

# OpenMP Application Program Interface Examples

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<b>Introduction</b>	<b>5</b>
<b>Examples</b>	<b>7</b>
1 A Simple Parallel Loop	7
2 The OpenMP Memory Model	9
3 Conditional Compilation	15
4 Internal Control Variables (ICVs)	16
5 The <b>parallel</b> Construct	19
6 Controlling the Number of Threads on Multiple Nesting Levels	21
7 Interaction Between the <b>num_threads</b> Clause and <b>omp_set_dynamic</b>	24
8 Fortran Restrictions on the <b>do</b> Construct	26
9 Fortran Private Loop Iteration Variables	28
10 The <b>nowait</b> clause	30
11 The <b>collapse</b> clause	33
12 The <b>parallel sections</b> Construct	38
13 The <b>firstprivate</b> Clause and the <b>sections</b> Construct	39
14 The <b>single</b> Construct	41
15 Tasking Constructs	43
16 The <b>taskyield</b> Directive	67
17 The <b>workshare</b> Construct	69
18 The <b>master</b> Construct	73
19 The <b>critical</b> Construct	75
20 worksharing Constructs Inside a <b>critical</b> Construct	77
21 Binding of <b>barrier</b> Regions	79
22 The <b>atomic</b> Construct	82
23 Restrictions on the <b>atomic</b> Construct	88

24	The <b>flush</b> Construct without a List .....	92
25	Placement of <b>flush</b> , <b>barrier</b> , <b>taskwait</b> and <b>taskyield</b> Directives .....	95
26	The <b>ordered</b> Clause and the <b>ordered</b> Construct .....	99
27	Cancellation Constructs .....	103
28	The <b>threadprivate</b> Directive .....	108
29	Parallel Random Access Iterator Loop .....	114
30	Fortran Restrictions on <b>shared</b> and <b>private</b> Clauses with Common Blocks .....	115
31	The <b>default(none)</b> Clause .....	117
32	Race Conditions Caused by Implied Copies of Shared Variables in Fortran .....	119
33	The <b>private</b> Clause .....	120
34	Fortran Restrictions on Storage Association with the <b>private</b> Clause .....	124
35	C/C++ Arrays in a <b>firstprivate</b> Clause .....	127
36	The <b>lastprivate</b> Clause .....	129
37	The <b>reduction</b> Clause .....	130
38	The <b>copyin</b> Clause .....	136
39	The <b>copyprivate</b> Clause .....	138
40	Nested Loop Constructs .....	142
41	Restrictions on Nesting of Regions .....	146
42	The <b>omp_set_dynamic</b> and <b>omp_set_num_threads</b> Routines ..	152
43	The <b>omp_get_num_threads</b> Routine .....	154
44	The <b>omp_init_lock</b> Routine .....	156
45	Ownership of Locks .....	157
46	Simple Lock Routines .....	159
47	Nestable Lock Routines .....	161
48	<b>target</b> Construct .....	164
49	<b>target data</b> Construct .....	171
50	<b>target update</b> Construct .....	182

51	<b>declare target</b> Construct	186
52	<b>teams</b> Constructs	194
53	Asynchronous Execution of a <b>target</b> Region Using Tasks	202
54	Array Sections in Device Constructs	206
55	Device Routines	210
56	Fortran ASSOCIATE Construct	214



# Introduction

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This collection of programming examples supplements the OpenMP API for Shared Memory Parallelization specifications, and is not part of the formal specifications. It assumes familiarity with the OpenMP specifications, and shares the typographical conventions used in that document.

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**Note** – This first release of the OpenMP Examples reflects the OpenMP Version 4.0 specifications. Additional examples are being developed and will be published in future releases of this document.

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The OpenMP API specification provides a model for parallel programming that is portable across shared memory architectures from different vendors. Compilers from numerous vendors support the OpenMP API.

The directives, library routines, and environment variables demonstrated in this document allow users to create and manage parallel programs while permitting portability. The directives extend the C, C++ and Fortran base languages with single program multiple data (SPMD) constructs, tasking constructs, device constructs, worksharing constructs, and synchronization constructs, and they provide support for sharing and privatizing data. The functionality to control the runtime environment is provided by library routines and environment variables. Compilers that support the OpenMP API often include a command line option to the compiler that activates and allows interpretation of all OpenMP directives.

Complete information about the OpenMP API and a list of the compilers that support the OpenMP API can be found at the OpenMP.org web site

**`http://www.openmp.org`**

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# Examples

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The following are examples of the OpenMP API directives, constructs, and routines.

C/C++

A statement following a directive is compound only when necessary, and a non-compound statement is indented with respect to a directive preceding it.

C/C++

---

## 1

### A Simple Parallel Loop

The following example demonstrates how to parallelize a simple loop using the parallel loop construct . The loop iteration variable is private by default, so it is not necessary to specify it explicitly in a **private** clause.

C/C++

*Example 1.1c*

```
void simple(int n, float *a, float *b)
{
    int i;

    #pragma omp parallel for
        for (i=1; i<n; i++) /* i is private by default */
            b[i] = (a[i] + a[i-1]) / 2.0;
}
```

C/C++

*Example 1.1f*

```
SUBROUTINE SIMPLE(N, A, B)

  INTEGER I, N
  REAL B(N), A(N)

  !$OMP PARALLEL DO !I is private by default
    DO I=2,N
      B(I) = (A(I) + A(I-1)) / 2.0
    ENDDO
  !$OMP END PARALLEL DO

  END SUBROUTINE SIMPLE
```

## 2

# The OpenMP Memory Model

In the following example, at Print 1, the value of  $x$  could be either 2 or 5, depending on the timing of the threads, and the implementation of the assignment to  $x$ . There are two reasons that the value at Print 1 might not be 5. First, Print 1 might be executed before the assignment to  $x$  is executed. Second, even if Print 1 is executed after the assignment, the value 5 is not guaranteed to be seen by thread 1 because a flush may not have been executed by thread 0 since the assignment.

The barrier after Print 1 contains implicit flushes on all threads, as well as a thread synchronization, so the programmer is guaranteed that the value 5 will be printed by both Print 2 and Print 3.

C/C++

### *Example 2.1c*

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

int main(){
    int x;

    x = 2;
    #pragma omp parallel num_threads(2) shared(x)
    {

        if (omp_get_thread_num() == 0) {
            x = 5;
        } else {
            /* Print 1: the following read of x has a race */
            printf("1: Thread# %d: x = %d\n", omp_get_thread_num(), x );
        }

        #pragma omp barrier

        if (omp_get_thread_num() == 0) {
            /* Print 2 */
            printf("2: Thread# %d: x = %d\n", omp_get_thread_num(), x );
        } else {
            /* Print 3 */
            printf("3: Thread# %d: x = %d\n", omp_get_thread_num(), x );
        }
    }
    return 0;
}
```

C/C++

*Example 2.1f*

```

PROGRAM MEMMODEL
  INCLUDE "omp_lib.h"      ! or USE OMP_LIB
  INTEGER X

  X = 2
!$OMP PARALLEL NUM_THREADS(2) SHARED(X)

  IF (OMP_GET_THREAD_NUM() .EQ. 0) THEN
    X = 5
  ELSE
    ! PRINT 1: The following read of x has a race
    PRINT *, "1: THREAD# ", OMP_GET_THREAD_NUM(), "X = ", X
  ENDIF

!$OMP BARRIER

  IF (OMP_GET_THREAD_NUM() .EQ. 0) THEN
    ! PRINT 2
    PRINT *, "2: THREAD# ", OMP_GET_THREAD_NUM(), "X = ", X
  ELSE
    ! PRINT 3
    PRINT *, "3: THREAD# ", OMP_GET_THREAD_NUM(), "X = ", X
  ENDIF

!$OMP END PARALLEL

END PROGRAM MEMMODEL

```

The following example demonstrates why synchronization is difficult to perform correctly through variables. The value of flag is undefined in both prints on thread 1 and the value of data is only well-defined in the second print.

*Example 2.2c*

```

#include <omp.h>
#include <stdio.h>
int main()
{
    int data;
    int flag=0;
    #pragma omp parallel num_threads(2)
    {
        if (omp_get_thread_num()==0)
        {
            /* Write to the data buffer that will be
            read by thread */
            data = 42;
            /* Flush data to thread 1 and strictly order
            the write to data
            relative to the write to the flag */
            #pragma omp flush(flag, data)
            /* Set flag to release thread 1 */
            flag = 1;
            /* Flush flag to ensure that thread 1 sees
            the change */
            #pragma omp flush(flag)
        }
        else if(omp_get_thread_num()==1)
        {
            /* Loop until we see the update to the flag */
            #pragma omp flush(flag, data)
            while (flag < 1)
            {
                #pragma omp flush(flag, data)
            }
            /* Values of flag and data are undefined */
            printf("flag=%d data=%d\n", flag, data);
            #pragma omp flush(flag, data)
            /* Values data will be 42, value of flag
            still undefined */
            printf("flag=%d data=%d\n", flag, data);
        }
    }
    return 0;
}

```

*Example 2.2f*

```

PROGRAM EXAMPLE
INCLUDE "omp_lib.h" ! or USE OMP_LIB
INTEGER DATA
INTEGER FLAG

FLAG = 0
!$OMP PARALLEL NUM_THREADS(2)
  IF(OMP_GET_THREAD_NUM() .EQ. 0) THEN
    ! Write to the data buffer that will be read by thread 1
    DATA = 42
    ! Flush DATA to thread 1 and strictly order the write to DATA
    ! relative to the write to the FLAG
    !$OMP FLUSH(FLAG, DATA)
    ! Set FLAG to release thread 1
    FLAG = 1;
    ! Flush FLAG to ensure that thread 1 sees the change */
    !$OMP FLUSH(FLAG)
  ELSE IF(OMP_GET_THREAD_NUM() .EQ. 1) THEN
    ! Loop until we see the update to the FLAG
    !$OMP FLUSH(FLAG, DATA)
    DO WHILE(FLAG .LT. 1)
      !$OMP FLUSH(FLAG, DATA)
    ENDDO

    ! Values of FLAG and DATA are undefined
    PRINT *, 'FLAG=', FLAG, ' DATA=', DATA
    !$OMP FLUSH(FLAG, DATA)

    !Values DATA will be 42, value of FLAG still undefined */
    PRINT *, 'FLAG=', FLAG, ' DATA=', DATA
  ENDIF
!$OMP END PARALLEL
END

```

The next example demonstrates why synchronization is difficult to perform correctly through variables. Because the *write(1)-flush(1)-flush(2)-read(2)* sequence cannot be guaranteed in the example, the statements on thread 0 and thread 1 may execute in either order.

*Example 2.3c*

```

#include <omp.h>
#include <stdio.h>
int main()
{
    int flag=0;

    #pragma omp parallel num_threads(3)
    {
        if(omp_get_thread_num()==0)
        {
            /* Set flag to release thread 1 */
            #pragma omp atomic update
            flag++;
            /* Flush of flag is implied by the atomic directive */
        }
        else if(omp_get_thread_num()==1)
        {
            /* Loop until we see that flag reaches 1*/
            #pragma omp flush(flag)
            while(flag < 1)
            {
                #pragma omp flush(flag)
            }
            printf("Thread 1 awoken\n");

            /* Set flag to release thread 2 */
            #pragma omp atomic update
            flag++;
            /* Flush of flag is implied by the atomic directive */
        }
        else if(omp_get_thread_num()==2)
        {
            /* Loop until we see that flag reaches 2 */
            #pragma omp flush(flag)
            while(flag < 2)
            {
                #pragma omp flush(flag)
            }
            printf("Thread 2 awoken\n");
        }
    }
    return 0;
}

```

*Example 2.3f*

```

PROGRAM EXAMPLE
INCLUDE "omp_lib.h" ! or USE OMP_LIB
INTEGER FLAG

FLAG = 0
!$OMP PARALLEL NUM_THREADS(3)
  IF(OMP_GET_THREAD_NUM() .EQ. 0) THEN
    ! Set flag to release thread 1
    !$OMP ATOMIC UPDATE
    FLAG = FLAG + 1
    !Flush of FLAG is implied by the atomic directive
  ELSE IF(OMP_GET_THREAD_NUM() .EQ. 1) THEN
    ! Loop until we see that FLAG reaches 1
    !$OMP FLUSH(FLAG, DATA)
    DO WHILE(FLAG .LT. 1)
      !$OMP FLUSH(FLAG, DATA)
    ENDDO

    PRINT *, 'Thread 1 awoken'

    ! Set FLAG to release thread 2
    !$OMP ATOMIC UPDATE
    FLAG = FLAG + 1
    !Flush of FLAG is implied by the atomic directive
  ELSE IF(OMP_GET_THREAD_NUM() .EQ. 2) THEN
    ! Loop until we see that FLAG reaches 2
    !$OMP FLUSH(FLAG, DATA)
    DO WHILE(FLAG .LT. 2)
      !$OMP FLUSH(FLAG, DATA)
    ENDDO

    PRINT *, 'Thread 2 awoken'

  ENDIF
!$OMP END PARALLEL
END

```



## 3

# Conditional Compilation

### C/C++

The following example illustrates the use of conditional compilation using the OpenMP macro `_OPENMP`. With OpenMP compilation, the `_OPENMP` macro becomes defined.

#### *Example 3.1c*

```
#include <stdio.h>

int main()
{
    # ifdef _OPENMP
        printf("Compiled by an OpenMP-compliant implementation.\n");
    # endif

    return 0;
}
```

### C/C++

### Fortran

The following example illustrates the use of the conditional compilation sentinel. With OpenMP compilation, the conditional compilation sentinel `!$` is recognized and treated as two spaces. In fixed form source, statements guarded by the sentinel must start after column 6.

#### *Example 3.1f*

```
PROGRAM EXAMPLE

C234567890
!$   PRINT *, "Compiled by an OpenMP-compliant implementation."

END PROGRAM EXAMPLE
```

### Fortran

---

## 4 Internal Control Variables (ICVs)

According to §, an OpenMP implementation must act as if there are ICVs that control the behavior of the program. This example illustrates two ICVs, *nthreads-var* and *max-active-levels-var*. The *nthreads-var* ICV controls the number of threads requested for encountered parallel regions; there is one copy of this ICV per task. The *max-active-levels-var* ICV controls the maximum number of nested active parallel regions; there is one copy of this ICV for the whole program.

In the following example, the *nest-var*, *max-active-levels-var*, *dyn-var*, and *nthreads-var* ICVs are modified through calls to the runtime library routines `omp_set_nested`, `omp_set_max_active_levels`, `omp_set_dynamic`, and `omp_set_num_threads` respectively. These ICVs affect the operation of **parallel** regions. Each implicit task generated by a **parallel** region has its own copy of the *nest-var*, *dyn-var*, and *nthreads-var* ICVs.

In the following example, the new value of *nthreads-var* applies only to the implicit tasks that execute the call to `omp_set_num_threads`. There is one copy of the *max-active-levels-var* ICV for the whole program and its value is the same for all tasks. This example assumes that nested parallelism is supported.

The outer **parallel** region creates a team of two threads; each of the threads will execute one of the two implicit tasks generated by the outer **parallel** region.

Each implicit task generated by the outer **parallel** region calls `omp_set_num_threads(3)`, assigning the value 3 to its respective copy of *nthreads-var*. Then each implicit task encounters an inner **parallel** region that creates a team of three threads; each of the threads will execute one of the three implicit tasks generated by that inner **parallel** region.

Since the outer **parallel** region is executed by 2 threads, and the inner by 3, there will be a total of 6 implicit tasks generated by the two inner **parallel** regions.

Each implicit task generated by an inner **parallel** region will execute the call to `omp_set_num_threads(4)`, assigning the value 4 to its respective copy of *nthreads-var*.

The print statement in the outer **parallel** region is executed by only one of the threads in the team. So it will be executed only once.

The print statement in an inner **parallel** region is also executed by only one of the threads in the team. Since we have a total of two inner **parallel** regions, the print statement will be executed twice -- once per inner **parallel** region.

*Example 4.1c*

```

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

int main (void)
{
    omp_set_nested(1);
    omp_set_max_active_levels(8);
    omp_set_dynamic(0);
    omp_set_num_threads(2);
    #pragma omp parallel
    {
        omp_set_num_threads(3);

        #pragma omp parallel
        {
            omp_set_num_threads(4);
            #pragma omp single
            {
                /*
                 * The following should print:
                 * Inner: max_act_lev=8, num_thds=3, max_thds=4
                 * Inner: max_act_lev=8, num_thds=3, max_thds=4
                 */
                printf ("Inner: max_act_lev=%d, num_thds=%d, max_thds=%d\n",
                    omp_get_max_active_levels(), omp_get_num_threads(),
                    omp_get_max_threads());
            }
        }

        #pragma omp barrier
        #pragma omp single
        {
            /*
             * The following should print:
             * Outer: max_act_lev=8, num_thds=2, max_thds=3
             */
            printf ("Outer: max_act_lev=%d, num_thds=%d, max_thds=%d\n",
                omp_get_max_active_levels(), omp_get_num_threads(),
                omp_get_max_threads());
        }
    }
    return 0;
}

```

*Example 4.1f*

```

program icv
use omp_lib

call omp_set_nested(.true.)
call omp_set_max_active_levels(8)
call omp_set_dynamic(.false.)
call omp_set_num_threads(2)

!$omp parallel
  call omp_set_num_threads(3)

!$omp parallel
  call omp_set_num_threads(4)
!$omp single
!   The following should print:
!   Inner: max_act_lev= 8 , num_thds= 3 , max_thds= 4
!   Inner: max_act_lev= 8 , num_thds= 3 , max_thds= 4
  print *, "Inner: max_act_lev=", omp_get_max_active_levels(),
    &      " , num_thds=", omp_get_num_threads(),
    &      " , max_thds=", omp_get_max_threads()
!$omp end single
!$omp end parallel

!$omp barrier
!$omp single
!   The following should print:
!   Outer: max_act_lev= 8 , num_thds= 2 , max_thds= 3
  print *, "Outer: max_act_lev=", omp_get_max_active_levels(),
    &      " , num_thds=", omp_get_num_threads(),
    &      " , max_thds=", omp_get_max_threads()
!$omp end single
!$omp end parallel
end

```

## 5

# The parallel Construct

The `parallel` construct can be used in coarse-grain parallel programs. In the following example, each thread in the `parallel` region decides what part of the global array `x` to work on, based on the thread number:

C/C++

*Example 5.1c*

```
#include <omp.h>

void subdomain(float *x, int istart, int ipoints)
{
    int i;

    for (i = 0; i < ipoints; i++)
        x[istart+i] = 123.456;
}

void sub(float *x, int npoints)
{
    int iam, nt, ipoints, istart;

#pragma omp parallel default(shared) private(iam,nt,ipoints,istart)
    {
        iam = omp_get_thread_num();
        nt = omp_get_num_threads();
        ipoints = npoints / nt; /* size of partition */
        istart = iam * ipoints; /* starting array index */
        if (iam == nt-1) /* last thread may do more */
            ipoints = npoints - istart;
        subdomain(x, istart, ipoints);
    }
}

int main()
{
    float array[10000];

    sub(array, 10000);

    return 0;
}
```

C/C++

*Example 5.1f*

```

SUBROUTINE SUBDOMAIN(X, ISTART, IPOINTS)
  INTEGER ISTART, IPOINTS
  REAL X(*)

  INTEGER I

  DO 100 I=1,IPOINTS
    X(ISTART+I) = 123.456
100  CONTINUE

END SUBROUTINE SUBDOMAIN

SUBROUTINE SUB(X, NPOINTS)
  INCLUDE "omp_lib.h"      ! or USE OMP_LIB

  REAL X(*)
  INTEGER NPOINTS
  INTEGER IAM, NT, IPOINTS, ISTART

!$OMP PARALLEL DEFAULT(PRIVATE) SHARED(X,NPOINTS)

  IAM = OMP_GET_THREAD_NUM()
  NT = OMP_GET_NUM_THREADS()
  IPOINTS = NPOINTS/NT
  ISTART = IAM * IPOINTS
  IF (IAM .EQ. NT-1) THEN
    IPOINTS = NPOINTS - ISTART
  ENDIF
  CALL SUBDOMAIN(X,ISTART,IPOINTS)

!$OMP END PARALLEL
END SUBROUTINE SUB

PROGRAM PAREXAMPLE
  REAL ARRAY(10000)
  CALL SUB(ARRAY, 10000)
END PROGRAM PAREXAMPLE

```

## 6

# Controlling the Number of Threads on Multiple Nesting Levels

The following examples demonstrate how to use the `OMP_NUM_THREADS` environment variable to control the number of threads on multiple nesting levels:

C/C++

### *Example 6.1c*

```
#include <stdio.h>
#include <omp.h>
int main (void)
{
    omp_set_nested(1);
    omp_set_dynamic(0);
    #pragma omp parallel
    {
        #pragma omp parallel
        {
            #pragma omp single
            {
                /*
                 * If OMP_NUM_THREADS=2,3 was set, the following should print:
                 * Inner: num_thds=3
                 * Inner: num_thds=3
                 */
                /*
                 * If nesting is not supported, the following should print:
                 * Inner: num_thds=1
                 * Inner: num_thds=1
                 */
                printf ("Inner: num_thds=%d\n", omp_get_num_threads());
            }
        }
        #pragma omp barrier
        omp_set_nested(0);
        #pragma omp parallel
        {
            #pragma omp single
            {
                /*
                 * Even if OMP_NUM_THREADS=2,3 was set, the following should
                 * print, because nesting is disabled:
                 * Inner: num_thds=1
                 * Inner: num_thds=1
                 */
                printf ("Inner: num_thds=%d\n", omp_get_num_threads());
            }
        }
    }
}
```

```

        #pragma omp barrier
        #pragma omp single
        {
            /*
             * If OMP_NUM_THREADS=2,3 was set, the following should print:
             * Outer: num_thds=2
             */
            printf ("Outer: num_thds=%d\n", omp_get_num_threads());
        }
    }
    return 0;
}

```

C/C++

Fortran

### *Example 6.1f*

```

program icv
    use omp_lib
    call omp_set_nested(.true.)
    call omp_set_dynamic(.false.)
!$omp parallel
!$omp parallel
!$omp single
    ! If OMP_NUM_THREADS=2,3 was set, the following should print:
    ! Inner: num_thds= 3
    ! Inner: num_thds= 3
    ! If nesting is not supported, the following should print:
    ! Inner: num_thds= 1
    ! Inner: num_thds= 1
    print *, "Inner: num_thds=", omp_get_num_threads()
!$omp end single
!$omp end parallel
!$omp barrier
    call omp_set_nested(.false.)
!$omp parallel
!$omp single
    ! Even if OMP_NUM_THREADS=2,3 was set, the following should print,
    ! because nesting is disabled:
    ! Inner: num_thds= 1
    ! Inner: num_thds= 1
    print *, "Inner: num_thds=", omp_get_num_threads()
!$omp end single
!$omp end parallel
!$omp barrier
!$omp single
    ! If OMP_NUM_THREADS=2,3 was set, the following should print:
    ! Outer: num_thds= 2
    print *, "Outer: num_thds=", omp_get_num_threads()
!$omp end single
!$omp end parallel

```



end

Fortran

## 7 Interaction Between the `num_threads` Clause and `omp_set_dynamic`

The following example demonstrates the `num_threads` clause and the effect of the `omp_set_dynamic` routine on it.

The call to the `omp_set_dynamic` routine with argument `0` in C/C++, or `.FALSE.` in Fortran, disables the dynamic adjustment of the number of threads in OpenMP implementations that support it. In this case, 10 threads are provided. Note that in case of an error the OpenMP implementation is free to abort the program or to supply any number of threads available.

Example 7.1c

```
#include <omp.h>
int main()
{
    omp_set_dynamic(0);
    #pragma omp parallel num_threads(10)
    {
        /* do work here */
    }
    return 0;
}
```

C/C++

Example 7.1f

```
PROGRAM EXAMPLE
    INCLUDE "omp_lib.h"          ! or USE OMP_LIB
    CALL OMP_SET_DYNAMIC(.FALSE.)
!$OMP    PARALLEL NUM_THREADS(10)
        ! do work here
!$OMP    END PARALLEL
END PROGRAM EXAMPLE
```

Fortran

The call to the `omp_set_dynamic` routine with a non-zero argument in C/C++, or `.TRUE.` in Fortran, allows the OpenMP implementation to choose any number of threads between 1 and 10.

C/C++

*Example 7.2c*

```
#include <omp.h>
int main()
{
    omp_set_dynamic(1);
    #pragma omp parallel num_threads(10)
    {
        /* do work here */
    }
    return 0;
}
```

C/C++

Fortran

*Example 7.2f*

```
PROGRAM EXAMPLE
    INCLUDE "omp_lib.h"          ! or USE OMP_LIB
    CALL OMP_SET_DYNAMIC(.TRUE.)
!$OMP    PARALLEL NUM_THREADS(10)
        ! do work here
!$OMP    END PARALLEL
END PROGRAM EXAMPLE
```

Fortran

It is good practice to set the *dyn-var* ICV explicitly by calling the `omp_set_dynamic` routine, as its default setting is implementation defined.

## 8 Fortran Restrictions on the `do` Construct

If an `end do` directive follows a *do-construct* in which several `DO` statements share a `DO` termination statement, then a `do` directive can only be specified for the outermost of these `DO` statements. The following example contains correct usages of loop constructs:

*Example 8.1f*

```

SUBROUTINE WORK(I, J)
  INTEGER I,J
  END SUBROUTINE WORK

SUBROUTINE DO_GOOD()
  INTEGER I, J
  REAL A(1000)

