

Introduction to OpenMP

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Loops with Tasks

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Introduction to OpenMP



- Task generating construct: decompose a loop into chunks, create a task for each loop chunk

```
#pragma omp taskloop [clause[[,] clause]...]  
{structured-for-loops}
```

- Where clause is one of:

→ shared(list) → private(list) → firstprivate(list) → lastprivate(list) → default(sh <u>pr</u> / <u>fp</u> none) → reduction(r-id: list) → in_reduction(r-id: list)	Data Environment
→ grainsize(grain-size) → num_tasks(num-tasks)	Chunks/Grain

→ if(scalar-expression) → final(scalar-expression) → mergeable	Cutoff Strategies
→ untied → priority(priority-value)	Scheduler (R/H)
→ collapse(n) → nogroup → allocate([allocator:] list)	Miscellaneous



Worksharing vs. taskloop constructs (1/2)

```
subroutine worksharing
  integer :: x
  integer :: i
  integer, parameter :: T = 16
  integer, parameter :: N = 1024

  x = 0
  !$omp parallel shared(x) num_threads(T)

  !$omp do
    do i = 1,N
      !$omp atomic
        x = x + 1
      !$omp end atomic
    end do
  !$omp end do

  !$omp end parallel
  write (*, '(A,I0)') 'x = ', x
end subroutine
```

Result: x = 1024

```
subroutine taskloop
  integer :: x
  integer :: i
  integer, parameter :: T = 16
  integer, parameter :: N = 1024

  x = 0
  !$omp parallel shared(x) num_threads(T)

  !$omp taskloop
    do i = 1,N
      !$omp atomic
        x = x + 1
      !$omp end atomic
    end do
  !$omp end taskloop

  !$omp end parallel
  write (*, '(A,I0)') 'x = ', x
end subroutine
```

Result: x = 16384

Worksharing vs. taskloop constructs (2/2)

```
subroutine worksharing
  integer :: x
  integer :: i
  integer, parameter :: T = 16
  integer, parameter :: N = 1024

  x = 0
  !$omp parallel shared(x) num_threads(T)

  !$omp do
    do i = 1,N
  !$omp atomic
      x = x + 1
  !$omp end atomic
    end do
  !$omp end do

  !$omp end parallel
  write (*, '(A,I0)') 'x = ', x
end subroutine
```

Result: x = 1024

```
subroutine taskloop
  integer :: x
  integer :: i
  integer, parameter :: T = 16
  integer, parameter :: N = 1024

  x = 0
  !$omp parallel shared(x) num_threads(T)
  !$omp single
  !$omp taskloop
    do i = 1,N
  !$omp atomic
      x = x + 1
  !$omp end atomic
    end do
  !$omp end taskloop
  !$omp end single
  !$omp end parallel
  write (*, '(A,I0)') 'x = ', x
end subroutine
```

Result: x = 1024

Clause: grainsize(grain-size)

- Chunks have at least grain-size iterations
- Chunks have maximum 2x grain-size iterations

```
int TS = 4 * 1024;
#pragma omp taskloop grainsize(TS)
for ( i = 0; i<SIZE; i+=1) {
    A[i]=A[i]*B[i]*S;
}
```

Clause: num_tasks(num-tasks)

- Create num-tasks chunks
- Each chunk must have at least one iteration

```
int NT = 4 * omp_get_num_threads();
#pragma omp taskloop num_tasks(NT)
for ( i = 0; i<SIZE; i+=1) {
    A[i]=A[i]*B[i]*S;
}
```

- If none of previous clauses is present, the *number of chunks* and the *number of iterations per chunk* is implementation defined
- Additional considerations:
 - The order of the creation of the loop tasks is unspecified
 - Taskloop creates an implicit taskgroup region; **nogroup** → no implicit taskgroup region is created

Questions?

