Parallel programming principles: Task decomposition

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- ► Atenea: Unit 3 Task decomposition
  - ▶ Video lesson 4 (partially covering slides 5–20)
  - Questions after video lesson 4
  - Going further: cut-off based on number of tasks pending to be executed (OPTIONAL)
- Collection of Exercises: problems in Chapter 3



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- From a specification that solves the original problem, find a decomposition of the problem
  - Identify pieces of work (tasks) that can be done concurrently
  - Identify data structures (input, output and/or intermediate) or parts of them that can be managed concurrently
  - ... ensuring that the same result is produced
    - ► Identify dependencies that impose ordering constraints (synchronizations) and data sharing



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Task ordering

# Objective: problem decomposition

- Having in mind two productivity goals:
  - Performance: maximize concurrency and reduce overheads (maximize potential speedup)
  - Programmability: readability and portability, target architecture independency
- ► Two usual approaches to problem decomposition
  - Task decomposition (Chapter 3)
  - Data decomposition (Chapter 5)

Either way, you'll look at both tasks and data; difference is in which you look at first, and then the other follows



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### Outline

#### Introduction to problem decomposition



- #pragma omp parallel: One implicit task is created for each thread in the team (and immediately executed)
  - #pragma omp for worksharing: iterations in a loop are distributed among the implicit tasks in the parallel region
- #pragma omp task: One explicit task is created, packaging code and data for (possible) deferred execution
  - Tasks can be nested



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- ► Linear task decomposition
  - ► Task = code block or procedure invocation
- ▶ (Linear) Iterative task decomposition
  - Tasks = body of iterative constructs, such as loops (countable or uncountable)
  - ► Examples: Pi computation, Mandelbrot and heat diffusion equation in lab sessions, vector and matrix operations, ...
- Recursive task decomposition
  - Tasks = recursive procedure invocations, for example in divide-and-conquer problems
  - Examples: Fibonacci, multisort in lab session, branch and bound problems, ...



A task is a sequence of instructions, as for example in sum of two vectors artificially divided in two parts:

```
void vector_add(int *A, int *B, int *C, int n) {
    #pragma omp task
    for (int i=0; i< n/2; i++)
        C[i] = A[i] + B[i];

    #pragma omp task
    for (int i=n/2; i< n; i++)
        C[i] = A[i] + B[i];
}

void main() {
    ....
    #pragma omp parallel
    #pragma omp single
    vector_add(a, b, c, N);
    ....
}</pre>
```



## Example 2: (linear) iterative task decomposition

But more naturally, a task can be a single loop iteration:

```
void vector_add(int *A, int *B, int *C, int n) {
  for (int i=0; i< n; ii++)
       #pragma omp task
      C[i] = A[i] + B[i];
7
void main() {
  #pragma omp parallel
  #pragma omp single
  vector_add(a, b, c, N);
}
```

Each explicit task executes a single iteration of the i loop, large task creation overhead, very fine granularity!

# Example 2: (linear) iterative task decomposition (cont.)

#### Chunk of loop iterations, requires loop transformation:

```
void vector_add(int *A, int *B, int *C, int n) {
  for (int ii=0; ii< n; ii+=BS)
       #pragma omp task
      for (int i = ii; i < min(ii+BS, n), i++)
           C[i] = A[i] + B[i];
void main() {
  #pragma omp parallel
  #pragma omp single
  vector_add(a, b, c, N);
```

Outer loop jumps over chunks of BS iterations, inner loop traverses each chunk

# Example 2: (linear) iterative task decomposition (cont.)

#### OpenMP provides two alternative constructs, one using taskloop:

```
void vector add(int *A, int *B, int *C, int n) {
  #pragma omp taskloop grainsize(BS) // or alternatively num_tasks(n/BS)
  for (int i=0; i< n; i++)
      C[i] = A[i] + B[i]:
void main() {
  #pragma omp parallel
  #pragma omp single
   ... vector_add(a, b, c, N); ...
7
```

#### and another using implicit tasks and worksharing:

```
void vector add(int *A, int *B, int *C, int n) {
   #pragma omp parallel for schedule(dynamic,BS)
  for (int i=0: i< n: i++)
       C[i] = A[i] + B[i]:
void main() {
   ... vector_add(a, b, c, N); ...
7
```