  DO 100 I = 1,10
!$OMP DO
    DO 100 J = 1,10
      CALL WORK(I,J)
100  CONTINUE      ! !$OMP ENDDO implied here

!$OMP DO
  DO 200 J = 1,10
200  A(I) = I + 1
!$OMP ENDDO

!$OMP DO
  DO 300 I = 1,10
    DO 300 J = 1,10
      CALL WORK(I,J)
300  CONTINUE
!$OMP ENDDO
  END SUBROUTINE DO_GOOD

```

The following example is non-conforming because the matching `do` directive for the `end do` does not precede the outermost loop:

*Example 8.2f*

```

SUBROUTINE WORK(I, J)
  INTEGER I,J
  END SUBROUTINE WORK

SUBROUTINE DO_WRONG
  INTEGER I, J

```

```
      DO 100 I = 1,10
!$OMP      DO
      DO 100 J = 1,10
      CALL WORK(I,J)
100      CONTINUE
!$OMP      ENDDO
      END SUBROUTINE DO_WRONG
```

Fortran

## 9 Fortran Private Loop Iteration Variables

In general loop iteration variables will be private, when used in the *do-loop* of a **do** and **parallel do** construct or in sequential loops in a **parallel** construct (see § and §). In the following example of a sequential loop in a **parallel** construct the loop iteration variable *I* will be private.

### *Example 9.1f*

```
SUBROUTINE PLOOP_1(A,N)
  INCLUDE "omp_lib.h"      ! or USE OMP_LIB

  REAL A(*)
  INTEGER I, MYOFFSET, N

  !$OMP PARALLEL PRIVATE(MYOFFSET)
    MYOFFSET = OMP_GET_THREAD_NUM()*N
    DO I = 1, N
      A(MYOFFSET+I) = FLOAT(I)
    ENDDO
  !$OMP END PARALLEL

END SUBROUTINE PLOOP_1
```

In exceptional cases, loop iteration variables can be made shared, as in the following example:

### *Example 9.2f*

```
SUBROUTINE PLOOP_2(A,B,N,I1,I2)
  REAL A(*), B(*)
  INTEGER I1, I2, N

  !$OMP PARALLEL SHARED(A,B,I1,I2)
  !$OMP SECTIONS
  !$OMP SECTION
    DO I1 = 1, N
      IF (A(I1).NE.0.0) EXIT
    ENDDO
  !$OMP SECTION
    DO I2 = 1, N
      IF (B(I2).NE.0.0) EXIT
    ENDDO
  !$OMP END SECTIONS
  !$OMP SINGLE
    IF (I1.LE.N) PRINT *, 'ITEMS IN A UP TO ', I1, 'ARE ALL ZERO.'
    IF (I2.LE.N) PRINT *, 'ITEMS IN B UP TO ', I2, 'ARE ALL ZERO.'
  !$OMP END SINGLE
```

```
!$OMP END PARALLEL
```

```
END SUBROUTINE PLOOP_2
```

Note however that the use of shared loop iteration variables can easily lead to race conditions.

---

Fortran

## 10 The `nowait` clause

If there are multiple independent loops within a `parallel` region, you can use the `nowait` clause to avoid the implied barrier at the end of the loop construct, as follows:

▼ C/C++ ▼  
*Example 10.1c*

```
#include <math.h>

void nowait_example(int n, int m, float *a, float *b, float *y, float *z)
{
    int i;
    #pragma omp parallel
    {
        #pragma omp for nowait
        for (i=1; i<n; i++)
            b[i] = (a[i] + a[i-1]) / 2.0;

        #pragma omp for nowait
        for (i=0; i<m; i++)
            y[i] = sqrt(z[i]);
    }
}
```

▲ C/C++ ▲

▼ Fortran ▼  
*Example 10.1f*

```
SUBROUTINE NOWAIT_EXAMPLE(N, M, A, B, Y, Z)

    INTEGER N, M
    REAL A(*), B(*), Y(*), Z(*)

    INTEGER I

    !$OMP PARALLEL

    !$OMP DO
        DO I=2,N
            B(I) = (A(I) + A(I-1)) / 2.0
        ENDDO
    !$OMP END DO NOWAIT

    !$OMP DO
        DO I=1,M
```



```

        Y(I) = SQRT(Z(I))
      ENDDO
!$OMP END DO NOWAIT

!$OMP END PARALLEL

END SUBROUTINE NOWAIT_EXAMPLE

```

## Fortran

In the following example, static scheduling distributes the same logical iteration numbers to the threads that execute the three loop regions. This allows the **nowait** clause to be used, even though there is a data dependence between the loops. The dependence is satisfied as long the same thread executes the same logical iteration numbers in each loop.

Note that the iteration count of the loops must be the same. The example satisfies this requirement, since the iteration space of the first two loops is from 0 to  $n-1$  (from 1 to  $N$  in the Fortran version), while the iteration space of the last loop is from 1 to  $n$  (2 to  $N+1$  in the Fortran version).

## C/C++

### Example 10.2c

```

#include <math.h>
void nowait_example2(int n, float *a, float *b, float *c, float *y, float *z)
{
    int i;
    #pragma omp parallel
    {
        #pragma omp for schedule(static) nowait
        for (i=0; i<n; i++)
            c[i] = (a[i] + b[i]) / 2.0f;
        #pragma omp for schedule(static) nowait
        for (i=0; i<n; i++)
            z[i] = sqrtf(c[i]);
        #pragma omp for schedule(static) nowait
        for (i=1; i<=n; i++)
            y[i] = z[i-1] + a[i];
    }
}

```

## C/C++

## Fortran

### Example 10.2f

```

SUBROUTINE NOWAIT_EXAMPLE2(N, A, B, C, Y, Z)
  INTEGER N
  REAL A(*), B(*), C(*), Y(*), Z(*)
  INTEGER I
!$OMP PARALLEL
!$OMP DO SCHEDULE(STATIC)

```

```

      DO I=1,N
        C(I) = (A(I) + B(I)) / 2.0
      ENDDO
!$OMP END DO NOWAIT
!$OMP DO SCHEDULE(STATIC)
      DO I=1,N
        Z(I) = SQRT(C(I))
      ENDDO
!$OMP END DO NOWAIT
!$OMP DO SCHEDULE(STATIC)
      DO I=2,N+1
        Y(I) = Z(I-1) + A(I)
      ENDDO
!$OMP END DO NOWAIT
!$OMP END PARALLEL
      END SUBROUTINE NOWAIT_EXAMPLE2

```

Fortran

## 11

# The collapse clause

In the following example, the **k** and **j** loops are associated with the loop construct. So the iterations of the **k** and **j** loops are collapsed into one loop with a larger iteration space, and that loop is then divided among the threads in the current team. Since the **i** loop is not associated with the loop construct, it is not collapsed, and the **i** loop is executed sequentially in its entirety in every iteration of the collapsed **k** and **j** loop.

### C/C++

The variable **j** can be omitted from the **private** clause when the **collapse** clause is used since it is implicitly private. However, if the **collapse** clause is omitted then **j** will be shared if it is omitted from the **private** clause. In either case, **k** is implicitly private and could be omitted from the **private** clause.

#### Example 11.1c

```
void bar(float *a, int i, int j, int k);
int kl, ku, ks, jl, ju, js, il, iu, is;
void sub(float *a)
{
    int i, j, k;
    #pragma omp for collapse(2) private(i, k, j)
    for (k=kl; k<=ku; k+=ks)
        for (j=jl; j<=ju; j+=js)
            for (i=il; i<=iu; i+=is)
                bar(a,i,j,k);
}
```

### C/C++

### Fortran

#### Example 11.1f

```
subroutine sub(a)
real a(*)
integer kl, ku, ks, jl, ju, js, il, iu, is
common /csub/ kl, ku, ks, jl, ju, js, il, iu, is
integer i, j, k
!$omp do collapse(2) private(i,j,k)
  do k = kl, ku, ks
    do j = jl, ju, js
      do i = il, iu, is
        call bar(a,i,j,k)
      enddo
    enddo
  enddo
```

```
!$omp end do  
    end subroutine
```

## Fortran

In the next example, the **k** and **j** loops are associated with the loop construct. So the iterations of the **k** and **j** loops are collapsed into one loop with a larger iteration space, and that loop is then divided among the threads in the current team.

The sequential execution of the iterations in the **k** and **j** loops determines the order of the iterations in the collapsed iteration space. This implies that in the sequentially last iteration of the collapsed iteration space, **k** will have the value **2** and **j** will have the value **3**. Since **klast** and **jlast** are **lastprivate**, their values are assigned by the sequentially last iteration of the collapsed **k** and **j** loop. This example prints: **2 3**.

*Example 11.2c*

```

#include <stdio.h>
void test()
{
    int j, k, jlast, klast;
    #pragma omp parallel
    {
        #pragma omp for collapse(2) lastprivate(jlast, klast)
        for (k=1; k<=2; k++)
            for (j=1; j<=3; j++)
            {
                jlast=j;
                klast=k;
            }
        #pragma omp single
        printf("%d %d\n", klast, jlast);
    }
}

```

*Example 11.2f*

```

program test
!$omp parallel
!$omp do private(j,k) collapse(2) lastprivate(jlast, klast)
do k = 1,2
do j = 1,3
jlast=j
klast=k
enddo
enddo
!$omp end do
!$omp single
print *, klast, jlast
!$omp end single
!$omp end parallel
end program test

```

The next example illustrates the interaction of the **collapse** and **ordered** clauses.

In the example, the loop construct has both a **collapse** clause and an **ordered** clause. The **collapse** clause causes the iterations of the **k** and **j** loops to be collapsed into one loop with a larger iteration space, and that loop is divided among the threads in the current team. An **ordered** clause is added to the loop construct, because an ordered region binds to the loop region arising from the loop construct.

According to §, a thread must not execute more than one ordered region that binds to the same loop region. So the **collapse** clause is required for the example to be conforming. With the **collapse** clause, the iterations of the **k** and **j** loops are collapsed into one loop, and therefore only one ordered region will bind to the collapsed **k** and **j** loop. Without the **collapse** clause, there would be two ordered regions that bind to each iteration of the **k** loop (one arising from the first iteration of the **j** loop, and the other arising from the second iteration of the **j** loop).

The code prints

```
0 1 1
0 1 2
0 2 1
1 2 2
1 3 1
1 3 2
```

C/C++

### Example 11.3c

```
#include <omp.h>
#include <stdio.h>
void work(int a, int j, int k);
void sub()
{
    int j, k, a;
    #pragma omp parallel num_threads(2)
    {
        #pragma omp for collapse(2) ordered private(j,k) schedule(static,3)
        for (k=1; k<=3; k++)
            for (j=1; j<=2; j++)
            {
                #pragma omp ordered
                printf("%d %d %d\n", omp_get_thread_num(), k, j);
                /* end ordered */
                work(a,j,k);
            }
    }
}
```

C/C++

*Example 11.3f*

```
program test
  include 'omp_lib.h'
!$omp parallel num_threads(2)
!$omp do collapse(2) ordered private(j,k) schedule(static,3)
  do k = 1,3
    do j = 1,2
!$omp ordered
      print *, omp_get_thread_num(), k, j
!$omp end ordered
      call work(a,j,k)
    enddo
  enddo
!$omp end do
!$omp end parallel
end program test
```

## 12

# The parallel sections Construct

In the following example routines **XAXIS**, **YAXIS**, and **ZAXIS** can be executed concurrently. The first **section** directive is optional. Note that all **section** directives need to appear in the **parallel sections** construct.

▼ C/C++ ▼  
*Example 12.1c*

```
void XAXIS();
void YAXIS();
void ZAXIS();

void sect_example()
{
    #pragma omp parallel sections
    {
        #pragma omp section
            XAXIS();

        #pragma omp section
            YAXIS();

        #pragma omp section
            ZAXIS();
    }
}
```

▲ C/C++ ▲

▼ Fortran ▼  
*Example 12.1f*

```
      SUBROUTINE SECT_EXAMPLE()
!$OMP PARALLEL SECTIONS
!$OMP SECTION
      CALL XAXIS()
!$OMP SECTION
      CALL YAXIS()

!$OMP SECTION
      CALL ZAXIS()

!$OMP END PARALLEL SECTIONS
      END SUBROUTINE SECT_EXAMPLE
```

▲ Fortran ▲



## 13

# The `firstprivate` Clause and the `sections` Construct

In the following example of the `sections` construct the `firstprivate` clause is used to initialize the private copy of `section_count` of each thread. The problem is that the `section` constructs modify `section_count`, which breaks the independence of the `section` constructs. When different threads execute each section, both sections will print the value 1. When the same thread executes the two sections, one section will print the value 1 and the other will print the value 2. Since the order of execution of the two sections in this case is unspecified, it is unspecified which section prints which value.

C/C++

*Example 13.1c*

```
#include <omp.h>
#include <stdio.h>
#define NT 4
int main( ) {
    int section_count = 0;
    omp_set_dynamic(0);
    omp_set_num_threads(NT);
    #pragma omp parallel
    #pragma omp sections firstprivate( section_count )
    {
        #pragma omp section
        {
            section_count++;
            /* may print the number one or two */
            printf( "section_count %d\n", section_count );
        }
        #pragma omp section
        {
            section_count++;
            /* may print the number one or two */
            printf( "section_count %d\n", section_count );
        }
    }
    return 1;
}
```

C/C++

*Example 13.1f*

```
program section
  use omp_lib
  integer :: section_count = 0
  integer, parameter :: NT = 4
  call omp_set_dynamic(.false.)
  call omp_set_num_threads(NT)
!$omp parallel
!$omp sections firstprivate ( section_count )
!$omp section
  section_count = section_count + 1
! may print the number one or two
  print *, 'section_count', section_count
!$omp section
  section_count = section_count + 1
! may print the number one or two
  print *, 'section_count', section_count
!$omp end sections
!$omp end parallel
end program section
```

## 14 The `single` Construct

The following example demonstrates the `single` construct . In the example, only one thread prints each of the progress messages. All other threads will skip the `single` region and stop at the barrier at the end of the `single` construct until all threads in the team have reached the barrier. If other threads can proceed without waiting for the thread executing the `single` region, a `nowait` clause can be specified, as is done in the third `single` construct in this example. The user must not make any assumptions as to which thread will execute a `single` region.

C/C++

*Example 14.1c*

```
#include <stdio.h>

void work1() {}
void work2() {}

void single_example()
{
    #pragma omp parallel
    {
        #pragma omp single
        printf("Beginning work1.\n");

        work1();

        #pragma omp single
        printf("Finishing work1.\n");

        #pragma omp single nowait
        printf("Finished work1 and beginning work2.\n");

        work2();
    }
}
```

C/C++

Fortran

*Example 14.1f*

```
SUBROUTINE WORK1()
END SUBROUTINE WORK1

SUBROUTINE WORK2()
```

```

        END SUBROUTINE WORK2

        PROGRAM SINGLE_EXAMPLE
!$OMP PARALLEL

!$OMP SINGLE
        print *, "Beginning work1."
!$OMP END SINGLE

        CALL WORK1()

!$OMP SINGLE
        print *, "Finishing work1."
!$OMP END SINGLE

!$OMP SINGLE
        print *, "Finished work1 and beginning work2."
!$OMP END SINGLE NOWAIT

        CALL WORK2()

!$OMP END PARALLEL

        END PROGRAM SINGLE_EXAMPLE

```

Fortran

## 15 Tasking Constructs

The following example shows how to traverse a tree-like structure using explicit tasks. Note that the **traverse** function should be called from within a parallel region for the different specified tasks to be executed in parallel. Also note that the tasks will be executed in no specified order because there are no synchronization directives. Thus, assuming that the traversal will be done in post order, as in the sequential code, is wrong.

C/C++

*Example 15.1c*

```
struct node {
    struct node *left;
    struct node *right;
};
extern void process(struct node *);
void traverse( struct node *p ) {
    if (p->left)
#pragma omp task    // p is firstprivate by default
        traverse(p->left);
    if (p->right)
#pragma omp task    // p is firstprivate by default
        traverse(p->right);
    process(p);
}
```

C/C++

Fortran

*Example 15.1f*

```
RECURSIVE SUBROUTINE traverse ( P )
    TYPE Node
        TYPE(Node), POINTER :: left, right
    END TYPE Node
    TYPE(Node) :: P
    IF (associated(P%left)) THEN
        !$OMP TASK    ! P is firstprivate by default
        call traverse(P%left)
        !$OMP END TASK
    ENDIF
    IF (associated(P%right)) THEN
        !$OMP TASK    ! P is firstprivate by default
        call traverse(P%right)
        !$OMP END TASK
    ENDIF
END SUBROUTINE
```

```

ENDIF
CALL process ( P )
END SUBROUTINE

```

## Fortran

In the next example, we force a postorder traversal of the tree by adding a **taskwait** directive. Now, we can safely assume that the left and right sons have been executed before we process the current node.

## C/C++

### Example 15.2c

```

struct node {
    struct node *left;
    struct node *right;
};
extern void process(struct node *);
void postorder_traverse( struct node *p ) {
    if (p->left)
        #pragma omp task    // p is firstprivate by default
        postorder_traverse(p->left);
    if (p->right)
        #pragma omp task    // p is firstprivate by default
        postorder_traverse(p->right);
    #pragma omp taskwait
    process(p);
}

```

## C/C++

## Fortran

### Example 15.2f

```

RECURSIVE SUBROUTINE traverse ( P )
    TYPE Node
        TYPE(Node), POINTER :: left, right
    END TYPE Node
    TYPE(Node) :: P
    IF (associated(P%left)) THEN
        !$OMP TASK    ! P is firstprivate by default
        call traverse(P%left)
        !$OMP END TASK
    ENDIF
    IF (associated(P%right)) THEN
        !$OMP TASK    ! P is firstprivate by default
        call traverse(P%right)
        !$OMP END TASK
    ENDIF
    !$OMP TASKWAIT

```

```

        CALL process ( P )
    END SUBROUTINE

```

## Fortran

The following example demonstrates how to use the **task** construct to process elements of a linked list in parallel. The thread executing the **single** region generates all of the explicit tasks, which are then executed by the threads in the current team. The pointer *p* is **firstprivate** by default on the **task** construct so it is not necessary to specify it in a **firstprivate** clause.

## C/C++

### Example 15.3c

```

typedef struct node node;
struct node {
    int data;
    node * next;
};

void process(node * p)
{
    /* do work here */
}

void increment_list_items(node * head)
{
    #pragma omp parallel
    {
        #pragma omp single
        {
            node * p = head;
            while (p) {
                #pragma omp task
                // p is firstprivate by default
                process(p);
                p = p->next;
            }
        }
    }
}

```

## C/C++

## Fortran

### Example 15.3f

```

MODULE LIST
  TYPE NODE
    INTEGER :: PAYLOAD
    TYPE (NODE), POINTER :: NEXT

```

```

        END TYPE NODE
CONTAINS
    SUBROUTINE PROCESS(p)
        TYPE (NODE), POINTER :: P
        ! do work here
    END SUBROUTINE
    SUBROUTINE INCREMENT_LIST_ITEMS (HEAD)
        TYPE (NODE), POINTER :: HEAD
        TYPE (NODE), POINTER :: P
        !$OMP PARALLEL PRIVATE(P)
            !$OMP SINGLE
                P => HEAD
                DO
                    !$OMP TASK
                        ! P is firstprivate by default
                        CALL PROCESS(P)
                    !$OMP END TASK
                    P => P%NEXT
                    IF ( .NOT. ASSOCIATED (P) ) EXIT
                END DO
            !$OMP END SINGLE
        !$OMP END PARALLEL
    END SUBROUTINE
END MODULE

```

## Fortran

The `fib()` function should be called from within a **parallel** region for the different specified tasks to be executed in parallel. Also, only one thread of the **parallel** region should call `fib()` unless multiple concurrent Fibonacci computations are desired.

## C/C++

### Example 15.4c

```

int fib(int n) {
    int i, j;
    if (n<2)
        return n;
    else {
        #pragma omp task shared(i)
        i=fib(n-1);
        #pragma omp task shared(j)
        j=fib(n-2);
        #pragma omp taskwait
        return i+j;
    }
}

```

## C/C++



*Example 15.4f*

```

      RECURSIVE INTEGER FUNCTION fib(n) RESULT(res)
      INTEGER n, i, j
      IF ( n .LT. 2) THEN
        res = n
      ELSE
!$OMP TASK SHARED(i)
        i = fib( n-1 )
!$OMP END TASK
!$OMP TASK SHARED(j)
        j = fib( n-2 )
!$OMP END TASK
!$OMP TASKWAIT
        res = i+j
      END IF
      END FUNCTION

```

Note: There are more efficient algorithms for computing Fibonacci numbers. This classic recursion algorithm is for illustrative purposes.

The following example demonstrates a way to generate a large number of tasks with one thread and execute them with the threads in the team. While generating these tasks, the implementation may reach its limit on unassigned tasks. If it does, the implementation is allowed to cause the thread executing the task generating loop to suspend its task at the task scheduling point in the **task** directive, and start executing unassigned tasks. Once the number of unassigned tasks is sufficiently low, the thread may resume execution of the task generating loop.

*Example 15.5c*

```

#define LARGE_NUMBER 10000000
double item[LARGE_NUMBER];
extern void process(double);

int main() {
#pragma omp parallel
{
  #pragma omp single
  {
    int i;
    for (i=0; i<LARGE_NUMBER; i++)
      #pragma omp task // i is firstprivate, item is shared
        process(item[i]);
  }
}

```

```
}
}
```

C/C++

Fortran

### Example 15.5f

```
real*8 item(10000000)
integer i

!$omp parallel
!$omp single ! loop iteration variable i is private
do i=1,10000000
!$omp task
    ! i is firstprivate, item is shared
    call process(item(i))
!$omp end task
end do
!$omp end single
!$omp end parallel
end
```

Fortran

The following example is the same as the previous one, except that the tasks are generated in an untied task. While generating the tasks, the implementation may reach its limit on unassigned tasks. If it does, the implementation is allowed to cause the thread executing the task generating loop to suspend its task at the task scheduling point in the **task** directive, and start executing unassigned tasks. If that thread begins execution of a task that takes a long time to complete, the other threads may complete all the other tasks before it is finished.

In this case, since the loop is in an untied task, any other thread is eligible to resume the task generating loop. In the previous examples, the other threads would be forced to idle until the generating thread finishes its long task, since the task generating loop was in a tied task.

C/C++

### Example 15.6c

```
#define LARGE_NUMBER 10000000
double item[LARGE_NUMBER];
extern void process(double);
int main() {
#pragma omp parallel
{
    #pragma omp single
    {
```

```

        int i;
        #pragma omp task untied
        // i is firstprivate, item is shared
        {
            for (i=0; i<LARGE_NUMBER; i++)
                #pragma omp task
                process(item[i]);
        }
    }
}
return 0;
}

```

C/C++

Fortran

### Example 15.6f

```

        real*8 item(10000000)
!$omp parallel
!$omp single
!$omp task untied
! loop iteration variable i is private
do i=1,10000000
!$omp task ! i is firstprivate, item is shared
call process(item(i))
!$omp end task
end do
!$omp end task
!$omp end single
!$omp end parallel
end

```

Fortran

The following two examples demonstrate how the scheduling rules illustrated in § affect the usage of **threadprivate** variables in tasks. A **threadprivate** variable can be modified by another task that is executed by the same thread. Thus, the value of a **threadprivate** variable cannot be assumed to be unchanged across a task scheduling point. In untied tasks, task scheduling points may be added in any place by the implementation.

A task switch may occur at a task scheduling point. A single thread may execute both of the task regions that modify **tp**. The parts of these task regions in which **tp** is modified may be executed in any order so the resulting value of **var** can be either 1 or 2.

C/C++

### Example 15.7c

```

int tp;
#pragma omp threadprivate(tp)

```

```

int var;
void work()
{
#pragma omp task
{
    /* do work here */
#pragma omp task
{
    tp = 1;
    /* do work here */
#pragma omp task
{
    /* no modification of tp */
}
    var = tp; //value of tp can be 1 or 2
}
    tp = 2;
}
}

```

C/C++

Fortran

### *Example 15.7f*

```

module example
integer tp
!$omp threadprivate(tp)
integer var
contains
subroutine work
use globals
!$omp task
! do work here
!$omp task
tp = 1
! do work here
!$omp task
! no modification of tp
!$omp end task
var = tp ! value of var can be 1 or 2
!$omp end task
tp = 2
!$omp end task
end subroutine
end module

```

Fortran

In this example, scheduling constraints prohibit a thread in the team from executing a new task that modifies **tp** while another such task region tied to the same thread is suspended. Therefore, the value written will persist across the task scheduling point.

*Example 15.8c*

```

int tp;
#pragma omp threadprivate(tp)
int var;
void work()
{
#pragma omp parallel
{
    /* do work here */
#pragma omp task
{
    tp++;
    /* do work here */
#pragma omp task
{
    /* do work here but don't modify tp */
}
    var = tp; //Value does not change after write above
}
}
}

```

*Example 15.8f*

```

module example
integer tp
!$omp threadprivate(tp)
integer var
contains
subroutine work
!$omp parallel
! do work here
!$omp task
tp = tp + 1
! do work here
!$omp task
! do work here but don't modify tp
!$omp end task
var = tp ! value does not change after write above
!$omp end task
!$omp end parallel
end subroutine
end module

```

The following two examples demonstrate how the scheduling rules illustrated in § affect the usage of locks and critical sections in tasks. If a lock is held across a task scheduling point, no attempt should be made to acquire the same lock in any code that may be interleaved. Otherwise, a deadlock is possible.

In the example below, suppose the thread executing task 1 defers task 2. When it encounters the task scheduling point at task 3, it could suspend task 1 and begin task 2 which will result in a deadlock when it tries to enter critical region 1.

### Example 15.9c

