assistant Director

2009 Keynote: Ed Seidel Director NSF Office of Cyberinfrastructure



2004 Keynote: Sangtae Kim NSF Shared Cyberinfrastructure Division Director





2005 Keynote: Walt Brooks NASA Advanced Supercomputing Division Director



2006 Keynote: Dan Atkins Head of NSF's Office of Cyberinfrastructure



kins 2007 Keynote: NSF's Jay Boisseau of Director er- Texas Advanced cture Computing Center U. Texas Austin



2008 Keynote: José Munoz Deputy Office Director/ Senior Scientific Advisor Office of Cyberinfrastructure National Science Foundation

FREE! Wed Oct 7 2009 @ OU http://symposium2009.oscer.ou.edu/

Parallel Programming Workshop FREE! Tue Oct 6 2009 @ OU Sponsored by SC09 Education Program FREE! Symposium Wed Oct 7 2009 @ OU

Supercomputing in Plain English: Shared Memory Multithreading

Tuesday March 3 2009



### **SC09 Summer Workshops**

- This coming summer, the SC09 Education Program, part of the SC09 (Supercomputing 2009) conference, is planning to hold two weeklong supercomputing-related workshops in Oklahoma, for **FREE** (except you pay your own travel):
- <u>At OU</u>: Parallel Programming & Cluster Computing, date to be decided, weeklong, for <u>FREE</u>
- <u>At OSU</u>: Computational Chemistry (tentative), date to be decided, weeklong, for <u>FREE</u>
- We'll alert everyone when the details have been ironed out and the registration webpage opens.
- Please note that you must apply for a seat, and acceptance <u>CANNOT</u> be guaranteed.





## Outline

- Parallelism
- Shared Memory Parallelism
- OpenMP





# Parallelism



### Parallelism

**Parallelism** means doing multiple things at the same time: you can get more work done in the same amount of time.





























More fish!





# What Is Parallelism?

<u>Parallelism</u> is the use of multiple processing units – either processors or parts of an individual processor – to solve a problem, and in particular the use of multiple processing units operating concurrently on different parts of a problem.

The different parts could be different tasks, or the same task on different pieces of the problem's data.





# **Kinds of Parallelism**

- Instruction Level Parallelism (the past two topics)
- Shared Memory Multithreading (our topic today)
- Distributed Memory Multiprocessing (next time)
- Hybrid Parallelism (Shared + Distributed)





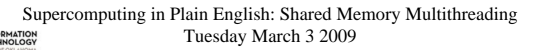
- <u>The Trees</u>: We like parallelism because, as the number of processing units working on a problem grows, we can solve <u>the same problem in less time</u>.
- <u>The Forest</u>: We like parallelism because, as the number of processing units working on a problem grows, we can solve <u>bigger problems</u>.





# **Parallelism Jargon**

- <u>*Threads*</u> are execution sequences that share a single memory area ("<u>*address space*</u>")
- <u>Processes</u> are execution sequences with their own independent, private memory areas
- ... and thus:
- *Multithreading*: parallelism via multiple <u>threads</u>
- *Multiprocessing*: parallelism via multiple processes
   Generally:
- Shared Memory Parallelism is concerned with <u>threads</u>, and
- Distributed Parallelism is concerned with **processes**.





# **Jargon Alert!**

In principle:

- "shared memory parallelism" → "multithreading"
- "distributed parallelism" → "multiprocessing"
- In practice, sadly, these terms are often used interchangeably:
- Parallelism
- *Concurrency* (not as popular these days)
- Multithreading
- Multiprocessing
- Typically, you have to figure out what is meant based on the context.





## **Amdahl's Law**

In 1967, Gene Amdahl came up with an idea so crucial to our understanding of parallelism that they named a <u>Law</u> for him:

$$S = \frac{1}{(1 - F_p) + \frac{F_p}{S_p}}$$

where *S* is the overall speedup achieved by parallelizing a code,  $F_p$  is the fraction of the code that's parallelizable, and  $S_p$  is the speedup achieved in the parallel part.<sup>[1]</sup>



What does Amdahl's Law tell us?

Imagine that you run your code on a zillion processors. The parallel part of the code could speed up by as much as a factor of a zillion.

For sufficiently large values of a zillion, the <u>parallel part</u> <u>would take zero time</u>!

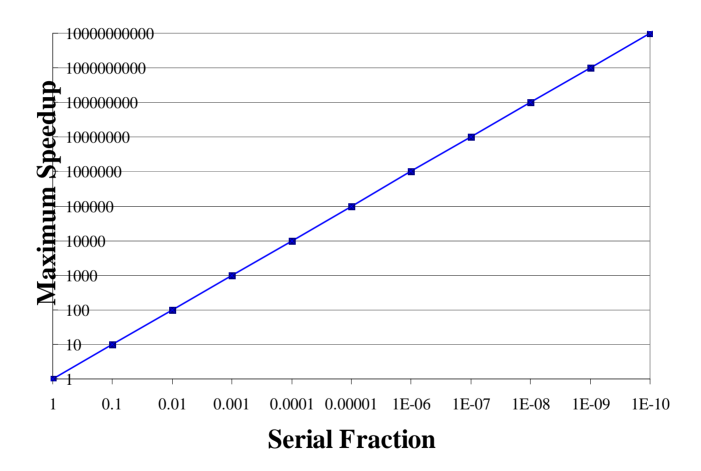
But, the <u>serial</u> (non-parallel) part would take the <u>same</u> <u>amount of time as on a single processor</u>.

So running your code on infinitely many processors would still take at least as much time as it takes to run just the serial part.





