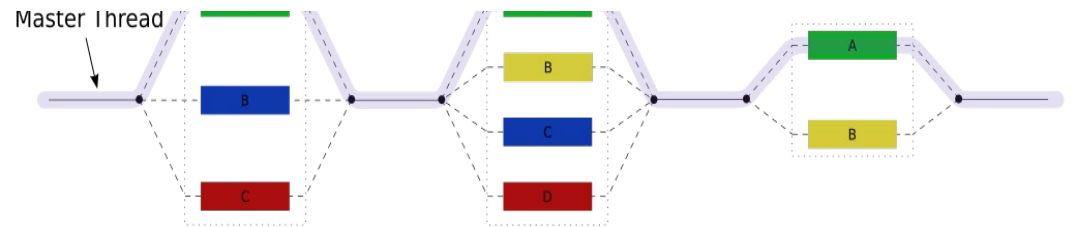


CS 470 Spring 2025

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OpenMP

OpenMP

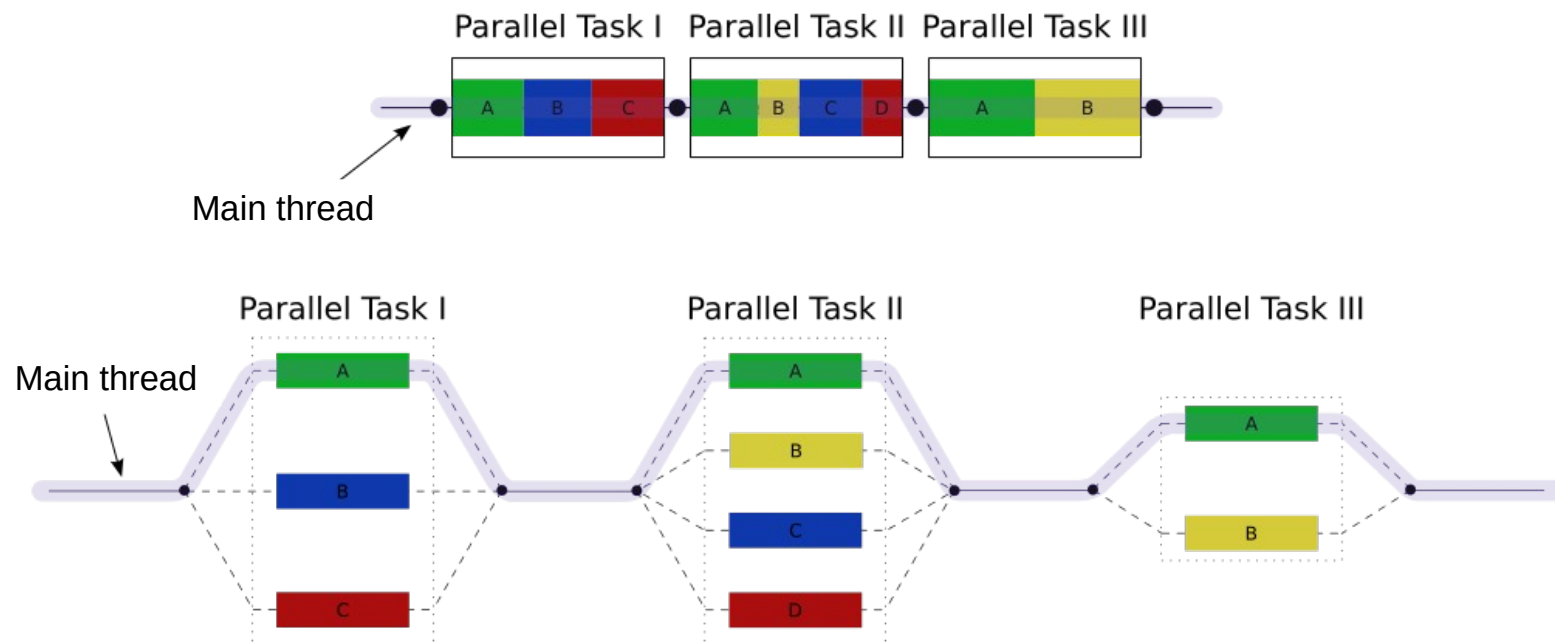
- Programming language extension
 - Compiler support required
 - "**Open Multi-Processing**"
 - Open standard: latest version is 5.1 (released Nov 2020)
 - Managed by a consortium: openmp.org
- “Automatic” thread-level parallelism
 - Guided by programmer-supplied **directives** (**pragmas**)
 - Does NOT verify correctness of parallel transformations
 - Targets shared-memory systems
 - Used in distributed systems for on-node parallelism