```
void work()
{
    #pragma omp task
    { //Task 1
        #pragma omp task
        { //Task 2
            #pragma omp critical //Critical region 1
            { /*do work here */ }
        }
        #pragma omp critical //Critical Region 2
        {
            //Capture data for the following task
            #pragma omp task
            { /* do work here */ } //Task 3
        }
    }
}
```

### Example 15.9f

```
module example
contains
subroutine work
!$omp task
! Task 1
!$omp task
! Task 2
!$omp critical
! Critical region 1
! do work here
!$omp end critical
!$omp end task
!$omp critical
! Critical region 2
```

```
        ! Capture data for the following task
!$omp task
        !Task 3
        ! do work here
!$omp end task
!$omp end critical
!$omp end task
        end subroutine
    end module
```

Fortran

In the following example, **lock** is held across a task scheduling point. However, according to the scheduling restrictions, the executing thread can't begin executing one of the non-descendant tasks that also acquires **lock** before the task region is complete. Therefore, no deadlock is possible.

## C/C++

### Example 15.10c

```
#include <omp.h>
void work() {
    omp_lock_t lock;
    omp_init_lock(&lock);
#pragma omp parallel
    {
        int i;
#pragma omp for
        for (i = 0; i < 100; i++) {
#pragma omp task
            {
                // lock is shared by default in the task
                omp_set_lock(&lock);
                // Capture data for the following task
#pragma omp task
                    {
                        // Task Scheduling Point 1
                        { /* do work here */ }
                        omp_unset_lock(&lock);
                    }
            }
        }
    }
    omp_destroy_lock(&lock);
}
```

## C/C++

## Fortran

### Example 15.10f

```
module example
include 'omp_lib.h'
integer (kind=omp_lock_kind) lock
integer i
contains
subroutine work
call omp_init_lock(lock)
!$omp parallel
!$omp do
do i=1,100
!$omp task
! Outer task
```



```

        call omp_set_lock(lock)      ! lock is shared by
                                     ! default in the task
        ! Capture data for the following task
        !$omp task      ! Task Scheduling Point 1
            ! do work here
        !$omp end task
        call omp_unset_lock(lock)
    !$omp end task
end do
!$omp end parallel
call omp_destroy_lock(lock)
end subroutine
end module

```

### Fortran

The following examples illustrate the use of the **mergeable** clause in the **task** construct. In this first example, the **task** construct has been annotated with the **mergeable** clause. The addition of this clause allows the implementation to reuse the data environment (including the ICVs) of the parent task for the task inside **foo** if the task is included or undeferred. Thus, the result of the execution may differ depending on whether the task is merged or not. Therefore the mergeable clause needs to be used with caution. In this example, the use of the mergeable clause is safe. As **x** is a shared variable the outcome does not depend on whether or not the task is merged (that is, the task will always increment the same variable and will always compute the same value for **x**).

### C/C++

#### *Example 15.11c*

```

#include <stdio.h>
void foo ( )
{
    int x = 2;
    #pragma omp task shared(x) mergeable
    {
        x++;
    }
    #pragma omp taskwait
    printf("%d\n",x); // prints 3
}

```

### C/C++

### Fortran

#### *Example 15.11f*

```

subroutine foo()
    integer :: x

```

```

    x = 2
!$omp task shared(x) mergeable
    x = x + 1
!$omp end task
!$omp taskwait
    print *, x      ! prints 3
end subroutine

```

Fortran

This second example shows an incorrect use of the **mergeable** clause. In this example, the created task will access different instances of the variable **x** if the task is not merged, as **x** is **firstprivate**, but it will access the same variable **x** if the task is merged. As a result, the behavior of the program is unspecified and it can print two different values for **x** depending on the decisions taken by the implementation.

C/C++

*Example 15.12c*

```

#include <stdio.h>
void foo ( )
{
    int x = 2;
    #pragma omp task mergeable
    {
        x++;
    }
    #pragma omp taskwait
    printf("%d\n",x); // prints 2 or 3
}

```

C/C++

Fortran

*Example 15.12f*

```

subroutine foo()
    integer :: x
    x = 2
!$omp task mergeable
    x = x + 1
!$omp end task
!$omp taskwait
    print *, x      ! prints 2 or 3
end subroutine

```

Fortran

The following example shows the use of the **final** clause and the **omp\_in\_final** API call in a recursive binary search program. To reduce overhead, once a certain depth of recursion is reached the program uses the **final** clause to create only included tasks, which allow additional optimizations.

The use of the **omp\_in\_final** API call allows programmers to optimize their code by specifying which parts of the program are not necessary when a task can create only included tasks (that is, the code is inside a **final** task). In this example, the use of a different state variable is not necessary so once the program reaches the part of the computation that is finalized and copying from the parent state to the new state is eliminated. The allocation of **new\_state** in the stack could also be avoided but it would make this example less clear. The **final** clause is most effective when used in conjunction with the **mergeable** clause since all tasks created in a **final** task region are included tasks that can be merged if the **mergeable** clause is present.

## C/C++

### *Example 15.13c*

```
#include <string.h>
#include <omp.h>
#define LIMIT 3 /* arbitrary limit on recursion depth */
void check_solution(char *);
void bin_search (int pos, int n, char *state)
{
    if ( pos == n ) {
        check_solution(state);
        return;
    }
    #pragma omp task final( pos > LIMIT ) mergeable
    {
        char new_state[n];
        if (!omp_in_final() ) {
            memcpy(new_state, state, pos );
            state = new_state;
        }
        state[pos] = 0;
        bin_search(pos+1, n, state );
    }
    #pragma omp task final( pos > LIMIT ) mergeable
    {
        char new_state[n];
        if (! omp_in_final() ) {
            memcpy(new_state, state, pos );
            state = new_state;
        }
        state[pos] = 1;
        bin_search(pos+1, n, state );
    }
}
#pragma omp taskwait
```

}

C/C++

Fortran

### *Example 15.13f*

```
recursive subroutine bin_search(pos, n, state)
  use omp_lib
  integer :: pos, n
  character, pointer :: state(:)
  character, target, dimension(n) :: new_state1, new_state2
  integer, parameter :: LIMIT = 3
  if (pos .eq. n) then
    call check_solution(state)
    return
  endif
!$omp task final(pos > LIMIT) mergeable
  if (.not. omp_in_final()) then
    new_state1(1:pos) = state(1:pos)
    state => new_state1
  endif
  state(pos+1) = 'z'
  call bin_search(pos+1, n, state)
!$omp end task
!$omp task final(pos > LIMIT) mergeable
  if (.not. omp_in_final()) then
    new_state2(1:pos) = state(1:pos)
    state => new_state2
  endif
  state(pos+1) = 'y'
  call bin_search(pos+1, n, state)
!$omp end task
!$omp taskwait
end subroutine
```

Fortran

The following example illustrates the difference between the **if** and the **final** clauses. The **if** clause has a local effect. In the first nest of tasks, the one that has the **if** clause will be undeferred but the task nested inside that task will not be affected by the **if** clause and will be created as usual. Alternatively, the **final** clause affects all **task** constructs in the **final** task region but not the **final** task itself. In the second nest of tasks, the nested tasks will be created as included tasks. Note also that the conditions for the **if** and **final** clauses are usually the opposite.

*Example 15.14c*

```
void foo ( )
{
    int i;
    #pragma omp task if(0) // This task is undeferred
    {
        #pragma omp task // This task is a regular task
        for (i = 0; i < 3; i++) {
            #pragma omp task // This task is a regular task
            bar();
        }
    }
    #pragma omp task final(1) // This task is a regular task
    {
        #pragma omp task // This task is included
        for (i = 0; i < 3; i++) {
            #pragma omp task // This task is also included
            bar();
        }
    }
}
```

*Example 15.14f*

```

subroutine foo()
integer i
!$omp task if(.FALSE.) ! This task is undeferred
!$omp task             ! This task is a regular task
  do i = 1, 3
    !$omp task          ! This task is a regular task
      call bar()
    !$omp end task
  enddo
!$omp end task
!$omp end task
!$omp task final(.TRUE.) ! This task is a regular task
!$omp task              ! This task is included
  do i = 1, 3
    !$omp task          ! This task is also included
      call bar()
    !$omp end task
  enddo
!$omp end task
!$omp end task
end subroutine

```

## Task Dependences

### Flow Dependence

In this example we show a simple flow dependence expressed using the **depend** clause on the **task** construct.

*Example 15.15c*

```

#include <stdio.h>
int main()
{
  int x = 1;
  #pragma omp parallel
  #pragma omp single
  {
    #pragma omp task shared(x) depend(out: x)
    x = 2;
  }
}

```

```

        #pragma omp task shared(x) depend(in: x)
        printf("x = %d\n", x);
    }
    return 0;

```

C/C++

Fortran

### Example 15.15f

```

program example
  integer :: x
  x = 1
  !$omp parallel
  !$omp single
    !$omp task shared(x) depend(out: x)
    x = 2
  !$omp end task
  !$omp task shared(x) depend(in: x)
  print*, "x = ", x
  !$omp end task
  !$omp end single
  !$omp end parallel
end program

```

Fortran

The program will always print "x = 2", because the **depend** clauses enforce the ordering of the tasks. If the **depend** clauses had been omitted, then the tasks could execute in any order and the program would have a race condition.

## Anti-dependence

In this example we show an anti-dependence expressed using the **depend** clause on the **task** construct.

C/C++

### Example 15.16c

```

#include <stdio.h>
int main()
{
    int x = 1;
    #pragma omp parallel
    #pragma omp single
    {
        #pragma omp task shared(x) depend(in: x)
        printf("x = %d\n", x);
    }
}

```

```

        #pragma omp task shared(x) depend(out: x)
        x = 2;
    }
    return 0;
}

```

C/C++

Fortran

### Example 15.16f

```

program example
  integer :: x
  x = 1
  !$omp parallel
  !$omp single
    !$omp task shared(x) depend(in: x)
    print*, "x = ", x
    !$omp end task
    !$omp task shared(x) depend(out: x)
    x = 2
    !$omp end task
  !$omp end single
  !$omp end parallel
end program

```

Fortran

The program will always print "x = 1", because the **depend** clauses enforce the ordering of the tasks. If the **depend** clauses had been omitted, then the tasks could execute in any order and the program would have a race condition.

## Output Dependence

In this example we show an output dependence expressed using the **depend** clause on the **task** construct.

C/C++

### Example 15.17c

```

#include <stdio.h>
int main()
{
    int x;
    #pragma omp parallel
    #pragma omp single
    {
        #pragma omp task shared(x) depend(out: x)

```



```

        x = 1;
        #pragma omp task shared(x) depend(out: x)
        x = 2;
        #pragma omp taskwait
        printf("x = %d\n", x);
    }
    return 0;
}

```

C/C++

Fortran

### *Example 15.17f*

```

program example
    integer :: x
    !$omp parallel
    !$omp single
        !$omp task shared(x) depend(out: x)
        x = 1
    !$omp end task
        !$omp task shared(x) depend(out: x)
        x = 2
    !$omp end task
    !$omp taskwait
    print*, "x = ", x
    !$omp end single
    !$omp end parallel
end program

```

Fortran

The program will always print "x = 2", because the **depend** clauses enforce the ordering of the tasks. If the **depend** clauses had been omitted, then the tasks could execute in any order and the program would have a race condition.

## Concurrent Execution with Dependences

In this example we show potentially concurrent execution of tasks using multiple flow dependences expressed using the **depend** clause on the **task** construct.

C/C++

### *Example 15.18c*

```

#include <stdio.h>
int main()
{
    int x = 1;

```

```

#pragma omp parallel
#pragma omp single
{
    #pragma omp task shared(x) depend(out: x)
    x = 2;
    #pragma omp task shared(x) depend(in: x)
    printf("x + 1 = %d. ", x+1);
    #pragma omp task shared(x) depend(in: x)
    printf("x + 2 = %d\n", x+2);
}
return 0;
}

```

▲ C/C++ ▲

▼ Fortran ▼

### *Example 15.18f*

```

program example
integer :: x
x = 1
!$omp parallel
!$omp single
    !$omp task shared(x) depend(out: x)
    x = 2
    !$omp end task
    !$omp task shared(x) depend(in: x)
    print*, "x + 1 = ", x+1, "."
    !$omp end task
    !$omp task shared(x) depend(in: x)
    print*, "x + 2 = ", x+2, "."
    !$omp end task
!$omp end single
!$omp end parallel
end program

```

▲ Fortran ▲

The last two tasks are dependent on the first task. However there is no dependence between the last two tasks, which may execute in any order (or concurrently if more than one thread is available). Thus, the possible outputs are "x + 1 = 3. x + 2 = 4. " and "x + 2 = 4. x + 1 = 3. ". If the **depend** clauses had been omitted, then all of the tasks could execute in any order and the program would have a race condition.

## Matrix multiplication

This example shows a task-based blocked matrix multiplication. Matrices are of  $N \times N$  elements, and the multiplication is implemented using blocks of  $BS \times BS$  elements.

C/C++

*Example 15.19c*

```
// Assume BS divides N perfectly
void matmul_depend(int N, int BS, float A[N][N], float B[N][N], float C[N][N] )
{
    int i, j, k, ii, jj, kk;
    for (i = 0; i < N; i+=BS) {
        for (j = 0; j < N; j+=BS) {
            for (k = 0; k < N; k+=BS) {
#pragma omp task depend ( in: A[i:BS][k:BS], B[k:BS][j:BS] ) \
                        depend ( inout: C[i:BS][j:BS] )
                for (ii = i; ii < i+BS; ii++ )
                    for (jj = j; jj < j+BS; jj++ )
                        for (kk = k; kk < k+BS; kk++ )
                            C[ii][jj] = C[ii][jj] + A[ii][kk] * B[kk][jj];
            }
        }
    }
}
```

C/C++

*Example 15.19f*

```

! Assume BS divides N perfectly
subroutine matmul_depend (N, BS, A, B, C)
  integer :: N, BS, BM
  real, dimension(N, N) :: A, B, C
  integer :: i, j, k, ii, jj, kk
  BM = BS - 1
  do i = 1, N, BS
    do j = 1, N, BS
      do k = 1, N, BS
!$omp task depend ( in: A(i:i+BM, k:k+BM), B(k:k+BM, j:j+BM) ) &
!$omp depend ( inout: C(i:i+BM, j:j+BM) )
        do ii = i, i+BS
          do jj = j, j+BS
            do kk = k, k+BS
              C(jj,ii) = C(jj,ii) + A(kk,ii) * B(jj,kk)
            end do
          end do
        end do
      end do
    end do
  !$omp end task
  end do
end do
end subroutine

```

## 16 The `taskyield` Directive

The following example illustrates the use of the `taskyield` directive. The tasks in the example compute something useful and then do some computation that must be done in a critical region. By using `taskyield` when a task cannot get access to the `critical` region the implementation can suspend the current task and schedule some other task that can do something useful.

C/C++

*Example 16.1c*

```
#include <omp.h>

void something_useful ( void );
void something_critical ( void );
void foo ( omp_lock_t * lock, int n )
{
    int i;

    for ( i = 0; i < n; i++ )
        #pragma omp task
        {
            something_useful();
            while ( !omp_test_lock(lock) ) {
                #pragma omp taskyield
            }
            something_critical();
            omp_unset_lock(lock);
        }
}
```

C/C++

Fortran

*Example 16.1f*

```
subroutine foo ( lock, n )
    use omp_lib
    integer (kind=omp_lock_kind) :: lock
    integer n
    integer i

    do i = 1, n
        !$omp task
        call something_useful()
        do while ( .not. omp_test_lock(lock) )

```

```
        !$omp taskyield
    end do
    call something_critical()
    call omp_unset_lock(lock)
    !$omp end task
end do

end subroutine
```

Fortran

## 17

## The workshare Construct

The following are examples of the **workshare** construct.

In the following example, **workshare** spreads work across the threads executing the **parallel** region, and there is a barrier after the last statement. Implementations must enforce Fortran execution rules inside of the **workshare** block.

*Example 17.1f*

```

SUBROUTINE WSHARE1(AA, BB, CC, DD, EE, FF, N)
  INTEGER N
  REAL AA(N,N), BB(N,N), CC(N,N), DD(N,N), EE(N,N), FF(N,N)

  !$OMP PARALLEL
  !$OMP WORKSHARE
    AA = BB
    CC = DD
    EE = FF
  !$OMP END WORKSHARE
  !$OMP END PARALLEL

  END SUBROUTINE WSHARE1

```

In the following example, the barrier at the end of the first **workshare** region is eliminated with a **nowait** clause. Threads doing **CC = DD** immediately begin work on **EE = FF** when they are done with **CC = DD**.

## Fortran (cont.)

*Example 17.2f*

```

SUBROUTINE WSHARE2(AA, BB, CC, DD, EE, FF, N)
  INTEGER N
  REAL AA(N,N), BB(N,N), CC(N,N)
  REAL DD(N,N), EE(N,N), FF(N,N)

  !$OMP PARALLEL
  !$OMP WORKSHARE
    AA = BB
    CC = DD
  !$OMP END WORKSHARE NOWAIT
  !$OMP WORKSHARE
    EE = FF
  !$OMP END WORKSHARE
  !$OMP END PARALLEL

  END SUBROUTINE WSHARE2

```

The following example shows the use of an **atomic** directive inside a **workshare** construct. The computation of **SUM(AA)** is workshared, but the update to **R** is atomic.

### Example 17.3f

```

SUBROUTINE WSHARE3(AA, BB, CC, DD, N)
  INTEGER N
  REAL AA(N,N), BB(N,N), CC(N,N), DD(N,N)
  REAL R
  R=0
!$OMP PARALLEL
!$OMP WORKSHARE
  AA = BB
!$OMP ATOMIC UPDATE
  R = R + SUM(AA)
  CC = DD
!$OMP END WORKSHARE
!$OMP END PARALLEL
END SUBROUTINE WSHARE3

```

Fortran **WHERE** and **FORALL** statements are *compound statements*, made up of a *control* part and a *statement* part. When **workshare** is applied to one of these compound statements, both the control and the statement parts are workshared. The following example shows the use of a **WHERE** statement in a **workshare** construct.

Each task gets worked on in order by the threads:

```

AA = BB then
CC = DD then
EE .ne. 0 then
FF = 1 / EE then
GG = HH

```

▼----- Fortran (cont.) -----▼

### Example 17.4f

```

SUBROUTINE WSHARE4(AA, BB, CC, DD, EE, FF, GG, HH, N)
  INTEGER N
  REAL AA(N,N), BB(N,N), CC(N,N)
  REAL DD(N,N), EE(N,N), FF(N,N)
  REAL GG(N,N), HH(N,N)

!$OMP PARALLEL
!$OMP WORKSHARE
  AA = BB
  CC = DD
  WHERE (EE .ne. 0) FF = 1 / EE
  GG = HH
!$OMP END WORKSHARE
!$OMP END PARALLEL

```



```
END SUBROUTINE WSHARE4
```

In the following example, an assignment to a shared scalar variable is performed by one thread in a **workshare** while all other threads in the team wait.

### *Example 17.5f*

```

SUBROUTINE WSHARE5 (AA, BB, CC, DD, N)
  INTEGER N
  REAL AA(N,N), BB(N,N), CC(N,N), DD(N,N)

  INTEGER SHR

  !$OMP PARALLEL SHARED(SHR)
  !$OMP WORKSHARE
    AA = BB
    SHR = 1
    CC = DD * SHR
  !$OMP END WORKSHARE
  !$OMP END PARALLEL

END SUBROUTINE WSHARE5

```

The following example contains an assignment to a private scalar variable, which is performed by one thread in a **workshare** while all other threads wait. It is non-conforming because the private scalar variable is undefined after the assignment statement.

### *Example 17.6f*

```

SUBROUTINE WSHARE6_WRONG (AA, BB, CC, DD, N)
  INTEGER N
  REAL AA(N,N), BB(N,N), CC(N,N), DD(N,N)

  INTEGER PRI

  !$OMP PARALLEL PRIVATE(PRI)
  !$OMP WORKSHARE
    AA = BB
    PRI = 1
    CC = DD * PRI
  !$OMP END WORKSHARE
  !$OMP END PARALLEL

END SUBROUTINE WSHARE6_WRONG

```

Fortran execution rules must be enforced inside a **workshare** construct. In the following example, the same result is produced in the following program fragment regardless of whether the code is executed sequentially or inside an OpenMP program with multiple threads:

*Example 17.7f*

```
SUBROUTINE WSHARE7(AA, BB, CC, N)
  INTEGER N
  REAL AA(N), BB(N), CC(N)

!$OMP  PARALLEL
!$OMP    WORKSHARE
      AA(1:50) = BB(11:60)
      CC(11:20) = AA(1:10)
!$OMP    END WORKSHARE
!$OMP  END PARALLEL

END SUBROUTINE WSHARE7
```

Fortran

## 18 The master Construct

The following example demonstrates the master construct . In the example, the master keeps track of how many iterations have been executed and prints out a progress report. The other threads skip the master region without waiting.

C/C++

*Example 18.1c*

```
#include <stdio.h>

extern float average(float,float,float);

void master_example( float* x, float* xold, int n, float tol )
{
    int c, i, toobig;
    float error, y;
    c = 0;
    #pragma omp parallel
    {
        do{
            #pragma omp for private(i)
            for( i = 1; i < n-1; ++i ){
                xold[i] = x[i];
            }
            #pragma omp single
            {
                toobig = 0;
            }
            #pragma omp for private(i,y,error) reduction(+:toobig)
            for( i = 1; i < n-1; ++i ){
                y = x[i];
                x[i] = average( xold[i-1], x[i], xold[i+1] );
                error = y - x[i];
                if( error > tol || error < -tol ) ++toobig;
            }
            #pragma omp master
            {
                ++c;
                printf( "iteration %d, toobig=%d\n", c, toobig );
            }
        }while( toobig > 0 );
    }
}
```

C/C++

*Example 18.1f*

```

SUBROUTINE MASTER_EXAMPLE( X, XOLD, N, TOL )
REAL X(*), XOLD(*), TOL
INTEGER N
INTEGER C, I, TOOBIG
REAL ERROR, Y, AVERAGE
EXTERNAL AVERAGE
C = 0
TOOBIG = 1
!$OMP PARALLEL
    DO WHILE( TOOBIG > 0 )
!$OMP      DO PRIVATE(I)
            DO I = 2, N-1
                XOLD(I) = X(I)
            ENDDO
!$OMP      SINGLE
            TOOBIG = 0
!$OMP      END SINGLE
!$OMP      DO PRIVATE(I,Y,ERROR), REDUCTION(+:TOOBIG)
            DO I = 2, N-1
                Y = X(I)
                X(I) = AVERAGE( XOLD(I-1), X(I), XOLD(I+1) )
                ERROR = Y-X(I)
                IF( ERROR > TOL .OR. ERROR < -TOL ) TOOBIG = TOOBIG+1
            ENDDO
!$OMP      MASTER
            C = C + 1
            PRINT *, 'Iteration ', C, 'TOOBIG=', TOOBIG
!$OMP      END MASTER
    ENDDO
!$OMP END PARALLEL
END SUBROUTINE MASTER_EXAMPLE

```

## 19

# The critical Construct

The following example includes several **critical** constructs . The example illustrates a queuing model in which a task is dequeued and worked on. To guard against multiple threads dequeuing the same task, the dequeuing operation must be in a **critical** region. Because the two queues in this example are independent, they are protected by **critical** constructs with different names, *xaxis* and *yaxis*.

C/C++

### Example 19.1c

```
int dequeue(float *a);
void work(int i, float *a);

void critical_example(float *x, float *y)
{
    int ix_next, iy_next;

    #pragma omp parallel shared(x, y) private(ix_next, iy_next)
    {
        #pragma omp critical (xaxis)
        ix_next = dequeue(x);
        work(ix_next, x);

        #pragma omp critical (yaxis)
        iy_next = dequeue(y);
        work(iy_next, y);
    }
}
```

C/C++

Fortran

### Example 19.1f

```
SUBROUTINE CRITICAL_EXAMPLE(X, Y)

    REAL X(*), Y(*)
    INTEGER IX_NEXT, IY_NEXT

    !$OMP PARALLEL SHARED(X, Y) PRIVATE(IX_NEXT, IY_NEXT)

    !$OMP CRITICAL(XAXIS)
        CALL DEQUEUE(IX_NEXT, X)
    !$OMP END CRITICAL(XAXIS)
```

```
        CALL WORK (IX_NEXT, X)

!$OMP CRITICAL (YAXIS)
        CALL DEQUEUE (IY_NEXT, Y)
!$OMP END CRITICAL (YAXIS)
        CALL WORK (IY_NEXT, Y)

!$OMP END PARALLEL

END SUBROUTINE CRITICAL_EXAMPLE
```



Fortran

## worksharing Constructs Inside a critical Construct

The following example demonstrates using a worksharing construct inside a **critical** construct. This example is conforming because the worksharing **single** region is not closely nested inside the **critical** region. A single thread executes the one and only section in the **sections** region, and executes the **critical** region. The same thread encounters the nested **parallel** region, creates a new team of threads, and becomes the master of the new team. One of the threads in the new team enters the **single** region and increments **i** by 1. At the end of this example **i** is equal to 2.

C/C++

*Example 20.1c*

```
void critical_work()
{
    int i = 1;
    #pragma omp parallel sections
    {
        #pragma omp section
        {
            #pragma omp critical (name)
            {
                #pragma omp parallel
                {
                    #pragma omp single
                    {
                        i++;
                    }
                }
            }
        }
    }
}
```

C/C++

Fortran

*Example 20.1f*