# Example 3: (linear) iterative task decomposition

List of elements, traversed using an uncountable (while) loop

```
int main() {
    struct node *p;

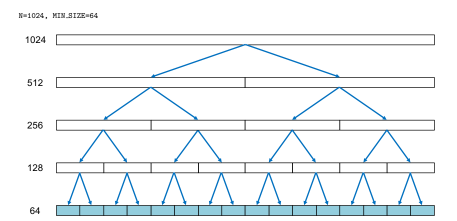
    p = init.list(n);
    ...

#pragma omp parallel
#pragma omp single
while (p != NULL) {
        #pragma omp task firstprivate(p)
        process.work(p);
        p = p->next;
     }
    ...
}
```

# Example 4: "Divide-and-conquer" task decomposition

Sum of two vectors by recursively dividing the problem into smaller sub-problems

```
#define N 1024
#define MIN SIZE 64
void vector_add(int *A, int *B, int *C, int n) {
  for (int i=0; i< n; i++) C[i] = A[i] + B[i];
void rec_vector_add(int *A, int *B, int *C, int n) {
  if (n>MIN_SIZE) {
      int n2 = n / 2:
      rec_vector_add(A, B, C, n2):
      rec_vector_add(A+n2, B+n2, C+n2, n-n2);
   else vector_add(A, B, C, n):
void main() {
  rec_vector_add(a, b, c, N);
```



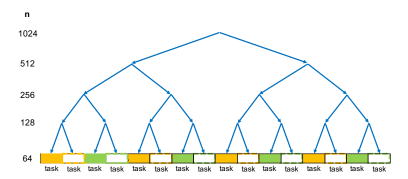


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### Two possible decomposition strategies

▶ **Leaf strategy**: a task corresponds with each invocation of vector\_add once the recursive invocations stop



Sequential generation of tasks



# Implementing the decomposition strategies

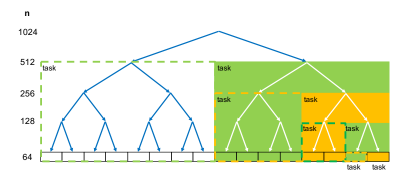
### Leaf parallelization

```
#define N 1024
#define MIN SIZE 64
void vector_add(int *A, int *B, int *C, int n) {
  for (int i=0: i< n: i++) C[i] = A[i] + B[i]:
}
void rec_vector_add(int *A, int *B, int *C, int n) {
  if (n>MIN SIZE) {
      int n2 = n / 2:
      rec_vector_add(A, B, C, n2);
      rec vector add(A+n2, B+n2, C+n2, n-n2);
  else
       #pragma omp task
       vector add(A, B, C, n):
}
void main() {
   . . . .
   #pragma omp parallel
  #pragma omp single
  rec_vector_add(a, b, c, N);
   . . .
```

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▶ Tree strategy: a task corresponds with each invocation of rec vector add



- Parallel generation of tasks
- Granularity: some tasks simply generate new tasks



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# Implementing the decomposition strategies (cont.)

#### Tree parallelization

```
#define N 1024
#define MIN SIZE 64
void vector_add(int *A, int *B, int *C, int n) {
  for (int i=0; i< n; i++) C[i] = A[i] + B[i];
}
void rec_vector_add(int *A, int *B, int *C, int n) {
  if (n>MIN SIZE) {
       int n2 = n / 2:
       #pragma omp task
       rec_vector_add(A, B, C, n2);
       #pragma omp task
      rec_vector_add(A+n2, B+n2, C+n2, n-n2);
  else vector add(A. B. C. n):
}
void main() {
   . . . .
   #pragma omp parallel
  #pragma omp single
  rec_vector_add(a, b, c, N);
   . . .
```

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Introduction to problem decomposition

Task decomposition strategies

Task generation contro

Task ordering constraints

Data sharing constraints



# Task generation control

Excessive task generation if recursion depth is very high

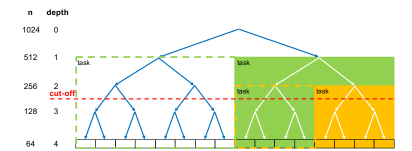
- May cause excessive overhead
- It may be not necessary for certain problems
- ▶ Need to stop task generation ... cut-off control
  - after certain number of recursive calls (static control)
  - when the size of the vector is too small (static control)
  - when the number of generated tasks is too much (dynamic control)



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### Cut-off control

#### Tree parallelization with depth recursion control





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# Cut-off control (cont.)