# Max Speedup by Serial %





## Amdahl's Law Example (F90)

```
PROGRAM amdahl_test
IMPLICIT NONE
REAL,DIMENSION(a_lot) :: array
REAL :: scalar
INTEGER :: index
READ *, scalar !! Serial part
DO index = 1, a_lot !! Parallel part
array(index) = scalar * index
END DO
END PROGRAM amdahl_test
```

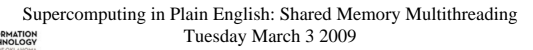
If we run this program on infinitely many CPUs, then the total run time will still be at least as much as the time it takes to perform the **READ**.





## **Amdahl's Law Example (C)**

```
int main ()
     float array[a lot];
     float scalar;
     int index;
     scanf("%f", scalar); /* Serial part */
     /* Parallel part */
     for (index = 0; index < a_lot; index++) {</pre>
        array(index) = scalar * index
If we run this program on infinitely many CPUs, then the total
run time will still be at least as much as the time it takes to
perform the scanf.
```





**<u>Rule of Thumb</u>**: When you write a parallel code, try to make as much of the code parallel as possible, because the <u>serial</u> <u>part will be the limiting factor</u> on parallel speedup.

Note that this rule will not hold when the *overhead* cost of parallelizing exceeds the parallel speedup. More on this presently.







The goal in parallelism is *linear speedup*: getting the speed of the job to increase by a factor equal to the number of processors.

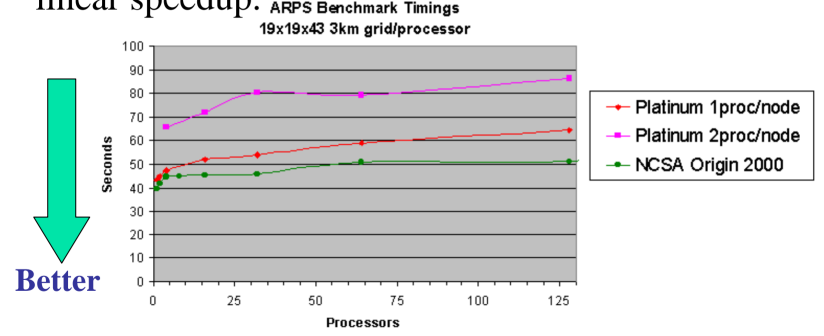
Very few programs actually exhibit linear speedup, but some come close.





## **Scalability**

### <u>Scalable</u> means "performs just as well regardless of how big the problem is." A scalable code has near linear speedup. ARPS Benchmark Timings



Platinum = NCSA 1024 processor PIII/1GHZ Linux Cluster Note: NCSA Origin timings are scaled from 19x19x53 domains.

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## **Strong vs Weak Scalability**

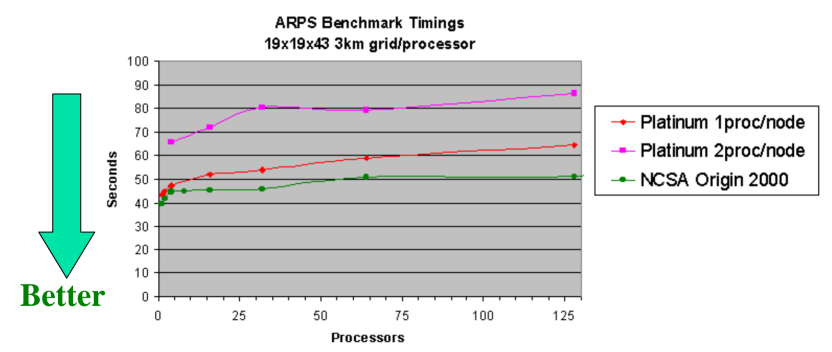
- Strong Scalability: If you double the number of processors, but you keep the problem size constant, then the problem takes half as long to complete.
- <u>Weak Scalability</u>: If you double the number of processors, and <u>double the problem size</u>, then the problem takes the <u>same amount of time</u> to complete.







#### This benchmark shows <u>weak</u> scalability.



Platinum = NCSA 1024 processor PIII/1GHZ Linux Cluster Note: NCSA Origin timings are scaled from 19x19x53 domains.

Supervised a second and the University of Oklahoma



# Granularity

<u>Granularity</u> is the size of the subproblem that each thread or process works on, and in particular the size that it works on between communicating or synchronizing with the others.
Some codes are <u>coarse grain</u> (a few very big parallel parts) and some are <u>fine grain</u> (many little parallel parts).
Usually, coarse grain codes are more scalable than fine

grain codes, because less of the runtime is spent managing the parallelism, so a higher proportion of the runtime is spent getting the work done.





# **Parallel Overhead**

**Parallelism isn't free**. Behind the scenes, the compiler and the hardware have to do a lot of *overhead* work to make parallelism happen.

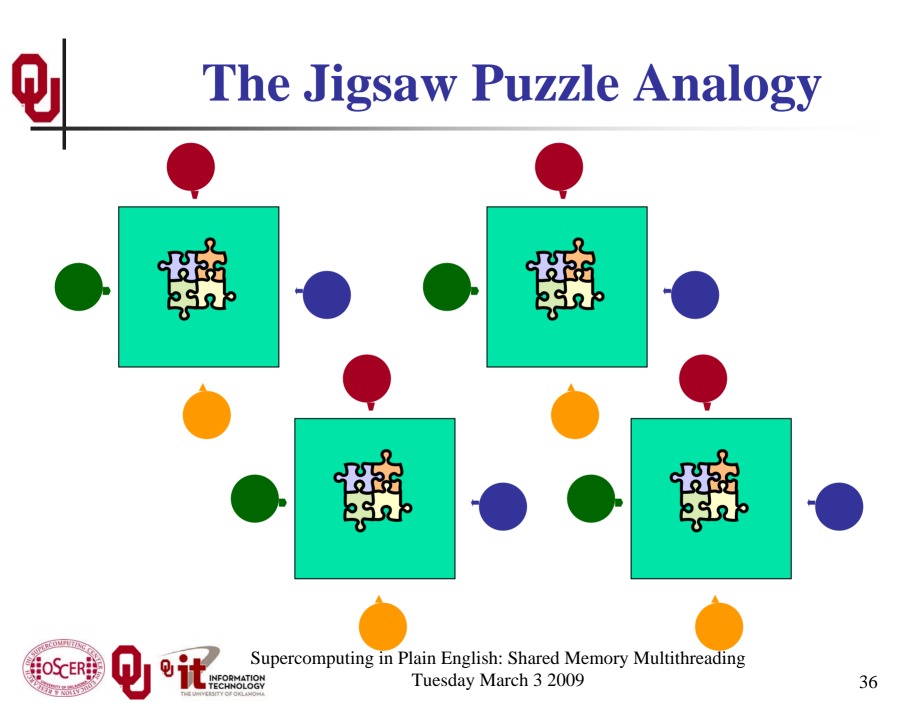
The overhead typically includes:

- **Managing** the multiple threads/processes
- **<u>Communication</u>** among threads/processes
- <u>Synchronization</u> (described later)



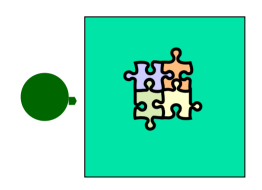








# **Serial Computing**

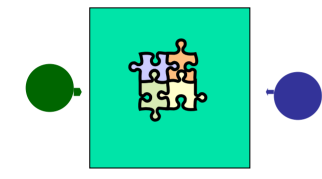


Suppose you want to do a jigsaw puzzle that has, say, a thousand pieces.

We can imagine that it'll take you a certain amount of time. Let's say that you can put the puzzle together in an hour.



### **Shared Memory Parallelism**

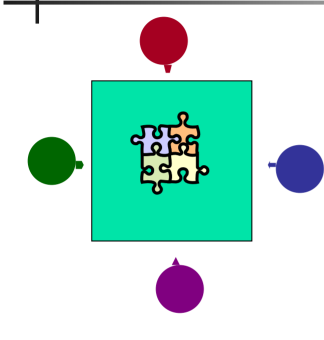


If Scott sits across the table from you, then he can work on his half of the puzzle and you can work on yours. Once in a while, you'll both reach into the pile of pieces at the same time (you'll *contend* for the same resource), which will cause a little bit of slowdown. And from time to time you'll have to work together (communicate) at the interface between his half and yours. The speedup will be nearly 2-to-1: y'all might take 35 minutes instead of 30.



# Q

### **The More the Merrier?**



Now let's put Paul and Charlie on the other two sides of the table. Each of you can work on a part of the puzzle, but there'll be a lot more contention for the shared resource (the pile of puzzle pieces) and a lot more communication at the interfaces. So y'all will get noticeably less than a 4-to-1 speedup, but you'll still have an improvement, maybe something like 3-to-1: the four of you can get it done in 20 minutes instead of an hour.

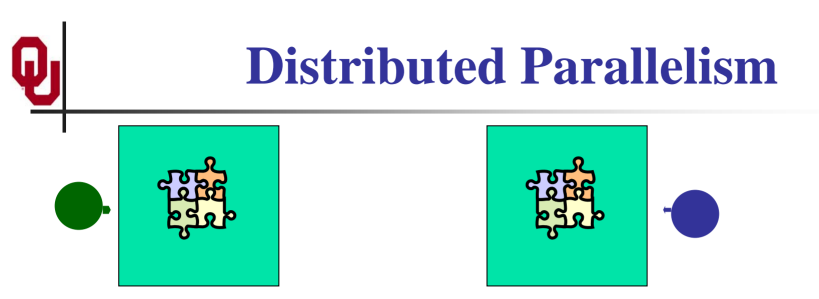


### **Diminishing Returns**

If we now put Dave and Tom and Horst and Brandon on the corners of the table, there's going to be a whole lot of contention for the shared resource, and a lot of communication at the many interfaces. So the speedup y'all get will be much less than we'd like; you'll be lucky to get 5-to-1.

So we can see that adding more and more workers onto a shared resource is eventually going to have a diminishing return.



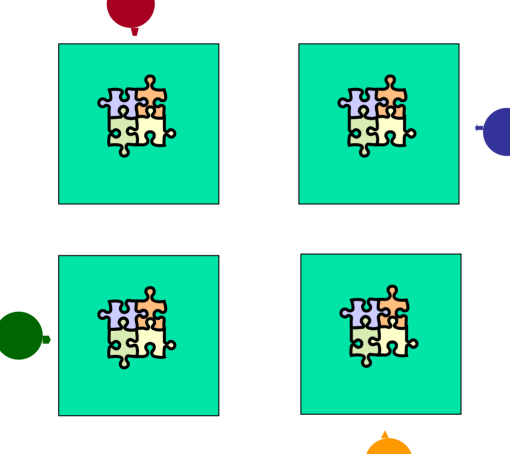


Now let's try something a little different. Let's set up two tables, and let's put you at one of them and Scott at the other. Let's put half of the puzzle pieces on your table and the other half of the pieces on Scott's. Now y'all can work completely independently, without any contention for a shared resource. **<u>BUT</u>**, the cost per communication is <u>**MUCH**</u> higher (you have to scootch your tables together), and you need the ability to split up (*decompose*) the puzzle pieces reasonably evenly, which may be tricky to do for some puzzles.