```
SUBROUTINE CRITICAL_WORK()

    INTEGER I
    I = 1
```

```
!$OMP    PARALLEL SECTIONS
!$OMP      SECTION
!$OMP        CRITICAL (NAME)
!$OMP          PARALLEL
!$OMP            SINGLE
!$OMP              I = I + 1
!$OMP            END SINGLE
!$OMP          END PARALLEL
!$OMP        END CRITICAL (NAME)
!$OMP      END PARALLEL SECTIONS
!$OMP    END SUBROUTINE CRITICAL_WORK
```

Fortran



---

## Binding of **barrier** Regions

The binding rules call for a **barrier** region to bind to the closest enclosing **parallel** region.

In the following example, the call from the main program to *sub2* is conforming because the **barrier** region (in *sub3*) binds to the **parallel** region in *sub2*. The call from the main program to *sub1* is conforming because the **barrier** region binds to the **parallel** region in subroutine *sub2*.

The call from the main program to *sub3* is conforming because the **barrier** region binds to the implicit inactive **parallel** region enclosing the sequential part. Also note that the **barrier** region in *sub3* when called from *sub2* only synchronizes the team of threads in the enclosing **parallel** region and not all the threads created in *sub1*.

*Example 21.1c*

```

void work(int n) {}

void sub3(int n)
{
    work(n);
    #pragma omp barrier
    work(n);
}

void sub2(int k)
{
    #pragma omp parallel shared(k)
        sub3(k);
}

void sub1(int n)
{
    int i;
    #pragma omp parallel private(i) shared(n)
    {
        #pragma omp for
        for (i=0; i<n; i++)
            sub2(i);
    }
}

int main()
{
    sub1(2);
    sub2(2);
    sub3(2);
    return 0;
}

```

*Example 21.1f*

```

      SUBROUTINE WORK(N)
        INTEGER N
      END SUBROUTINE WORK

      SUBROUTINE SUB3(N)
        INTEGER N
        CALL WORK(N)
!$OMP BARRIER
        CALL WORK(N)

```

```

        END SUBROUTINE SUB3

        SUBROUTINE SUB2(K)
        INTEGER K
!$OMP   PARALLEL SHARED(K)
            CALL SUB3(K)
!$OMP   END PARALLEL
        END SUBROUTINE SUB2

        SUBROUTINE SUB1(N)
        INTEGER N
        INTEGER I
!$OMP   PARALLEL PRIVATE(I) SHARED(N)
!$OMP   DO
            DO I = 1, N
                CALL SUB2(I)
            END DO
!$OMP   END PARALLEL
        END SUBROUTINE SUB1

        PROGRAM EXAMPLE
        CALL SUB1(2)
        CALL SUB2(2)
        CALL SUB3(2)
        END PROGRAM EXAMPLE

```

Fortran

## 22 The `atomic` Construct

The following example avoids race conditions (simultaneous updates of an element of  $x$  by multiple threads) by using the `atomic` construct .

The advantage of using the `atomic` construct in this example is that it allows updates of two different elements of  $x$  to occur in parallel. If a `critical` construct were used instead, then all updates to elements of  $x$  would be executed serially (though not in any guaranteed order).

Note that the `atomic` directive applies only to the statement immediately following it. As a result, elements of  $y$  are not updated atomically in this example.

C/C++

*Example 22.1c*

```
float work1(int i)
{
    return 1.0 * i;
}

float work2(int i)
{
    return 2.0 * i;
}

void atomic_example(float *x, float *y, int *index, int n)
{
    int i;

    #pragma omp parallel for shared(x, y, index, n)
    for (i=0; i<n; i++) {
        #pragma omp atomic update
        x[index[i]] += work1(i);
        y[i] += work2(i);
    }
}

int main()
{
    float x[1000];
    float y[10000];
    int index[10000];
    int i;

    for (i = 0; i < 10000; i++) {
        index[i] = i % 1000;
        y[i]=0.0;
    }
}
```

```

    }
    for (i = 0; i < 1000; i++)
        x[i] = 0.0;
    atomic_example(x, y, index, 10000);
    return 0;
}

```

C/C++

Fortran

### *Example 22.1f*

```

REAL FUNCTION WORK1(I)
  INTEGER I
  WORK1 = 1.0 * I
  RETURN
END FUNCTION WORK1

REAL FUNCTION WORK2(I)
  INTEGER I
  WORK2 = 2.0 * I
  RETURN
END FUNCTION WORK2

SUBROUTINE SUB(X, Y, INDEX, N)
  REAL X(*), Y(*)
  INTEGER INDEX(*), N

  INTEGER I

!$OMP PARALLEL DO SHARED(X, Y, INDEX, N)
  DO I=1,N
!$OMP ATOMIC UPDATE
    X(INDEX(I)) = X(INDEX(I)) + WORK1(I)
    Y(I) = Y(I) + WORK2(I)
  ENDDO

END SUBROUTINE SUB

PROGRAM ATOMIC_EXAMPLE
  REAL X(1000), Y(10000)
  INTEGER INDEX(10000)
  INTEGER I

  DO I=1,10000
    INDEX(I) = MOD(I, 1000) + 1
    Y(I) = 0.0
  ENDDO

  DO I = 1,1000
    X(I) = 0.0
  ENDDO

```

```

        CALL SUB(X, Y, INDEX, 10000)

END PROGRAM ATOMIC_EXAMPLE

```

## Fortran

The following example illustrates the **read** and **write** clauses for the **atomic** directive. These clauses ensure that the given variable is read or written, respectively, as a whole. Otherwise, some other thread might read or write part of the variable while the current thread was reading or writing another part of the variable. Note that most hardware provides atomic reads and writes for some set of properly aligned variables of specific sizes, but not necessarily for all the variable types supported by the OpenMP API.

## C/C++

### Example 22.2c

```

int atomic_read(const int *p)
{
    int value;
    /* Guarantee that the entire value of *p is read atomically. No part of
     * *p can change during the read operation.
     */
    #pragma omp atomic read
        value = *p;
    return value;
}

void atomic_write(int *p, int value)
{
    /* Guarantee that value is stored atomically into *p. No part of *p can change
     * until after the entire write operation is completed.
     */
    #pragma omp atomic write
        *p = value;
}

```

## C/C++

## Fortran

### Example 22.2f

```

function atomic_read(p)
    integer :: atomic_read
    integer, intent(in) :: p
    ! Guarantee that the entire value of p is read atomically. No part of
    ! p can change during the read operation.

```

```

!$omp atomic read
    atomic_read = p
    return
end function atomic_read

subroutine atomic_write(p, value)
    integer, intent(out) :: p
    integer, intent(in) :: value
! Guarantee that value is stored atomically into p. No part of p can change
! until after the entire write operation is completed.
!$omp atomic write
    p = value
end subroutine atomic_write

```

## Fortran

The following example illustrates the **capture** clause for the **atomic** directive. In this case the value of a variable is captured, and then the variable is incremented. These operations occur atomically. This particular example could be implemented using the fetch-and-add instruction available on many kinds of hardware. The example also shows a way to implement a spin lock using the **capture** and **read** clauses.

## C/C++

### *Example 22.3c*

```

int fetch_and_add(int *p)
{
    /* Atomically read the value of *p and then increment it. The previous value is
     * returned. This can be used to implement a simple lock as shown below.
     */
    int old;
#pragma omp atomic capture
    { old = *p; (*p)++; }
    return old;
}

/*
 * Use fetch_and_add to implement a lock
 */
struct locktype {
    int ticketnumber;
    int turn;
};
void do_locked_work(struct locktype *lock)
{
    int atomic_read(const int *p);
    void work();

    // Obtain the lock
    int myturn = fetch_and_add(&lock->ticketnumber);
}

```

```

while (atomic_read(&lock->turn) != myturn)
;
// Do some work. The flush is needed to ensure visibility of
// variables not involved in atomic directives

#pragma omp flush
work();
#pragma omp flush
// Release the lock
fetch_and_add(&lock->turn);
}

```

C/C++

Fortran

### Example 22.3f

```

function fetch_and_add(p)
    integer :: fetch_and_add
    integer, intent(inout) :: p

! Atomically read the value of p and then increment it. The previous value is
! returned. This can be used to implement a simple lock as shown below.
!$omp atomic capture
    fetch_and_add = p
    p = p + 1
!$omp end atomic
end function fetch_and_add

! Use fetch_and_add to implement a lock
module m
    interface
        function fetch_and_add(p)
            integer :: fetch_and_add
            integer, intent(inout) :: p
        end function
        function atomic_read(p)
            integer :: atomic_read
            integer, intent(in) :: p
        end function
    end interface
    type locktype
        integer ticketnumber
        integer turn
    end type
    contains
        subroutine do_locked work(lock)
            type(locktype), intent(inout) :: lock
            integer myturn
            integer junk
! obtain the lock
            myturn = fetch_and_add(lock%ticketnumber)
            do while (atomic_read(lock%turn) .ne. myturn)

```



```

        continue
    enddo
! Do some work. The flush is needed to ensure visibility of variables
! not involved in atomic directives
!$omp flush
    call work
!$omp flush
! Release the lock
    junk = fetch_and_add(lock%turn)
end subroutine
end module

```

Fortran

## 23

# Restrictions on the `atomic` Construct

The following non-conforming examples illustrate the restrictions on the `atomic` construct.

### Example 23.1c

C/C++

```
void atomic_wrong ()
{
    union {int n; float x;} u;

    #pragma omp parallel
    {
        #pragma omp atomic update
        u.n++;

        #pragma omp atomic update
        u.x += 1.0;

        /* Incorrect because the atomic constructs reference the same location
           through incompatible types */
    }
}
```

C/C++

### Example 23.1f

Fortran

```
SUBROUTINE ATOMIC_WRONG()
    INTEGER:: I
    REAL:: R
    EQUIVALENCE(I,R)

    !$OMP PARALLEL
    !$OMP ATOMIC UPDATE
        I = I + 1
    !$OMP ATOMIC UPDATE
        R = R + 1.0
    ! incorrect because I and R reference the same location
    ! but have different types
    !$OMP END PARALLEL
END SUBROUTINE ATOMIC_WRONG
```

Fortran

*Example 23.2c*

```
void atomic_wrong2 ()
{
    int x;
    int *i;
    float *r;

    i = &x;
    r = (float *)&x;

    #pragma omp parallel
    {
        #pragma omp atomic update
            *i += 1;

        #pragma omp atomic update
            *r += 1.0;

        /* Incorrect because the atomic constructs reference the same location
           through incompatible types */

    }
}
```

The following example is non-conforming because **I** and **R** reference the same location but have different types.

### *Example 23.2f*

```

SUBROUTINE SUB()
  COMMON /BLK/ R
  REAL R

!$OMP  ATOMIC UPDATE
      R = R + 1.0
END SUBROUTINE SUB

SUBROUTINE ATOMIC_WRONG2()
  COMMON /BLK/ I
  INTEGER I

!$OMP  PARALLEL

!$OMP  ATOMIC UPDATE
      I = I + 1
      CALL SUB()
!$OMP  END PARALLEL
END SUBROUTINE ATOMIC_WRONG2

```

Although the following example might work on some implementations, this is also non-conforming:

### *Example 23.3f*

```

SUBROUTINE ATOMIC_WRONG3
  INTEGER:: I
  REAL:: R
  EQUIVALENCE(I,R)

!$OMP  PARALLEL
!$OMP  ATOMIC UPDATE
      I = I + 1
! incorrect because I and R reference the same location
! but have different types
!$OMP  END PARALLEL

!$OMP  PARALLEL
!$OMP  ATOMIC UPDATE
      R = R + 1.0
! incorrect because I and R reference the same location
! but have different types
!$OMP  END PARALLEL

```

```
END SUBROUTINE ATOMIC_WRONG3
```

Fortran

## 24

# The `flush` Construct without a List

The following example distinguishes the shared variables affected by a `flush` construct with no list from the shared objects that are not affected:

C/C++

*Example 24.1c*

```
int x, *p = &x;

void f1(int *q)
{
    *q = 1;
    #pragma omp flush
    /* x, p, and *q are flushed */
    /* because they are shared and accessible */
    /* q is not flushed because it is not shared. */
}

void f2(int *q)
{
    #pragma omp barrier
    *q = 2;
    #pragma omp barrier

    /* a barrier implies a flush */
    /* x, p, and *q are flushed */
    /* because they are shared and accessible */
    /* q is not flushed because it is not shared. */
}

int g(int n)
{
    int i = 1, j, sum = 0;
    *p = 1;
    #pragma omp parallel reduction(+: sum) num_threads(10)
    {
        f1(&j);

        /* i, n and sum were not flushed */
        /* because they were not accessible in f1 */
        /* j was flushed because it was accessible */
        sum += j;

        f2(&j);

        /* i, n, and sum were not flushed */
        /* because they were not accessible in f2 */
        /* j was flushed because it was accessible */
    }
}
```

```

        sum += i + j + *p + n;
    }
    return sum;
}

int main()
{
    int result = g(7);
    return result;
}

```

C/C++

Fortran

### *Example 24.1f*

```

SUBROUTINE F1(Q)
    COMMON /DATA/ X, P
    INTEGER, TARGET :: X
    INTEGER, POINTER :: P
    INTEGER Q

    Q = 1
!$OMP FLUSH
    ! X, P and Q are flushed
    ! because they are shared and accessible
END SUBROUTINE F1

SUBROUTINE F2(Q)
    COMMON /DATA/ X, P
    INTEGER, TARGET :: X
    INTEGER, POINTER :: P
    INTEGER Q

!$OMP BARRIER
    Q = 2
!$OMP BARRIER
    ! a barrier implies a flush
    ! X, P and Q are flushed
    ! because they are shared and accessible
END SUBROUTINE F2

INTEGER FUNCTION G(N)
    COMMON /DATA/ X, P
    INTEGER, TARGET :: X
    INTEGER, POINTER :: P
    INTEGER N
    INTEGER I, J, SUM

    I = 1
    SUM = 0
    P = 1

```

```

!$OMP  PARALLEL REDUCTION(+: SUM) NUM_THREADS(10)
      CALL F1(J)
        ! I, N and SUM were not flushed
        !   because they were not accessible in F1
        ! J was flushed because it was accessible
      SUM = SUM + J

      CALL F2(J)
        ! I, N, and SUM were not flushed
        !   because they were not accessible in f2
        ! J was flushed because it was accessible
      SUM = SUM + I + J + P + N
!$OMP  END PARALLEL

      G = SUM
END FUNCTION G

PROGRAM FLUSH_NOLIST
  COMMON /DATA/ X, P
  INTEGER, TARGET  :: X
  INTEGER, POINTER :: P
  INTEGER RESULT, G

  P => X
  RESULT = G(7)
  PRINT *, RESULT
END PROGRAM FLUSH_NOLIST

```

Fortran



## 25

# Placement of `flush`, `barrier`, `taskwait` and `taskyield` Directives

The following example is non-conforming, because the `flush`, `barrier`, `taskwait`, and `taskyield` directives are stand-alone directives and cannot be the immediate substatement of an `if` statement.

C/C++

*Example 25.1c*

```
void standalone_wrong()
{
    int a = 1;

    if (a != 0)
        #pragma omp flush(a)
    /* incorrect as flush cannot be immediate substatement
       of if statement */

    if (a != 0)
        #pragma omp barrier
    /* incorrect as barrier cannot be immediate substatement
       of if statement */

    if (a!=0)
        #pragma omp taskyield
    /* incorrect as taskyield cannot be immediate substatement of if statement */

    if (a != 0)
        #pragma omp taskwait
    /* incorrect as taskwait cannot be immediate substatement
       of if statement */
}
```

C/C++

The following example is non-conforming, because the `flush`, `barrier`, `taskwait`, and `taskyield` directives are stand-alone directives and cannot be the action statement of an `if` statement or a labeled branch target.

*Example 25.1f*

```

SUBROUTINE STANDALONE_WRONG()
  INTEGER A
  A = 1
  ! the FLUSH directive must not be the action statement
  ! in an IF statement
  IF (A .NE. 0) !$OMP FLUSH(A)

  ! the BARRIER directive must not be the action statement
  ! in an IF statement
  IF (A .NE. 0) !$OMP BARRIER

  ! the TASKWAIT directive must not be the action statement
  ! in an IF statement
  IF (A .NE. 0) !$OMP TASKWAIT

  ! the TASKYIELD directive must not be the action statement
  ! in an IF statement
  IF (A .NE. 0) !$OMP TASKYIELD

  GOTO 100

  ! the FLUSH directive must not be a labeled branch target
  ! statement
  100 !$OMP FLUSH(A)
  GOTO 200

  ! the BARRIER directive must not be a labeled branch target
  ! statement
  200 !$OMP BARRIER
  GOTO 300

  ! the TASKWAIT directive must not be a labeled branch target
  ! statement
  300 !$OMP TASKWAIT
  GOTO 400

  ! the TASKYIELD directive must not be a labeled branch target
  ! statement
  400 !$OMP TASKYIELD

END SUBROUTINE

```

The following version of the above example is conforming because the **flush**, **barrier**, **taskwait**, and **taskyield** directives are enclosed in a compound statement.

*Example 25.2c*

```

void standalone_ok()
{
    int a = 1;

    #pragma omp parallel
    {
        if (a != 0) {
            #pragma omp flush(a)
        }
        if (a != 0) {
            #pragma omp barrier
        }
        if (a != 0) {
            #pragma omp taskwait
        }
        if (a != 0) {
            #pragma omp taskyield
        }
    }
}

```

The following example is conforming because the **flush**, **barrier**, **taskwait**, and **taskyield** directives are enclosed in an **if** construct or follow the labeled branch target.

*Example 25.2f*

```

SUBROUTINE STANDALONE_OK()
    INTEGER A
    A = 1
    IF (A .NE. 0) THEN
        !$OMP FLUSH(A)
    ENDIF
    IF (A .NE. 0) THEN
        !$OMP BARRIER
    ENDIF
    IF (A .NE. 0) THEN
        !$OMP TASKWAIT
    ENDIF
    IF (A .NE. 0) THEN
        !$OMP TASKYIELD
    ENDIF
    GOTO 100
100 CONTINUE

```

```
!$OMP FLUSH(A)
GOTO 200
200 CONTINUE
!$OMP BARRIER
GOTO 300
300 CONTINUE
!$OMP TASKWAIT
GOTO 400
400 CONTINUE
!$OMP TASKYIELD
END SUBROUTINE
```



Fortran

## 26

# The ordered Clause and the ordered Construct

Ordered constructs are useful for sequentially ordering the output from work that is done in parallel. The following program prints out the indices in sequential order:

C/C++

*Example 26.1c*

```
#include <stdio.h>

void work(int k)
{
    #pragma omp ordered
    printf(" %d\n", k);
}

void ordered_example(int lb, int ub, int stride)
{
    int i;

    #pragma omp parallel for ordered schedule(dynamic)
    for (i=lb; i<ub; i+=stride)
        work(i);
}

int main()
{
    ordered_example(0, 100, 5);
    return 0;
}
```

C/C++

Fortran

*Example 26.1f*

```
SUBROUTINE WORK(K)
    INTEGER k

    !$OMP ORDERED
        WRITE(*,*) K
    !$OMP END ORDERED

END SUBROUTINE WORK
```

```

SUBROUTINE SUB(LB, UB, STRIDE)
  INTEGER LB, UB, STRIDE
  INTEGER I

!$OMP PARALLEL DO ORDERED SCHEDULE(DYNAMIC)
  DO I=LB,UB,STRIDE
    CALL WORK(I)
  END DO
!$OMP END PARALLEL DO

END SUBROUTINE SUB

PROGRAM ORDERED_EXAMPLE
  CALL SUB(1,100,5)
END PROGRAM ORDERED_EXAMPLE

```

## Fortran

It is possible to have multiple **ordered** constructs within a loop region with the **ordered** clause specified. The first example is non-conforming because all iterations execute two **ordered** regions. An iteration of a loop must not execute more than one **ordered** region:

*Example 26.2c*

```

void work(int i) {}

void ordered_wrong(int n)
{
    int i;
    #pragma omp for ordered
    for (i=0; i<n; i++) {
/* incorrect because an iteration may not execute more than one
   ordered region */
        #pragma omp ordered
        work(i);
        #pragma omp ordered
        work(i+1);
    }
}

```

*Example 26.2f*

```

      SUBROUTINE WORK(I)
      INTEGER I
      END SUBROUTINE WORK

      SUBROUTINE ORDERED_WRONG(N)
      INTEGER N

      INTEGER I
!$OMP DO ORDERED
      DO I = 1, N
! incorrect because an iteration may not execute more than one
! ordered region
!$OMP ORDERED
          CALL WORK(I)
!$OMP END ORDERED

!$OMP ORDERED
          CALL WORK(I+1)
!$OMP END ORDERED
      END DO
      END SUBROUTINE ORDERED_WRONG

```

The following is a conforming example with more than one **ordered** construct. Each iteration will execute only one **ordered** region:

*Example 26.3c*

```

void work(int i) {}
void ordered_good(int n)
{
    int i;
    #pragma omp for ordered
    for (i=0; i<n; i++) {
        if (i <= 10) {
            #pragma omp ordered
            work(i);
        }
        if (i > 10) {
            #pragma omp ordered
            work(i+1);
        }
    }
}

```

*Example 26.3f*

```

      SUBROUTINE ORDERED_GOOD(N)
      INTEGER N

      !$OMP DO ORDERED
      DO I = 1,N
          IF (I <= 10) THEN
      !$OMP ORDERED
              CALL WORK(I)
      !$OMP END ORDERED
          ENDIF

          IF (I > 10) THEN
      !$OMP ORDERED
              CALL WORK(I+1)
      !$OMP END ORDERED
          ENDIF
      ENDDO
      END SUBROUTINE ORDERED_GOOD

```



C/C++

*Example 27.1c*

The following example shows how the **cancel** directive can be used to terminate an OpenMP region. Although the **cancel** construct terminates the OpenMP worksharing region, programmers must still track the exception through the pointer `ex` and issue a cancellation for the **parallel** region if an exception has been raised. The master thread checks the exception pointer to make sure that the exception is properly handled in the sequential part. If cancellation of the **parallel** region has been requested, some threads might have executed `phase_1()`. However, it is guaranteed that none of the threads executed `phase_2()`.

```
void example() {
    std::exception *ex = NULL;
#pragma omp parallel shared(ex)
    {
#pragma omp for
        for (int i = 0; i < N; i++) {
            // no 'if' that prevents compiler optimizations
            try {
                causes_an_exception();
            }
            catch (const std::exception *e) {
                // still must remember exception for later handling
#pragma omp atomic write
                ex = e;

                // cancel worksharing construct
#pragma omp cancel for
            }
            // if an exception has been raised, cancel parallel region
            if (ex) {
#pragma omp cancel parallel
            }
            phase_1();
#pragma omp barrier
            phase_2();
        }
        // continue here if an exception has been thrown in the worksharing loop
        if (ex) {
            // handle exception stored in ex
        }
    }
}
```

C/C++

*Example 27.1f*

The following example illustrates the use of the **cancel** construct in error handling. If there is an error condition from the **allocate** statement, the cancellation is activated. The encountering thread sets the shared variable **err** and other threads of the binding thread set proceed to the end of the worksharing construct after the cancellation has been activated.

```

subroutine example(n, dim)
  integer, intent(in) :: n, dim(n)
  integer :: i, s, err
  real, allocatable :: B(:)
  err = 0
!$omp parallel shared(err)
! ...
!$omp do private(s, B)
  do i=1, n
!$omp cancellation point
    allocate(B(dim(i)), stat=s)
    if (s .gt. 0) then
!$omp atomic write
      err = s
!$omp cancel do
    endif
! ...
! deallocate private array B
    if (allocated(B)) then
      deallocate(B)
    endif
  enddo
!$omp end parallel
end subroutine

```

*Example 27.2c*

The following example shows how to cancel a parallel search on a binary tree as soon as the search value has been detected. The code creates a task to descend into the child nodes of the current tree node. If the search value has been found, the code remembers the tree node with the found value through an **atomic** write to the result variable and then cancels execution of all search tasks. The function **search\_tree\_parallel** groups all search tasks into a single task group to control the effect of the **cancel taskgroup** directive. The *level* argument is used to create undeferred tasks after the first ten levels of the tree.

```

binary_tree_t *search_tree(binary_tree_t *tree, int value, int level) {
    binary_tree_t *found = NULL;
    if (tree) {
        if (tree->value == value) {
            found = tree;
        }
        else {
#pragma omp task shared(found) if(level < 10)
            {
                binary_tree_t *found_left = NULL;
                found_left = search_tree(tree->left, value, level + 1);
                if (found_left) {
#pragma omp atomic write
                    found = found_left;
#pragma omp cancel taskgroup
                }
            }
#pragma omp task shared(found) if(level < 10)
            {
                binary_tree_t *found_right = NULL;
                found_right = search_tree(tree->right, value, level + 1);
                if (found_right) {
#pragma omp atomic write
                    found = found_right;
#pragma omp cancel taskgroup
                }
            }
#pragma omp taskwait
        }
    }
    return found;
}

binary_tree_t *search_tree_parallel(binary_tree_t *tree, int value) {
    binary_tree_t *found = NULL;
#pragma omp parallel shared(found, tree, value)
    {
#pragma omp master
        {
#pragma omp taskgroup
            {
                found = search_tree(tree, value, 0);
            }
        }
    }
    return found;
}

```

C/C++

*Example 27.2f*

The following is the equivalent parallel search example in Fortran.