### Tree parallelization with depth recursion control

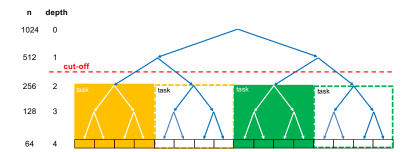
```
#define CUTOFF 3
void rec vector add(int *A. int *B. int *C. int n. int depth) {
    if (n>MIN_SIZE) {
        int n2 = n / 2:
       if (depth < CUTOFF) {
            #pragma omp task
            rec_vector_add(A, B, C, n2, depth+1);
            #pragma omp task
            rec_vector_add(A+n2, B+n2, C+n2, n-n2, depth+1);
       else {
            rec_vector_add(A, B, C, n2, depth+1);
            rec_vector_add(A+n2, B+n2, C+n2, n-n2, depth+1);
    } else vector add(A, B, C, n):
}
void main() {
   #pragma omp parallel
   #pragma omp single
   rec_vector_add(a, b, c, N, 0);
```

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#### Cut-off control

#### Leaf parallelization with depth recursion control





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#### Leaf parallelization with depth recursion control

```
#define CUTOFF 2
void rec_vector_add(int *A, int *B, int *C, int n, int depth) {
    if (n>MIN_SIZE) {
        int n2 = n / 2:
        if (depth == CUTOFF) {
            #pragma omp task
            rec_vector_add(A, B, C, n2, depth+1);
            #pragma omp task
            rec_vector_add(A+n2, B+n2, C+n2, n-n2, depth+1);
        else {
            rec_vector_add(A, B, C, n2, depth+1);
            rec_vector_add(A+n2, B+n2, C+n2, n-n2, depth+1);
    7
   else
        if (depth <= CUTOFF)
            #pragma omp task
            vector_add(A, B, C, n);
        else
            vector add(A, B, C, n):
. . .
```



- ▶ if clause: If the expression of an if clause evaluates to false the encountering parent task is suspended and the new one undeferred with respect it (but not necessarily executed by same thread). The parent resumes when the undeferred task finishes
- ▶ final clause: If the expression of a final clause evaluates to *true* the generated task and **all of its descendent tasks** will be final. The execution of a final task is sequentially **included** in the generating task



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mergeable clause: when a mergeable clause is present on a task construct, and the generated task is an undeferred task or an included task, then the compiler may choose to generate a merged task instead. If a merged task is generated, then the behavior is as though there was no task directive at all

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```
#define N 1024
#define MIN SIZE 64
#define CUTOFF 3
void vector_add(int *A, int *B, int *C, int n) {
  for (int i=0: i< n: i++) C[i] = A[i] + B[i]:
7
void rec vector add(int *A. int *B. int *C. int n. int depth) {
   if (n>MIN SIZE) {
       int n2 = n / 2:
       #pragma omp task final(depth >= CUTOFF) mergeable
       rec_vector_add(A, B, C, n2, depth+1);
       #pragma omp task final(depth >= CUTOFF) mergeable
       rec_vector_add(A+n2, B+n2, C+n2, n-n2, depth+1);
   else vector_add(A, B, C, n);
void main() {
  #pragma omp parallel
  #pragma omp single
  rec vector add(a, b, c, N, 0):
   . . .
```

### Outline

Introduction to problem decomposition

Task decomposition strategies

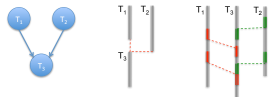
Task generation control

Task ordering constraints

Data sharing constraints



- Constraints in the parallel execution of tasks
  - ► Task ordering constraints: they force the execution of (groups of) tasks in a required order
  - Data sharing constraints: they force the access to data to fulfil certain properties (write-after-read, exclusive, commutative, ...)



Cut-off

- ▶ If no constraints are defined during the parallel execution of the tasks, the algorithm is called "embarrassingly" parallel. There are still challenges in this case:
  - Load balancing: how to distribute the work to perform so the load is evenly balanced among tasks (e.g. pi computation vs. Mandelbrot set)
  - ▶ Data locality: how to assign work to tasks so that data resides in the memory hierarchy levels close to the processor



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Task ordering

# Task ordering constraints

- ► Control flow constraints: the creation of a task depends on the outcome (decision) of one or more previous tasks
- ▶ Data flow constraints: the execution of a task can not start until one or more previous tasks have computed some data

► Task ordering constraints are easily imposed by sequentially composing tasks and/or using global synchronizations.



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- ► Thread barriers (#pragma omp barrier): wait for all threads to finish previous work
- Task barriers:
  - taskwait: Suspends the current task waiting on the completion of child tasks of the current task. The taskwait construct is a stand-alone directive.
  - taskgroup: Suspends the current task at the end of structured block waiting on completion of child tasks of the current task and their descendent tasks.
- Task dependences



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# Fibonnaci series example

#### Sequential code

