#### Lecture 4:

## Parallel Programming Basics

Parallel Computer Architecture and Programming CMU 15-418/15-618, Spring 2016

#### Tunes

## Gorillaz

"Clint Eastwood"
(Gorillaz)

I'm happy, I'm feeling glad
I got a new toolbox, in my bag
My code's useless, but not for long
I know what my tasks de-pend on
I'm happy, I'm feeling glad
I used the 418 skills, in my bag
My code is per-formant
and structured for parallelization
parallelization...
parallelization...

#### Quiz

```
export void sinx(
   uniform int N,
   uniform int terms,
   uniform float* x,
   uniform float* result)
   // assume N % programCount = 0
   for (uniform int i=0; i<N; i+=programCount)</pre>
      int idx = i + programIndex;
      float value = x[idx];
      float numer = x[idx] * x[idx] * x[idx];
      uniform int denom = 6; // 3!
      uniform int sign = -1;
      for (uniform int j=1; j<=terms; j++)</pre>
         value += sign * numer / denom
         numer *= x[idx] * x[idx];
         denom *= (2*j+2) * (2*j+3);
         sign *= -1;
      result[idx] = value;
```

This is an ISPC function.

It contains a loop nest.

Which iterations of the loop(s) are parallelized by ISPC? Which are not?

**Answer: none** 

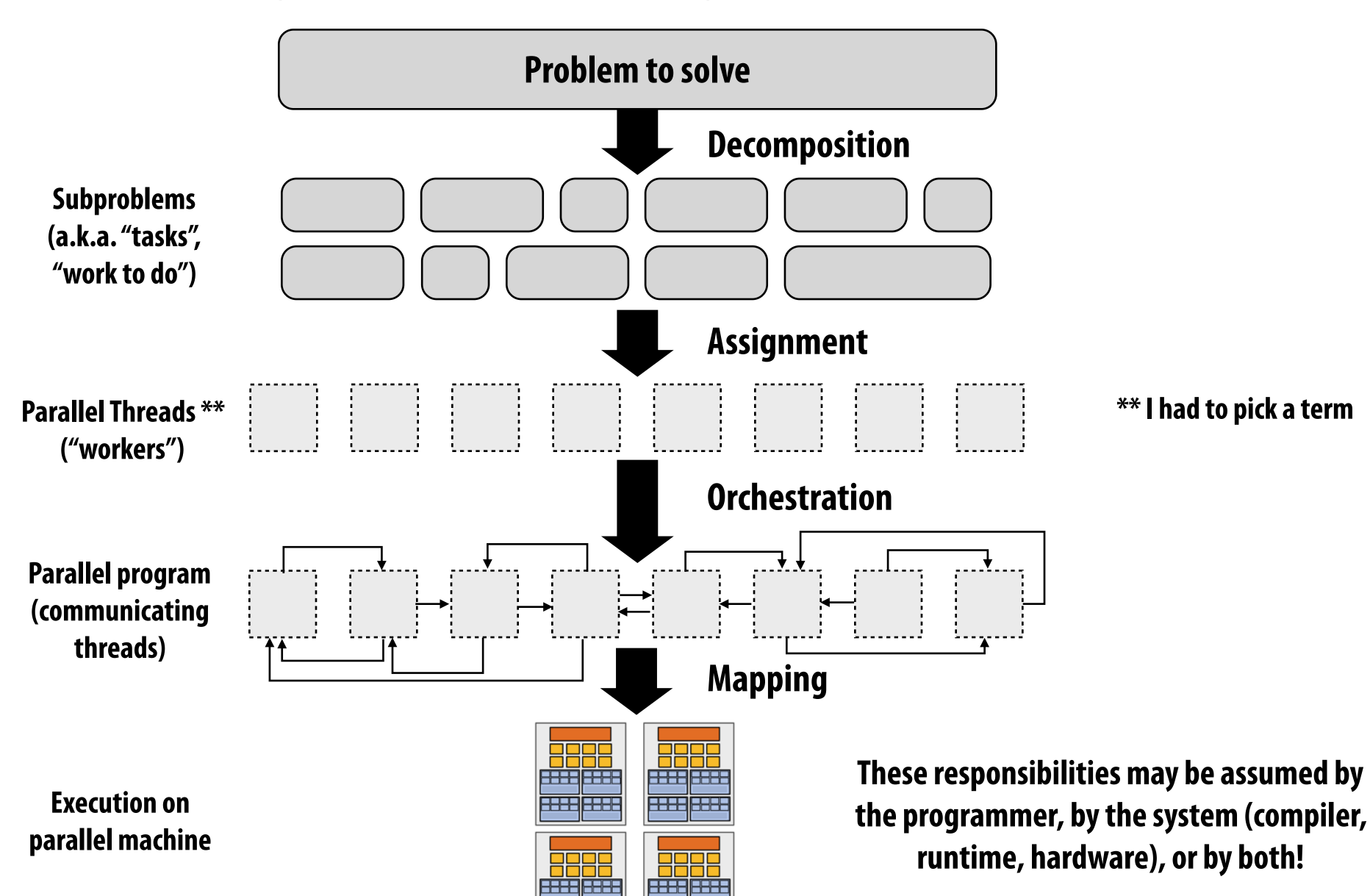
#### Creating a parallel program

- Thought process:
  - 1. Identify work that can be performed in parallel
  - 2. Partition work (and also data associated with the work)
  - 3. Manage data access, communication, and synchronization

Recall one of our main goals is speedup \*
For a fixed computation:

<sup>\*</sup> Other goals include high efficiency (cost, area, power, etc.) or working on bigger problems than can fit on one machine

#### Creating a parallel program



Adopted from: Culler, Singh, and Gupta

#### Decomposition

- Break up problem into tasks that <u>can</u> be carried out in parallel
  - Decomposition need not happen statically
  - New tasks can be identified as program executes
- Main idea: create at least enough tasks to keep all execution units on a machine busy

## Key aspect of decomposition: identifying dependencies (or... a lack of dependencies)

# Amdahl's Law: dependencies limit maximum speedup due to parallelism

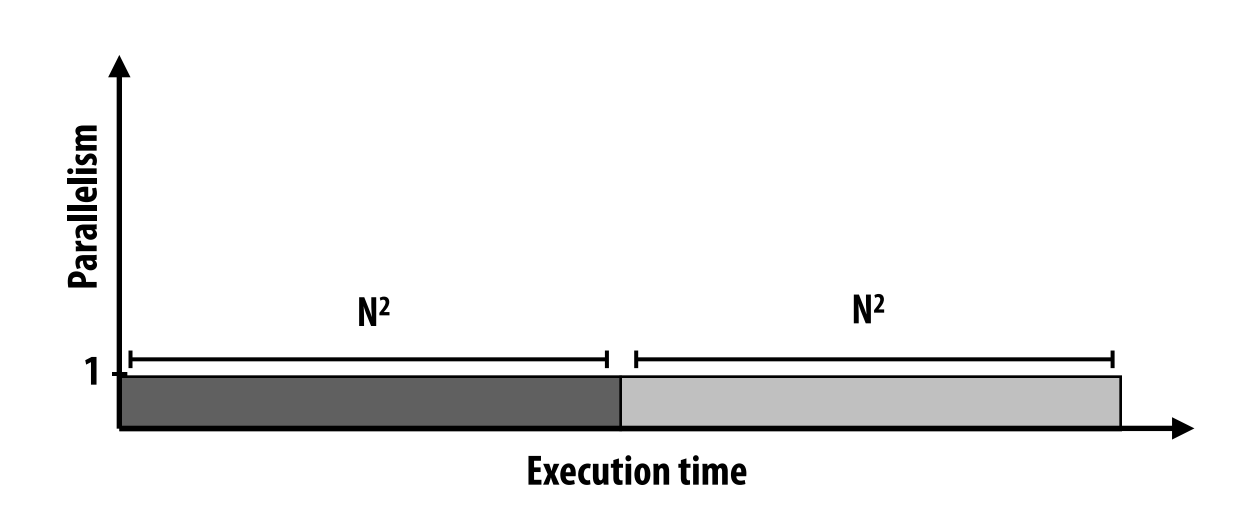
■ You run your favorite sequential program...

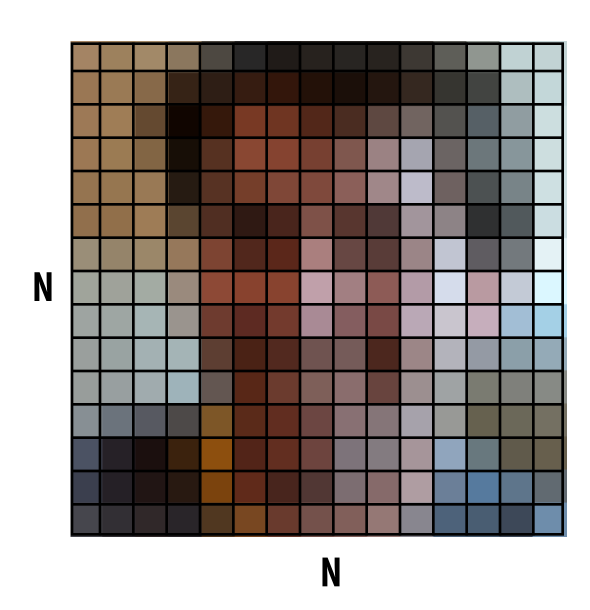
Let S = the fraction of sequential execution that is inherently sequential (dependencies prevent parallel execution)

■ Then maximum speedup due to parallel execution  $\leq 1/S$ 

#### A simple example

- Consider a two-step computation on a N x N image
  - Step 1: double brightness of all pixels (independent computation on each grid element)
  - Step 2: compute average of all pixel values
- Sequential implementation of program
  - Both steps take  $\sim N^2$  time, so total time is  $\sim 2N^2$





