



EPSRC

# Introduction to OpenMP

Lecture 6: Further topics in OpenMP



| epcc |



# Nested parallelism

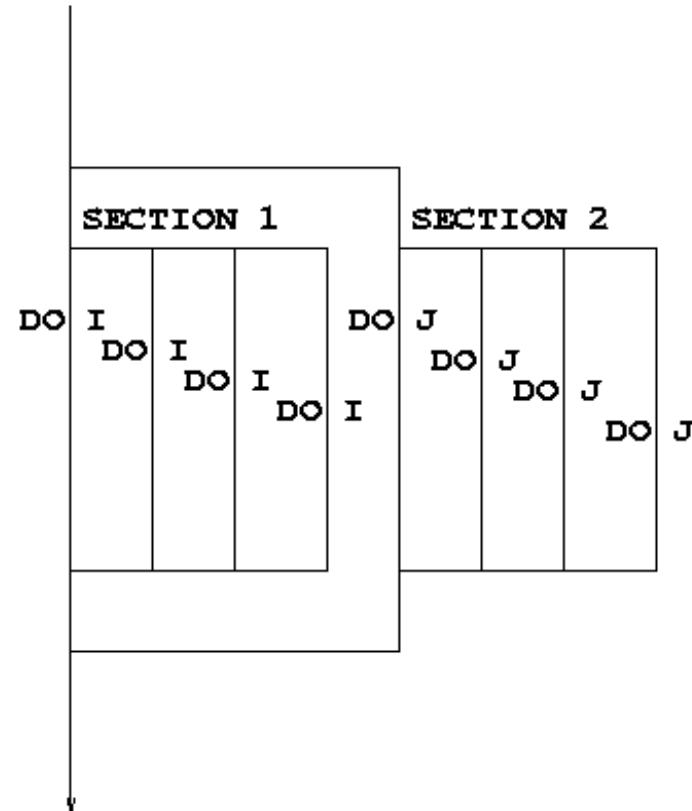
- Unlike most previous directive systems, nested parallelism is permitted in OpenMP.
- This is enabled with the **OMP\_NESTED** environment variable or the **OMP\_SET\_NESTED** routine.
- If a **PARALLEL** directive is encountered within another **PARALLEL** directive, a new team of threads will be created.
- The new team will contain only one thread unless nested parallelism is enabled.



# Nested parallelism (cont)

Example:

```
!$OMP PARALLEL
!$OMP SECTIONS
!$OMP SECTION
!$OMP PARALLEL DO
  do i = 1,n
    x(i) = 1.0
  end do
!$OMP SECTION
!$OMP PARALLEL DO
  do j = 1,n
    y(j) = 2.0
  end do
!$OMP END SECTIONS
!$OMP END PARALLEL
```



# Nested parallelism (cont)

- Not often needed, but can be useful to exploit non-scalable parallelism (SECTIONS).
- Note: nested parallelism isn't supported in some implementations (the code will execute, but as if OMP\_NESTED is set to FALSE).
  - turns out to be hard to do correctly without impacting performance significantly.



# NUMTHREADS clause

- One way to control the number of threads used at each level is with the NUM\_THREADS clause:

```
!$OMP PARALLEL DO NUM_THREADS (4)
    DO I = 1,4
!$OMP PARALLEL DO NUM_THREADS (TOTALTHREADS/4)
    DO J = 1,N
        A(I,J) = B(I,J)
    END DO
END DO
```

- The value set in the clause supersedes the value in the environment variable OMP\_NUM\_THREADS (or that set by `omp_set_num_threads()`)



# Orphaned directives

- Directives are active in the *dynamic* scope of a parallel region, not just its *lexical* scope.
- Example:

```
!$OMP PARALLEL
    call fred()
!$OMP END PARALLEL

subroutine fred()
!$OMP DO
    do i = 1,n
        a(i) = a(i) + 23.5
    end do
    return
end
```



# Orphaned directives (cont)

- This is very useful, as it allows a modular programming style....
- But it can also be rather confusing if the call tree is complicated (what happens if `fred` is also called from outside a parallel region?)
- There are some extra rules about data scope attributes....