Technology comparison

- [Cilk / Cilk Plus / Open Cilk](#)
 - Language extension - new keywords: `spawn`, `sync`, `cilk_for`
 - Purchased by Intel in 2009; lost support in 2017; now maintained by MIT
- [Intel Thread Building Blocks](#) (TBB)
 - Template library (C++ only)
 - Gaining popularity, but fairly complicated to use
- [OpenMP](#)
 - Directive-based; supported by most major compilers
 - Currently the most popular CPU-based technology
- [OpenACC](#)
 - Directive-based; similar to OpenMP
 - Primarily aimed at GPU parallelism (driven by NVIDIA)

Fork-join threading

- OpenMP provides **directives** to control threading
 - General **fork-join** threading model w/ **teams** of threads
 - One **main** (or “master”) thread and multiple **worker** threads



C preprocessor (261 review)

- Text-based processing phase of compilation
 - Can be run individually with “cpp”
 - Performs textual modifications on program code

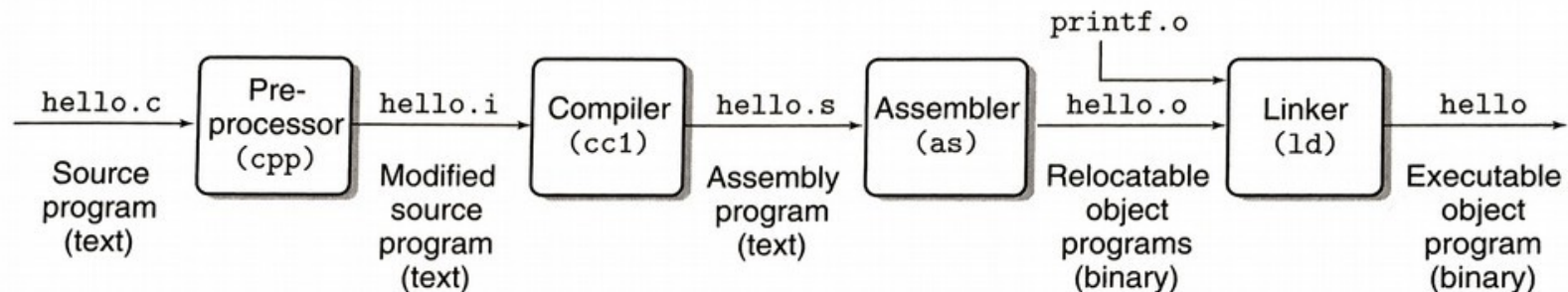


Figure 1.3 The compilation system.

```
#define MAX_LEN 128
```

```
char name[MAX_LEN];  
snprintf(name, MAX_LEN, "%s", argv[1]);
```

Before

```
char name[128];  
snprintf(name, 128, "%s", argv[1]);
```

After

C preprocessor (261 review)

- Controlled by **directives** on lines beginning with “#”
 - Must be the first non-whitespace character
 - Alignment is a matter of personal style

```
#include <stdio.h>
#define F00
#define BAR 5

int main() {
#   ifdef F00
    printf("Hello!\n");
#   else
    printf("Goodbye!\n");
#   endif
    printf("%d\n", BAR);
    return 0;
}
```

my preference

```
#include <stdio.h>
#define F00
#define BAR 5

int main() {
    #ifdef F00
        printf("Hello!\n");
    #else
        printf("Goodbye!\n");
    #endif
    printf("%d\n", BAR);
    return 0;
}
```