#### First attempt at parallelism (P processors)

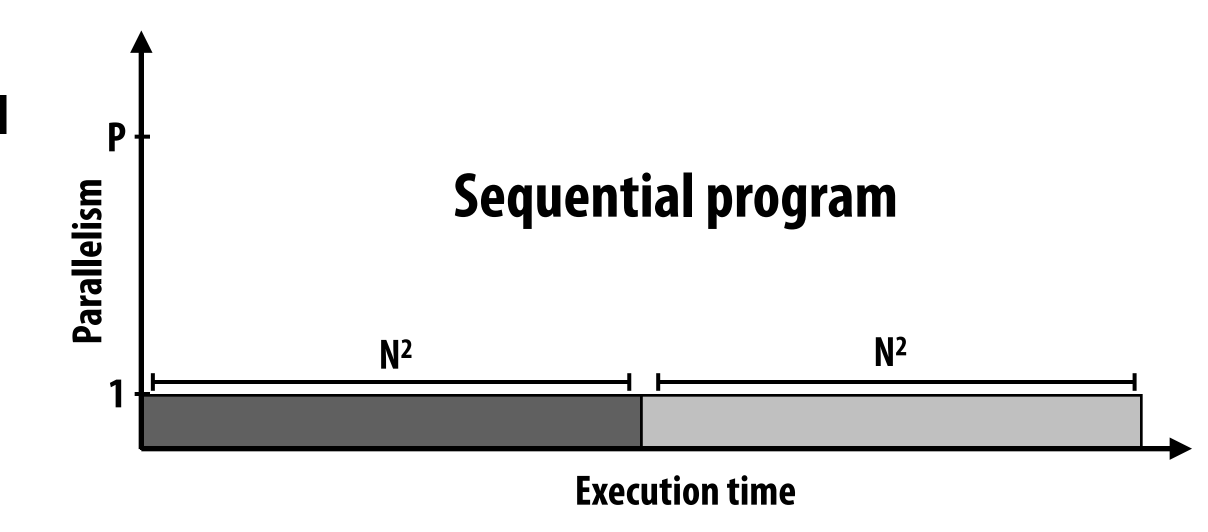
#### Strategy:

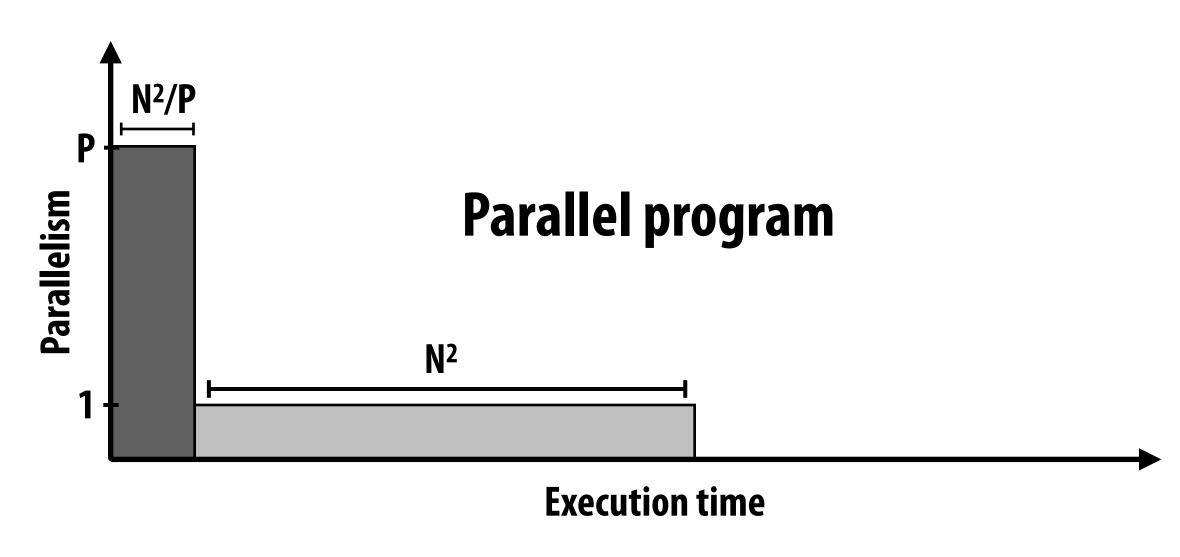
- Step 1: execute in parallel
  - time for phase 1: N<sup>2</sup>/P
- Step 2: execute serially
  - time for phase 2: N<sup>2</sup>

#### Overall performance:

Speedup 
$$\leq \frac{2n^2}{\frac{n^2}{p} + n^2}$$

Speedup ≤ 2





### Parallelizing step 2

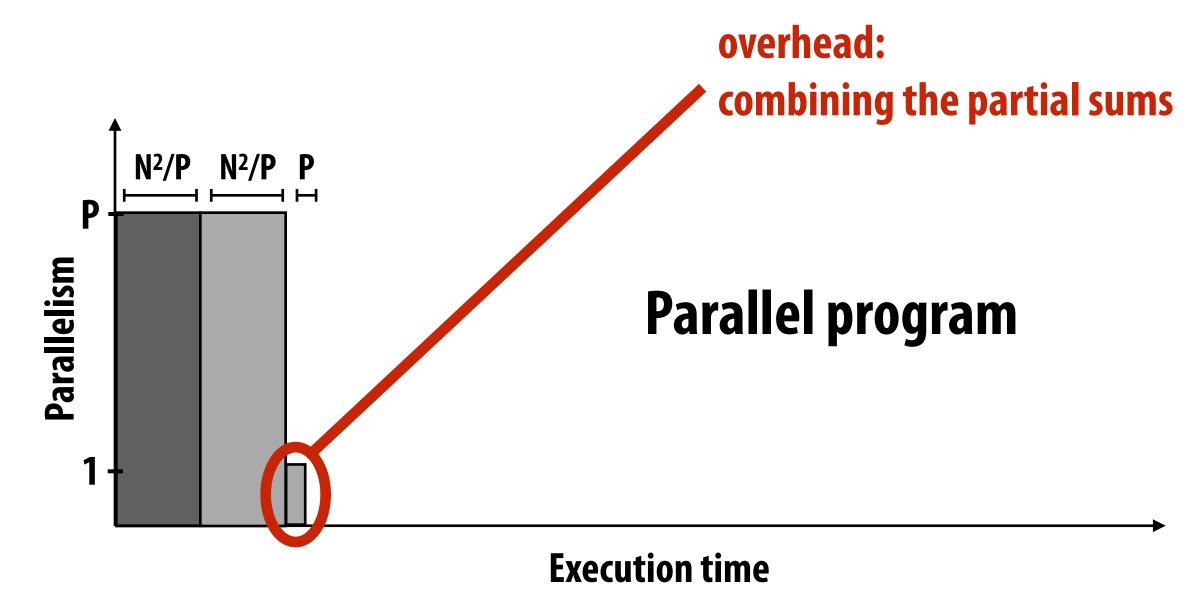
#### Strategy:

- Step 1: execute in parallel
  - time for phase 1: N<sup>2</sup>/P
- Step 2: compute partial sums in parallel, combine results serially
  - time for phase 2:  $N^{2}/P + P$

#### Overall performance:

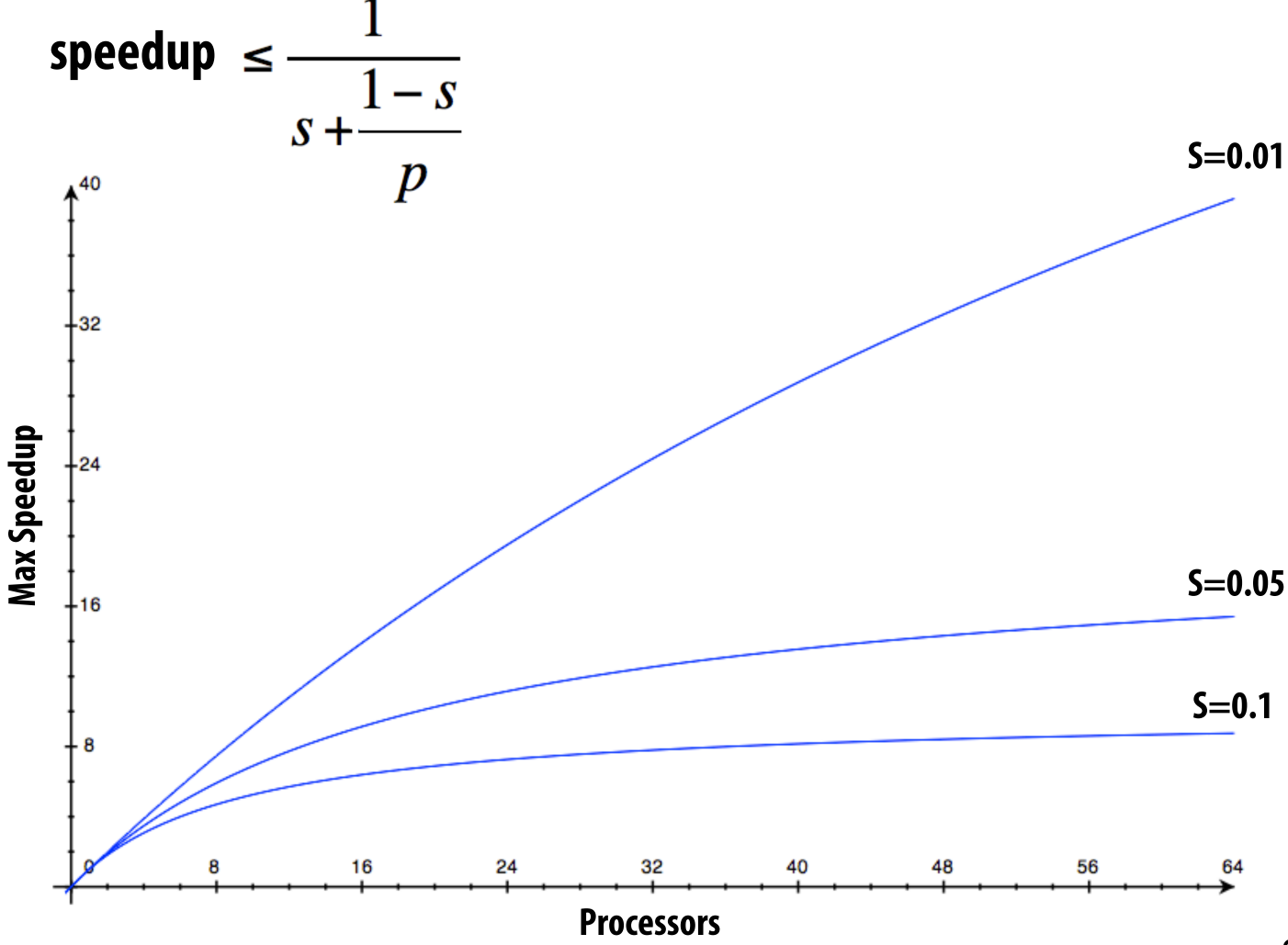
- Speedup 
$$\leq \frac{2n^2}{2n^2 + p}$$