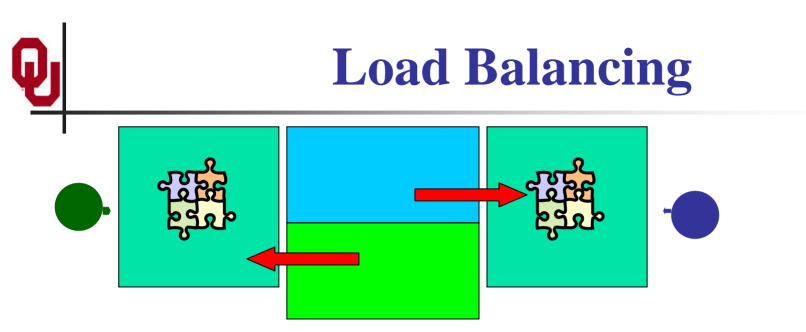




### **More Distributed Processors**



It's a lot easier to add more processors in distributed parallelism. But, you always have to be aware of the need to decompose the problem and to communicate among the processors. Also, as you add more processors, it may be harder to *load balance* the amount of work that each processor gets.



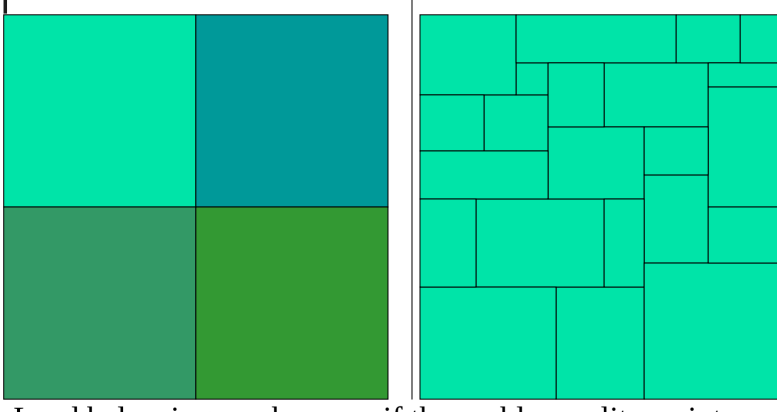
*Load balancing* means ensuring that everyone completes their workload at roughly the same time.

For example, if the jigsaw puzzle is half grass and half sky, then you can do the grass and Scott can do the sky, and then y'all only have to communicate at the horizon – and the amount of work that each of you does on your own is roughly equal. So you'll get pretty good speedup.





## Load Balancing

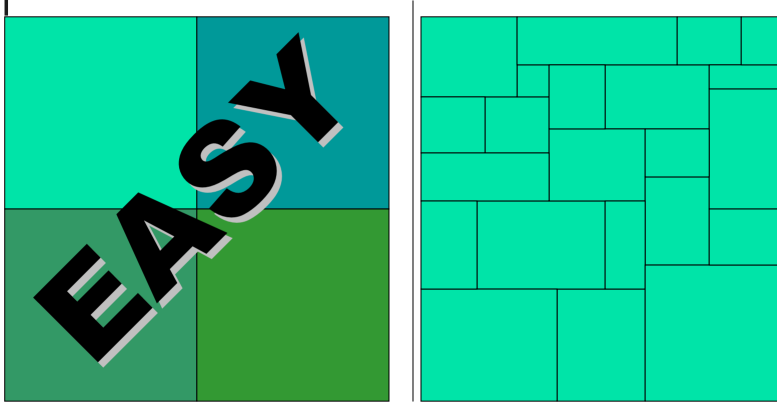


Load balancing can be easy, if the problem splits up into chunks of roughly equal size, with one chunk per processor. Or load balancing can be very hard.





## **Load Balancing**

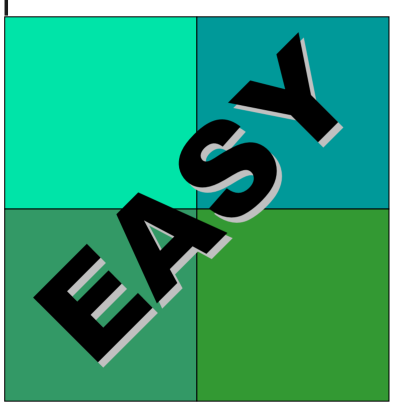


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## **Load Balancing**





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# How Shared Memory Parallelism Behaves

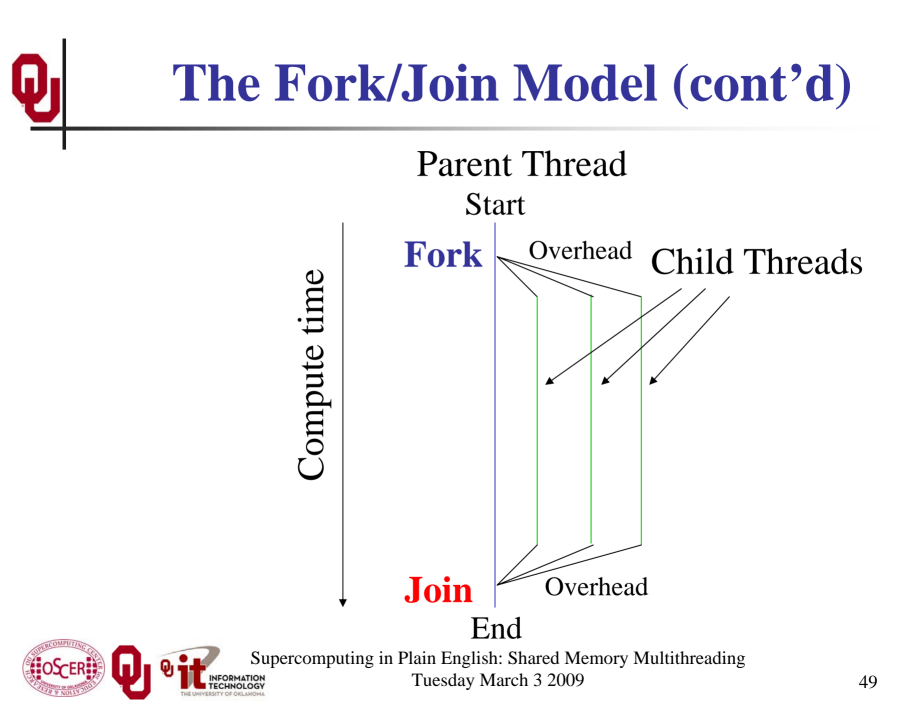


# **The Fork/Join Model**

Many shared memory parallel systems use a programming model called *Fork/Join*. Each program begins executing on just a single thread, called the *parent*.