Pragmas

- `#pragma` - generic preprocessor directive
 - Provides direction or info to later compiler phases
 - Ignored by compilers that don't support it
 - All OpenMP pragma directives begin with "`omp`"
 - Basic threading directive: "`parallel`"
 - Runs the following code construct in fork/join parallel threads
 - Implicit barrier at end of construct

```
#pragma play(global_thermonuclear_war)  
do_something();
```

```
#pragma omp parallel  
do_something_else();
```

Compiling and running w/ OpenMP

- Must `#include <omp.h>`
- Must compile with "`-fopenmp`" flag

```
gcc -g -std=c99 -Wall -fopenmp -o omp omp.c  
srun ./omp
```

- Use `OMP_NUM_THREADS` environment variable to set thread count
 - Default value is core count (w/ hyper-threads)

```
OMP_NUM_THREADS=4 srun ./omp
```


"Hello World" example

```
#include <stdio.h>
#include <stdlib.h>
#include <omp.h>

int main(int argc, char *argv[])
{
    #    pragma omp parallel
    printf("Hello!\n");

    printf("Goodbye!\n");

    return EXIT_SUCCESS;
}
```

Mutual exclusion

- Use "**critical**" directive to enforce mutual exclusion
 - Only one thread at a time can execute the following construct
 - A critical section can optionally be **named**
 - Sections that share a name share exclusivity
 - *CAUTION: all unnamed sections "share" a name!*

```
#    pragma omp critical(gres)  
    global_result += my_result ;
```

Pragma scope

- Most OpenMP pragmas apply to the immediately-following **statement** or **block**
 - Not necessarily just the next line!

```
# pragma omp parallel
printf("hello!\n");
```

```
# pragma omp parallel
total += a * b + c;
```

```
# pragma omp parallel
{
    int a = 0;
    ...
    global_var += a;
}
```