Note: speedup  $\rightarrow$  P when N >> P



#### Amdahl's law

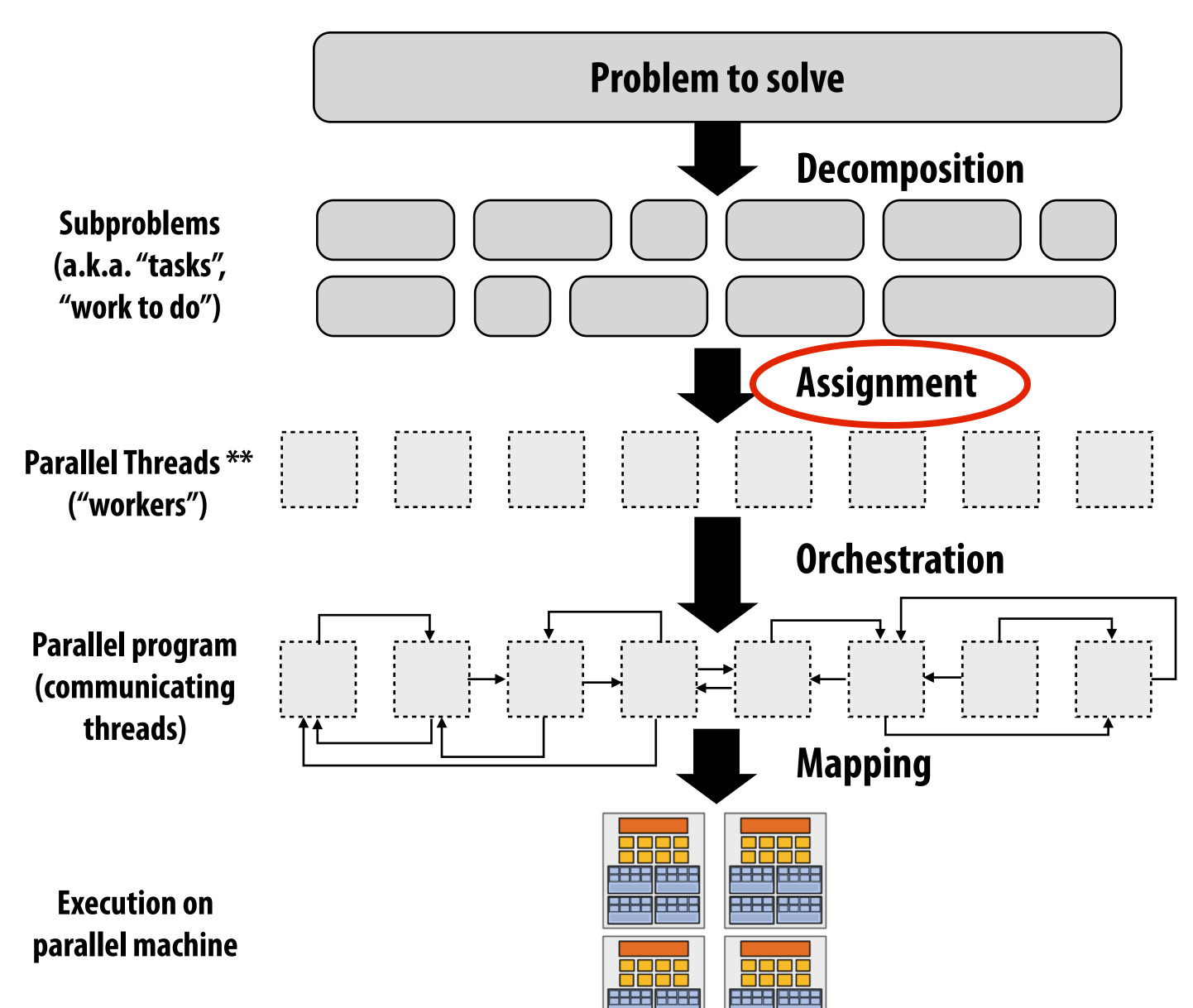
- Let S = the fraction of total work that is inherently sequential
- Max speedup on P processors given by:



#### Decomposition

- Who is responsible for performing decomposition?
  - In most cases: the programmer
- Automatic decomposition of sequential programs continues to be a challenging research problem (very difficult in general case)
  - Compiler must analyze program, identify dependencies
    - What if dependencies are data dependent (not known at compile time)?
  - Researchers have had modest success with simple loop nests
  - The "magic parallelizing compiler" for complex, general-purpose code has not yet been achieved

## Assignment



\*\* I had to pick a term

#### Assignment

Assigning tasks to threads \*\*\*

\*\* I had to pick a term (will explain in a second)

- Think of "tasks" as things to do
- Think of threads as "workers"
- Goals: balance workload, reduce communication costs
- Can be performed statically, or dynamically during execution
- While programmer often responsible for decomposition, many languages/runtimes take responsibility for assignment.

#### Assignment examples in ISPC

```
export void sinx(
   uniform int N,
   uniform int terms,
   uniform float* x,
   uniform float* result)
   // assumes N % programCount = 0
   for (uniform int i=0; i<N; i+=programCount)</pre>
      int idx = i + programIndex;
      float value = x[idx];
      float numer = x[idx] * x[idx] * x[idx];
      uniform int denom = 6; // 3!
      uniform int sign = -1;
      for (uniform int j=1; j<=terms; j++)</pre>
         value += sign * numer / denom;
         numer *= x[idx] * x[idx];
         denom *= (2*j+2) * (2*j+3);
         sign *= -1;
      result[i] = value;
```

**Decomposition of work by loop iteration** 

**Programmer-managed assignment:** 

Static assignment
Assign iterations to ISPC program instances in interleaved fashion

```
export void sinx(
   uniform int N,
   uniform int terms,
   uniform float* x,
   uniform float* result)
  foreach (i = 0 ... N)
      float value = x[i];
      float numer = x[i] * x[i] * x[i];
      uniform int denom = 6; // 3!
      uniform int sign = -1;
      for (uniform int j=1; j<=terms; j++)</pre>
         value += sign * numer / denom;
         numer *= x[i] * x[i];
         denom *= (2*j+2) * (2*j+3);
         sign *= -1;
      result[i] = value;
```

#### **Decomposition of work by loop iteration**

foreach construct exposes independent work to system System-manages assignment of iterations (work) to ISPC program instances (abstraction leaves room for dynamic assignment, but current ISPC implementation is static)

## Static assignment example using pthreads

```
typedef struct {
   int N, terms;
  float* x, *result;
} my_args;
void parallel_sinx(int N, int terms, float* x, float* result)
    pthread_t thread_id;
    my_args args;
    args.N = N/2;
    args.terms = terms;
    args.x = x;
    args.result = result;
    // launch second thread, do work on first half of array
    pthread_create(&thread_id, NULL, my_thread_start, &args);
    // do work on second half of array in main thread
    sinx(N - args.N, terms, x + args.N, result + args.N);
    pthread_join(thread_id, NULL);
void my_thread_start(void* thread_arg)
   my_args* thread_args = (my_args*)thread_arg;
   sinx(args->N, args->terms, args->x, args->result); // do work
```

**Decomposition of work by loop iteration** 

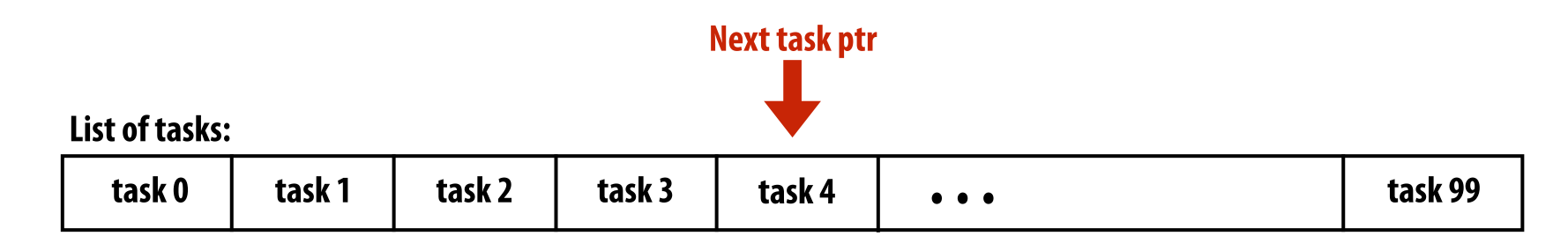
**Programmer-managed assignment:** 

Static assignment

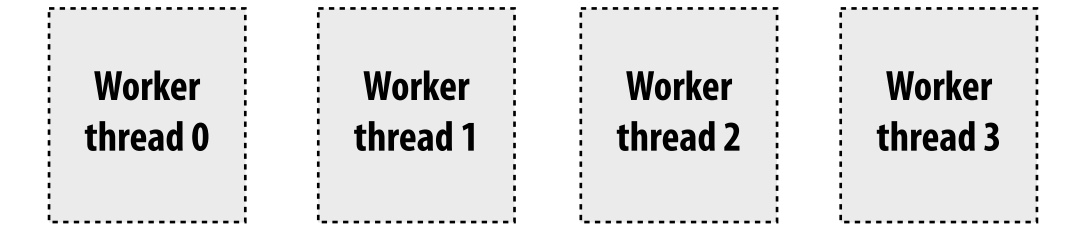
Assign iterations to pthreads in blocked fashion (first half of array to spawned thread, second half to main thread)

### Dynamic assignment using ISPC tasks

ISPC runtime assign tasks to worker threads



Assignment policy: after completing current task, worker thread inspects list and assigns itself the next uncompleted task.