- *Fork*: When a parallel region is reached, the *parent* thread *spawns* additional *child* threads as needed.
- *Join*: When the parallel region ends, the <u>child threads shut</u> <u>down</u>, leaving only the parent still running.







In principle, as a parallel section completes, the child threads shut down (join the parent), forking off again when the parent reaches another parallel section.

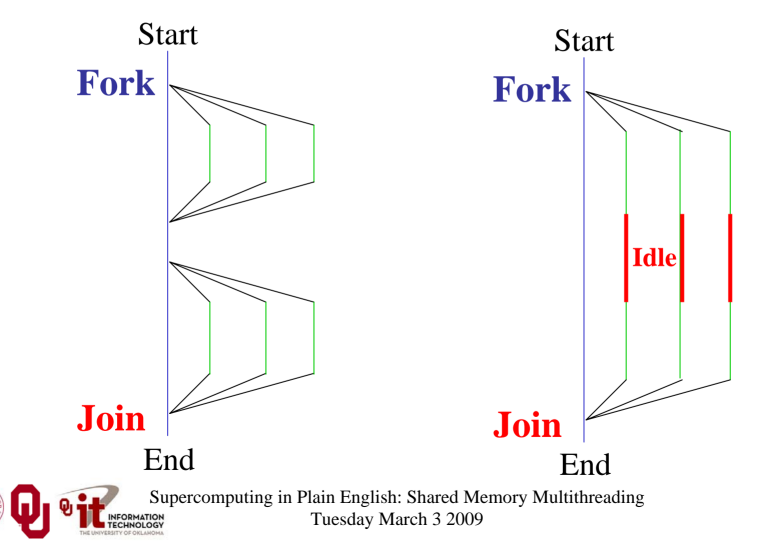
In practice, the child threads often continue to exist but are <u>idle</u>.

Why?





### **Principle vs. Practice**





# Why Idle?

- On some shared memory multithreading computers, the <u>overhead cost</u> of forking and joining is <u>high</u> compared to the cost of computing, so rather than waste time on overhead, the <u>children sit idle</u> until the next parallel section.
- On some computers, joining threads releases a program's control over the child processors, so they may not be available for more parallel work later in the run. <u>Gang</u> <u>scheduling</u> is preferable, because then all of the processors are guaranteed to be available for the whole run.







Most of this discussion is from [2], with a little bit from [3].



# What Is OpenMP?

OpenMP is a standardized way of expressing shared memory parallelism.

# OpenMP consists of *compiler directives*, *functions* and *environment variables*.

- When you compile a program that has OpenMP in it, if your compiler knows OpenMP, then you get an executable that can run in parallel; otherwise, the compiler ignores the OpenMP stuff and you get a purely serial executable.
- OpenMP can be used in Fortran, C and C++, but only if your preferred compiler explicitly supports it.





# **Compiler Directives**

- A <u>compiler directive</u> is a line of source code that gives the compiler special information about the statement or block of code that immediately follows.
- C++ and C programmers already know about compiler directives:

#### #include "MyClass.h"

- Many Fortran programmers already have seen at least one compiler directive:
- INCLUDE 'mycommon.inc'
- OR

#### INCLUDE "mycommon.inc"





OpenMP compiler directives in Fortran look like this: **!\$OMP** ....*stuff*...

In C++ and C, OpenMP directives look like:

- #pragma omp ...stuff...
- Both directive forms mean "the rest of this line contains OpenMP information."

Aside: "*pragma*" is the Greek word for "thing." Go figure.





Fortran

- **!\$OMP PARALLEL DO**
- **!\$OMP CRITICAL**
- **!\$OMP MASTER**
- **!\$OMP BARRIER**
- **!\$OMP SINGLE**
- **!\$OMP ATOMIC**
- **!\$OMP SECTION**
- **!\$OMP FLUSH**
- **!\$OMP ORDERED**

C++/C

#pragma omp parallel for #pragma omp critical #pragma omp master #pragma omp barrier #pragma omp single #pragma omp atomic #pragma omp section #pragma omp flush #pragma omp ordered

#### Note that we won't cover all of these.



# A First OpenMP Program (F90)

```
PROGRAM hello world
  IMPLICIT NONE
  INTEGER :: number of threads, this thread, iteration
  INTEGER, EXTERNAL :: omp get max threads,
                      omp get thread num
  number of threads = omp get_max_threads()
  WRITE (0,"(I2,A)") number_of_threads, " threads"
!SOMP PARALLEL DO DEFAULT(PRIVATE) &
! SOMP
                  SHARED(number of threads)
  DO iteration = 0, number of threads - 1
    this thread = omp get thread num()
    WRITE (0,"(A,I2,A,I2,A) ")"Iteration ", &
      iteration, ", thread ", this thread, &
 Se .
 æ
      ": Hello, world!"
  END DO
END PROGRAM hello world
```



# A First OpenMP Program (C)

```
int main ()
{
  int number of threads, this thread, iteration;
  int omp get max threads(), omp get thread num();
 number of threads = omp get max threads();
  fprintf(stderr, "%2d threads\n", number of threads);
\# pragma omp parallel for default(private) \setminus
                            shared(number of threads)
  for (iteration = 0;
       iteration < number_of_threads; iteration++) {</pre>
    this_thread = omp_get_thread_num();
    fprintf(stderr, "Iteration %2d, thread %2d: Hello, world!\n",
        iteration, this thread);
```