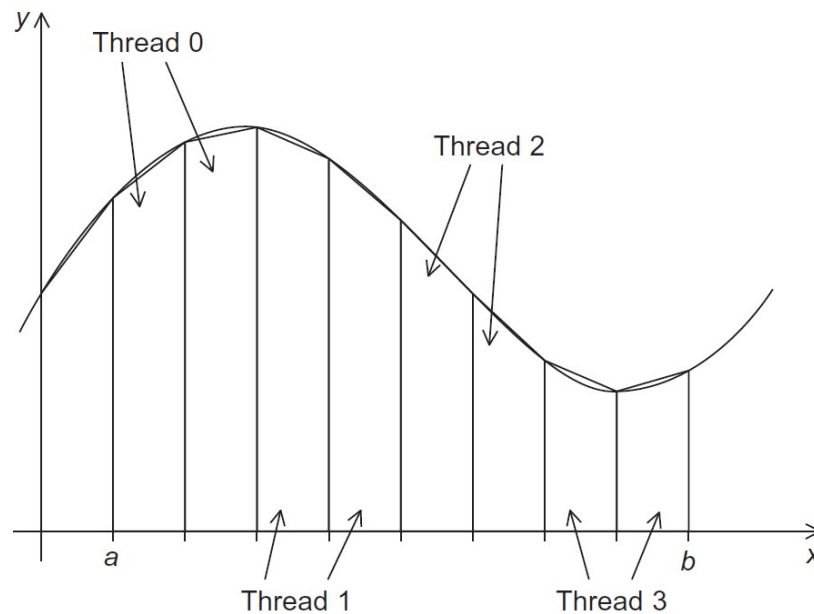
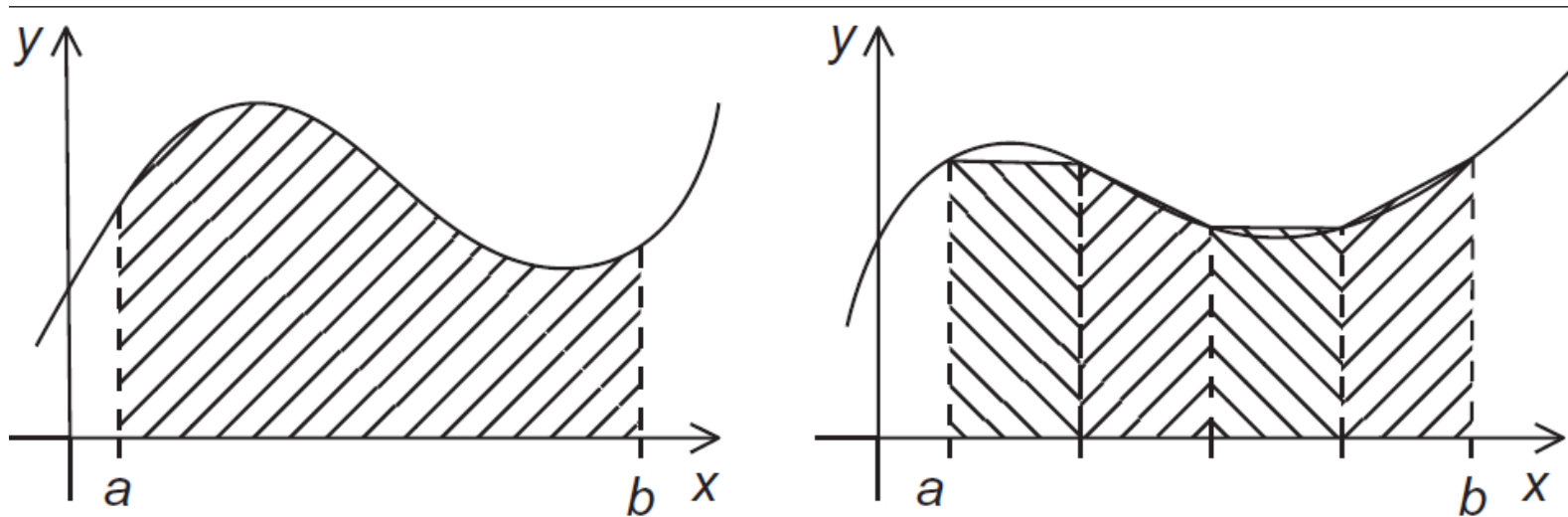
```
# pragma omp parallel
for (i = 0; i < n; i++) {
    sum += i;
}
```

Warning: different semantics from
#pragma omp parallel for

Functions

- Built-in functions:
 - **omp_get_num_threads()**
 - Returns the number of threads in the current team
 - **omp_get_max_threads()**
 - Returns the maximum number of threads in a team
 - Can be used outside a parallel region
 - **omp_get_thread_num()**
 - Returns the caller's thread ID within the current team
 - **omp_get_wtime()**
 - Returns the elapsed wall time in seconds

Trapezoid example (from textbook)



Is this task or data parallelism?

What problem(s) might we run into?

Trapezoid example (from textbook)

- Solution: *(w/ some non-OMP code omitted for brevity)*

```
int main() {
    double global_result = 0.0;

    # pragma omp parallel
        trapezoid(&global_result);

    printf("Estimated area: %.14e\n", global_result);
}

void trapezoid(double* global_result) {
    double my_result;

    int my_tid = omp_get_thread_num();
    int num_threads = omp_get_num_threads()
    // calculate my_result based on my_tid and num_threads

    # pragma omp critical
        *global_result += my_result;
}
```

Incremental parallelization

- Pragas allow **incremental parallelization**
 - Gradually add parallel constructs
 - OpenMP programs can be correct serial programs when compiled without "-fopenmp"
 - Silence the pragma warnings with "-Wno-unknown-pragmas"
 - Still need to guard the `#include` and function calls
 - Use `"_OPENMP"` preprocessor variable to test
 - If defined, it is safe to call OpenMP functions

```
#ifndef _OPENMP
#include <omp.h>
#endif
```

```
#   ifdef _OPENMP
        int my_rank = omp_get_thread_num();
        int thread_count = omp_get_num_threads();
#   else
        int my_rank = 0;
        int thread_count = 1;
#   endif
```

Barriers

- Explicit barrier: "`barrier`" directive
 - All threads must sync

```
#    pragma omp barrier
```


Clauses

- Directives can be modified by **clauses**
 - Text that follows the directive
 - Some clauses take parameters
 - E.g., "**num_threads**"

```
# pragma omp parallel num_threads(thread_count)
```

WARNING: Only use the "num_threads" clause if you wish to hard-code the number of threads (this is not considered best practice for OpenMP!)