#### Orchestration

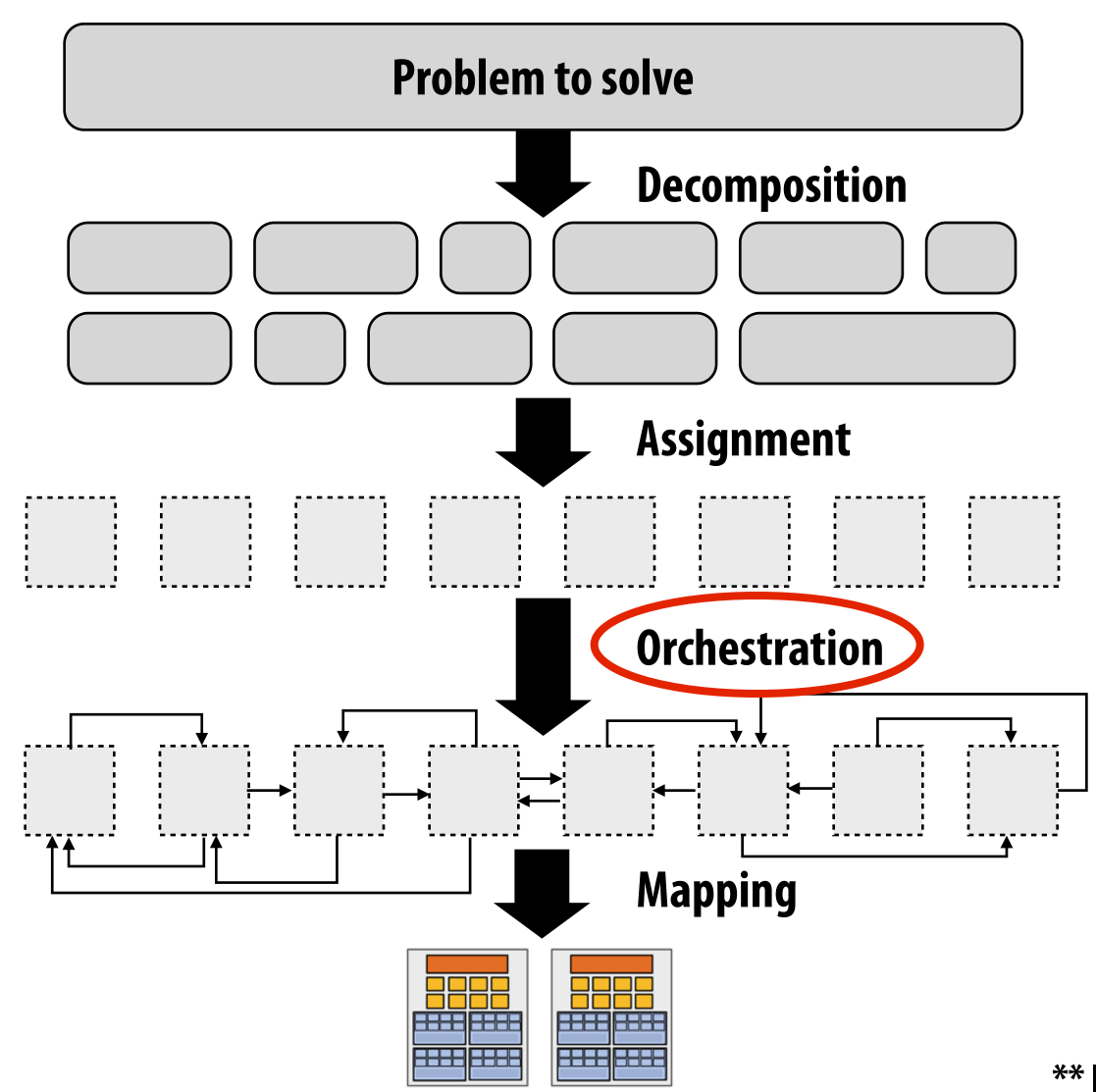
Subproblems (a.k.a. "tasks", "work to do")

Parallel Threads \*\*

("workers")

Parallel program (communicating threads)

Execution on parallel machine



\*\* I had to pick a term

#### Orchestration

#### Involves:

- Structuring communication
- Adding synchronization to preserve dependencies if necessary
- Organizing data structures in memory
- Scheduling tasks
- Goals: reduce costs of communication/sync, preserve locality of data reference, reduce overhead, etc.

- Machine details impact many of these decisions
  - If synchronization is expensive, might use it more sparsely

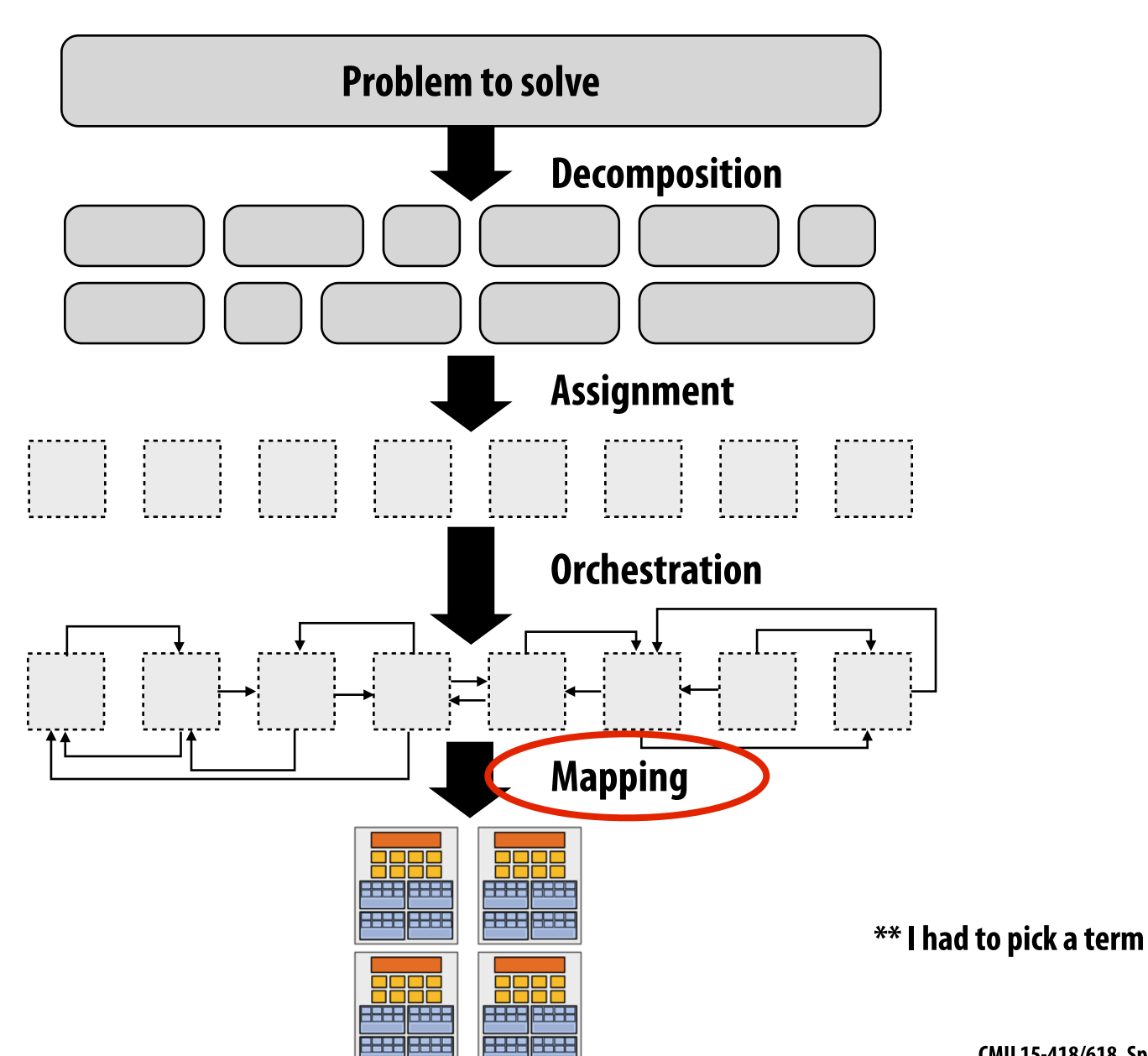
#### Mapping to hardware

Subproblems (a.k.a. "tasks", "work to do")

Parallel Threads \*\*
("workers")

Parallel program (communicating threads)