0,

### Running hello\_world

Л

<pre>% setenv OMP_NUM_THREADS 4</pre>						
% hello_world						
4 threads						
Iteration	Ο,	thread	0:	Hello,	world!	
Iteration	1,	thread	1:	Hello,	world!	
Iteration	З,	thread	3:	Hello,	world!	
Iteration	2,	thread	2:	Hello,	world!	
% hello_world						
4 threads						
Iteration	2,	thread	2:	Hello,	world!	
Iteration	1,	thread	1:	Hello,	world!	
Iteration	Ο,	thread	0:	Hello,	world!	
Iteration	З,	thread	3:	Hello,	world!	
% hello_world						
4 threads						
Iteration	1,	thread	1:	Hello,	world!	
Iteration	2,	thread	2:	Hello,	world!	
Iteration	0,	thread	0:	Hello,	world!	
Iteration	3,	thread	3:	Hello,	world!	

OND NUM BUDDENDO



From the **hello\_world** program, we learn that:

• At some point before running an OpenMP program, you must set an environment variable

OMP\_NUM\_THREADS

that represents the number of threads to use.

• The order in which the threads execute is **nondeterministic**.





The **PARALLEL DO** directive tells the compiler that the **DO** loop immediately after the directive should be executed in parallel; for example:

```
!$OMP PARALLEL DO
DO index = 1, length
array(index) = index * index
END DO
```

The iterations of the loop will be computed in parallel (note that they are independent of one another).

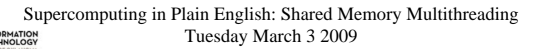




The **parallel for** directive tells the compiler that the **for** loop immediately after the directive should be executed in parallel; for example:

# pragma omp parallel for for (index = 0; index < length; index++) { array[index] = index \* index; }

The iterations of the loop will be computed in parallel (note that they are independent of one another).





### A Change to hello\_world

Suppose we do 3 loop iterations per thread:

#### DO iteration = 0, number\_of\_threads \* 3 - 1

<pre>% hello_world 4 threads</pre>						
Iteration	9,	thread	3: Hello, world!			
Iteration	Ο,	thread	0: Hello, world!			
Iteration	10,	thread	3: Hello, world!			
Iteration	11,	thread	3: Hello, world!			
Iteration	1,	thread	0: Hello, world!			
Iteration	2,	thread	0: Hello, world!			
Iteration	3,	thread	1: Hello, world!			
Iteration	6,	thread	2: Hello, world!			
Iteration	7,	thread	2: Hello, world!			
Iteration	8,	thread	2: Hello, world!			
Iteration	4,	thread	1: Hello, world!			
Iteration	5,	thread	1: Hello, world!			

Notice that the iterations are split into contiguous <u>*chunks*</u>, and each thread gets one chunk of iterations.





### Chunks

- By default, OpenMP splits the iterations of a loop into chunks of equal (or roughly equal) size, assigns each chunk to a thread, and lets each thread loop through its subset of the iterations.
- So, for example, if you have 4 threads and 12 iterations, then each thread gets three iterations:
- Thread 0: iterations 0, 1, 2
- Thread 1: iterations 3, 4, 5
- Thread 2: iterations 6, 7, 8
- Thread 3: iterations 9, 10, 11
- Notice that each thread performs its own chunk in deterministic order, but that the overall order is nondeterministic.



### **Private and Shared Data**

<u>*Private*</u> data are data that are owned by, and only visible to, a single individual thread.

<u>Shared</u> data are data that are owned by and visible to all threads.

(Note: In distributed parallelism, all data are private, as we'll see next time.)





# **Should All Data Be Shared?**

In our example program, we saw this:

**!\$OMP PARALLEL DO DEFAULT(PRIVATE) SHARED(number\_of\_threads)** 

What do **DEFAULT ( PRIVATE )** and **SHARED** mean?

- We said that OpenMP uses shared memory parallelism. So **PRIVATE** and **SHARED** refer to memory.
- Would it make sense for all data within a parallel loop to be shared?





## A Private Variable

#### Consider this loop: !\$OMP PARALLEL DO ... DO iteration = 0, number\_of\_threads - 1 this\_thread = omp\_get\_thread\_num() WRITE (0,"(A,I2,A,I2,A) ") "Iteration ", iteration, & & ", thread ", this\_thread, ": Hello, world!" END DO

Notice that, if the iterations of the loop are executed concurrently, then the loop index variable named **iteration** will be wrong for all but one of the threads.

Each thread should get its own copy of the variable named **iteration**.





### **Another Private Variable**

```
!$OMP PARALLEL DO ...
DO iteration = 0, number_of_threads - 1
this_thread = omp_get_thread_num()
WRITE (0,"(A,I2,A,I2,A)") "Iteration ", iteration, &
& ", thread ", this_thread, ": Hello, world!"
END DO
```

Notice that, if the iterations of the loop are executed concurrently, then **this\_thread** will be wrong for all but one of the threads.

Each thread should get its own copy of the variable named **this\_thread**.





### **A Shared Variable**

#### !\$OMP PARALLEL DO ... DO iteration = 0, number\_of\_threads - 1 this\_thread = omp\_get\_thread\_num() WRITE (0,"(A,I2,A,I2,A)"") "Iteration ", iteration, & & ", thread ", this\_thread, ": Hello, world!" END DO

Notice that, regardless of whether the iterations of the loop are executed serially or in parallel, **number\_of\_threads** will be correct for all of the threads.

All threads should share a single instance of **number\_of\_threads**.





### SHARED & PRIVATE Clauses

The **PARALLEL DO** directive allows extra <u>*clauses*</u> to be appended that tell the compiler which variables are shared and which are private:

!\$OMP PARALLEL DO PRIVATE(iteration,this\_thread) &

!\$OMP SHARED (number\_of\_threads)

This tells that compiler that **iteration** and **this\_thread** are private but that **number\_of\_threads** is shared.