```
long fib(long n)
{
    if (n < 2) return n;
        return(fib(n-1) + fib(n-2));
}

void main (int argc, char *argv[])
{
        n = atoi(argv[1]);
        res = fib(n);
        printf("Fibonacci for %d is %d", n, res);
}</pre>
```

- ▶ The invocations of fib for n-1 and n-2 can be a task
- We need to guarantee that both instances of fib finish before returning the result



#### OpenMP code using task and taskwait, with cut-off

```
long fib_parallel(long n, int d)
   long x, y;
   if (n < 2) return n:
    if (d<CUTOFF) {
        #pragma omp task shared(x) // firstprivate(n) by default
        x = fib_parallel(n-1, d+1):
        #pragma omp task shared(y) // firstprivate(n) by default
        v = fib_parallel(n-2, d+1);
        #pragma omp taskwait
    } else {
        x = fib_parallel(n-1, d); // or fib(n-1)
       y = fib_parallel(n-2, d); // or fib(n-2)
   return (x+y);
void main (int argc, char *argv∏)
     n = atoi(argv[1]);
     #pragma omp parallel
     #pragma omp single
     res = fib_parallel(n,0);
     printf("Fibonacci for %d is %d", n, res);
```

 OpenMP allows the specification of dependences (through argument directionality) between sibling tasks (i.e. from the same parent task)

Task dependences are derived from the dependence type (in, out or inout) and its items in var\_list. This list may include array sections

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- ▶ The in dependence-type: the generated task will be a dependent task of all previously generated sibling tasks that reference at least one of the list items in an out or inout dependence-type list
- ► The out and inout dependence-types: the generated task will be a dependent task of all previously generated sibling tasks that reference at least one of the list items in an in. out, or inout dependence-type list.



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## Task dependences

- Example:
  - Function foo(i, j) processes block(i, j)
  - ► Wave-front execution: the execution of foo(i,j) depends on foo(i-1, j) and foo(i, j-1)

```
#pragma omp parallel private(i, j)
#pragma omp single
  for (i=1; i<n i++) {
       for (j=1; j<n;j++) {
           #pragma omp task // firstprivate(i, j) by default
                            depend(in : block[i-1][j], block[i][j-1])
                            depend(out: block[i][j])
           foo(i,j);
```

A **doacross** loop is a loop nest where cross-iteration dependences exist

- ► The ordered(n) clause with an integer argument n is used to define the number of loops within the doacross nest
- depend clauses on ordered constructs within an ordered loop describe the dependences of the doacross loops
  - depend(sink:expr) defines the wait point for the completion of computation in a previous iteration defined by expr
  - depend(source) indicates the completion of computation from the current iteration



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Task ordering

## The doacross loop nest: first example

```
#pragma omp for ordered(1)
for ( i = 1; i < N; i++ ) {
     A[i] = foo(i);
     #pragma omp ordered depend(sink: i-1)
     B[i] = goo(A[i], B[i-1]);
     #pragma omp ordered depend(source)
     C[i] = too(B[i]);
```

- ▶ In this example an i-1 to i cross-iteration dependence is defined, but only for the statement computing B.
- ▶ The computation of A and C can go in parallel across multiple iterations.



#### The doacross loop nest: wavefront example

- Example:
  - Function foo(i, j) processes block(i, j)
  - ► Wave-front execution: the execution of foo(i,j) depends on foo(i-1, j) and foo(i, j-1)

```
#pragma omp for schedule(static.1) ordered(2)
for ( i = 1; i < N; i++ )
  for (j = 1; j < N; j++) {
      #pragma omp ordered depend(sink: i-1,j) depend(sink: i,j-1)
     foo (i, j);
      #pragma omp ordered depend(source)
```



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#### The doacross loop nest: last example