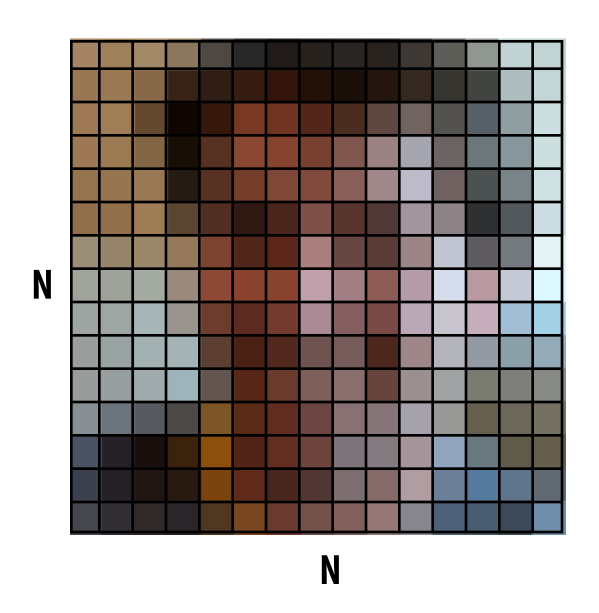
Execution on parallel machine



#### Mapping to hardware

- Mapping "threads" ("workers") to hardware execution units
- **Example 1: mapping by the operating system** 
  - e.g., map pthread to HW execution context on a CPU core
- Example 2: mapping by the compiler
  - Map ISPC program instances to vector instruction lanes
- Example 3: mapping by the hardware
  - Map CUDA thread blocks to GPU cores (future lecture)
- Some interesting mapping decisions:
  - Place <u>related</u> threads (cooperating threads) on the same processor (maximize locality, data sharing, minimize costs of comm/sync)
  - Place <u>unrelated</u> threads on the same processor (one might be bandwidth limited and another might be compute limited) to use machine more efficiently

#### Decomposing computation or data?



Often, the reason a problem requires lots of computation (and needs to be parallelized) is that it involves manipulating a lot of data.

I've described the process of parallelizing programs as an act of partitioning computation (work).

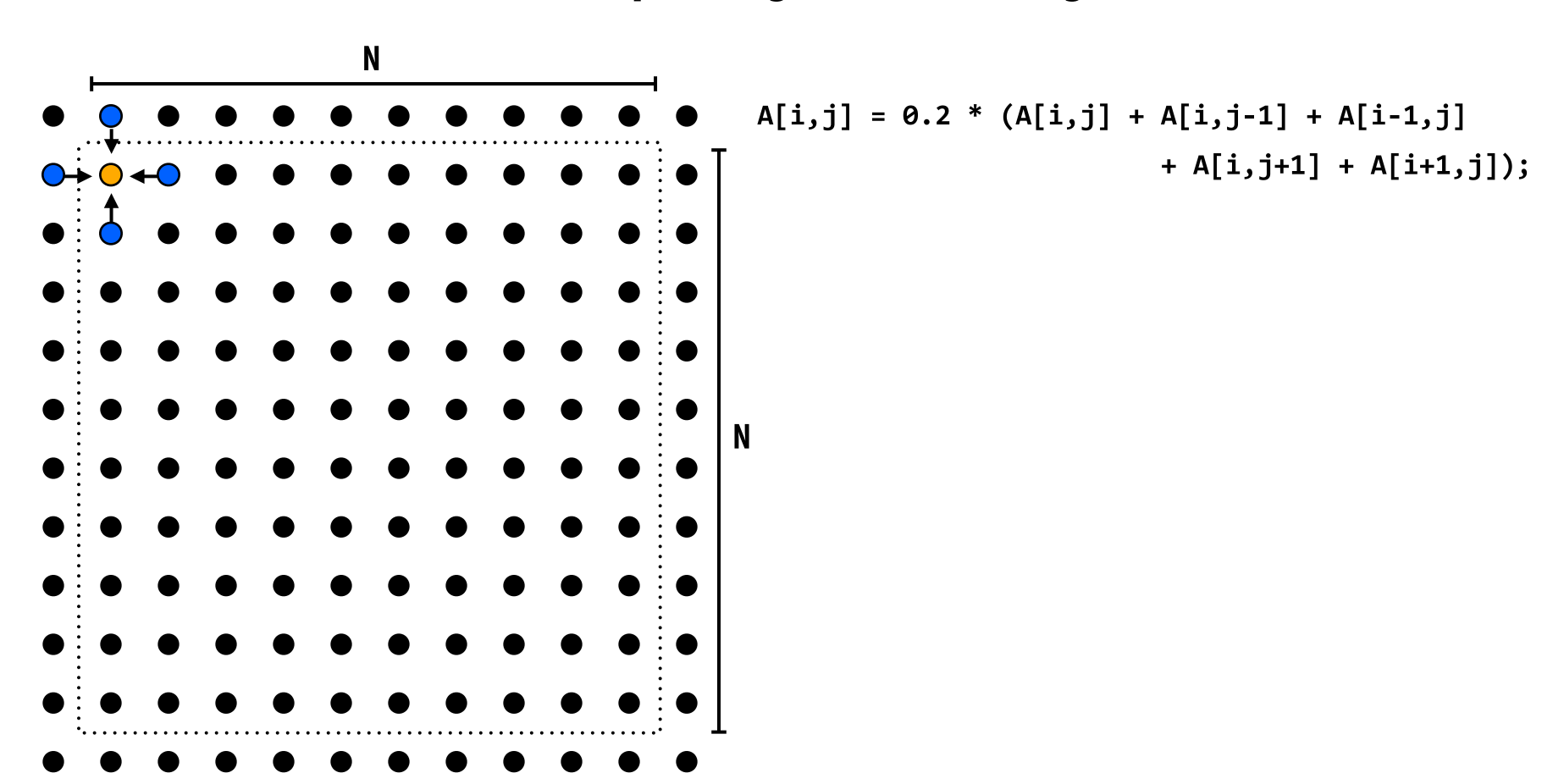
Often, it's equally valid to think of <u>partitioning data</u>. (computations go with the data)

But there are many computations where the correspondence between work-to-do ("tasks") and data is less clear. In these cases it's natural to think of partitioning computation.

## A parallel programming example

#### A 2D-grid based solver

- Solve partial differential equation (PDE) on  $(N+2) \times (N+2)$  grid
- Iterative solution
  - Perform Gauss-Seidel sweeps over grid until convergence

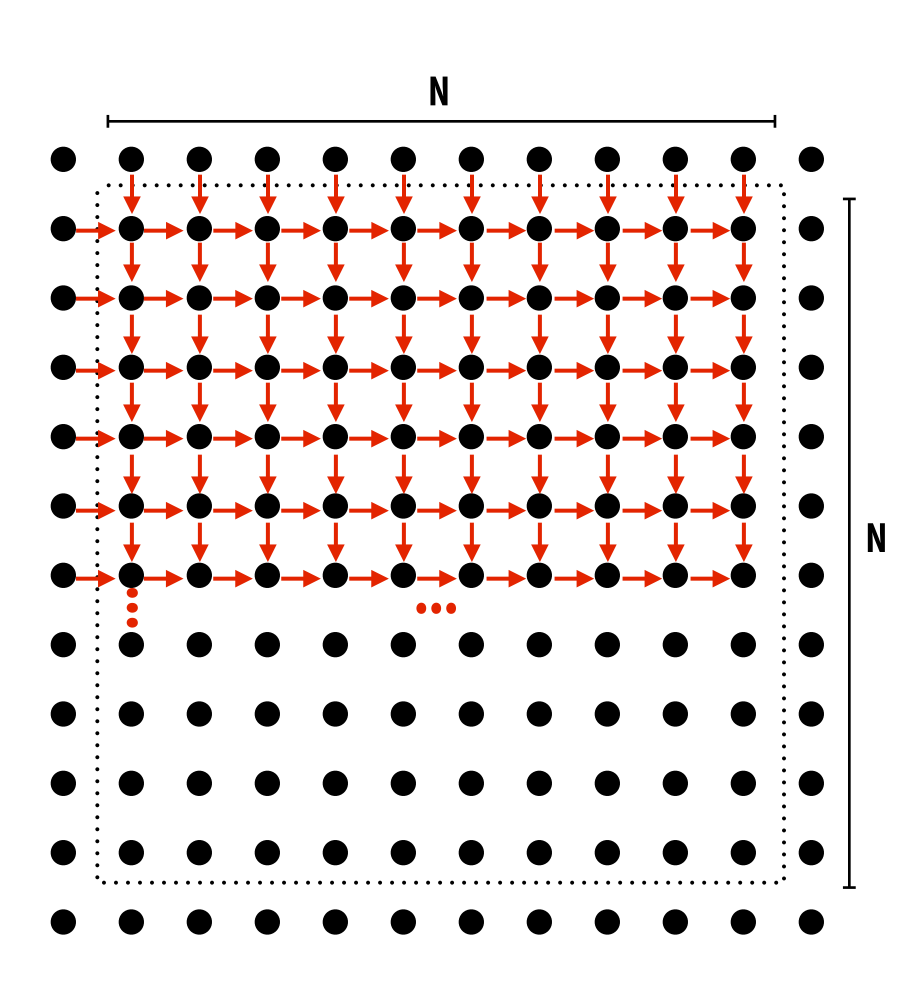


#### Grid solver algorithm

C-like pseudocode for sequential algorithm is provided below

```
const int n;
float* A;
                             // assume allocated to grid of N+2 x N+2 elements
void solve(float* A) {
  float diff, prev;
  bool done = false;
 while (!done) {
                                        // outermost loop: iterations
   diff = 0.f;
    for (int i=1; i<n i++) {
                                        // iterate over non-border points of grid
      for (int j=1; j<n; j++) {
        prev = A[i,j];
       A[i,j] = 0.2f * (A[i,j] + A[i,j-1] + A[i-1,j] +
                                  A[i,j+1] + A[i+1,j]);
        diff += fabs(A[i,j] - prev);  // compute amount of change
    if (diff/(n*n) < TOLERANCE)</pre>
                                 // quit if converged
      done = true;
```