Single-thread regions

- Implicit barrier: "**single**" directive
 - Only one thread executes the following construct
 - Could be any thread; don't assume it's the first/main thread
 - For main-thread-only, use "**master**" directive
 - All threads must sync at end of directive
 - Use "**nowait**" clause to prevent this implicit barrier

```
#    pragma omp single  
    global_result /= 2;
```

```
#    pragma omp single nowait  
    global_iter_count++;
```

Reductions

- The `reduction(op:var)` clause applies an operator to a sequence of operands to get a single result
 - Designates a shared-memory **reduction variable** (`var`)
 - OpenMP manages per-thread intermediate results
 - OpenMP handles synchronization (w/ implicit mutex) and stores final results in the reduction variable
 - Supported operations (`op`): `+`, `-`, `*`, `&`, `|`, `^`, `&&`, `||`, `min`, `max`

```
double foo = 0.0;
# pragma omp parallel reduction(+:foo)
foo += (do_calc() * PI)/2.0;
```

Scope of variables

- In OpenMP, each variable has a thread "scope"
 - **Shared** scope: accessible by all threads in team
 - Default for variables declared **before** a parallel block
 - **Private** scope: accessible by only a single thread
 - Default for variables declared **inside** a parallel block

```
double foo = 0.0;                                // shared
# pragma omp parallel
{
    double bar = do_calc() * PI;                  // private
# pragma omp critical
    foo = foo + bar/2.0;
}
```

Default scoping

- The "**default**" clause changes the default scope for variables declared outside the parallel block
 - **default(none)** mandates explicit scope declaration
 - Use "**shared**", "**reduction**", and "**private**" clauses
 - Compiler will check that you declared all variables
 - This is good programming practice and **required in CS 470**

```
double sum = 0.0;
# pragma omp parallel for num_threads(thread_count) \
    default(none) reduction(+:sum) private(k, factor) \
    shared(n)
for (k = 0; k < n; k++) {
    if (k % 2 == 0)
        factor = 1.0;
    else
        factor = -1.0;
    sum += factor/(2*k+1);
}
```

Private variable nuances

- Sometimes it is useful to have a variable that is neither completely shared nor completely private
- Use `firstprivate` to initialize with the value before parallel region
 - Useful if all threads need to start with the same value but later diverge
- Use `lastprivate` to save last value after parallel region

```
int i;  
#pragma omp parallel  
{  
#   pragma omp for lastprivate(i)  
    for (i = 0; i < n-1; i++)  
        a[i] = b[i] + b[i+1];  
}  
a[i] = b[i];
```

Parallel for loops

- The "`parallel for`" directive parallelizes a loop
 - Probably the most powerful and most-used directive
 - Divides loop iterations among a team of threads
 - **CAVEAT**: the for-loop must have a very particular form

```
for (
    index = start ; index < end      index++
    index <= end   index--          ++index
    index >= end   --index
    index > end    index += incr
                  index -= incr
                  index = index + incr
                  index = incr + index
                  index = index - incr
)
```

Parallel for loops

- The compiler must be able to determine the number of iterations *prior to the execution of the loop*
- Implications/restrictions:
 - The number of iterations must be finite (no "**for** (;;)")
 - The **break** statement cannot be used (although **exit()** is ok)
 - The **index** variable must have an integer or pointer type
 - The **index** variable must only be modified by the "increment" part of the loop declaration
 - The **index**, **start**, **end**, and **incr** expressions/variables must all have compatible types
 - The **start**, **end**, and **incr** expressions must not change during execution of the loop