(Note the syntax for continuing a directive in Fortran90.)





### **DEFAULT** Clause

- If your loop has lots of variables, it may be cumbersome to put all of them into **SHARED** and **PRIVATE** clauses.
- So, OpenMP allows you to declare one kind of data to be the default, and then you only need to explicitly declare variables of the other kind:
- !\$OMP PARALLEL DO DEFAULT(PRIVATE) &
   SHARED(number\_of\_threads)
  The default DEFAULT (so to speak) is SHARED, except for the
  - loop index variable, which by default is **PRIVATE**.



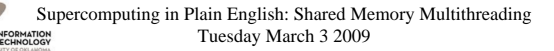


## **Different Workloads**

What happens if the threads have different amounts of work to do?

```
!$OMP PARALLEL DO
DO index = 1, length
    x(index) = index / 3.0
    IF (x(index) < 0) THEN
        y(index) = LOG(x(index))
    ELSE
        y(index) = 1.0 - x(index)
        END IF
END DO
The threads that finish early have to use
```

The threads that finish early have to wait.





## Chunks

- By default, OpenMP splits the iterations of a loop into chunks of equal (or roughly equal) size, assigns each chunk to a thread, and lets each thread loop through its subset of the iterations.
- So, for example, if you have 4 threads and 12 iterations, then each thread gets three iterations:
- Thread 0: iterations 0, 1, 2
- Thread 1: iterations 3, 4, 5
- Thread 2: iterations 6, 7, 8
- Thread 3: iterations 9, 10, 11

Notice that each thread performs its own chunk in deterministic order, but that the overall order is nondeterministic.





# **Scheduling Strategies**

OpenMP supports three scheduling strategies:

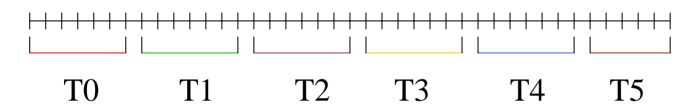
- <u>Static</u>: The default, as described in the previous slides good for iterations that are inherently load balanced.
- <u>Dynamic</u>: Each thread gets a chunk of a few iterations, and when it finishes that chunk it goes back for more, and so on until all of the iterations are done – good when iterations aren't load balanced at all.
- <u>Guided</u>: Each thread gets smaller and smaller chunks over time – a compromise.





# **Static Scheduling**

For  $N_i$  iterations and  $N_t$  threads, each thread gets one chunk of  $N_i/N_t$  loop iterations:

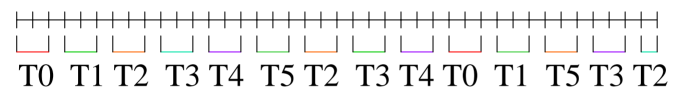


- Thread #0: iterations 0 through  $N_i/N_t-1$
- Thread #1: iterations  $N_i/N_t$  through  $2N_i/N_t-1$
- Thread #2: iterations  $2N_i/N_t$  through  $3N_i/N_t-1$
- Thread  $\#N_t$ -1: iterations  $(N_t$ -1) $N_i/N_t$  through  $N_i$ -1

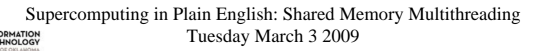


# **Dynamic Scheduling**

For  $N_i$  iterations and  $N_t$  threads, each thread gets a fixed-size chunk of *k* loop iterations:



- When a particular thread finishes its chunk of iterations, it gets assigned a new chunk. So, the relationship between iterations and threads is nondeterministic.
- Advantage: very flexible
- Disadvantage: high overhead lots of decision making about which thread gets each chunk





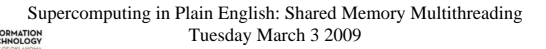
# **Guided Scheduling**

For  $N_i$  iterations and  $N_t$  threads, initially each thread gets a fixed-size chunk of  $k < N_i/N_t$  loop iterations:



#### T0 T1 T2 T3 T4 T5 2 3 4 1 0 2 5 4 2 3 1

- After each thread finishes its chunk of k iterations, it gets a chunk of k/2 iterations, then k/4, etc. Chunks are assigned dynamically, as threads finish their previous chunks.
- Advantage over static: can handle imbalanced load
- Advantage over dynamic: fewer decisions, so less overhead





# How to Know Which Schedule?

Test all three using a typical case as a *benchmark*.

Whichever wins is probably the one you want to use most of the time on that particular platform.

This may vary depending on problem size, new versions of the compiler, who's on the machine, what day of the week it is, etc, so you may want to benchmark the three schedules from time to time.





#### **SCHEDULE** Clause

The **PARALLEL DO** directive allows a **SCHEDULE** clause to be appended that tell the compiler which variables are shared and which are private:

**!**\$OMP PARALLEL DO ... SCHEDULE(STATIC)

This tells that compiler that the schedule will be static.

Likewise, the schedule could be **GUIDED** or **DYNAMIC**.

However, the very best schedule to put in the **SCHEDULE** clause is **RUNTIME**.

You can then set the environment variable **OMP\_SCHEDULE** to **STATIC** or **GUIDED** or **DYNAMIC** at runtime – great for benchmarking!





## **Synchronization**

**Jargon**: Waiting for other threads to finish a parallel loop (or other parallel section) before going on to the work after the parallel section is called *synchronization*.

Synchronization is <u>**BAD**</u>, because when a thread is waiting for the others to finish, it isn't getting any work done, so it isn't contributing to speedup.

So why would anyone ever synchronize?