# Data scoping rules

When we call a subroutine from inside a parallel region:

- Variables in the argument list inherit their data scope attribute from the calling routine.
- Global variables in C++ and COMMON blocks or module variables in Fortran are shared, unless declared THREADPRIVATE (see later).
- **static** local variables in C/C++ and **SAVE** variables in Fortran are shared.
- All other local variables are private.





# Binding rules

- There could be ambiguity about which parallel region directives refer to, so we need a rule....
- DO/FOR, SECTIONS, SINGLE, MASTER and BARRIER directives always bind to the nearest enclosing PARALLEL directive.



# Thread private global variables

- It can be convenient for each thread to have its own copy of variables with global scope (e.g. COMMON blocks and module data in Fortran, or file-scope and namespace-scope variables in C/C++).
- Outside parallel regions and in MASTER directives, accesses to these variables refer to the master thread's copy.



# Thread private globals (cont)

Syntax:

Fortran: **!\$OMP THREADPRIVATE** (*list*)

where *list* contains named common blocks (enclosed in slashes), module variables and SAVEd variables..

This directive must come after all the declarations for the common blocks or variables.

C/C++: **#pragma omp threadprivate** (*list*)

This directive must be at file or namespace scope, after all declarations of variables in *list* and before any references to variables in *list*. See standard document for other restrictions.



# COPYIN clause

- Allows the values of the master thread's THREADPRIVATE data to be copied to all other threads at the start of a parallel region.

Syntax:

Fortran: **COPYIN** (*list*)

C/C++: **copyin** (*list*)

In Fortran the list can contain variables in THREADPRIVATE COMMON blocks.



# COPYIN clause

Example:

```
common /junk/ nx
common /stuff/ a,b,c
!$OMP THREADPRIVATE (/JUNK/,/STUFF/)
nx = 32
c = 17.9
. . .
!$OMP PARALLEL PRIVATE(NX2,CSQ) COPYIN(/JUNK/,C)
nx2 = nx * 2
csq = c*c
. . .
```



# if clause

- Can add `if` clause to:
  - `parallel`
  - `for/do`
  - `sections`
- `if` clause takes scalar expression (C/C++) or scalar logical expression (Fortran)
  - `if(i)`
  - `if(i<100)`
  - `logical :: mylogical`  
...  
`if(mylogical)`



# if clause

```
!$OMP PARALLEL shared(b,n) private(i) if(n>100)
!$OMP DO
do i=1,n
    b(i) = b(i) * 2
end do
if(omp_in_parallel()) then
    write(*,*) `done the work in parallel`
else
    write(*,*) `done the work in serial`
end if
!$OMP END PARALLEL
```



# Timing routines

OpenMP supports a portable timer:

- return current wall clock time (relative to arbitrary origin) with:

```
DOUBLE PRECISION FUNCTION OMP_GET_WTIME ()
```

```
double omp_get_wtime(void);
```

- return clock precision with

```
DOUBLE PRECISION FUNCTION OMP_GET_WTICK ()
```

```
double omp_get_wtick(void);
```





# Using timers

```
DOUBLE PRECISION STARTTIME, TIME
```

```
STARTTIME = OMP_GET_WTIME()
```

```
.....(work to be timed)
```

```
TIME = OMP_GET_WTIME() - STARTTIME
```

Note: timers are local to a thread: must make both calls on the same thread.

Also note: no guarantees about resolution!



# Exercise

## Molecular dynamics again

- Aim: use of orphaned directives.
- Modify the molecular dynamics code so by placing a parallel region directive around the iteration loop in the main program, and making all code within this sequential except for the forces loop.
- Modify the code further so that each thread accumulates the forces into a local copy of the force array, and reduce these copies into the main array at the end of the loop.