Issue: correctness

```
fib[0] = fib[1] = 1;  
for (i = 2; i < n; i++)  
    fib[i] = fib[i-1] + fib[i-2];
```



```
fib[0] = fib[1] = 1;  
# pragma omp parallel for  
for (i = 2; i < n; i++)  
    fib[i] = fib[i-1] + fib[i-2];
```

1 1 2 3 5 8 13 21 34 55

this is correct

but sometimes we
get this (w/ 2 threads)

1 1 2 3 5 8 0 0 0 0

Loop dependencies

- A loop has a **data dependence** if one iteration depends on another iteration
 - Explicitly (as in Fibonacci example) or implicitly
 - Includes side effects!
 - Sometimes called **loop-carried dependence**
- A loop with dependencies cannot (usually) be parallelized correctly by OpenMP
 - Identifying dependencies is very important!
 - OpenMP does not check for them

Loop dependencies

- Examples:

```
for (i = 0; i < n; i++) {  
    a[i] = b[i] * c[i];  
}
```

```
for (i = 1; i < n; i++) {  
    a[i] += a[i-1]  
}
```

```
for (i = 0; i < n; i++) {  
    a[i] += b[i]  
}
```

```
for (i = 1; i < n; i += 2) {  
    a[i] += a[i-1]  
}
```

```
for (i = 0; i < n; i++) {  
    a[i] += a[i]  
}
```

```
for (i = 1; i < n; i++) {  
    a[i] += b[i-1]  
}
```

Loop dependencies

- Examples:

```
for (i = 0; i < n; i++) {  
    a[i] = b[i] * c[i];  
}
```

OK!

```
for (i = 0; i < n; i++) {  
    a[i] += b[i]  
}
```

OK!

```
for (i = 0; i < n; i++) {  
    a[i] += a[i]  
}
```

OK!

```
for (i = 1; i < n; i++) {  
    a[i] += a[i-1]  
}
```

BAD! (iteration i depends on i-1)

```
for (i = 1; i < n; i += 2) {  
    a[i] += a[i-1]  
}
```

OK!

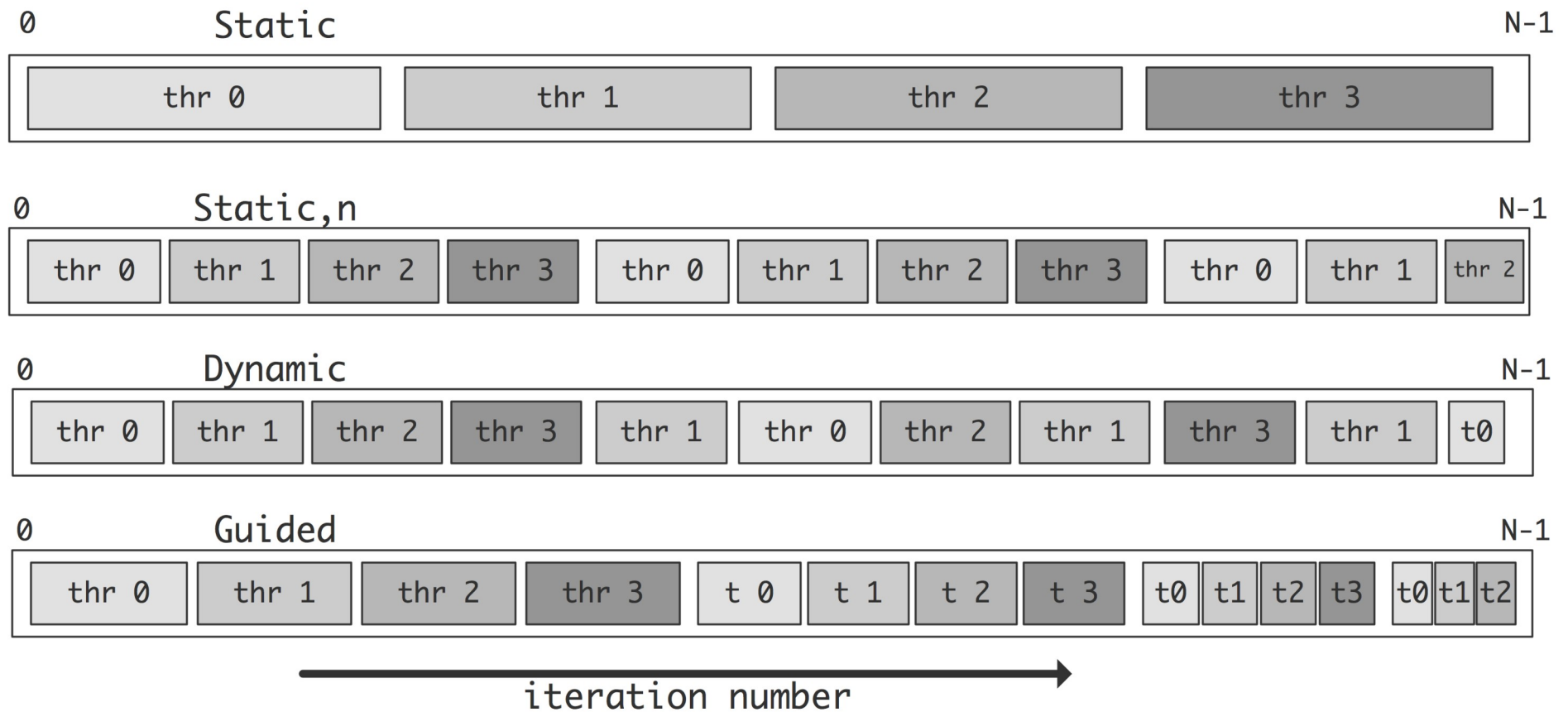
```
for (i = 1; i < n; i++) {  
    a[i] += b[i-1]  
}
```