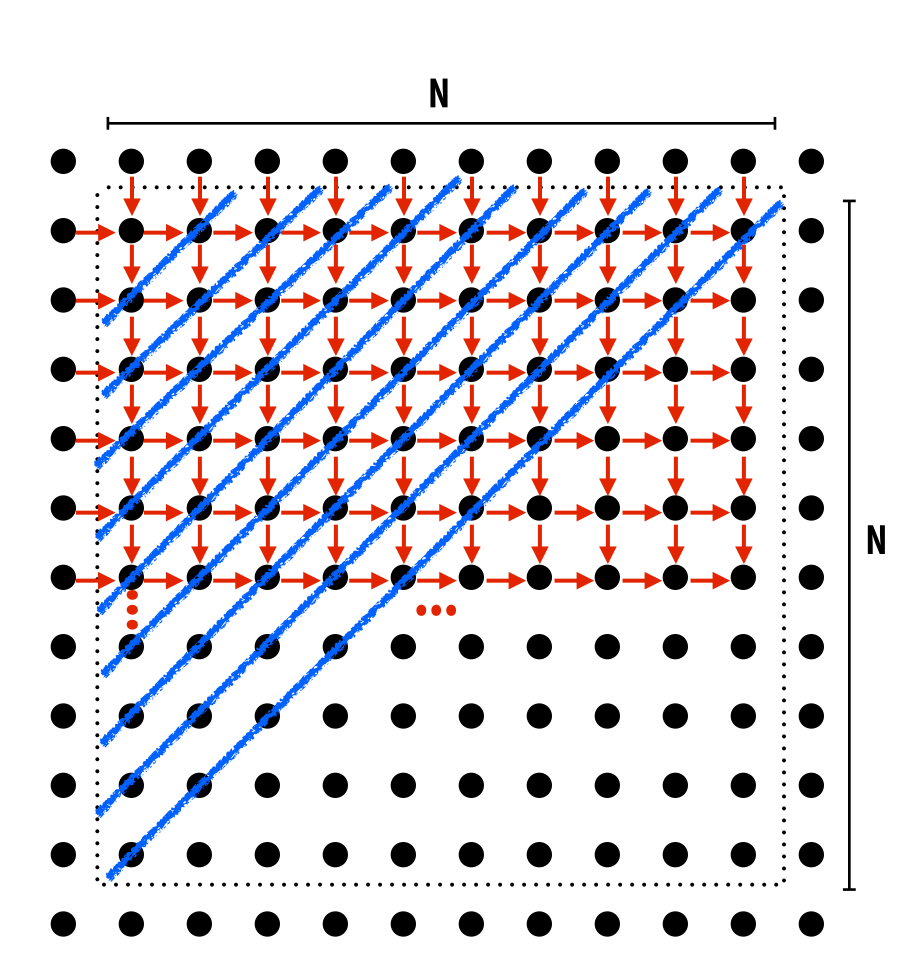
# Step 1: identify dependencies (problem decomposition phase)



Each row element depends on element to left.

Each column depends on previous column.

# Step 1: identify dependencies (problem decomposition phase)



There is independent work along the diagonals!

**Good: parallelism exists!** 

Possible implementation strategy:

- 1. Partition grid cells on a diagonal into tasks
- 2. Update values in parallel
- 3. When complete, move to next diagonal

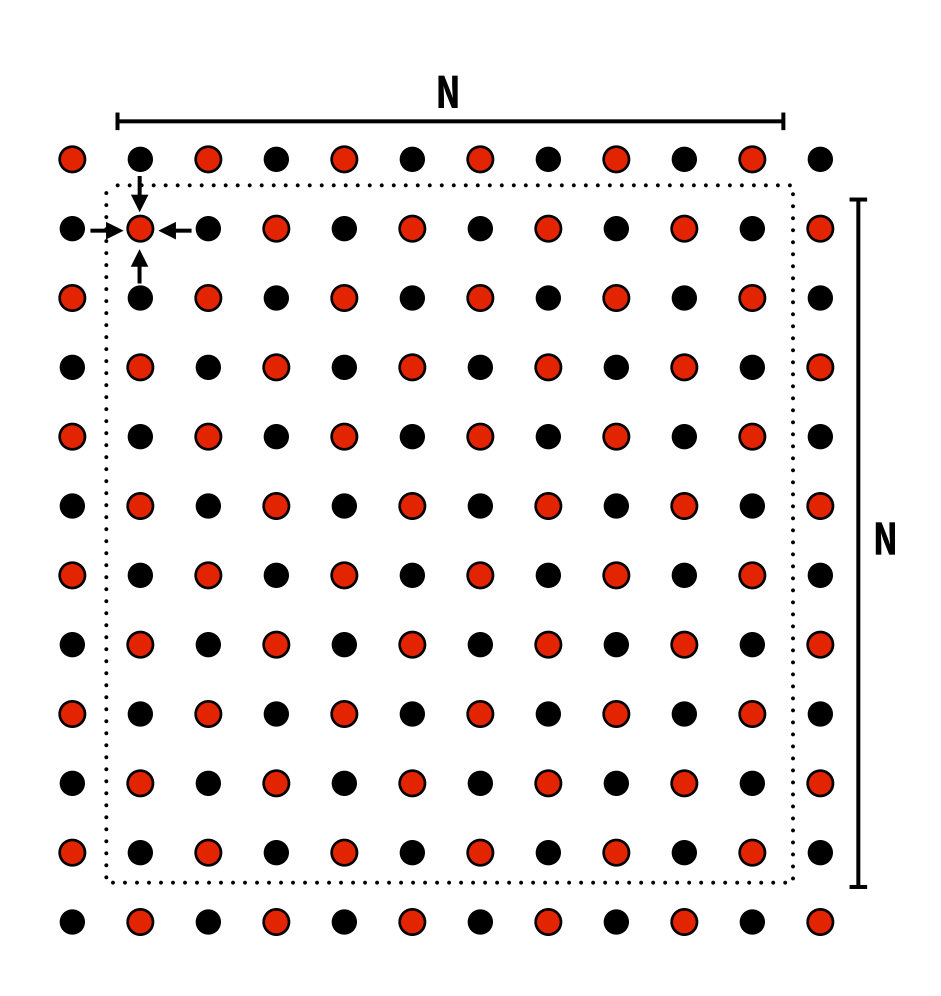
Bad: independent work is hard to exploit Not much parallelism at beginning and end of computation.

Frequent synchronization (after completing each diagonal)

#### Let's make life easier on ourselves

- Idea: improve performance by changing the algorithm to one that is more amenable to parallelism
  - Change the order grid cell cells are updated
  - New algorithm iterates to same solution (approximately),
     but converges to solution differently
    - Note: floating-point values computed are different, but solution still converges to within error threshold
  - Yes, we needed domain knowledge of Gauss-Seidel method for solving a linear system to realize this change is permissible for the application

## New approach: reorder grid cell update via red-black coloring

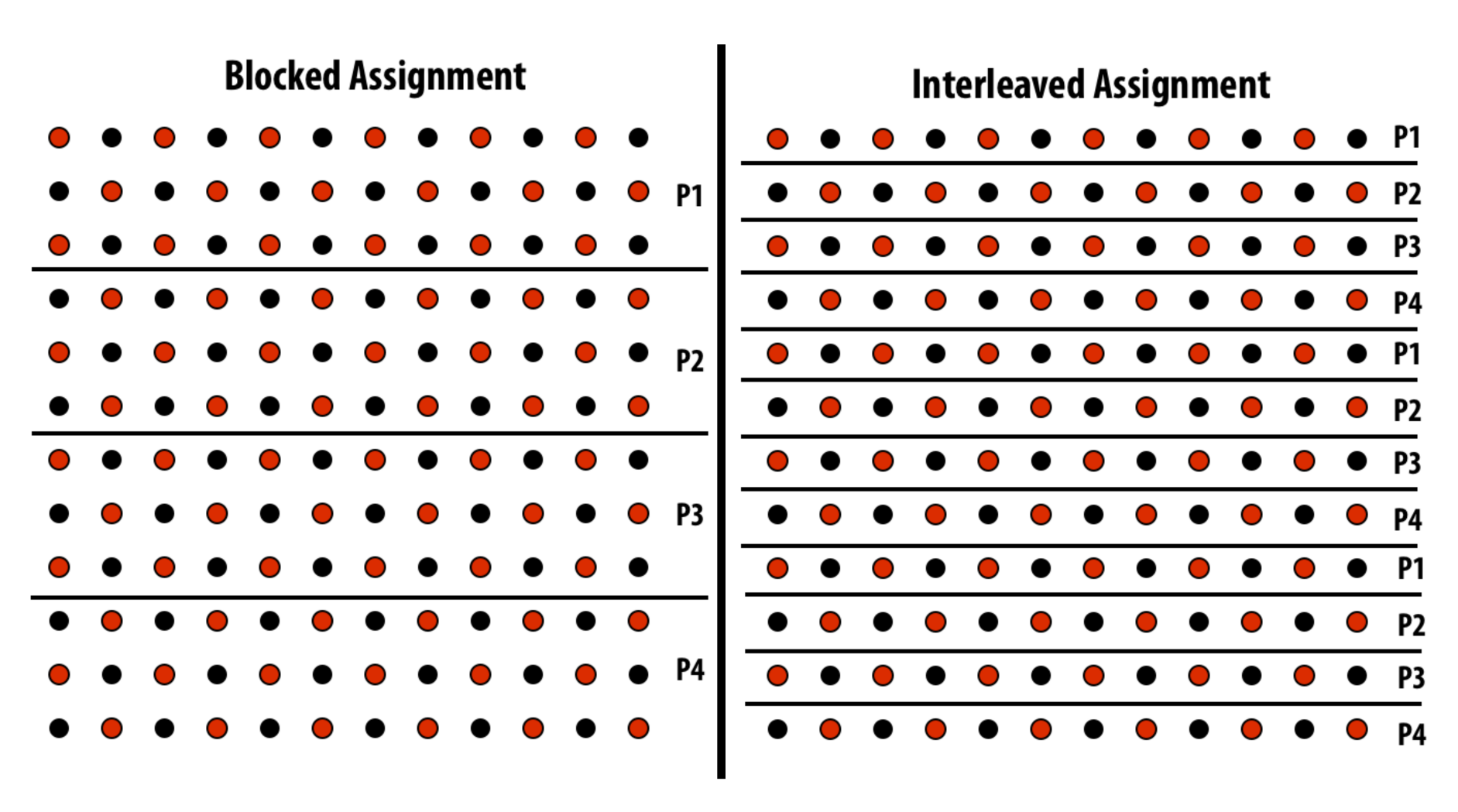


Update all red cells in parallel

When done updating red cells, update all black cells in parallel (respect dependency on red cells)