Synchronizing is necessary when the code that follows a parallel section needs all threads to have their final answers.

```
!$OMP PARALLEL DO
DO index = 1, length
    x(index) = index / 1024.0
    IF ((index / 1000) < 1) THEN
        y(index) = LOG(x(index))
    ELSE
        y(index) = x(index) + 2
    END IF
END DO
! Need to synchronize here!
DO index = 1, length
    z(index) = y(index) + y(length - index + 1)
END DO
```



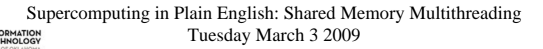


A *barrier* is a place where synchronization is forced to occur; that is, where faster threads have to wait for slower ones.

The **PARALLEL DO** directive automatically puts an invisible, implied barrier at the end of its **DO** loop:

- **!\$OMP PARALLEL DO** 
  - DO index = 1, length ... parallel stuff ... END DO
- ! Implied barrier
  - ... serial stuff ...

OpenMP also has an explicit **BARRIER** directive, but most people don't need it.





#### **Critical Sections**

A *critical section* is a piece of code that any thread can execute, but that only one thread can execute at a time. **!**SOMP PARALLEL DO DO index = 1, length ... parallel stuff ... !\$OMP CRITICAL(summing) sum = sum + x(index) \* y(index)**!\$OMP END CRITICAL(summing)** ... more parallel stuff ... END DO

What's the point?





# **Why Have Critical Sections?**

If only one thread at a time can execute a critical section, that slows the code down, because the other threads may be waiting to enter the critical section.

But, for certain statements, if you don't ensure *mutual exclusion*, then you can get nondeterministic results.





# **If No Critical Section**

```
!$OMP CRITICAL(summing)
```

```
sum = sum + x(index) * y(index)
!$OMP END CRITICAL(summing)
```

Suppose for thread #0, index is 27, and for thread #1, index is 92.

- If the two threads execute the above statement at the same time, **sum** could be
- the value after adding x(27) \* y(27), or
- the value after adding x(92) \* y(92), or
- garbage!

This is called a *race condition*: the result depends on who wins the race.





#### **Pen Game #1: Take the Pen**

We need two volunteers for this game.

- 1. I'll hold a pen in my hand.
- 2. You win by taking the pen from my hand.
- 3. One, two, three, go!
- Can we predict the outcome? Therefore, can we guarantee that we get the correct outcome?





We need two volunteers for this game.

- 1. I'll hold a pen in my hand.
- 2. You win by looking at the pen.
- 3. One, two, three, go!
- Can we predict the outcome? Therefore, can we guarantee that we get the correct outcome?





#### **Race Conditions**

A *race condition* is a situation in which multiple processes can change the value of a variable at the same time.

As in Pen Game #1 (Take the Pen), a race condition can lead to unpredictable results.

So, race conditions are **<u>BAD</u>**.





#### Reductions

- A <u>*reduction*</u> converts an array to a scalar: sum, product, minimum value, maximum value, location of minimum value, location of maximum value, Boolean AND, Boolean OR, number of occurrences, etc.
- Reductions are so common, and so important, that OpenMP has a specific construct to handle them: the **REDUCTION** clause in a **PARALLEL DO** directive.





#### **Reduction Clause**

```
total_mass = 0
!$OMP PARALLEL DO REDUCTION(+:total_mass)
D0 index = 1, length
   total_mass = total_mass + mass(index)
END D0 !! index = 1, length
```

#### This is equivalent to:

```
total_mass = 0
D0 thread = 0, number_of_threads - 1
    thread_mass(thread) = 0
END D0
$OMP PARALLEL D0
D0 index = 1, length
    thread = omp_get_thread_num()
    thread_mass(thread) = thread_mass(thread) + mass(index)
END D0 !! index = 1, length
D0 thread = 0, number_of_threads - 1
    total_mass = total_mass + thread_mass(thread)
END D0
```



## **Parallelizing a Serial Code #1**

#### **PROGRAM big\_science** ... declarations ...

DO ... ... parallelizable work ... END DO ... serial work ...



DO ... ... more parallelizable work ... END DO ... serial work ... ... etc ... END PROGRAM big\_science PROGRAM big\_science ... declarations ... **!\$OMP PARALLEL DO ...** DO ... ... parallelizable work ... END DO ... serial work ... **!**\$OMP PARALLEL DO ... DO ... ... more parallelizable work ... END DO ... serial work ... ... etc ... END PROGRAM big science

#### This way may have lots of **<u>synchronization</u>** overhead.





## **Parallelizing a Serial Code #2**

#### **PROGRAM** big\_science ... declarations ...

DO task = 1, numtasks CALL science\_task(...) END DO END PROGRAM big\_science

SUBROUTINE science\_task (...)
... parallelizable work ...

... serial work ...

... more parallelizable work ...

... serial work ...

... etc ... END PROGRAM big\_science **PROGRAM** big science ... declarations ... **!**\$OMP PARALLEL DO ... DO task = 1, numtasks CALL science task(...) END DO END PROGRAM big science SUBROUTINE science task ( ... ) ... parallelizable work ... **!**\$OMP MASTER ... serial work ... **!**\$OMP END MASTER ... more parallelizable work ... **!**SOMP MASTER ... serial work ... **!**SOMP END MASTER ... etc ... END PROGRAM big science



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#### **Thanks for helping!**

- OSCER operations staff (Brandon George, Dave Akin, Brett Zimmerman, Josh Alexander)
- OU Research Campus staff (Patrick Calhoun, Josh Maxey, Gabe Wingfield)
- Kevin Blake, OU IT (videographer)
- Katherine Kantardjieff, CSU Fullerton
- John Chapman and Amy Apon, U Arkansas
- Andy Fleming, KanREN/Kan-ed
- This material is based upon work supported by the National Science Foundation under Grant No. OCI-0636427, "CI-TEAM Demonstration: Cyberinfrastructure Education for Bioinformatics and Beyond."





# Thanks for your attention!





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