OK!

Loop scheduling

- Use the `schedule` clause to control how parallel for-loop iterations are allocated to threads
 - Modified by `chunksize` parameter
 - `static`: split into chunks before loop is executed
 - `dynamic`: split into chunks, dynamically allocated to threads (similar to thread pool or tasks)
 - `guided`: like dynamic, but chunk sizes decrease
 - The specified chunksize is the minimum
 - `auto`: allows the compiler or runtime to choose
 - `runtime`: allows specification using `OMP_SCHEDULE`

Loop scheduling



Parallel regions

- Often useful: multiple for-loops inside a `parallel` region
 - Many pragmas bind dynamically to any active `parallel` region
 - Less thread creation/joining overhead
 - Private variables can be re-used across multiple loops

```
# pragma omp parallel for
for (int i = 0; i < n; i++) {
    do_something_parallel();
}

do_something_serial();

# pragma omp parallel for
for (int j = 0; j < m; j++) {
    do_something_else_parallel();
}
```

Original

```
# pragma omp parallel
{
#     pragma omp for
    for (int i = 0; i < n; i++) {
        do_something_parallel();
    }

#     pragma omp single
    do_something_serial();

#     pragma omp for
    for (int j = 0; j < m; j++) {
        do_something_else_parallel();
    }
}
```

Faster

Nested loops

- The parallel for loop only applies to the loop layer that you specify
 - For nested loops, use the `collapse` clause to combine iteration spaces
 - Spaces must be “square”
 - i.e., inner loop iteration count should not depend on outer loop value
 - Project note: this is NOT the case for many loops in P3!

```
#pragma omp parallel for collapse(2)
for (i = 0; i < n; i++) {           // row
    for (j = 0; j < n; j++) {       // column
        a[i*n + j] = 1.0;
    }
}
```

Atomics

- OpenMP provides access to highly-efficient hardware synchronization mechanisms
 - Use the `atomic` pragma to annotate a single statement
 - Statement must be a single increment/decrement or in the following form:
 - `x <op>= <expr>;` `// <op> can be +, -, *, /, &, |, ^, <<, >>`
 - Many ISAs provide an atomic load/modify/store instruction
 - In x86-64, specified using the LOCK prefix
 - Far more efficient than using a mutex (i.e., `critical`)
 - This requires multiple function calls!