```

module parallel_search
  type binary_tree
    integer :: value
    type(binary_tree), pointer :: right
    type(binary_tree), pointer :: left
  end type
!
contains
  recursive subroutine search_tree(tree, value, level, found)
    type(binary_tree), intent(in), pointer :: tree
    integer, intent(in) :: value, level
    type(binary_tree), pointer :: found
    type(binary_tree), pointer :: found_left => NULL(), found_right => NULL()
!
    if (.not. associated(found)) then
      allocate(found)
    endif
!
    if (associated(tree)) then
      if (tree%value .eq. value) then
        found = tree
      else
!$omp task shared(found) if(level<10)
        call search_tree(tree%left, value, level+1, found_left)
        if (associated(found_left)) then
!$omp atomic write
          found = found_left
!
!$omp cancel taskgroup
        endif
!$omp end task
!
!$omp task shared(found) if(level<10)
        call search_tree(tree%right, value, level+1, found_right)
        if (associated(found_right)) then
!$omp atomic write
          found = found_right
!
!$omp cancel taskgroup
        endif
!$omp end task
!
!$omp taskwait
        endif
      endif
    end subroutine
!

```

```

subroutine search_tree_parallel(tree, value, found)
  type(binary_tree), intent(in), pointer :: tree
  integer, intent(in) :: value
  type(binary_tree), pointer :: found
!
  if (associated(found)) then
    allocate(found)
  endif
!$omp parallel shared(found, tree, value)
!$omp master
!$omp taskgroup
  call search_tree(tree, value, 0, found)
!$omp end taskgroup
!$omp end master
!$omp end parallel
  end subroutine
!
end module parallel_search

```

Fortran

## 28

# The threadprivate Directive

The following examples demonstrate how to use the **threadprivate** directive to give each thread a separate counter.

Example 28.1c

```
int counter = 0;
#pragma omp threadprivate(counter)

int increment_counter()
{
    counter++;
    return(counter);
}
```

C/C++

Example 28.1f

```
INTEGER FUNCTION INCREMENT_COUNTER()
COMMON/INC_COMMON/COUNTER
!$OMP THREADPRIVATE (/INC_COMMON/)

COUNTER = COUNTER +1
INCREMENT_COUNTER = COUNTER
RETURN
END FUNCTION INCREMENT_COUNTER
```

Fortran

The following example uses **threadprivate** on a static variable:

Example 28.2c

```
int increment_counter_2()
{
    static int counter = 0;
    #pragma omp threadprivate(counter)
    counter++;
    return(counter);
}
```

C/C++

The following example demonstrates unspecified behavior for the initialization of a **threadprivate** variable. A **threadprivate** variable is initialized once at an unspecified point before its first reference. Because **a** is constructed using the value of **x** (which is modified by the statement **x++**), the value of **a.val** at the start of the **parallel** region could be either 1 or 2. This problem is avoided for **b**, which uses an auxiliary **const** variable and a copy-constructor.

### *Example 28.3c*

```
class T {
public:
    int val;
    T (int);
    T (const T&);
};

T :: T (int v){
    val = v;
}

T :: T (const T& t) {
    val = t.val;
}

void g(T a, T b){
    a.val += b.val;
}

int x = 1;
T a(x);
const T b_aux(x); /* Capture value of x = 1 */
T b(b_aux);
#pragma omp threadprivate(a, b)

void f(int n) {
    x++;
    #pragma omp parallel for
    /* In each thread:
     * a is constructed from x (with value 1 or 2?)
     * b is copy-constructed from b_aux
     */

    for (int i=0; i<n; i++) {
        g(a, b); /* Value of a is unspecified. */
    }
}
```

C/C++

The following examples show non-conforming uses and correct uses of the **threadprivate** directive.

The following example is non-conforming because the common block is not declared local to the subroutine that refers to it:

*Example 28.2f*

```

MODULE INC_MODULE
  COMMON /T/ A
END MODULE INC_MODULE

SUBROUTINE INC_MODULE_WRONG()
  USE INC_MODULE
!$OMP  THREADPRIVATE(/T/)
!non-conforming because /T/ not declared in INC_MODULE_WRONG
END SUBROUTINE INC_MODULE_WRONG

```

The following example is also non-conforming because the common block is not declared local to the subroutine that refers to it:

*Example 28.3f*

```

SUBROUTINE INC_WRONG()
  COMMON /T/ A
!$OMP  THREADPRIVATE(/T/)

  CONTAINS
    SUBROUTINE INC_WRONG_SUB()
!$OMP    PARALLEL COPYIN(/T/)
!non-conforming because /T/ not declared in INC_WRONG_SUB
!$OMP    END PARALLEL
    END SUBROUTINE INC_WRONG_SUB
END SUBROUTINE INC_WRONG

```



The following example is a correct rewrite of the previous example:

*Example 28.4f*

```

SUBROUTINE INC_GOOD()
COMMON /T/ A
!$OMP THREADPRIVATE(/T/)

CONTAINS
SUBROUTINE INC_GOOD_SUB()
COMMON /T/ A
!$OMP THREADPRIVATE(/T/)

!$OMP PARALLEL COPYIN(/T/)
!$OMP END PARALLEL
END SUBROUTINE INC_GOOD_SUB
END SUBROUTINE INC_GOOD

```

The following is an example of the use of **threadprivate** for local variables:

*Example 28.5f*

```

PROGRAM INC_GOOD2
INTEGER, ALLOCATABLE, SAVE :: A(:)
INTEGER, POINTER, SAVE :: PTR
INTEGER, SAVE :: I
INTEGER, TARGET :: TARG
LOGICAL :: FIRSTIN = .TRUE.
!$OMP THREADPRIVATE(A, I, PTR)

ALLOCATE (A(3))
A = (/1,2,3/)
PTR => TARG
I = 5

!$OMP PARALLEL COPYIN(I, PTR)
!$OMP CRITICAL
IF (FIRSTIN) THEN
TARG = 4 ! Update target of ptr
I = I + 10
IF (ALLOCATED(A)) A = A + 10
FIRSTIN = .FALSE.
END IF

IF (ALLOCATED(A)) THEN
PRINT *, 'a = ', A
ELSE

```

```

        PRINT *, 'A is not allocated'
    END IF

    PRINT *, 'ptr = ', PTR
    PRINT *, 'i = ', I
    PRINT *

!$OMP    END CRITICAL
!$OMP    END PARALLEL
END PROGRAM INC_GOOD2

```

The above program, if executed by two threads, will print one of the following two sets of output:

```

a = 11 12 13
ptr = 4
i = 15

A is not allocated
ptr = 4
i = 5

```

or

```

A is not allocated
ptr = 4
i = 15

a = 1 2 3
ptr = 4
i = 5

```

The following is an example of the use of **threadprivate** for module variables:

### *Example 28.6f*

```

MODULE INC_MODULE_GOOD3
    REAL, POINTER :: WORK(:)
    SAVE WORK
!$OMP    THREADPRIVATE(WORK)
END MODULE INC_MODULE_GOOD3

SUBROUTINE SUB1(N)
    USE INC_MODULE_GOOD3
!$OMP    PARALLEL PRIVATE (THE_SUM)
        ALLOCATE (WORK(N))

```

```

        CALL SUB2 (THE_SUM)
        WRITE (*,*) THE_SUM
!$OMP   END PARALLEL
END SUBROUTINE SUB1

SUBROUTINE SUB2 (THE_SUM)
    USE INC_MODULE_GOOD3
    WORK(:) = 10
    THE_SUM=SUM(WORK)
END SUBROUTINE SUB2

PROGRAM INC_GOOD3
    N = 10
    CALL SUB1 (N)
END PROGRAM INC_GOOD3

```

Fortran

C/C++

The following example illustrates initialization of **threadprivate** variables for class-type **T**. **t1** is default constructed, **t2** is constructed taking a constructor accepting one argument of integer type, **t3** is copy constructed with argument **f()**:

#### *Example 28.4c*

```

static T t1;
#pragma omp threadprivate(t1)
static T t2( 23 );
#pragma omp threadprivate(t2)
static T t3 = f();
#pragma omp threadprivate(t3)

```

The following example illustrates the use of **threadprivate** for static class members. The **threadprivate** directive for a static class member must be placed inside the class definition.

#### *Example 28.5c*

```

class T {
public:
    static int i;
#pragma omp threadprivate(i)
};

```

C/C++

## Parallel Random Access Iterator Loop

The following example shows a parallel random access iterator loop.

*Example 29.1c*

```
#include <vector>
void iterator_example()
{
    std::vector<int> vec(23);
    std::vector<int>::iterator it;
    #pragma omp parallel for default(none) shared(vec)
    for (it = vec.begin(); it < vec.end(); it++)
    {
        // do work with *it //
    }
}
```

## 30

## Fortran Restrictions on shared and private Clauses with Common Blocks

When a named common block is specified in a **private**, **firstprivate**, or **lastprivate** clause of a construct, none of its members may be declared in another data-sharing attribute clause on that construct. The following examples illustrate this point.

The following example is conforming:

### *Example 30.1f*

```

SUBROUTINE COMMON_GOOD()
  COMMON /C/ X,Y
  REAL X, Y

!$OMP  PARALLEL PRIVATE (/C/)
      ! do work here
!$OMP  END PARALLEL
!$OMP  PARALLEL SHARED (X,Y)
      ! do work here
!$OMP  END PARALLEL
END SUBROUTINE COMMON_GOOD

```

The following example is also conforming:

### *Example 30.2f*

```

SUBROUTINE COMMON_GOOD2()
  COMMON /C/ X,Y
  REAL X, Y
  INTEGER I

!$OMP  PARALLEL
!$OMP    DO PRIVATE(/C/)
        DO I=1,1000
          ! do work here
        ENDDO
!$OMP    END DO
!$OMP    DO PRIVATE(X)
        DO I=1,1000
          ! do work here
        ENDDO
!$OMP    END DO
!$OMP  END PARALLEL
END SUBROUTINE COMMON_GOOD2

```

The following example is conforming:

*Example 30.3f*

```
SUBROUTINE COMMON_GOOD3()
  COMMON /C/ X,Y
!$OMP  PARALLEL PRIVATE (/C/)
      ! do work here
!$OMP  END PARALLEL
!$OMP  PARALLEL SHARED (/C/)
      ! do work here
!$OMP  END PARALLEL
END SUBROUTINE COMMON_GOOD3
```

The following example is non-conforming because **x** is a constituent element of **c**:

*Example 30.4f*

```
SUBROUTINE COMMON_WRONG()
  COMMON /C/ X,Y
! Incorrect because X is a constituent element of C
!$OMP  PARALLEL PRIVATE(/C/), SHARED(X)
      ! do work here
!$OMP  END PARALLEL
END SUBROUTINE COMMON_WRONG
```

The following example is non-conforming because a common block may not be declared both shared and private:

*Example 30.5f*

```
SUBROUTINE COMMON_WRONG2()
  COMMON /C/ X,Y
! Incorrect: common block C cannot be declared both
! shared and private
!$OMP  PARALLEL PRIVATE (/C/), SHARED(/C/)
      ! do work here
!$OMP  END PARALLEL

END SUBROUTINE COMMON_WRONG2
```

Fortran

## 31

# The default (none) Clause

The following example distinguishes the variables that are affected by the **default (none)** clause from those that are not.

C/C++

*Example 31.1c*

```
#include <omp.h>
int x, y, z[1000];
#pragma omp threadprivate(x)

void default_none(int a) {
    const int c = 1;
    int i = 0;

    #pragma omp parallel default(none) private(a) shared(z)
    {
        int j = omp_get_num_threads();
        /* O.K. - j is declared within parallel region */
        a = z[j]; /* O.K. - a is listed in private clause */
        /*      - z is listed in shared clause */
        x = c;    /* O.K. - x is threadprivate */
        /*      - c has const-qualified type */
        z[i] = y; /* Error - cannot reference i or y here */

        #pragma omp for firstprivate(y)
        /* Error - Cannot reference y in the firstprivate clause */
        for (i=0; i<10 ; i++) {
            z[i] = i; /* O.K. - i is the loop iteration variable */
        }

        z[i] = y; /* Error - cannot reference i or y here */
    }
}
```

C/C++

Fortran

*Example 31.1f*

```
SUBROUTINE DEFAULT_NONE(A)
  INCLUDE "omp_lib.h"      ! or USE OMP_LIB

  INTEGER A
```

```

        INTEGER X, Y, Z(1000)
        COMMON/BLOCKX/X
        COMMON/BLOCKY/Y
        COMMON/BLOCKZ/Z
!$OMP THREADPRIVATE (/BLOCKX/)

        INTEGER I, J
        i = 1

!$OMP  PARALLEL DEFAULT(NONE) PRIVATE(A) SHARED(Z) PRIVATE(J)
        J = OMP_GET_NUM_THREADS();
            ! O.K. - J is listed in PRIVATE clause
        A = Z(J) ! O.K. - A is listed in PRIVATE clause
            ! - Z is listed in SHARED clause
        X = 1    ! O.K. - X is THREADPRIVATE
        Z(I) = Y ! Error - cannot reference I or Y here

!$OMP DO firstprivate(y)
    ! Error - Cannot reference y in the firstprivate clause
    DO I = 1,10
        Z(I) = I ! O.K. - I is the loop iteration variable
    END DO

        Z(I) = Y    ! Error - cannot reference I or Y here
!$OMP  END PARALLEL
        END SUBROUTINE DEFAULT_NONE

```

Fortran



## 32

## Race Conditions Caused by Implied Copies of Shared Variables in Fortran

The following example contains a race condition, because the shared variable, which is an array section, is passed as an actual argument to a routine that has an assumed-size array as its dummy argument. The subroutine call passing an array section argument may cause the compiler to copy the argument into a temporary location prior to the call and copy from the temporary location into the original variable when the subroutine returns. This copying would cause races in the **parallel** region.

### *Example 32.1f*

```
SUBROUTINE SHARED_RACE

    INCLUDE "omp_lib.h"      ! or USE OMP_LIB

    REAL A(20)
    INTEGER MYTHREAD

    !$OMP PARALLEL SHARED(A) PRIVATE(MYTHREAD)

        MYTHREAD = OMP_GET_THREAD_NUM()
        IF (MYTHREAD .EQ. 0) THEN
            CALL SUB(A(1:10)) ! compiler may introduce writes to A(6:10)
        ELSE
            A(6:10) = 12
        ENDIF

    !$OMP END PARALLEL

END SUBROUTINE SHARED_RACE

SUBROUTINE SUB(X)
    REAL X(*)
    X(1:5) = 4
END SUBROUTINE SUB
```

## 33 The private Clause

In the following example, the values of original list items *i* and *j* are retained on exit from the **parallel** region, while the private list items *i* and *j* are modified within the **parallel** construct.

Example 33.1c

```
#include <stdio.h>
#include <assert.h>

int main()
{
    int i, j;
    int *ptr_i, *ptr_j;

    i = 1;
    j = 2;

    ptr_i = &i;
    ptr_j = &j;

    #pragma omp parallel private(i) firstprivate(j)
    {
        i = 3;
        j = j + 2;
        assert (*ptr_i == 1 && *ptr_j == 2);
    }

    assert(i == 1 && j == 2);

    return 0;
}
```

C/C++

Fortran

Example 33.1f

```
PROGRAM PRIV_EXAMPLE
    INTEGER I, J

    I = 1
    J = 2

    !$OMP PARALLEL PRIVATE(I) FIRSTPRIVATE(J)
```

```

        I = 3
        J = J + 2
!$OMP   END PARALLEL

        PRINT *, I, J ! I .eq. 1 .and. J .eq. 2
END PROGRAM PRIV_EXAMPLE

```

### Fortran

In the following example, all uses of the variable *a* within the loop construct in the routine *f* refer to a private list item *a*, while it is unspecified whether references to *a* in the routine *g* are to a private list item or the original list item.

### C/C++

#### Example 33.2c

```

int a;

void g(int k) {
    a = k; /* Accessed in the region but outside of the construct;
           * therefore unspecified whether original or private list
           * item is modified. */
}

void f(int n) {
    int a = 0;

    #pragma omp parallel for private(a)
    for (int i=1; i<n; i++) {
        a = i;
        g(a*2); /* Private copy of "a" */
    }
}

```

### C/C++

### Fortran

#### Example 33.2f

```

MODULE PRIV_EXAMPLE2
  REAL A

  CONTAINS

  SUBROUTINE G(K)
    REAL K
    A = K ! Accessed in the region but outside of the
          ! construct; therefore unspecified whether
          ! original or private list item is modified.
  END SUBROUTINE G

```

```

        END SUBROUTINE G

        SUBROUTINE F(N)
        INTEGER N
        REAL A

            INTEGER I
!$OMP    PARALLEL DO PRIVATE(A)
            DO I = 1,N
                A = I
                CALL G(A*2)
            ENDDO
!$OMP    END PARALLEL DO
        END SUBROUTINE F

    END MODULE PRIV_EXAMPLE2

```

## Fortran

The following example demonstrates that a list item that appears in a **private** clause in a **parallel** construct may also appear in a **private** clause in an enclosed worksharing construct, which results in an additional private copy.

*Example 33.3c*

```

#include <assert.h>
void priv_example3()
{
    int i, a;

    #pragma omp parallel private(a)
    {
        a = 1;
        #pragma omp parallel for private(a)
        for (i=0; i<10; i++)
        {
            a = 2;
        }
        assert(a == 1);
    }
}

```

*Example 33.3f*

```

SUBROUTINE PRIV_EXAMPLE3()
    INTEGER I, A

    !$OMP PARALLEL PRIVATE(A)
        A = 1
    !$OMP PARALLEL DO PRIVATE(A)
        DO I = 1, 10
            A = 2
        END DO
    !$OMP END PARALLEL DO
        PRINT *, A ! Outer A still has value 1
    !$OMP END PARALLEL
END SUBROUTINE PRIV_EXAMPLE3

```

## 34

## Fortran Restrictions on Storage Association with the `private` Clause

The following non-conforming examples illustrate the implications of the `private` clause rules with regard to storage association.

### *Example 34.1f*

```

SUBROUTINE SUB()
COMMON /BLOCK/ X
PRINT *,X           ! X is undefined
END SUBROUTINE SUB

PROGRAM PRIV_RESTRICT
COMMON /BLOCK/ X
X = 1.0
!$OMP PARALLEL PRIVATE (X)
X = 2.0
CALL SUB()
!$OMP END PARALLEL
END PROGRAM PRIV_RESTRICT

```

### *Example 34.2f*

```

PROGRAM PRIV_RESTRICT2
COMMON /BLOCK2/ X
X = 1.0

!$OMP PARALLEL PRIVATE (X)
X = 2.0
CALL SUB()
!$OMP END PARALLEL

CONTAINS

SUBROUTINE SUB()
COMMON /BLOCK2/ Y

PRINT *,X           ! X is undefined
PRINT *,Y           ! Y is undefined
END SUBROUTINE SUB

END PROGRAM PRIV_RESTRICT2

```

*Example 34.3f*

```

PROGRAM PRIV_RESTRICT3
EQUIVALENCE (X,Y)
X = 1.0

!$OMP PARALLEL PRIVATE(X)
  PRINT *,Y                      ! Y is undefined
  Y = 10
  PRINT *,X                      ! X is undefined
!$OMP END PARALLEL
END PROGRAM PRIV_RESTRICT3

```

*Example 34.4f*

```

PROGRAM PRIV_RESTRICT4
INTEGER I, J
INTEGER A(100), B(100)
EQUIVALENCE (A(51), B(1))

!$OMP PARALLEL DO DEFAULT(PRIVATE) PRIVATE(I,J) LASTPRIVATE(A)
  DO I=1,100
    DO J=1,100
      B(J) = J - 1
    ENDDO

    DO J=1,100
      A(J) = J    ! B becomes undefined at this point
    ENDDO

    DO J=1,50
      B(J) = B(J) + 1 ! B is undefined
                      ! A becomes undefined at this point
    ENDDO
  ENDDO
!$OMP END PARALLEL DO          ! The LASTPRIVATE write for A has
                              ! undefined results

  PRINT *, B    ! B is undefined since the LASTPRIVATE
                ! write of A was not defined
END PROGRAM PRIV_RESTRICT4

```

### Example 34.5f

```
SUBROUTINE SUB1(X)
  DIMENSION X(10)

  ! This use of X does not conform to the
  ! specification. It would be legal Fortran 90,
  ! but the OpenMP private directive allows the
  ! compiler to break the sequence association that
  ! A had with the rest of the common block.

  FORALL (I = 1:10) X(I) = I
END SUBROUTINE SUB1

PROGRAM PRIV_RESTRICT5
  COMMON /BLOCK5/ A

  DIMENSION B(10)
  EQUIVALENCE (A,B(1))

  ! the common block has to be at least 10 words
  A = 0

!$OMP  PARALLEL PRIVATE(/BLOCK5/)

  ! Without the private clause,
  ! we would be passing a member of a sequence
  ! that is at least ten elements long.
  ! With the private clause, A may no longer be
  ! sequence-associated.

  CALL SUB1(A)
!$OMP  MASTER
  PRINT *, A
!$OMP  END MASTER

!$OMP  END PARALLEL
END PROGRAM PRIV_RESTRICT5
```

Fortran



## C/C++ Arrays in a **firstprivate** Clause

The following example illustrates the size and value of list items of array or pointer type in a **firstprivate** clause. The size of new list items is based on the type of the corresponding original list item, as determined by the base language.

In this example:

- The type of **A** is array of two arrays of two ints.
- The type of **B** is adjusted to pointer to array of **n** ints, because it is a function parameter.
- The type of **C** is adjusted to pointer to int, because it is a function parameter.
- The type of **D** is array of two arrays of two ints.
- The type of **E** is array of **n** arrays of **n** ints.

Note that **B** and **E** involve variable length array types.

The new items of array type are initialized as if each integer element of the original array is assigned to the corresponding element of the new array. Those of pointer type are initialized as if by assignment from the original item to the new item.

### Example 35.1c

```
#include <assert.h>

int A[2][2] = {1, 2, 3, 4};

void f(int n, int B[n][n], int C[])
{
    int D[2][2] = {1, 2, 3, 4};
    int E[n][n];

    assert(n >= 2);
    E[1][1] = 4;

    #pragma omp parallel firstprivate(B, C, D, E)
    {
        assert(sizeof(B) == sizeof(int (*)[n]));
        assert(sizeof(C) == sizeof(int*));
        assert(sizeof(D) == 4 * sizeof(int));
        assert(sizeof(E) == n * n * sizeof(int));

        /* Private B and C have values of original B and C. */
        assert(&B[1][1] == &A[1][1]);
        assert(&C[3] == &A[1][1]);
        assert(D[1][1] == 4);
        assert(E[1][1] == 4);
    }
}

int main() {
    f(2, A, A[0]);
    return 0;
}
```

C/C++

## 36

# The lastprivate Clause

Correct execution sometimes depends on the value that the last iteration of a loop assigns to a variable. Such programs must list all such variables in a **lastprivate** clause so that the values of the variables are the same as when the loop is executed sequentially.

C/C++

*Example 36.1c*

```
void lastpriv (int n, float *a, float *b)
{
    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];      /* i == n-1 here */
}
```

C/C++

Fortran

*Example 36.1f*

```
SUBROUTINE LASTPRIV(N, A, B)

    INTEGER N
    REAL A(*), B(*)
    INTEGER I
!$OMP PARALLEL
!$OMP DO LASTPRIVATE(I)

    DO I=1,N-1
        A(I) = B(I) + B(I+1)
    ENDDO

!$OMP END PARALLEL
    A(I) = B(I)      ! I has the value of N here

END SUBROUTINE LASTPRIV
```

Fortran

## 37 The reduction Clause

The following example demonstrates the **reduction** clause ; note that some reductions can be expressed in the loop in several ways, as shown for the **max** and **min** reductions below:

C/C++

*Example 37.1c*

```
#include <math.h>
void reduction1(float *x, int *y, int n)
{
    int i, b, c;
    float a, d;
    a = 0.0;
    b = 0;
    c = y[0];
    d = x[0];
    #pragma omp parallel for private(i) shared(x, y, n) \
        reduction(+:a) reduction(^:b) \
        reduction(min:c) reduction(max:d)
    for (i=0; i<n; i++) {
        a += x[i];
        b ^= y[i];
        if (c > y[i]) c = y[i];
        d = fmaxf(d,x[i]);
    }
}
```

C/C++

Fortran

*Example 37.1f*

```
SUBROUTINE REDUCTION1(A, B, C, D, X, Y, N)
    REAL :: X(*), A, D
    INTEGER :: Y(*), N, B, C
    INTEGER :: I
    A = 0
    B = 0
    C = Y(1)
    D = X(1)
    !$OMP PARALLEL DO PRIVATE(I) SHARED(X, Y, N) REDUCTION(+:A) &
    !$OMP& REDUCTION(IEOR:B) REDUCTION(MIN:C) REDUCTION(MAX:D)
    DO I=1,N
        A = A + X(I)
        B = IEOR(B, Y(I))
    
```

```

        C = MIN(C, Y(I))
        IF (D < X(I)) D = X(I)
    END DO

END SUBROUTINE REDUCTION1

```

## Fortran

A common implementation of the preceding example is to treat it as if it had been written as follows:

## C/C++

### *Example 37.2c*

```

#include <limits.h>
#include <math.h>
void reduction2(float *x, int *y, int n)
{
    int i, b, b_p, c, c_p;
    float a, a_p, d, d_p;
    a = 0.0f;
    b = 0;
    c = y[0];
    d = x[0];
    #pragma omp parallel shared(a, b, c, d, x, y, n) \
        private(a_p, b_p, c_p, d_p)
    {
        a_p = 0.0f;
        b_p = 0;
        c_p = INT_MAX;
        d_p = -HUGE_VALF;
        #pragma omp for private(i)
        for (i=0; i<n; i++) {
            a_p += x[i];
            b_p ^= y[i];
            if (c_p > y[i]) c_p = y[i];
            d_p = fmaxf(d_p, x[i]);
        }
        #pragma omp critical
        {
            a += a_p;
            b ^= b_p;
            if (c > c_p) c = c_p;
            d = fmaxf(d, d_p);
        }
    }
}

```

## C/C++

## Example 37.2f