Repeat until convergence

### Possible assignments of work to processors

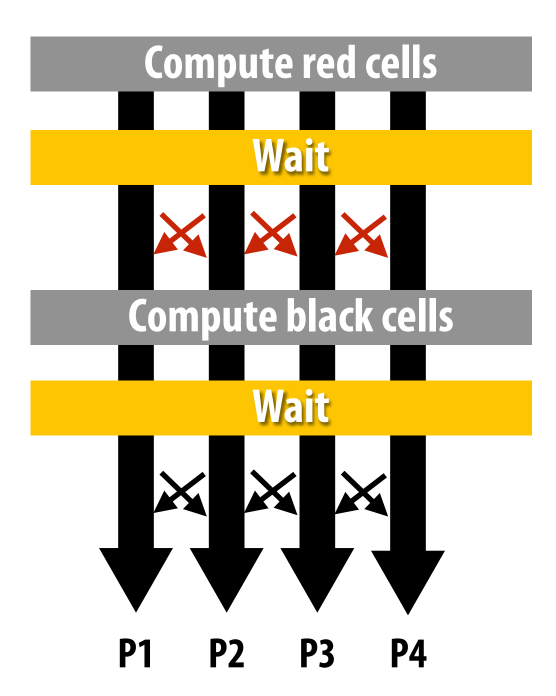


**Question: Which is better? Does it matter?** 

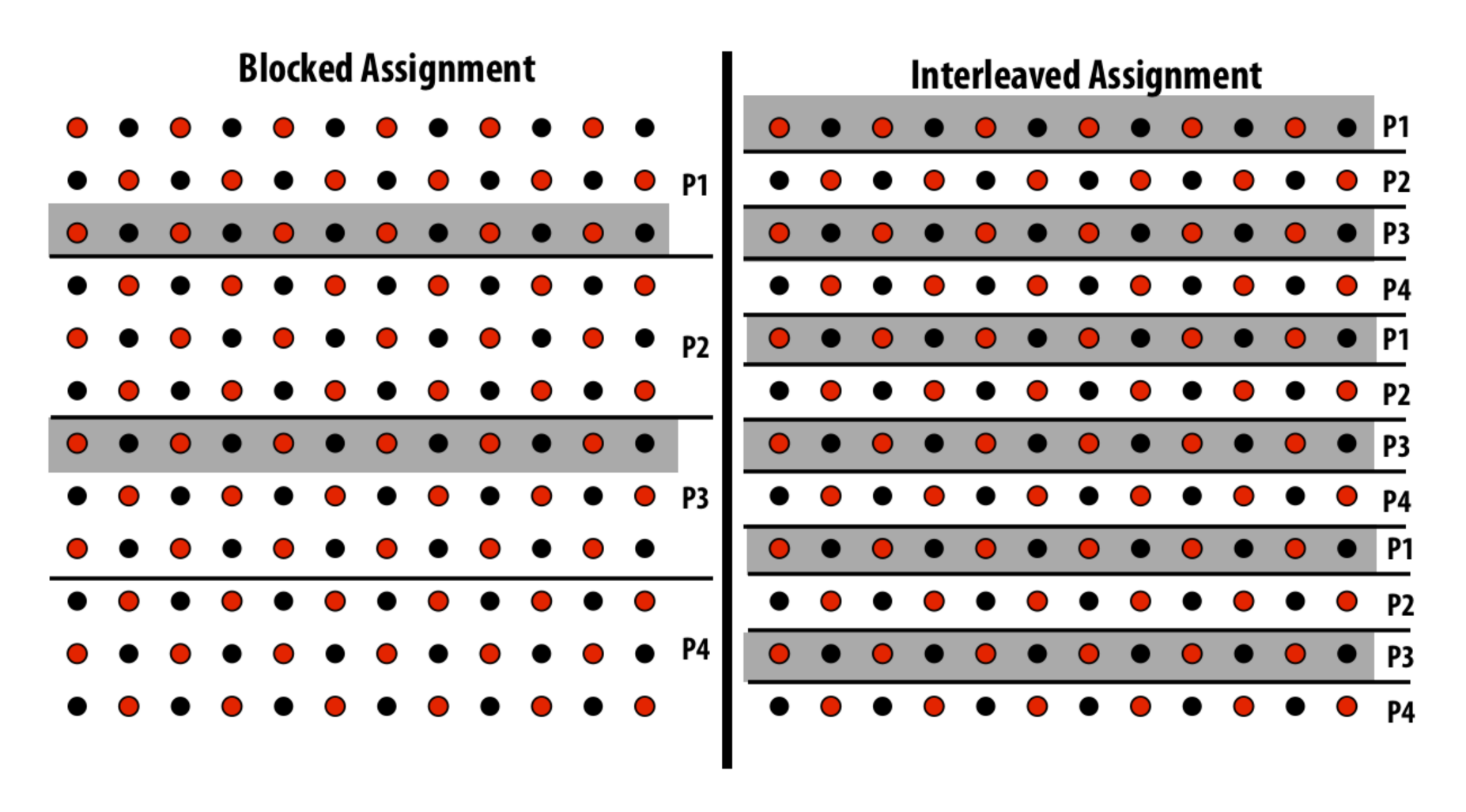
Answer: it depends on the system this program is running on

#### Consider dependencies (data flow)

- 1. Perform red update in parallel
- 2. Wait until all processors done with update
- 3. Communicate updated red cells to other processors
- 4. Perform black update in parallel
- 5. Wait until all processors done with update
- 6. Communicate updated black cells to other processors
- 7. Repeat



## Communication resulting from assignment



= data that must be sent to P2 each iteration

Blocked assignment requires less data to be communicated between processors

## Data-parallel expression of solver

#### Data-parallel expression of grid solver

Note: to simplify pseudocode: just showing red-cell update

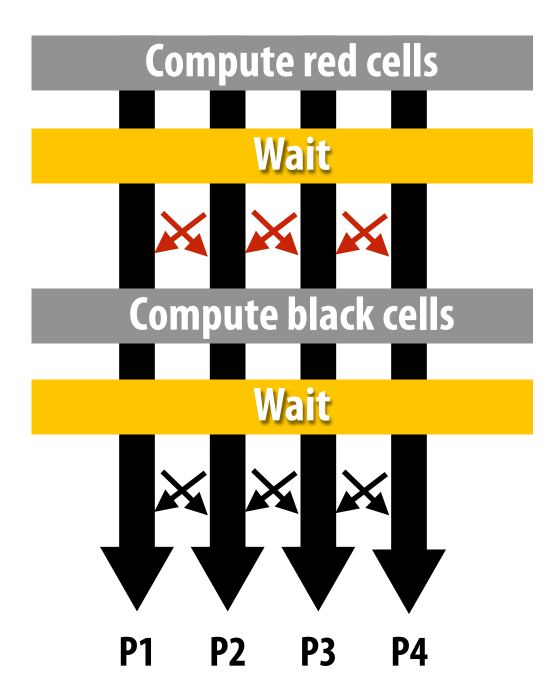
```
const int n;
                                                                                 Assignment: ???
float* A = allocate(n+2, n+2)); // allocate grid
void solve(float* A) {
   bool done = false;
   float diff = 0.f;
   while (!done) {
     for_all (red cells (i,j)) {
                                                                                 decomposition:
          +loat prev = A[i,j];
                                                                                 individual grid
          A[i,j] = 0.2f * (A[i-1,j] + A[i,j-1] + A[i,j] +
                                                                                 elements constitute
                              A[i+1,j] + A[i,j+1]);
                                                                                 independent work
          reduceAdd(diff, abs(A[i,j] - prev));
                                                               Orchestration: handled by system
                                                               (builtin communication primitive: reduceAdd)
     if (diff/(n*n) < TOLERANCE)
          done = true;
                                                                      Orchestration:
                                                                      handled by system
                                                                      (End of for_all block is implicit wait for all
                                                                      workers before returning to sequential control)
```

# Shared address space (with SPMD threads) expression of solver

## Shared address space expression of solver

SPMD execution model

- Programmer is responsible for synchronization
- Common synchronization primitives:
  - Locks (provide mutual exclusion): only one thread in the critical region at a time
  - Barriers: wait for threads to reach this point



## Shared address space solver (pseudocode in SPMD execution model)

```
Assume these are global variables
                                                                                        (accessible to all threads)
                               // grid size
int
        n;
        done = false;
bool
                                                                                       Assume solve function is executed by
        diff = 0.0;
float
LOCK
        myLock;
                                                                                       all threads. (SPMD-style)
BARRIER myBarrier;
// allocate grid
float* A = allocate(n+2, n+2);
                                                                                       Value of threadId is different for
void solve(float* A) {
                                                                                       each SPMD instance: use value to
                                                                                       compute region of grid to work on
   int threadId = getThreadId();
   int myMin = 1 + (threadId * n / NUM PROCESSORS);
   int myMax = myMin + (n / NUM_PROCESSORS)
   while (!done) {
     diff = 0.f;
                                                                                       Each thread computes the rows it is
     barrier(myBarrier, NUM_PROCESSORS);
                                                                                       responsible for updating
     for (j=myMin to myMax) {
        for (i = red cells in this row) {
            float prev = A[i,j];
            A[i,j] = 0.2f * (A[i-1,j] + A[i,j-1] + A[i,j] +
                              A[i+1,j], A[i,j+1]);
            lock(myLock)
            diff += abs(A[i,j] - prev));
            unlock(myLock);
     barrier(myBarrier, NUM PROCESSORS);
     if (diff/(n*n) < TOLERANCE)</pre>
                                                 // check convergence, all threads get same answer
          done = true;
     barrier(myBarrier, NUM_PROCESSORS);
```