```
#pragma omp for collapse(2) ordered(2)
for (i = 1; i < N-1; i++) {
   for (j = 1; j < M-1; j++) {
        A[i][j] = foo(i, j);
        #pragma omp ordered depend(source)
        B[i][j] = alpha * A[i][j];
        #pragma omp ordered depend(sink: i-1,j) depend(sink: i,j-1)
        C[i][j] = 0.2 * (A[i-1][j] + A[i+1][j] + A[i][j-1] + A[i][j+1]);
        }
}</pre>
```

- ▶ In this example the i and j loops are the associated loops for the collapsed loop as well as for the doacross loop nest.
- ► The dependence source directive is placed before the corresponding sink directive.



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Task ordering

Introduction to problem decomposition

Task decomposition strategies

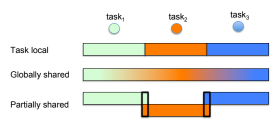
Task generation control

Task ordering constraints

Data sharing constraints



- Task-local data: variables (or chunks of them) only used by single task
- Globally shared data: variables (or chunks of them) not associated with any particular task
- Partially shared data: variables (or chunks of them) shared among smaller groups of tasks (e.g. boundary elements or halos as in Gauss-Seidel)





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- ► Task interactions are needed to guarantee proper access to shared data, without adding too much overhead
  - With shared memory architectures, all processors have access to all data, but must use synchronisation to prevent 'race conditions'
  - With distributed memory architectures, each processor has its own data, so race conditions are not possible, but must use communication to (in effect) share data (next chapter)
- ▶ Basic approach: first identify what data is shared, second figure out how it is accessed, and finally add the appropriate interactions to guarantee correctness



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#### Protecting task interactions

Sequence of statements in a task that may conflict with a sequence of statements in another task, creating a possible data race

Two sequences of statements conflict if both access the same data and at least one of them modifies the data

#### Two mechanisms:

- Atomic accesses: mechanism to guarantee atomicity in load/store instruction pairs
- Mutual exclusion: mechanism to ensure that only one task at a time executes the code within the critical section



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#### Task interactions: atomic construct

```
#pragma omp atomic [update | read | write]
     expression
```

- Ensures that a specific storage location is accessed atomically, avoiding the possibility of multiple, simultaneous reading and writing threads
  - Atomic updates: x += 1, x = x foo(), x[index[i]]++
  - Atomic reads: value = \*p
  - ► Atomic writes: \*p = value
- Only protects the read/operation/write
- Usually more efficient than a critical construct



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In OpenMP two alternatives for mutual exclusion: critical and low-level synchronization functions

- critical (name) pragma: if a thread arrives at the top of the critical-section block while another thread is processing the block, it waits until the prior thread exits
  - ► A critical section in OpenMP implies a memory flush on entry to and on exit from it (memory consistency)
  - name is an identifier that can be used to support disjoint sets of critical sections

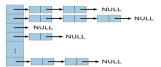


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linked lists

# ► Inserting elements in a hash table, defined as a collection of



```
for (i = 0; i < elements; i++) {
  index = hash(element[i]);
  insert_element (element[i], index);
}</pre>
```

▶ Updates to the list in any particular slot must be protected to prevent a race condition

```
#pragma omp parallel for private (index)
for (i = 0; i < elements; i++) {
   index = hash(element[i]);
   #pragma omp critical
   insert.element (element[i], index);
}</pre>
```



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 Low-level synchronization functions that provide a lock capability

```
void omp_init_lock(omp_lock_t *lock)
void omp_destroy_lock(omp_lock_t *lock)

void omp_set_lock(omp_lock_t *lock)
void omp_unset_lock(omp_lock_t *lock)
int omp_test_lock(omp_lock_t *lock)
```

- ► The lock functions guarantee that the lock variable itself is consistently updated between threads, ...
- ... but do not imply a flush of other variables
- Nested locks are possible → locking anomalies (deadlock)



Associate a lock variable with each slot in the hash table, protecting the chain of elements in an slot

```
/* hash.lock declared as type omp.lock.t */
omp_lock.t hash.lock[HASH.TABLE_SIZE];
/* locks initialed in function main */
for (i = 0; i < HASH.TABLE.SIZE; i++)
    omp.init.lock(&hash.lock[i]);

#pragma omp parallel for private (index)
for (i = 0; i < elements; i++) {
    index = hash(element[i]);
    omp.set.lock (&hash.lock[index]);
    insert.element (element[i], index);
    #pragma omp flush
    omp.unset.lock (&hash_lock[index]);
}
/* locks destroyed in function main */
for (i = 0; i < HASH.TABLE.SIZE; i++)
    omp.destroy.lock(&hash.lock[i]);</pre>
```

► Threads may be inserting elements into the hash table in parallel, as long as these elements hash to different slots

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#### Reducing task interactions

- Group tasks that have strong data interactions. This also has an effect in increasing task granularity
- Replicate computations into local structures to avoid data sharing
- ▶ Separable dependencies (reductions): the dependencies between tasks can be managed by replicating key data structures and then accumulating results into these local structures. The tasks then execute according to the embarrassingly parallel pattern and the local replicated data structures are combined into the final global result

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