```

SUBROUTINE REDUCTION2(A, B, C, D, X, Y, N)
  REAL :: X(*), A, D
  INTEGER :: Y(*), N, B, C
  REAL :: A_P, D_P
  INTEGER :: I, B_P, C_P
  A = 0
  B = 0
  C = Y(1)
  D = X(1)
  !$OMP PARALLEL SHARED(X, Y, A, B, C, D, N) &
  !$OMP&          PRIVATE(A_P, B_P, C_P, D_P)
    A_P = 0.0
    B_P = 0
    C_P = HUGE(C_P)
    D_P = -HUGE(D_P)
    !$OMP DO PRIVATE(I)
    DO I=1,N
      A_P = A_P + X(I)
      B_P = IEOR(B_P, Y(I))
      C_P = MIN(C_P, Y(I))
      IF (D_P < X(I)) D_P = X(I)
    END DO
    !$OMP CRITICAL
      A = A + A_P
      B = IEOR(B, B_P)
      C = MIN(C, C_P)
      D = MAX(D, D_P)
    !$OMP END CRITICAL
  !$OMP END PARALLEL
END SUBROUTINE REDUCTION2

```

The following program is non-conforming because the reduction is on the *intrinsic procedure name* **MAX** but that name has been redefined to be the variable named **MAX**.

## Fortran (cont.)

## Example 37.3f

```

PROGRAM REDUCTION_WRONG
  MAX = HUGE(0)
  M = 0

  !$OMP PARALLEL DO REDUCTION(MAX: M)
  ! MAX is no longer the intrinsic so this is non-conforming
  DO I = 1, 100
    CALL SUB(M, I)

```

```

END DO

END PROGRAM REDUCTION_WRONG

SUBROUTINE SUB(M,I)
  M = MAX(M,I)
END SUBROUTINE SUB

```

The following conforming program performs the reduction using the *intrinsic procedure name* **MAX** even though the intrinsic **MAX** has been renamed to **REN**.

### Example 37.4f

```

MODULE M
  INTRINSIC MAX
END MODULE M

PROGRAM REDUCTION3
  USE M, REN => MAX
  N = 0
  !$OMP PARALLEL DO REDUCTION(REN: N)      ! still does MAX
  DO I = 1, 100
    N = MAX(N,I)
  END DO
END PROGRAM REDUCTION3

```

The following conforming program performs the reduction using *intrinsic procedure name* **MAX** even though the intrinsic **MAX** has been renamed to **MIN**.

### Example 37.5f

```

MODULE MOD
  INTRINSIC MAX, MIN
END MODULE MOD

PROGRAM REDUCTION4
  USE MOD, MIN=>MAX, MAX=>MIN
  REAL :: R
  R = -HUGE(0.0)

  !$OMP PARALLEL DO REDUCTION(MIN: R)      ! still does MAX
  DO I = 1, 1000
    R = MIN(R, SIN(REAL(I)))
  END DO
  PRINT *, R
END PROGRAM REDUCTION4

```

The following example is non-conforming because the initialization (**a = 0**) of the original list item **a** is not synchronized with the update of **a** as a result of the reduction computation in the **for** loop. Therefore, the example may print an incorrect value for **a**.

To avoid this problem, the initialization of the original list item **a** should complete before any update of **a** as a result of the **reduction** clause. This can be achieved by adding an explicit barrier after the assignment **a = 0**, or by enclosing the assignment **a = 0** in a **single** directive (which has an implied barrier), or by initializing **a** before the start of the **parallel** region.

### Example 37.3c C/C++

```
#include <stdio.h>

int main (void)
{
    int a, i;

    #pragma omp parallel shared(a) private(i)
    {
        #pragma omp master
        a = 0;

        // To avoid race conditions, add a barrier here.

        #pragma omp for reduction(+:a)
        for (i = 0; i < 10; i++) {
            a += i;
        }

        #pragma omp single
        printf ("Sum is %d\n", a);
    }
}
```

### C/C++

### Example 37.6f Fortran

```
INTEGER A, I

!$OMP PARALLEL SHARED(A) PRIVATE(I)

!$OMP MASTER
    A = 0
```



```
!$OMP END MASTER

      ! To avoid race conditions, add a barrier here.

!$OMP DO REDUCTION(+:A)
  DO I= 0, 9
    A = A + I
  END DO

!$OMP SINGLE
  PRINT *, "Sum is ", A
!$OMP END SINGLE

!$OMP END PARALLEL
END
```

Fortran

## 38 The copyin Clause

The **copyin** clause is used to initialize threadprivate data upon entry to a **parallel** region. The value of the threadprivate variable in the master thread is copied to the threadprivate variable of each other team member.

C/C++

*Example 38.1c*

```
#include <stdlib.h>

float* work;
int size;
float tol;

#pragma omp threadprivate(work,size,tol)

void build()
{
    int i;
    work = (float*)malloc( sizeof(float)*size );
    for( i = 0; i < size; ++i ) work[i] = tol;
}

void copyin_example( float t, int n )
{
    tol = t;
    size = n;
    #pragma omp parallel copyin(tol,size)
    {
        build();
    }
}
```

C/C++

Fortran

*Example 38.1f*

```
MODULE M
  REAL, POINTER, SAVE :: WORK(:)
  INTEGER :: SIZE
  REAL :: TOL
!$OMP THREADPRIVATE(WORK,SIZE,TOL)
END MODULE M
```

```

SUBROUTINE COPYIN_EXAMPLE( T, N )
    USE M
    REAL :: T
    INTEGER :: N
    TOL = T
    SIZE = N
!$OMP    PARALLEL COPYIN(TOL,SIZE)
        CALL BUILD
!$OMP    END PARALLEL
END SUBROUTINE COPYIN_EXAMPLE

SUBROUTINE BUILD
    USE M
    ALLOCATE(WORK(SIZE))
    WORK = TOL
END SUBROUTINE BUILD

```

Fortran

## 39

# The copyprivate Clause

The **copyprivate** clause can be used to broadcast values acquired by a single thread directly to all instances of the private variables in the other threads. In this example, if the routine is called from the sequential part, its behavior is not affected by the presence of the directives. If it is called from a **parallel** region, then the actual arguments with which **a** and **b** are associated must be private.

The thread that executes the structured block associated with the **single** construct broadcasts the values of the private variables **a**, **b**, **x**, and **y** from its implicit task's data environment to the data environments of the other implicit tasks in the thread team. The broadcast completes before any of the threads have left the barrier at the end of the construct.

C/C++

*Example 39.1c*

```
#include <stdio.h>
float x, y;
#pragma omp threadprivate(x, y)

void init(float a, float b ) {
    #pragma omp single copyprivate(a,b,x,y)
    {
        scanf("%f %f %f %f", &a, &b, &x, &y);
    }
}
```

C/C++

Fortran

*Example 39.1f*

```
SUBROUTINE INIT(A,B)
  REAL A, B
  COMMON /XY/ X,Y
  !$OMP THREADPRIVATE (/XY/)

  !$OMP SINGLE
    READ (11) A,B,X,Y
  !$OMP END SINGLE COPYPRIVATE (A,B,/XY/)

  END SUBROUTINE INIT
```

Fortran

In this example, assume that the input must be performed by the master thread. Since the **master** construct does not support the **copyprivate** clause, it cannot broadcast the input value that is read. However, **copyprivate** is used to broadcast an address where the input value is stored.

## C/C++

### Example 39.2c

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

float read_next( ) {
    float * tmp;
    float return_val;

    #pragma omp single copyprivate(tmp)
    {
        tmp = (float *) malloc(sizeof(float));
    } /* copies the pointer only */

    #pragma omp master
    {
        scanf("%f", tmp);
    }

    #pragma omp barrier
    return_val = *tmp;
    #pragma omp barrier

    #pragma omp single nowait
    {
        free(tmp);
    }

    return return_val;
}
```

## C/C++

## Fortran

### Example 39.2f

```
REAL FUNCTION READ_NEXT()
REAL, POINTER :: TMP

!$OMP SINGLE
    ALLOCATE (TMP)
!$OMP END SINGLE COPYPRIVATE (TMP) ! copies the pointer only
```

```

!$OMP  MASTER
      READ (11) TMP
!$OMP  END MASTER

!$OMP  BARRIER
      READ_NEXT = TMP
!$OMP  BARRIER

!$OMP  SINGLE
      DEALLOCATE (TMP)
!$OMP  END SINGLE NOWAIT
      END FUNCTION READ_NEXT

```

### Fortran

Suppose that the number of lock variables required within a **parallel** region cannot easily be determined prior to entering it. The **copyprivate** clause can be used to provide access to shared lock variables that are allocated within that **parallel** region.

### C/C++

#### Example 39.3c

```

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

omp_lock_t *new_lock()
{
    omp_lock_t *lock_ptr;

    #pragma omp single copyprivate(lock_ptr)
    {
        lock_ptr = (omp_lock_t *) malloc(sizeof(omp_lock_t));
        omp_init_lock( lock_ptr );
    }

    return lock_ptr;
}

```

### C/C++

### Fortran

#### Example 39.3f

```

      FUNCTION NEW_LOCK()
      USE OMP_LIB      ! or INCLUDE "omp_lib.h"
      INTEGER(OMP_LOCK_KIND), POINTER :: NEW_LOCK

!$OMP  SINGLE

```

```

        ALLOCATE (NEW_LOCK)
        CALL OMP_INIT_LOCK (NEW_LOCK)
!$OMP   END SINGLE COPYPRIVATE (NEW_LOCK)
      END FUNCTION NEW_LOCK

```

Note that the effect of the **copyprivate** clause on a variable with the **allocatable** attribute is different than on a variable with the **pointer** attribute. The value of **A** is copied (as if by intrinsic assignment) and the pointer **B** is copied (as if by pointer assignment) to the corresponding list items in the other implicit tasks belonging to the **parallel** region.

### *Example 39.4f*

```

SUBROUTINE S(N)
  INTEGER N

  REAL, DIMENSION(:), ALLOCATABLE :: A
  REAL, DIMENSION(:), POINTER :: B

  ALLOCATE (A(N))
!$OMP   SINGLE
      ALLOCATE (B(N))
      READ (11) A,B
!$OMP   END SINGLE COPYPRIVATE(A,B)
      ! Variable A is private and is
      ! assigned the same value in each thread
      ! Variable B is shared

!$OMP   BARRIER
!$OMP   SINGLE
      DEALLOCATE (B)
!$OMP   END SINGLE NOWAIT
END SUBROUTINE S

```

Fortran

## 40 Nested Loop Constructs

The following example of loop construct nesting is conforming because the inner and outer loop regions bind to different **parallel** regions:

▼ C/C++ ▼  
*Example 40.1c*

```
void work(int i, int j) {}

void good_nesting(int n)
{
    int i, j;
    #pragma omp parallel default(shared)
    {
        #pragma omp for
        for (i=0; i<n; i++) {
            #pragma omp parallel shared(i, n)
            {
                #pragma omp for
                for (j=0; j < n; j++)
                    work(i, j);
            }
        }
    }
}
```

▲ C/C++ ▲

▼ Fortran ▼  
*Example 40.1f*

```
SUBROUTINE WORK(I, J)
  INTEGER I, J
END SUBROUTINE WORK

SUBROUTINE GOOD_NESTING(N)
  INTEGER N

  INTEGER I
  !$OMP PARALLEL DEFAULT(SHARED)
  !$OMP DO
    DO I = 1, N
      !$OMP PARALLEL SHARED(I,N)
      !$OMP DO
        DO J = 1, N
          CALL WORK(I,J)
        
```



```
                END DO
!$OMP          END PARALLEL
                END DO
!$OMP          END PARALLEL
END SUBROUTINE GOOD_NESTING
```

Fortran

The following variation of the preceding example is also conforming:

#### C/C++

##### *Example 40.2c*

```
void work(int i, int j) {}

void work1(int i, int n)
{
    int j;
    #pragma omp parallel default(shared)
    {
        #pragma omp for
        for (j=0; j<n; j++)
            work(i, j);
    }
}

void good_nesting2(int n)
{
    int i;
    #pragma omp parallel default(shared)
    {
        #pragma omp for
        for (i=0; i<n; i++)
            work1(i, n);
    }
}
```

#### C/C++

#### Fortran

##### *Example 40.2f*

```
      SUBROUTINE WORK(I, J)
      INTEGER I, J
      END SUBROUTINE WORK

      SUBROUTINE WORK1(I, N)
      INTEGER J
      !$OMP PARALLEL DEFAULT(SHARED)
      !$OMP DO
          DO J = 1, N
              CALL WORK(I,J)
          END DO
      !$OMP END PARALLEL
      END SUBROUTINE WORK1
```

```
        SUBROUTINE GOOD_NESTING2(N)
        INTEGER N
!$OMP PARALLEL DEFAULT(SHARED)
!$OMP DO
        DO I = 1, N
            CALL WORK1(I, N)
        END DO
!$OMP END PARALLEL
        END SUBROUTINE GOOD_NESTING2
```

Fortran

# 41

## Restrictions on Nesting of Regions

The examples in this section illustrate the region nesting rules.

The following example is non-conforming because the inner and outer loop regions are closely nested:

C/C++

*Example 41.1c*

```
void work(int i, int j) {}
void wrong1(int n)
{
    #pragma omp parallel default(shared)
    {
        int i, j;
        #pragma omp for
        for (i=0; i<n; i++) {
            /* incorrect nesting of loop regions */
            #pragma omp for
            for (j=0; j<n; j++)
                work(i, j);
        }
    }
}
```

C/C++

Fortran

*Example 41.1f*

```
SUBROUTINE WORK(I, J)
  INTEGER I, J
END SUBROUTINE WORK
SUBROUTINE WRONG1(N)
  INTEGER N
  INTEGER I,J
  !$OMP PARALLEL DEFAULT(SHARED)
  !$OMP DO
    DO I = 1, N
      !$OMP DO ! incorrect nesting of loop regions
        DO J = 1, N
          CALL WORK(I,J)
        END DO
      END DO
    END DO
  !$OMP END PARALLEL
```

END SUBROUTINE WRONG1

Fortran

The following orphaned version of the preceding example is also non-conforming:

C/C++

*Example 41.2c*

```
void work(int i, int j) {}
void work1(int i, int n)
{
    int j;
    /* incorrect nesting of loop regions */
    #pragma omp for
        for (j=0; j<n; j++)
            work(i, j);
}

void wrong2(int n)
{
    #pragma omp parallel default(shared)
    {
        int i;
        #pragma omp for
            for (i=0; i<n; i++)
                work1(i, n);
    }
}
```

C/C++

Fortran

*Example 41.2f*

```
SUBROUTINE WORK1(I,N)
  INTEGER I, N
  INTEGER J
!$OMP DO ! incorrect nesting of loop regions
  DO J = 1, N
    CALL WORK(I,J)
  END DO
END SUBROUTINE WORK1
SUBROUTINE WRONG2(N)
  INTEGER N
  INTEGER I
!$OMP PARALLEL DEFAULT(SHARED)
!$OMP DO
  DO I = 1, N
    CALL WORK1(I,N)
  END DO
!$OMP END PARALLEL
```

```
END SUBROUTINE WRONG2
```

## Fortran

The following example is non-conforming because the loop and **single** regions are closely nested:

## C/C++

### Example 41.3c

```
void work(int i, int j) {}
void wrong3(int n)
{
    #pragma omp parallel default(shared)
    {
        int i;
        #pragma omp for
        for (i=0; i<n; i++) {
/* incorrect nesting of regions */
            #pragma omp single
            work(i, 0);
        }
    }
}
```

## C/C++

## Fortran

### Example 41.3f

```
SUBROUTINE WRONG3 (N)
  INTEGER N

  INTEGER I
!$OMP PARALLEL DEFAULT (SHARED)
!$OMP DO
  DO I = 1, N
!$OMP SINGLE ! incorrect nesting of regions
    CALL WORK(I, 1)
!$OMP END SINGLE
  END DO
!$OMP END PARALLEL
END SUBROUTINE WRONG3
```

## Fortran

The following example is non-conforming because a **barrier** region cannot be closely nested inside a loop region:

*Example 41.4c*

```

void work(int i, int j) {}
void wrong4(int n)
{
    #pragma omp parallel default(shared)
    {
        int i;
        #pragma omp for
        for (i=0; i<n; i++) {
            work(i, 0);
        }
        /* incorrect nesting of barrier region in a loop region */
        #pragma omp barrier
        work(i, 1);
    }
}

```

*Example 41.4f*

```

SUBROUTINE WRONG4(N)
  INTEGER N

  INTEGER I
  !$OMP PARALLEL DEFAULT(SHARED)
  !$OMP DO
    DO I = 1, N
      CALL WORK(I, 1)
    ! incorrect nesting of barrier region in a loop region
  !$OMP BARRIER
    CALL WORK(I, 2)
    END DO
  !$OMP END PARALLEL
END SUBROUTINE WRONG4

```

The following example is non-conforming because the **barrier** region cannot be closely nested inside the **critical** region. If this were permitted, it would result in deadlock due to the fact that only one thread at a time can enter the **critical** region:

*Example 41.5c*

```

void work(int i, int j) {}
void wrong5(int n)
{
    #pragma omp parallel
    {
        #pragma omp critical
        {
            work(n, 0);
        }
        /* incorrect nesting of barrier region in a critical region */
        #pragma omp barrier
        work(n, 1);
    }
}

```

*Example 41.5f*

```

      SUBROUTINE WRONG5(N)
      INTEGER N

      !$OMP  PARALLEL DEFAULT(SHARED)
      !$OMP  CRITICAL
            CALL WORK(N,1)
      ! incorrect nesting of barrier region in a critical region
      !$OMP  BARRIER
            CALL WORK(N,2)
      !$OMP  END CRITICAL
      !$OMP  END PARALLEL
      END SUBROUTINE WRONG5

```

The following example is non-conforming because the **barrier** region cannot be closely nested inside the **single** region. If this were permitted, it would result in deadlock due to the fact that only one thread executes the **single** region:

*Example 41.6c*

```

void work(int i, int j) {}
void wrong6(int n)
{
    #pragma omp parallel

```



```

{
    #pragma omp single
    {
        work(n, 0);
    /* incorrect nesting of barrier region in a single region */
        #pragma omp barrier
        work(n, 1);
    }
}

```

C/C++

Fortran

### *Example 41.6f*

```

SUBROUTINE WRONG6 (N)
  INTEGER N

!$OMP  PARALLEL DEFAULT(SHARED)
!$OMP  SINGLE
      CALL WORK(N,1)
! incorrect nesting of barrier region in a single region
!$OMP  BARRIER
      CALL WORK(N,2)
!$OMP  END SINGLE
!$OMP  END PARALLEL
END SUBROUTINE WRONG6

```

Fortran

## The `omp_set_dynamic` and `omp_set_num_threads` Routines

Some programs rely on a fixed, prespecified number of threads to execute correctly. Because the default setting for the dynamic adjustment of the number of threads is implementation defined, such programs can choose to turn off the dynamic threads capability and set the number of threads explicitly to ensure portability. The following example shows how to do this using `omp_set_dynamic`, and `omp_set_num_threads`.

In this example, the program executes correctly only if it is executed by 16 threads. If the implementation is not capable of supporting 16 threads, the behavior of this example is implementation defined. Note that the number of threads executing a **parallel** region remains constant during the region, regardless of the dynamic threads setting. The dynamic threads mechanism determines the number of threads to use at the start of the **parallel** region and keeps it constant for the duration of the region.

C/C++

*Example 42.1c*

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

void do_by_16(float *x, int iam, int ipoints) {}

void dynthreads(float *x, int npoints)
{
    int iam, ipoints;

    omp_set_dynamic(0);
    omp_set_num_threads(16);

    #pragma omp parallel shared(x, npoints) private(iam, ipoints)
    {
        if (omp_get_num_threads() != 16)
            abort();

        iam = omp_get_thread_num();
        ipoints = npoints/16;
        do_by_16(x, iam, ipoints);
    }
}
```

C/C++

*Example 42.1f*

```

SUBROUTINE DO_BY_16(X, IAM, IPOINTS)
  REAL X(*)
  INTEGER IAM, IPOINTS
END SUBROUTINE DO_BY_16

SUBROUTINE DYNTHREADS(X, NPOINTS)

  INCLUDE "omp_lib.h"      ! or USE OMP_LIB

  INTEGER NPOINTS
  REAL X(NPOINTS)

  INTEGER IAM, IPOINTS

  CALL OMP_SET_DYNAMIC(.FALSE.)
  CALL OMP_SET_NUM_THREADS(16)

!$OMP  PARALLEL SHARED(X,NPOINTS) PRIVATE(IAM, IPOINTS)

  IF (OMP_GET_NUM_THREADS() .NE. 16) THEN
    STOP
  ENDIF

  IAM = OMP_GET_THREAD_NUM()
  IPOINTS = NPOINTS/16
  CALL DO_BY_16(X,IAM,IPOINTS)

!$OMP  END PARALLEL

END SUBROUTINE DYNTHREADS

```

## 43

# The `omp_get_num_threads` Routine

In the following example, the `omp_get_num_threads` call returns 1 in the sequential part of the code, so `np` will always be equal to 1. To determine the number of threads that will be deployed for the `parallel` region, the call should be inside the `parallel` region.

Example 43.1c

```
#include <omp.h>
void work(int i);

void incorrect()
{
    int np, i;

    np = omp_get_num_threads(); /* misplaced */

    #pragma omp parallel for schedule(static)
    for (i=0; i < np; i++)
        work(i);
}
```

C/C++

Example 43.1f

```
SUBROUTINE WORK(I)
  INTEGER I
  I = I + 1
END SUBROUTINE WORK

SUBROUTINE INCORRECT()
  INCLUDE "omp_lib.h"      ! or USE OMP_LIB
  INTEGER I, NP

  NP = OMP_GET_NUM_THREADS() !misplaced: will return 1
!$OMP PARALLEL DO SCHEDULE(STATIC)
  DO I = 0, NP-1
    CALL WORK(I)
  ENDDO
!$OMP END PARALLEL DO
END SUBROUTINE INCORRECT
```

Fortran

The following example shows how to rewrite this program without including a query for the number of threads:

C/C++

*Example 43.2c*

```
#include <omp.h>
void work(int i);

void correct()
{
    int i;

    #pragma omp parallel private(i)
    {
        i = omp_get_thread_num();
        work(i);
    }
}
```

C/C++

Fortran

*Example 43.2f*

```
SUBROUTINE WORK(I)
    INTEGER I

    I = I + 1

END SUBROUTINE WORK

SUBROUTINE CORRECT()
    INCLUDE "omp_lib.h"      ! or USE OMP_LIB
    INTEGER I

!$OMP    PARALLEL PRIVATE(I)
        I = OMP_GET_THREAD_NUM()
        CALL WORK(I)
!$OMP    END PARALLEL

END SUBROUTINE CORRECT
```

Fortran

## The `omp_init_lock` Routine

The following example demonstrates how to initialize an array of locks in a **parallel** region by using `omp_init_lock`.

C/C++

*Example 44.1c*

```
#include <omp.h>

omp_lock_t *new_locks()
{
    int i;
    omp_lock_t *lock = new omp_lock_t[1000];

    #pragma omp parallel for private(i)
    for (i=0; i<1000; i++)
    {
        omp_init_lock(&lock[i]);
    }
    return lock;
}
```

C/C++

Fortran

*Example 44.1f*

```
FUNCTION NEW_LOCKS()
    USE OMP_LIB          ! or INCLUDE "omp_lib.h"
    INTEGER(OMP_LOCK_KIND), DIMENSION(1000) :: NEW_LOCKS

    INTEGER I

    !$OMP PARALLEL DO PRIVATE(I)
        DO I=1,1000
            CALL OMP_INIT_LOCK(NEW_LOCKS(I))
        END DO
    !$OMP END PARALLEL DO

    END FUNCTION NEW_LOCKS
```

Fortran

---

## 45 Ownership of Locks

Ownership of locks has changed since OpenMP 2.5. In OpenMP 2.5, locks are owned by threads; so a lock released by the `omp_unset_lock` routine must be owned by the same thread executing the routine. With OpenMP 3.0, locks are owned by task regions; so a lock released by the `omp_unset_lock` routine in a task region must be owned by the same task region.

This change in ownership requires extra care when using locks. The following program is conforming in OpenMP 2.5 because the thread that releases the lock `lck` in the parallel region is the same thread that acquired the lock in the sequential part of the program (master thread of parallel region and the initial thread are the same). However, it is not conforming in OpenMP 3.0 and 3.1, because the task region that releases the lock `lck` is different from the task region that acquires the lock.

C/C++

*Example 45.1c*

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

int main()
{
    int x;
    omp_lock_t lck;

    omp_init_lock (&lck);
    omp_set_lock (&lck);
    x = 0;

#pragma omp parallel shared (x)
    {
        #pragma omp master
        {
            x = x + 1;
            omp_unset_lock (&lck);
        }

        /* Some more stuff. */
    }
    omp_destroy_lock (&lck);
    return 0;
}
```

C/C++

*Example 45.1f*

```
program lock
  use omp_lib
  integer :: x
  integer (kind=omp_lock_kind) :: lck

  call omp_init_lock (lck)
  call omp_set_lock(lck)
  x = 0

!$omp parallel shared (x)
!$omp master
  x = x + 1
  call omp_unset_lock(lck)
!$omp end master

!      Some more stuff.
!$omp end parallel

  call omp_destroy_lock(lck)
end
```



## Simple Lock Routines

In the following example, the lock routines cause the threads to be idle while waiting for entry to the first critical section, but to do other work while waiting for entry to the second. The `omp_set_lock` function blocks, but the `omp_test_lock` function does not, allowing the work in `skip` to be done.