#### Review: need for mutual exclusion

- Each thread executes
  - Load the value of diff into register r1
  - Add the register r2 to register r1
  - Store the value of register r1 into diff
- One possible interleaving: (let starting value of diff=0, r2=1)

	T0	<b>T1</b>	
<b>r1</b> ←	- diff		T0 reads value 0
		r1 ← diff	T1 reads value 0
r1 +	- r1 + r2		T0 sets value of its r1 to 1
		r1 ← r1 + r2	T1 sets value of its r1 to 1
diff	e ← r1		T0 stores 1 to diff
		diff ← r1	T1 stores 1 to diff

Need this set of three instructions to be atomic

# Mechanisms for preserving atomicity

Lock/unlock mutex around a critical section

```
LOCK(mylock);
// critical section
UNLOCK(mylock);
```

Some languages have first-class support for atomicity of code blocks

```
atomic {
   // critical section
}
```

Intrinsics for hardware-supported atomic read-modify-write operations

```
atomicAdd(x, 10);
```

## Shared address space solver

// grid size

(pseudocode in SPMD execution model)

```
bool done = false;
float diff = 0.0;
LOCK
       myLock;
BARRIER myBarrier;
                                                     Do you see a potential performance
// allocate grid
float* A = allocate(n+2, n+2);
                                                     problem with this implementation?
void solve(float* A) {
   int threadId = getThreadId();
   int myMin = 1 + (threadId * n / NUM_PROCESSORS);
   int myMax = myMin + (n / NUM_PROCESSORS)
   while (!done) {
    diff = 0.f;
     barrier(myBarrier, NUM_PROCESSORS);
     for (j=myMin to myMax) {
       for (i = red cells in this row) {
          float prev = A[i,j];
          A[i,j] = 0.2f * (A[i-1,j] + A[i,j-1] + A[i,j] +
                           A[i+1,j], A[i,j+1]);
          lock(myLock)
          diff += abs(A[i,j] - prev));
          unlock(myLock);
     barrier(myBarrier, NUM_PROCESSORS);
     if (diff/(n*n) < TOLERANCE)</pre>
                                           // check convergence, all threads get same answer
         done = true;
     barrier(myBarrier, NUM_PROCESSORS);
```

int

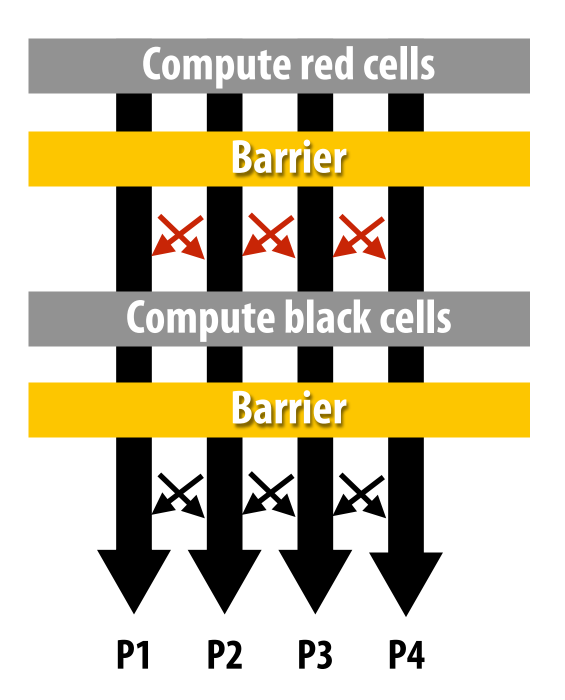
n;

#### Shared address space solver (SPMD execution model)

```
// grid size
int
       n;
      done = false;
bool
     diff = 0.0;
float
                                                     Improve performance by accumulating
LOCK
       myLock;
BARRIER myBarrier;
                                                     into partial sum locally, then complete
// allocate grid
                                                     reduction globally at the end of the
float* A = allocate(n+2, n+2);
void solve(float* A) {
                                                      iteration.
   float myDiff;
   int threadId = getThreadId();
   int myMin = 1 + (threadId * n / NUM_PROCESSORS);
   int myMax = myMin + (n / NUM PROCESSORS)
   while (!done) {
     float myDiff = 0.f;
     diff = 0.f;
     barrier(myBarrier, NUM_PROCESSORS);
     for (j=myMin to myMax) {
        for (i = red cells in this row) {
           float prev = A[i,j];
           A[i,j] = 0.2f * (A[i-1,j] + A[i,j-1] + A[i,j] +
                                                                                compute per worker partial sum
                            A[i+1,j], A[i,j+1]);
           myDiff += abs(A[i,j] - prev));
                                                              Now only only lock once per thread, not once
    lock(myLock);
                                                              per (i,j) loop iteration!
     diff += myDiff;
     unlock(myLock);
     barrier(myBarrier, NUM_PROCESSORS);
                                            // check convergence, all threads get same answer
     if (diff/(n*n) < TOLERANCE)</pre>
         done = true;
     barrier(myBarrier, NUM PROCESSORS);
```

#### Barrier synchronization primitive

- barrier(num\_threads)
- Barriers are a conservative way to express dependencies
- Barriers divide computation into phases
- All computations by all threads before the barrier complete before any computation in any thread after the barrier begins



#### Shared address space solver (SPMD execution model)

```
int
                         // grid size
       n;
bool done = false;
float diff = 0.0;
                                                          Why are there three barriers?
       myLock;
LOCK
BARRIER myBarrier;
// allocate grid
float* A = allocate(n+2, n+2);
void solve(float* A) {
   float myDiff;
   int threadId = getThreadId();
   int myMin = 1 + (threadId * n / NUM_PROCESSORS);
   int myMax = myMin + (n / NUM_PROCESSORS)
   while (!done) {
     float myDiff = 0.f;
     diff = 0.f:
     barrier(myBarrier, NUM PROCESSORS);
     for (j=myMin to myMax) {
        for (i = red cells in this row) {
           float prev = A[i,j];
           A[i,j] = 0.2f * (A[i-1,j] + A[i,j-1] + A[i,j] +
                            A[i+1,j], A[i,j+1]);
           myDiff += abs(A[i,j] - prev));
     lock(myLock);
     diff += myDiff;
     unlock(mvLock):
     barrier(myBarrier, NUM_PROCESSORS);
     if (diff/(n*n) < TOLERANCE)</pre>
                                            // check convergence, all threads get same answer
         done = true;
     barrier(myBarrier, NUM_PROCESSORS);
```

#### Shared address space solver: one barrier

```
Idea:
                        // grid size
int
bool done = false;
LOCK
       myLock;
                                                   Remove dependencies by using different diff
BARRIER myBarrier;
float diff[3]; // global diff, but now 3 copies
                                                   variables in successive loop iterations
float *A = allocate(n+2, n+2);
void solve(float* A) {
                                                   Trade off footprint for removing dependencies!
  float myDiff; // thread local variable
                                                   (a common parallel programming technique)
 int index = 0; // thread local variable
 diff[0] = 0.0f;
  barrier(myBarrier, NUM_PROCESSORS); // one-time only: just for init
 while (!done) {
   myDiff = 0.0f;
    // perform computation (accumulate locally into myDiff)
    //
    lock(myLock);
    diff[index] += myDiff; // atomically update global diff
    unlock(myLock);
    diff[(index+1) \% 3] = 0.0f;
   barrier(myBarrier, NUM_PROCESSORS);
    if (diff[index]/(n*n) < TOLERANCE)</pre>
      break;
    index = (index + 1) \% 3;
```

## More on specifying dependencies

- Barriers: simple, but conservative (coarse-granularity dependencies)
  - All work in program up until this point (for all threads) must finish before any thread begins next phase
- Specifying specific dependencies can increase performance (by revealing more parallelism)
  - Example: two threads. One produces a result, the other consumes it.

```
T0

// produce x, then let T1 know

x = 1;

flag = 1;

// do more work here...

T1

// do stuff independent

// of x here

while (flag == 0);

print x;
```

We just implemented a message queue (of length 1)

```
T0 \rightarrow \Box \Box \Box \rightarrow T1
```

#### Solver implementation in two programming models

#### Data-parallel programming model

- Synchronization:
  - Single logical thread of control, but iterations of forall loop <u>may</u> be parallelized by the system (implicit barrier at end of forall loop body)
- Communication
  - Implicit in loads and stores (like shared address space)
  - Special built-in primitives for more complex communication patterns:
     e.g., reduce

#### Shared address space

- Synchronization:
  - Mutual exclusion required for shared variables (e.g., via locks)
  - Barriers used to express dependencies (between phases of computation)
- Communication
  - Implicit in loads/stores to shared variables

# We will defer discussion of the message passing expression of solver to a later class.

#### Summary

- Amdahl's Law
  - Overall maximum speedup from parallelism is limited by amount of serial execution in a program
- Aspects of creating a parallel program
  - Decomposition to create independent work, assignment of work to workers, orchestration (to coordinate processing of work by workers), mapping to hardware
  - We'll talk a lot about making good decisions in each of these phases in the coming lectures (in practice, they are very inter-related)
- **■** Focus today: identifying dependencies
- Focus soon: identifying locality, reducing synchronization