Locks

- OpenMP provides a basic locking system
 - Useful for protecting a data structure rather than a region of code
 - `omp_lock_t`: lock variable
 - Similar to `pthread_mutex_t`
 - `omp_lock_init`: initialize lock
 - Similar to `pthread_mutex_init`
 - `omp_set_lock`: acquire lock
 - Similar to `pthread_mutex_lock`
 - `omp_unset_lock`: release lock
 - Similar to `pthread_mutex_unlock`
 - `omp_lock_destroy`: clean up a lock
 - Similar to `pthread_mutex_destroy`

Thread safety

- Don't **mix** mutual exclusion mechanisms
 - `#pragma omp critical`
 - `#pragma omp atomic`
 - `omp_set_lock()`
- Don't **nest** mutual exclusion mechanisms
 - Nesting unnamed `critical` sections *guarantees* deadlock!
 - The thread cannot enter the second section because it is still in the first section, and unnamed sections “share” a name
 - If you must, use **named** critical sections or **nested** locks

Nested locks

- Simple vs. nested locks
 - `omp_nest_lock_*` instead of `omp_lock_*`
 - A nested lock may be acquired multiple times
 - Must be in the same thread
 - Must be released the same number of times
 - Allows you to write functions that call each other but need to acquire the same lock

Sections

- OpenMP is most often used for **data parallelism** (`parallel for`)
- However, it also supports explicit **task parallelism**
- Pre-OpenMP 3.0 mechanism: `sections` directive
 - Contains multiple `section` blocks; each section runs on separate thread
 - Must list all sections in same location (cannot dynamically add new tasks)
 - Implicit barrier at end (unless `nowait` clause is specified)

```
#    pragma omp parallel sections
#    {
#        pragma omp section
#        producer();
#        pragma omp section
#        consumer();
#    }
```

Tasks

- Post-OpenMP 3.0 mechanism: `task` directive
 - Similar to thread pool task model
 - Tasks are assigned to available worker threads by the runtime
 - Tasks may be deferred if no workers available
 - No implicit barrier; use `taskwait` directive if needed
 - Use `single` region if only one thread should begin (e.g., recursion)
 - Use `nowait` clause to allow other threads to run tasks

```
main:
```

```
#  pragma omp parallel  
#  pragma omp single nowait  
    quick_sort(items, n);
```

```
quicksort:
```

```
    <select pivot and partition>
```

```
    // recursively sort each partition  
#  pragma omp task  
    quick_sort(items, p+1);  
#  pragma omp task  
    quick_sort(items+q, n-q);  
#  pragma omp taskwait
```

SIMD support

- Post-OpenMP 4.0 mechanism: `simd` directive
 - Enables generation of vector instructions (e.g., SSE or AVX)
 - Can encode some loop-carried dependencies using `safelen(X)` directive

```
#pragma omp simd
{
  for (i=0; i<N; i++) {
    a[i] = a[i] + b[i] * c[i];
  }
}
```

```
#pragma omp simd safelen(4)
{
  for (i=0; i<(N-4); i++) {
    a[i] = a[i+4] + b[i] * c[i];
  }
}
```


GPU support

- Post-OpenMP 4.0/4.5 mechanism: **target** directive
 - Offloads computation to a device (e.g., GPU)
 - Data management using **map** directive with **from** and **to** clauses

```
void vec_mult(int N)
{
    int i;
    float p[N], v1[N], v2[N];
    init(v1, v2, N);
    # pragma omp target map(to: v1, v2) map(from: p)
    # pragma omp parallel for
    for (i=0; i<N; i++) {
        p[i] = v1[i] * v2[i];
    }
    output(p, N);
}
```

More OpenMP examples

- Posted in `/shared/cs470`
 - Basic for-loop example (`omp-array`)
 - For-loop scheduling (`omp-sched`)
 - Critical sections and deadlock (`omp-deadlock`)
 - The 'atomic' directive (`omp-atomic`)
 - Tasks (`omp-qsort`)
 - Matrix multiplication (`omp-matmult`)