C/C++

Note that the argument to the lock routines should have type `omp_lock_t`, and that there is no need to flush it.

### *Example 46.1c*

```
#include <stdio.h>
#include <omp.h>
void skip(int i) {}
void work(int i) {}
int main()
{
    omp_lock_t lck;
    int id;
    omp_init_lock(&lck);

    #pragma omp parallel shared(lck) private(id)
    {
        id = omp_get_thread_num();

        omp_set_lock(&lck);
        /* only one thread at a time can execute this printf */
        printf("My thread id is %d.\n", id);
        omp_unset_lock(&lck);

        while (! omp_test_lock(&lck)) {
            skip(id); /* we do not yet have the lock,
                       so we must do something else */
        }

        work(id); /* we now have the lock
                   and can do the work */

        omp_unset_lock(&lck);
    }
    omp_destroy_lock(&lck);

    return 0;
}
```

C/C++

Note that there is no need to flush the lock variable.

*Example 46.1f*

```

SUBROUTINE SKIP(ID)
END SUBROUTINE SKIP

SUBROUTINE WORK(ID)
END SUBROUTINE WORK

PROGRAM SIMPLELOCK

    INCLUDE "omp_lib.h"      ! or USE OMP_LIB

    INTEGER(OMP_LOCK_KIND) LCK
    INTEGER ID

    CALL OMP_INIT_LOCK(LCK)

!$OMP PARALLEL SHARED(LCK) PRIVATE(ID)
    ID = OMP_GET_THREAD_NUM()
    CALL OMP_SET_LOCK(LCK)
    PRINT *, 'My thread id is ', ID
    CALL OMP_UNSET_LOCK(LCK)

    DO WHILE (.NOT. OMP_TEST_LOCK(LCK))
        CALL SKIP(ID)      ! We do not yet have the lock
                           ! so we must do something else
    END DO

    CALL WORK(ID)          ! We now have the lock
                           ! and can do the work

    CALL OMP_UNSET_LOCK( LCK )

!$OMP END PARALLEL

    CALL OMP_DESTROY_LOCK( LCK )

END PROGRAM SIMPLELOCK

```

## 47

# Nestable Lock Routines

The following example demonstrates how a nestable lock can be used to synchronize updates both to a whole structure and to one of its members.

C/C++

*Example 47.1c*

```
#include <omp.h>
typedef struct {
    int a,b;
    omp_nest_lock_t lck; } pair;

int work1();
int work2();
int work3();
void incr_a(pair *p, int a)
{
    /* Called only from incr_pair, no need to lock. */
    p->a += a;
}
void incr_b(pair *p, int b)
{
    /* Called both from incr_pair and elsewhere, */
    /* so need a nestable lock. */

    omp_set_nest_lock(&p->lck);
    p->b += b;
    omp_unset_nest_lock(&p->lck);
}
void incr_pair(pair *p, int a, int b)
{
    omp_set_nest_lock(&p->lck);
    incr_a(p, a);
    incr_b(p, b);
    omp_unset_nest_lock(&p->lck);
}
void nestlock(pair *p)
{
    #pragma omp parallel sections
    {
        #pragma omp section
        incr_pair(p, work1(), work2());
        #pragma omp section
        incr_b(p, work3());
    }
}
```

C/C++

*Example 47.1f*

```

MODULE DATA
  USE OMP_LIB, ONLY: OMP_NEST_LOCK_KIND
  TYPE LOCKED_PAIR
    INTEGER A
    INTEGER B
    INTEGER (OMP_NEST_LOCK_KIND) LCK
  END TYPE
END MODULE DATA

SUBROUTINE INCR_A(P, A)
  ! called only from INCR_PAIR, no need to lock
  USE DATA
  TYPE(LOCKED_PAIR) :: P
  INTEGER A
  P%A = P%A + A
END SUBROUTINE INCR_A

SUBROUTINE INCR_B(P, B)
  ! called from both INCR_PAIR and elsewhere,
  ! so we need a nestable lock
  USE OMP_LIB      ! or INCLUDE "omp_lib.h"
  USE DATA
  TYPE(LOCKED_PAIR) :: P
  INTEGER B
  CALL OMP_SET_NEST_LOCK(P%LCK)
  P%B = P%B + B
  CALL OMP_UNSET_NEST_LOCK(P%LCK)
END SUBROUTINE INCR_B

SUBROUTINE INCR_PAIR(P, A, B)
  USE OMP_LIB      ! or INCLUDE "omp_lib.h"
  USE DATA
  TYPE(LOCKED_PAIR) :: P
  INTEGER A
  INTEGER B

  CALL OMP_SET_NEST_LOCK(P%LCK)
  CALL INCR_A(P, A)
  CALL INCR_B(P, B)
  CALL OMP_UNSET_NEST_LOCK(P%LCK)
END SUBROUTINE INCR_PAIR

SUBROUTINE NESTLOCK(P)
  USE OMP_LIB      ! or INCLUDE "omp_lib.h"
  USE DATA
  TYPE(LOCKED_PAIR) :: P
  INTEGER WORK1, WORK2, WORK3
  EXTERNAL WORK1, WORK2, WORK3

```

```
!$OMP  PARALLEL SECTIONS

!$OMP  SECTION
      CALL INCR_PAIR(P, WORK1(), WORK2())
!$OMP  SECTION
      CALL INCR_B(P, WORK3())
!$OMP  END PARALLEL SECTIONS

      END SUBROUTINE NESTLOCK
```

Fortran

## 48 target Construct

### target Construct on parallel Construct

This following example shows how the **target** construct offloads a code region to a target device. The variables `p`, `v1`, `v2`, and `N` are implicitly mapped to the the target device.

Example 48.1c

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

C/C++

Example 48.1f

```
subroutine vec_mult(N)
    integer :: i,N
    real    :: p(N), v1(N), v2(N)
    call init(v1, v2, N)
    !$omp target
    !$omp parallel do
    do i=1,N
        p(i) = v1(i) * v2(i)
    end do
    !$omp end target
    call output(p, N)
end subroutine
```

Fortran

## target Construct with map Clause

This following example shows how the **target** construct offloads a code region to a target device. The variables `p`, `v1`, `v2`, are explicitly mapped to the the target device using the map clause. The variable `N` is implicitly mapped to the target device.

C/C++

### *Example 48.2c*

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

C/C++

Fortran

### *Example 48.2f*

```
subroutine vec_mult(N)
    integer :: i,N
    real    :: p(N), v1(N), v2(N)
    call init(v1, v2, N)
    !$omp target map(v1,v2,p)
    !$omp parallel do
    do i=1,N
        p(i) = v1(i) * v2(i)
    end do
    !$omp end target
    call output(p, N)
end subroutine
```

Fortran

## map Clause With to/from map-types

The following example shows how the **target** construct offloads a code region to a target device. In the **map** clause, the **to** and **from** map-types define the mapping between the original (host) data and the target (device) data. The **to** map-type specifies that the data will only be read on the device, and the **from** map-type specifies that the data will only be written to on the device. By specifying a guaranteed access on the device, data transfers can be reduced for the **target** region.

The **to** map-type indicates that at the start of the **target** region the variables `v1` and `v2` are initialized with the values of the corresponding variables on the host device, and at the end of the **target** region the variables `v1` and `v2` are not assigned to their corresponding variables on the host device.

The **from** map-type indicates that at the start of the **target** region the variable `p` is not initialized with the value of the corresponding variable on the host device, and at the end of the **target** region the variable `p` is assigned to the corresponding variable on the host device.

### Example 48.3c

```
extern void init(float*, float*, int);
extern void output(float*, int);
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);
}
```

### Example 48.3f

The **to** and **from** map-types allow programmers to optimize data motion. Since data for the `v` arrays are not returned, and data for the `p` array are not transferred to the device, only one-half of the data is moved, compared to the default behavior of an implicit mapping.



```

subroutine vec_mult(N)
  integer :: i,N
  real    :: p(N), v1(N), v2(N)
  call init(v1, v2, N)
  !$omp target map(to: v1,v2) map(from: p)
  !$omp parallel do
    do i=1,N
      p(i) = v1(i) * v2(i)
    end do
  !$omp end target
  call output(p, N)
end subroutine

```

Fortran

## map Clause with Array Sections

The following example shows how the **target** construct offloads a code region to a target device. In the **map** clause, map-types are used to optimize the mapping of variables to the target device. Because variables **p**, **v1** and **v2** are pointers, array section notation must be used to map the arrays. The notation **:N** is equivalent to **0:N**.

C/C++

*Example 48.4c*

```

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

```

C/C++

*Example 48.4f*

In C, the length of the pointed-to array must be specified. In Fortran the extent of the array is known and the length need not be specified. A section of the array can be specified with the usual Fortran syntax, as shown in the following example. The value 1 is assumed for the lower bound for array section `v2 (:N)`.

```
module mults
contains
subroutine vec_mult(p,v1,v2,N)
  real,pointer,dimension(:) :: p, v1, v2
  integer                    :: N,i
  call init(v1, v2, N)
  !$omp target map(to: v1(1:N), v2(:N)) map(from: p(1:N))
  !$omp parallel do
  do i=1,N
    p(i) = v1(i) * v2(i)
  end do
  !$omp end target
  call output(p, N)
end subroutine
end module
```

A more realistic situation in which an assumed-size array is passed to `vec_mult` requires that the length of the arrays be specified, because the compiler does not know the size of the storage. A section of the array must be specified with the usual Fortran syntax, as shown in the following example. The value 1 is assumed for the lower bound for array section `v2 (:N)`.

```
module mults
contains
subroutine vec_mult(p,v1,v2,N)
  real,dimension(*) :: p, v1, v2
  integer            :: N,i
  call init(v1, v2, N)
  !$omp target map(to: v1(1:N), v2(:N)) map(from: p(1:N))
  !$omp parallel do
  do i=1,N
    p(i) = v1(i) * v2(i)
  end do
  call output(p, N)
  !$omp end target
end subroutine
end module
```

## target Construct with if Clause

The following example shows how the **target** construct offloads a code region to a target device.

The **if** clause on the **target** construct indicates that if the variable *N* is smaller than a given threshold, then the **target** region will be executed by the host device.

The **if** clause on the **parallel** construct indicates that if the variable *N* is smaller than a second threshold then the **parallel** region is inactive.

C/C++

### *Example 48.5c*

```
#define THRESHOLD1 1000000
#define THRESHOLD2 1000
extern void init(float*, float*, int);
extern void output(float*, int);
void vec_mult(float *p, float *v1, float *v2, int N)
{
    int i;
    init(v1, v2, N);
    #pragma omp target if(N>THRESHOLD1) map(to: v1[0:N], v2[:N])\
        map(from: p[0:N])
    #pragma omp parallel for if(N>THRESHOLD2)
    for (i=0; i<N; i++)
        p[i] = v1[i] * v2[i];
    output(p, N);
}
```

C/C++

Fortran

### *Example 48.5f*

```
module params
integer,parameter :: THRESHOLD1=1000000, THRESHHOLD2=1000
end module
subroutine vec_mult(p, v1, v2, N)
    use params
    real :: p(N), v1(N), v2(N)
    integer :: i
    call init(v1, v2, N)
    !$omp target if(N>THRESHHOLD1) map(to: v1, v2 ) map(from: p)
        !$omp parallel do if(N>THRESHOLD2)
            do i=1,N
                p(i) = v1(i) * v2(i)
            end do
        end do
    !$omp end target
end subroutine
```

```
call output(p, N)
end subroutine
```

Fortran

## Simple target data Construct

This example shows how the **target data** construct maps variables to a device data environment. The **target data** construct creates a new device data environment and maps the variables `v1`, `v2`, and `p` to the new device data environment. The **target** construct enclosed in the **target data** region creates a new device data environment, which inherits the variables `v1`, `v2`, and `p` from the enclosing device data environment. The variable `N` is mapped into the new device data environment from the encountering task's data environment.

C/C++

*Example 49.1c*

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

C/C++

Fortran

*Example 49.1f*

The Fortran code passes a reference and specifies the extent of the arrays in the declaration. No length information is necessary in the map clause, as is required with C/C++ pointers.

```

subroutine vec_mult(p, v1, v2, N)
  real    :: p(N), v1(N), v2(N)
  integer :: i
  call init(v1, v2, N)
  !$omp target data map(to: v1, v2) map(from: p)
  !$omp target
  !$omp parallel do
    do i=1,N
      p(i) = v1(i) * v2(i)
    end do
  !$omp end target
  !$omp end target data
  call output(p, N)
end subroutine

```

Fortran

## target data Region Enclosing Multiple target Regions

The following examples show how the **target data** construct maps variables to a device data environment of a **target** region. The **target data** construct creates a device data environment and encloses **target** regions, which have their own device data environments. The device data environment of the **target data** region is inherited by the device data environment of an enclosed **target** region. The **target data** construct is used to create variables that will persist throughout the **target data** region.

In the following example the variables `v1` and `v2` are mapped at each **target** construct. Instead of mapping the variable `p` twice, once at each **target** construct, `p` is mapped once by the **target data** construct.

C/C++

### Example 49.2c

```

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

```

```

        p[i] = v1[i] * v2[i];
    init_again(v1, v2, N);
    #pragma omp target map(to: v1[:N], v2[:N])
    #pragma omp parallel for
    for (i=0; i<N; i++)
        p[i] = p[i] + (v1[i] * v2[i]);
    }
    output(p, N);
}

```

C/C++

Fortran

### Example 49.2f

The Fortran code uses reference and specifies the extent of the p, v1 and v2 arrays. No length information is necessary in the **map** clause, as is required with C/C++ pointers. The arrays v1 and v2 are mapped at each **target** construct. Instead of mapping the array p twice, once at each target construct, p is mapped once by the **target data** construct.

```

subroutine vec_mult(p, v1, v2, N)
    real    :: p(N), v1(N), v2(N)
    integer :: i
    call init(v1, v2, N)
    !$omp target data map(from: p)
    !$omp target map(to: v1, v2 )
    !$omp parallel do
    do i=1,N
        p(i) = v1(i) * v2(i)
    end do
    !$omp end target
    call init_again(v1, v2, N)
    !$omp target map(to: v1, v2 )
    !$omp parallel do
    do i=1,N
        p(i) = p(i) + v1(i) * v2(i)
    end do
    !$omp end target
    !$omp end target data
    call output(p, N)
end subroutine

```

Fortran

C/C++

In the following example, the variable tmp defaults to **tofrom** map-type and is mapped at each **target** construct. The array Q is mapped once at the enclosing **target data** region instead of at each **target** construct.

```

#include <math.h>
void gramSchmidt(restrict float Q[][COLS], const int rows, const int cols)
{
    #pragma omp target data map(Q[0:rows][0:cols])
    for(int k=0; k < cols; k++)
    {
        double tmp = 0.0;
        #pragma omp target
        #pragma omp parallel for reduction(+:tmp)
        for(int i=0; i < rows; i++)
            tmp += (Q[i][k] * Q[i][k]);
        tmp = 1/sqrt(tmp);
        #pragma omp target
        #pragma omp parallel for
        for(int i=0; i < rows; i++)
            Q[i][k] *= tmp;
    }
}

```

▲ C/C++ ▲

▼ Fortran ▼

### *Example 49.3f*

In the following example the arrays `v1` and `v2` are mapped at each **target** construct. Instead of mapping the array `Q` twice at each **target** construct, `Q` is mapped once by the **target data** construct. Note, the `tmp` variable is implicitly remapped for each **target** region, mapping the value from the device to the host at the end of the first **target** region, and from the host to the device for the second **target** region.

```

subroutine gramSchmidt(Q,rows,cols)
integer          :: rows,cols, i,k
double precision :: Q(rows,cols), tmp
!$omp target data map(Q)
do k=1,cols
    tmp = 0.0d0
    !$omp target
        !$omp parallel do reduction(+:tmp)
        do i=1,rows
            tmp = tmp + (Q(i,k) * Q(i,k))
        end do
    !$omp end target
    tmp = 1.0d0/sqrt(tmp)
    !$omp target
        !$omp parallel do
        do i=1,rows
            Q(i,k) = Q(i,k)*tmp
        enddo
    !$omp end target

```



```

        end do
        !$omp end target data
    end subroutine

```

## Fortran

# target data Construct with Orphaned Call

The following two examples show how the **target data** construct maps variables to a device data environment. The **target data** construct's device data environment encloses the **target** construct's device data environment in the function `vec_mult()`.

When the type of the variable appearing in an array section is pointer, the pointer variable and the storage location of the corresponding array section are mapped to the device data environment. The pointer variable is treated as if it had appeared in a **map** clause with a map-type of **alloc**. The array section's storage location is mapped according to the map-type in the **map** clause (the default map-type is **tofrom**).

The **target** construct's device data environment inherits the storage locations of the array sections `v1[0:N]`, `v2[:n]`, and `p0[0:N]` from the enclosing target data construct's device data environment. Neither initialization nor assignment is performed for the array sections in the new device data environment.

The pointer variables `p1`, `v3`, and `v4` are mapped into the target construct's device data environment with an implicit map-type of **alloc** and they are assigned the address of the storage location associated with their corresponding array sections. Note that the following pairs of array section storage locations are equivalent (`p0[:N]`, `p1[:N]`), (`v1[:N]`, `v3[:N]`), and (`v2[:N]`, `v4[:N]`).

## C/C++

### Example 49.3c

```

void vec_mult(float*, float*, float*, int);
extern void init(float*, float*, int);
extern void output(float*, int);
void foo(float *p0, float *v1, float *v2, int N)
{
    init(v1, v2, N);
    #pragma omp target data map(to: v1[0:N], v2[:N]) map(from: p0[0:N])
    {
        vec_mult(p0, v1, v2, N);
    }
    output(p0, N);
}
void vec_mult(float *p1, float *v3, float *v4, int N)
{

```

```

    #pragma omp target map(to: v3[0:N], v4[:N]) map(from: p1[0:N])
    #pragma omp parallel for
    for (i=0; i<N; i++)
        p1[i] = v3[i] * v4[i];
}

```

C/C++

Fortran

#### Example 49.4f

The Fortran code maps the pointers and storage in an identical manner (same extent, but uses indices from 1 to N).

The **target** construct's device data environment inherits the storage locations of the arrays **v1**, **v2** and **p0** from the enclosing **target data** constructs's device data environment. However, in Fortran the associated data of the pointer is known, and the shape is not required.

The pointer variables **p1**, **v3**, and **v4** are mapped into the **target** construct's device data environment with an implicit map-type of **alloc** and they are assigned the address of the storage location associated with their corresponding array sections. Note that the following pair of array storage locations are equivalent (**p0**, **p1**), (**v1**, **v3**), and (**v2**, **v4**).

```

module mults
contains
subroutine foo(p0,v1,v2,N)
real,pointer,dimension(:) :: p0, v1, v2
integer :: N,i
    call init(v1, v2, N)
    !$omp target data map(to: v1, v2) map(from: p0)
    call vec_mult(p0,v1,v2,N)
    !$omp end target data
    call output(p0, N)
end subroutine
subroutine vec_mult(p1,v3,v4,N)
real,pointer,dimension(:) :: p1, v3, v4
integer :: N,i
    !$omp target map(to: v3, v4) map(from: p1)
    !$omp parallel do
    do i=1,N
        p1(i) = v3(i) * v4(i)
    end do
    !$omp end target
end subroutine
end module

```

Fortran

*Example 49.4c*

In the following example, the variables `p1`, `v3`, and `v4` are references to the pointer variables `p0`, `v1` and `v2` respectively. The **target** construct's device data environment inherits the pointer variables `p0`, `v1`, and `v2` from the enclosing **target data** construct's device data environment. Thus, `p1`, `v3`, and `v4` are already present in the device data environment.

```
void vec_mult(float* &, float* &, float* &, int &);
extern void init(float*, float*, int);
extern void output(float*, int);
void foo(float *p0, float *v1, float *v2, int N)
{
    init(v1, v2, N);
    #pragma omp target data map(to: v1[0:N], v2[:N]) map(from: p0[0:N])
    {
        vec_mult(p0, v1, v2, N);
    }
    output(p0, N);
}
void vec_mult(float* &p1, float* &v3, float* &v4, int &N)
{
    int i;
    #pragma omp target map(to: v3[0:N], v4[:N]) map(from: p1[0:N])
    #pragma omp parallel for
    for (i=0; i<N; i++)
        p1[i] = v3[i] * v4[i];
}
```

*Example 49.5f*

In the following example, the usual Fortran approach is used for dynamic memory. The `p0`, `v1`, `v2` arrays are allocated in the main program and passed as references from one routine to another. In `vec_mult`, `p1`, `v3` and `v4` are references to the `p0`, `v1`, and `v2` arrays, respectively. The **target** construct's device data environment inherits the arrays `p0`, `v1`, and `v2` from the enclosing target data construct's device data environment. Thus, `p1`, `v3`, and `v4` are already present in the device data environment.

```
module my_mult
contains
subroutine foo(p0,v1,v2,N)
```

```

real,dimension(:) :: p0, v1, v2
integer            :: N,i
  call init(v1, v2, N)
  !$omp target data map(to: v1, v2) map(from: p0)
  call vec_mult(p0,v1,v2,N)
  !$omp end target data
  call output(p0, N)
end subroutine
subroutine vec_mult(p1,v3,v4,N)
real,dimension(:) :: p1, v3, v4
integer            :: N,i
  !$omp target map(to: v3, v4) map(from: p1)
  !$omp parallel do
    do i=1,N
      p1(i) = v3(i) * v4(i)
    end do
  !$omp end target
end subroutine
end module
!
program main
use my_mult
integer, parameter :: N=1024
real,allocatable, dimension(:) :: p, v1, v2
  allocate( p(N), v1(N), v2(N) )
  call foo(p,v1,v2,N)
end program

```

Fortran

## target data Construct With if Clause

The following two examples show how the **target data** construct maps variables to a device data environment.

In the following example, the **if** clause on the **target data** construct indicates that if the variable **N** is smaller than a given threshold, then the **target data** construct will not create a device data environment.

The **target** constructs enclosed in the **target data** region must also use an **if** clause on the same condition, otherwise the pointer variable **p** is implicitly mapped with a map-type of **tofrom**, but the storage location for the array section **p[0:N]** will not be mapped in the device data environments of the **target** constructs.

C/C++

*Example 49.5c*

```
#define THRESHOLD 1000000
```

```

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

```

C/C++

Fortran

### Example 49.6f

The **if** clauses work the same way for the following Fortran code. The **target** constructs enclosed in the **target data** region should also use an **if** clause with the same condition, so that the **target data** region and the **target** region are either both created for the device, or are both ignored.

```

module params
integer,parameter :: THRESHOLD=1000000
end module
subroutine vec_mult(p, v1, v2, N)
    use params
    real    :: p(N), v1(N), v2(N)
    integer :: i
    call init(v1, v2, N)
    !$omp target data if (N>THRESHOLD) map(from: p)
    !$omp target if (N>THRESHOLD) map(to: v1, v2)
    !$omp parallel do
    do i=1,N
        p(i) = v1(i) * v2(i)
    end do
    !$omp end target
    call init_again(v1, v2, N)
    !$omp target if (N>THRESHOLD) map(to: v1, v2)
    !$omp parallel do

```

```

        do i=1,N
            p(i) = p(i) + v1(i) * v2(i)
        end do
    !$omp end target
    !$omp end target data
    call output(p, N)
end subroutine

```

## Fortran

In the following example, when the **if** clause conditional expression on the **target** construct evaluates to *false*, the target region will execute on the host device. However, the **target data** construct created an enclosing device data environment that mapped `p[0:N]` to a device data environment on the default device. At the end of the **target data** region the array section `p[0:N]` will be assigned from the device data environment to the corresponding variable in the data environment of the task that encountered the **target data** construct, resulting in undefined values in `p[0:N]`.

## C/C++

### Example 49.6c

```

#define THRESHOLD 1000000
extern void init(float*, float*, int);
extern void output(float*, int);
void vec_mult(float *p, float *v1, float *v2, int N)
{
    int i;
    init(v1, v2, N);
    #pragma omp target data map(from: p[0:N])
    {
        #pragma omp target if (N>THRESHOLD) map(to: v1[:N], v2[:N])
        #pragma omp parallel for
        for (i=0; i<N; i++)
            p[i] = v1[i] * v2[i];
    } /* UNDEFINED behavior if N<=THRESHOLD */
    output(p, N);
}

```

## C/C++

## Fortran

### Example 49.7f

The **if** clauses work the same way for the following Fortran code. When the **if** clause conditional expression on the **target** construct evaluates to *false*, the **target** region will execute on the host device. However, the **target data** construct created an enclosing device data environment that mapped the `p` array (and `v1` and `v2`) to a device data environment on the default target device. At the end of the **target data** region

the p array will be assigned from the device data environment to the corresponding variable in the data environment of the task that encountered the **target data** construct, resulting in undefined values in p.

```
module params
integer, parameter :: THRESHOLD=1000000
end module
subroutine vec_mult(p, v1, v2, N)
  use params
  real    :: p(N), v1(N), v2(N)
  integer :: i
  call init(v1, v2, N)
  !$omp target data map(from: p)
    !$omp target if(N>THRESHOLD) map(to: v1, v2)
    !$omp parallel do
      do i=1,N
        p(i) = v1(i) * v2(i)
      end do
    !$omp end target
  !$omp end target data
  call output(p, N)  !*** UNDEFINED behavior if N<=THRESHOLD
end subroutine
```

Fortran

## Simple target data and target update Constructs

The following example shows how the **target update** construct updates variables in a device data environment.

The **target data** construct maps array sections `v1[:N]` and `v2[:N]` (arrays `v1` and `v2` in the Fortran code) into a device data environment.

The task executing on the host device encounters the first **target** region and waits for the completion of the region.

After the execution of the first **target** region, the task executing on the host device then assigns new values to `v1[:N]` and `v2[:N]` (`v1` and `v2` arrays in Fortran code) in the task's data environment by calling the function `init_again()`.

The **target update** construct assigns the new values of `v1` and `v2` from the task's data environment to the corresponding mapped array sections in the device data environment of the **target data** construct.

The task executing on the host device then encounters the second **target** region and waits for the completion of the region.

The second **target** region uses the updated values of `v1[:N]` and `v2[:N]`.

C/C++

*Example 50.1c*

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



```

        p[i] = v1[i] * v2[i];
    init_again(v1, v2, N);
    #pragma omp target update to(v1[:N], v2[:N])
    #pragma omp target
    #pragma omp parallel for
    for (i=0; i<N; i++)
        p[i] = p[i] + (v1[i] * v2[i]);
    }
    output(p, N);
}

```

C/C++

Fortran

### *Example 50.1f*

```

subroutine vec_mult(p, v1, v2, N)
    real    :: p(N), v1(N), v2(N)
    integer :: i
    call init(v1, v2, N)
    !$omp target data map(to: v1, v2) map(from: p)
        !$omp target
        !$omp parallel do
            do i=1,N
                p(i) = v1(i) * v2(i)
            end do
        !$omp end target
    call init_again(v1, v2, N)
    !$omp target update to(v1, v2)
    !$omp target
    !$omp parallel do
        do i=1,N
            p(i) = p(i) + v1(i) * v2(i)
        end do
    !$omp end target
    !$omp end target data
    call output(p, N)
end subroutine

```

Fortran

## target update Construct With if Clause

The following example shows how the **target update** construct updates variables in a device data environment.

The **target data** construct maps array sections `v1[:N]` and `v2[:N]` (arrays `v1` and `v2` in the Fortran code) into a device data environment. In between the two **target** regions, the task executing on the host device conditionally assigns new values to `v1` and `v2` in the task's data environment. The function `maybe_init_again()` returns true if new data is written.

When the conditional expression (the return value of `maybe_init_again()`) in the **if** clause is *true*, the **target update** construct assigns the new values of `v1` and `v2` from the task's data environment to the corresponding mapped array sections in the **target data** construct's device data environment.

### Example 50.2c

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

*Example 50.2f*

```

subroutine vec_mult(p, v1, v2, N)
  interface
    logical function maybe_init_again (v1, N)
      real :: v1(N)
      integer :: N
    end function
  end interface
  real :: p(N), v1(N), v2(N)
  integer :: i
  logical :: changed
  call init(v1, v2, N)
  !$omp target data map(to: v1, v2) map(from: p)
    !$omp target
    !$omp parallel do
      do i=1, N
        p(i) = v1(i) * v2(i)
      end do
    !$omp end target
    changed = maybe_init_again(v1, N)
    !$omp target if(changed) update to(v1(:N))
    changed = maybe_init_again(v2, N)
    !$omp target if(changed) update to(v2(:N))
    !$omp target
    !$omp parallel do
      do i=1, N
        p(i) = p(i) + v1(i) * v2(i)
      end do
    !$omp end target
  !$omp end target data
  call output(p, N)
end subroutine

```

## declare target and end declare target for a Function

The following example shows how the **declare target** directive is used to indicate that the corresponding call inside a **target** region is to a `fib` function that can execute on the default target device.

A version of the function is also available on the host device. When the **if** clause conditional expression on the **target** construct evaluates to *false*, the **target** region (thus `fib`) will execute on the host device.

For C/C++ codes the declaration of the function `fib` appears between the **declare target** and **end declare target** directives.

▼ C/C++ ▼  
*Example 51.1c*

```
#pragma omp declare target
extern void fib(int N);
#pragma omp end declare target
#define THRESHOLD 1000000
void fib_wrapper(int n)
{
    #pragma omp target if(n > THRESHOLD)
    {
        fib(n);
    }
}
```

▲ C/C++ ▲

▼ Fortran ▼  
*Example 51.1f*

The Fortran `fib` subroutine contains a **declare target** declaration to indicate to the compiler to create an device executable version of the procedure. The subroutine name has not been included on the **declare target** directive and is, therefore, implicitly assumed.

The program uses the `module_fib` module, which presents an explicit interface to the compiler with the **declare target** declarations for processing the `fib` call.

```
module module_fib
contains
  subroutine fib(N)
    integer :: N
    !$omp declare target
    !...
  end subroutine
end module
module params
integer :: THRESHOLD=1000000
end module
program my_fib
use params
use module_fib
  !$omp target if( N > THRESHOLD )
    call fib(N)
  !$omp end target
end program
```

The next Fortran example shows the use of an external subroutine. Without an explicit interface (through module use or an interface block) the **declare target** declarations within a external subroutine are unknown to the main program unit; therefore, a **declare target** must be provided within the program scope for the compiler to determine that a target binary should be available.

### *Example 51.2f*

```
program my_fib
integer :: N = 8
!$omp declare target(fib)
  !$omp target
    call fib(N)
  !$omp end target
end program
subroutine fib(N)
integer :: N
!$omp declare target
  print*,"hello from fib"
  !...
end subroutine
```

Fortran

## declare target Construct for Class Type

The following example shows how the **declare target** and **end declare target** directives are used to enclose the declaration of a variable `varY` with a class type `typeY`. The member function `typeY::foo()` cannot be accessed on a target device because its declaration did not appear between **declare target** and **end declare target** directives.

*Example 51.2c*

C/C++

```
struct typeX
{
    int a;
}
class typeY
{
    int foo() { return a^0x01;}
    int a;
}
#pragma omp declare target
struct typeX varX; // ok
class typeY varY; // ok if varY.foo() not called on target device
#pragma omp end declare target
void foo()
{
    #pragma omp target
    {
        varX.a = 100; // ok
        varY.foo(); // error foo() is not available on a target device
    }
}
```

C/C++

## declare target and end declare target for Variables

The following examples show how the **declare target** and **end declare target** directives are used to indicate that global variables are mapped to the implicit device data environment of each target device.

In the following example, the declarations of the variables `p`, `v1`, and `v2` appear between **declare target** and **end declare target** directives indicating that the variables are mapped to the implicit device data environment of each target device. The

**target update** directive is then used to manage the consistency of the variables `p`, `v1`, and `v2` between the data environment of the encountering host device task and the implicit device data environment of the default target device.

### C/C++

#### Example 51.3c

```
#define N 1000
#pragma omp declare target
float p[N], v1[N], v2[N];
#pragma omp end declare target
extern void init(float *, float *, int);
extern void output(float *, int);
void vec_mult()
{
    int i;
    init(v1, v2, N);
    #pragma omp target update to(v1, v2)
    #pragma omp target
    #pragma omp parallel for
    for (i=0; i<N; i++)
        p[i] = v1[i] * v2[i];
    #pragma omp target update from(p)
    output(p, N);
}
```

### C/C++

### Fortran

#### Example 51.3f

The Fortran version of the above C code uses a different syntax. Fortran modules use a list syntax on the **declare target** directive to declare mapped variables.

```
module my_arrays
!$omp declare target (N, p, v1, v2)
integer, parameter :: N=1000
real                :: p(N), v1(N), v2(N)
end module
subroutine vec_mult()
use my_arrays
integer :: i
call init(v1, v2, N);
!$omp target update to(v1, v2)
!$omp target
!$omp parallel do
do i = 1,N
    p(i) = v1(i) * v2(i)
end do
end subroutine
```

```

        end do
        !$omp end target
        !$omp target update from (p)
        call output(p, N)
    end subroutine

```

#### Fortran

The following example also indicates that the function `Pfun()` is available on the target device, as well as the variable `Q`, which is mapped to the implicit device data environment of each target device. The **target update** directive is then used to manage the consistency of the variable `Q` between the data environment of the encountering host device task and the implicit device data environment of the default target device.

#### C/C++

In the following example, the function and variable declarations appear between the **declare target** and **end declare target** directives.

#### *Example 51.4c*

```

#define N 10000
#pragma omp declare target
float Q[N][N];
float Pfun(const int i, const int k)
{ return Q[i][k] * Q[k][i]; }
#pragma omp end declare target
float accum(int k)
{
    float tmp = 0.0;
    #pragma omp target update to(Q)
    #pragma omp target
    #pragma omp parallel for reduction(+:tmp)
    for(int i=0; i < N; i++)
        tmp += Pfun(i,k);
    return tmp;
}

```

#### C/C++

#### Fortran

#### *Example 51.4f*

The Fortran version of the above C code uses a different syntax. In Fortran modules a list syntax on the **declare target** directive is used to declare mapped variables and procedures. The `N` and `Q` variables are declared as a comma separated list. When the **declare target** directive is used to declare just the procedure, the procedure name need not be listed -- it is implicitly assumed, as illustrated in the `Pfun()` function.



```

module my_global_array
!$omp declare target (N,Q)
integer, parameter :: N=10
real                :: Q(N,N)
contains
function Pfun(i,k)
!$omp declare target
real                :: Pfun
integer,intent(in) :: i,k
    Pfun=(Q(i,k) * Q(k,i))
end function
end module
function accum(k) result(tmp)
use my_global_array
real    :: tmp
integer :: i, k
    tmp = 0.0e0
    !$omp target
    !$omp parallel do reduction(+:tmp)
    do i=1,N
        tmp = tmp + Pfun(k,i)
    end do
    !$omp end target
end function

```

Fortran

## declare target and end declare target with declare simd

The following example shows how the **declare target** and **end declare target** directives are used to indicate that a function is available on a target device. The **declare simd** directive indicates that there is a SIMD version of the function `P()` that is available on the target device as well as one that is available on the host device.

C/C++

### *Example 51.5c*

```

#define N 10000
#define M 1024
#pragma omp declare target
float Q[N][N];
#pragma omp declare simd uniform(i) linear(k) notinbranch
float P(const int i, const int k)
{
    return Q[i][k] * Q[k][i];
}

```

```

#pragma omp end declare target
float accum(void)
{
    float tmp = 0.0;
    int i, k;
#pragma omp target
#pragma omp parallel for reduction(+:tmp)
    for (i=0; i < N; i++) {
        float tmp1 = 0.0;
#pragma omp simd reduction(+:tmp1)
        for (k=0; k < M; k++) {
            tmp1 += P(i,k);
        }
        tmp += tmp1;
    }
    return tmp;
}

```

C/C++

Fortran

### Example 51.5f

The Fortran version of the above C code uses a different syntax. Fortran modules use a list syntax of the **declare target** declaration for the mapping. Here the N and Q variables are declared in the list form as a comma separated list. The function declaration does not use a list and implicitly assumes the function name. In this Fortran example row and column indices are reversed relative to the C/C++ example, as is usual for codes optimized for memory access.

```

module my_global_array
!$omp declare target (N,Q)
integer, parameter :: N=10000, M=1024
real                :: Q(N,N)
contains
function P(k,i)
!$omp declare simd uniform(i) linear(k) notinbranch
!$omp declare target
real                :: P
integer,intent(in) :: k,i
    P=(Q(k,i) * Q(i,k))
end function
end module
function accum() result(tmp)
use my_global_array
real    :: tmp, tmp1
integer :: i
    tmp = 0.0e0
!$omp target
!$omp parallel do private(tmp1) reduction(+:tmp)

```

```
do i=1,N
  tmp1 = 0.0e0
  !$omp simd reduction(+:tmp1)
  do k = 1,M
    tmp1 = tmp1 + P(k,i)
  end do
  tmp = tmp + tmp1
end do
!$omp end target
end function
```

Fortran

## target and teams Constructs with omp\_get\_num\_teams and omp\_get\_team\_num Routines

The following example shows how the **target** and **teams** constructs are used to create a league of thread teams that execute a region. The **teams** construct creates a league of at most two teams where the master thread of each team executes the **teams** region.

The `omp_get_num_teams` routine returns the number of teams executing in a **teams** region. The `omp_get_team_num` routine returns the team number, which is an integer between 0 and one less than the value returned by `omp_get_num_teams`. The following example manually distributes a loop across two teams.

C/C++

*Example 52.1c*

```
#include <stdlib.h>
#include <omp.h>
float dotprod(float B[], float C[], int N)
{
    float sum0 = 0.0;
    float sum1 = 0.0;
    #pragma omp target map(to: B[:N], C[:N])
    #pragma omp teams num_teams(2)
    {
        int i;
        if (omp_get_num_teams() != 2)
            abort();
        if (omp_get_team_num() == 0)
        {
            #pragma omp parallel for reduction(+:sum0)
            for (i=0; i<N/2; i++)
                sum0 += B[i] * C[i];
        }
        else if (omp_get_team_num() == 1)
        {
            #pragma omp parallel for reduction(+:sum1)
            for (i=N/2; i<N; i++)
                sum1 += B[i] * C[i];
        }
    }
}
```

```

    }
    return sum0 + sum1;
}

```

C/C++

Fortran

### *Example 52.1f*

```

function dotprod(B,C,N) result(sum)
use omp_lib, ONLY : omp_get_num_teams, omp_get_team_num
real    :: B(N), C(N), sum,sum0, sum1
integer :: N, i
sum0 = 0.0e0
sum1 = 0.0e0
!$omp target map(to: B, C)
!$omp teams num_teams(2)
  if (omp_get_num_teams() /= 2) stop "2 teams required"
  if (omp_get_team_num() == 0) then
    !$omp parallel do reduction(+:sum0)
    do i=1,N/2
      sum0 = sum0 + B(i) * C(i)
    end do
  else if (omp_get_team_num() == 1) then
    !$omp parallel do reduction(+:sum1)
    do i=N/2+1,N
      sum1 = sum1 + B(i) * C(i)
    end do
  end if
!$omp end teams
!$omp end target
sum = sum0 + sum1
end function

```

Fortran

## target, teams, and distribute Constructs

The following example shows how the **target**, **teams**, and **distribute** constructs are used to execute a loop nest in a **target** region. The **teams** construct creates a league and the master thread of each team executes the **teams** region. The **distribute** construct schedules the subsequent loop iterations across the master threads of each team.

The number of teams in the league is less than or equal to the variable `num_blocks`. Each team in the league has a number of threads less than or equal to the variable `block_threads`. The iterations in the outer loop are distributed among the master threads of each team.

When a team's master thread encounters the parallel loop construct before the inner loop, the other threads in its team are activated. The team executes the **parallel** region and then workshares the execution of the loop.

Each master thread executing the **teams** region has a private copy of the variable `sum` that is created by the **reduction** clause on the teams construct. The master thread and all threads in its team have a private copy of the variable `sum` that is created by the **reduction** clause on the parallel loop construct. The second private `sum` is reduced into the master thread's private copy of `sum` created by the **teams** construct. At the end of the **teams** region, each master thread's private copy of `sum` is reduced into the final `sum` that is implicitly mapped into the **target** region.

### Example 52.2c C/C++

```
float dotprod(float B[], float C[], int N, int block_size,
             int num_teams, int block_threads)
{
    float sum = 0;
    int i, i0;
    #pragma omp target map(to: B[0:N], C[0:N])
    #pragma omp teams num_teams(num_teams) thread_limit(block_threads) \
        reduction(+:sum)
    #pragma omp distribute
    for (i0=0; i0<N; i0 += block_size)
        #pragma omp parallel for reduction(+:sum)
        for (i=i0; i< min(i0+block_size,N); i++)
            sum += B[i] * C[i];
    return sum;
}
```

### C/C++

### Example 52.2f Fortran

```
function dotprod(B,C,N, block_size, num_teams, block_threads) result(sum)
implicit none
    real    :: B(N), C(N), sum
    integer :: N, block_size, num_teams, block_threads, i, i0
    sum = 0.0e0
    !$omp target map(to: B, C)
```

```

!$omp teams num_teams(num_teams) thread_limit(block_threads) &
    reduction(+:sum)
!$omp distribute
do i0=1,N, block_size
    !$omp parallel do reduction(+:sum)
    do i = i0, min(i0+block_size,N)
        sum = sum + B(i) * C(i)
    end do
end do
!$omp end teams
!$omp end target
end function

```

Fortran

## target teams, and Distribute Parallel Loop Constructs

The following example shows how the **target teams** and distribute parallel loop constructs are used to execute a **target** region. The **target teams** construct creates a league of teams where the master thread of each team executes the **teams** region.

The distribute parallel loop construct schedules the loop iterations across the master threads of each team and then across the threads of each team.

C/C++

### *Example 52.3c*

```

float dotprod(float B[], float C[], int N)
{
    float sum = 0;
    int i;
    #pragma omp target teams map(to: B[0:N], C[0:N])
    #pragma omp distribute parallel for reduction(+:sum)
    for (i=0; i<N; i++)
        sum += B[i] * C[i];
    return sum;
}

```

C/C++

*Example 52.3f*

```

function dotprod(B,C,N) result(sum)
  real    :: B(N), C(N), sum
  integer :: N, i
  sum = 0.0e0
  !$omp target teams map(to: B, C)
  !$omp distribute parallel do reduction(+:sum)
    do i = 1,N
      sum = sum + B(i) * C(i)
    end do
  !$omp end teams
  !$omp end target
end function

```

## target teams and Distribute Parallel Loop Constructs with Scheduling Clauses

The following example shows how the **target teams** and distribute parallel loop constructs are used to execute a **target** region. The **teams** construct creates a league of at most eight teams where the master thread of each team executes the **teams** region. The number of threads in each team is less than or equal to 16.

The **distribute** parallel loop construct schedules the subsequent loop iterations across the master threads of each team and then across the threads of each team.

The **dist\_schedule** clause on the distribute parallel loop construct indicates that loop iterations are distributed to the master thread of each team in chunks of 1024 iterations.

The **schedule** clause indicates that the 1024 iterations distributed to a master thread are then assigned to the threads in its associated team in chunks of 64 iterations.

*Example 52.4c*

```

#define N 1024*1024
float dotprod(float B[], float C[], int N)
{
  float sum = 0;
  int i;
  #pragma omp target map(to: B[0:N], C[0:N])
  #pragma omp teams num_teams(8) thread_limit(16)

```



```

#pragma omp distribute parallel for reduction(+:sum) \
    dist_schedule(static, 1024) schedule(static, 64)
for (i=0; i<N; i++)
    sum += B[i] * C[i];
return sum;
}

```

C/C++

Fortran

### *Example 52.4f*

```

module arrays
integer,parameter :: N=1024*1024
real :: B(N), C(N)
end module
function dotprod() result(sum)
use arrays
    real :: sum
    integer :: i
    sum = 0.0e0
    !$omp target map(to: B, C)
    !$omp teams num_teams(8) thread_limit(16)
    !$omp distribute parallel do reduction(+:sum) &
    !$omp& dist_schedule(static, 1024) schedule(static, 64)
        do i = 1,N
            sum = sum + B(i) * C(i)
        end do
    !$omp end teams
    !$omp end target
end function

```

Fortran

## target teams and distribute simd Constructs

The following example shows how the **target teams** and **distribute simd** constructs are used to execute a loop in a **target** region. The **target teams** construct creates a league of teams where the master thread of each team executes the **teams** region.

The **distribute simd** construct schedules the loop iterations across the master thread of each team and then uses SIMD parallelism to execute the iterations.

*Example 52.5c*

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

*Example 52.5f*

```
subroutine vec_mult(p, v1, v2, N)
    real    :: p(N), v1(N), v2(N)
    integer :: i
    call init(v1, v2, N)
    !$omp target teams map(to: v1, v2) map(from: p)
        !$omp distribute simd
        do i=1,N
            p(i) = v1(i) * v2(i)
        end do
    !$omp end target teams
    call output(p, N)
end subroutine
```

## target teams and Distribute Parallel Loop SIMD Constructs

The following example shows how the **target teams** and the distribute parallel loop SIMD constructs are used to execute a loop in a **target teams** region. The **target teams** construct creates a league of teams where the master thread of each team executes the **teams** region.

The distribute parallel loop SIMD construct schedules the loop iterations across the master thread of each team and then across the threads of each team where each thread uses SIMD parallelism.

C/C++

*Example 52.6c*

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

C/C++

Fortran

*Example 52.6f*

```
subroutine vec_mult(p, v1, v2, N)
    real    :: p(N), v1(N), v2(N)
    integer :: i
    call init(v1, v2, N)
    !$omp target teams map(to: v1, v2) map(from: p)
    !$omp distribute parallel do simd
    do i=1,N
        p(i) = v1(i) * v2(i)
    end do
    !$omp end target teams
    call output(p, N)
end subroutine
```

Fortran

## Asynchronous Execution of a target Region Using Tasks

The following example shows how the **task** and **target** constructs are used to execute multiple **target** regions asynchronously. The task that encounters the **task** construct generates an explicit task that contains a **target** region. The thread executing the explicit task encounters a task scheduling point while waiting for the execution of the **target** region to complete, allowing the thread to switch back to the execution of the encountering task or one of the previously generated explicit tasks.

C/C++

*Example 53.1c*

```
#pragma omp declare target
float F(float);
#pragma omp end declare target
#define N 1000000000
#define CHUNKSZ 1000000
void init(float *, int);
float Z[N];
void pipedF()
{
    int C, i;
    init(Z, N);
    for (C=0; C<N; C+=CHUNKSZ)
    {
        #pragma omp task
        #pragma omp target map(Z[C:CHUNKSZ])
        #pragma omp parallel for
        for (i=0; i<CHUNKSZ; i++)
            Z[i] = F(Z[i]);
    }
    #pragma omp taskwait
}
```

C/C++

Fortran

*Example 53.1f*

The Fortran version has an interface block that contains the **declare target**. An identical statement exists in the function declaration (not shown here).

```
module parameters
```

```

integer, parameter :: N=1000000000, CHUNKSZ=1000000
end module

subroutine pipedF()
use parameters, ONLY: N, CHUNKSZ
integer          :: C, i
real             :: z(N)

interface
  function F(z)
    !$omp declare target
    real, intent(IN) :: z
    real             :: F
  end function F
end interface

call init(z,N)

do C=1,N,CHUNKSZ

  !$omp task
  !$omp target map(z(C:C+CHUNKSZ-1))
  !$omp parallel do
    do i=C,C+CHUNKSZ-1
      z(i) = F(z(i))
    end do
  !$omp end target
  !$omp end task

end do
print*, z

end subroutine pipedF

```

## Fortran

The following example shows how the **task** and **target** constructs are used to execute multiple **target** regions asynchronously. The task dependence ensures that the storage is allocated and initialized on the device before it is accessed.

## C/C++

### Example 53.2c

```

#include <stdlib.h>
extern void init(float *, float *, int);
extern void output(float *, int);
void vec_mult(float *p, float *v1, float *v2, int N, int dev)
{
  int i;
  init(p, N);
  #pragma omp task depend(out: v1, v2)
  #pragma omp target device(dev) map(v1, v2)

```

```

{
    // check whether on device dev
    if (omp_is_initial_device())
        abort();
    v1 = malloc(N*sizeof(float));
    v2 = malloc(N*sizeof(float));
    init(v1,v2);
}
foo(); // execute other work asynchronously
#pragma omp task depend(in: v1, v2)
#pragma omp target device(dev) map(to: v1, v2) map(from: p[0:N])
{
    // check whether on device dev
    if (omp_is_initial_device())
        abort();
    #pragma omp parallel for
    for (i=0; i<N; i++)
        p[i] = v1[i] * v2[i];
    output(p, N);
    free(v1);
    free(v2);
}
}

```

C/C++

Fortran

### Example 53.2f

The Fortran example uses allocatable arrays for dynamic memory on the device.

```

subroutine mult(p, N, idev)
    use omp_lib, ONLY: omp_is_initial_device
    real                :: p(N)
    real,allocatable :: v1(:), v2(:)
    integer :: i, idev
    !$omp declare target (init)

    !$omp task depend(out: v1,v2)
    !$omp target device(idev) map(v1,v2)
        if( omp_is_initial_device() ) &
            stop "not executing on target device"
        allocate(v1(N), v2(N))
        call init(v1,v2,N)
    !$omp end target
    !$omp end task

    call foo() ! execute other work asynchronously

    !$omp task depend(in: v1,v2)
    !$omp target device(idev) map(to: v1,v2) map(from: p)

```

```

        if( omp_is_initial_device() ) &
            stop "not executing on target device"
        !$omp parallel do
            do i = 1,N
                p(i) = v1(i) * v2(i)
            end do
            deallocate(v1,v2)
                                                    !!

        !$omp end target
        !$omp end task
                                                    !!

        call output(p, N)
                                                    !!
    end subroutine

```

Fortran

## Array Sections in Device Constructs

The following examples show the usage of array sections in **map** clauses on **target** and **target data** constructs.

This example shows the invalid usage of two separate sections of the same array inside of a **target** construct.

C/C++

*Example 54.1c*

```
void foo ()
{
    int A[30];
    #pragma omp target data map( A[0:4] )
    {
        /* Cannot map distinct parts of the same array */
        #pragma omp target map( A[7:20] )
        {
            A[2] = 0;
        }
    }
}
```

C/C++

Fortran

*Example 54.1f*

```
subroutine foo()
integer :: A(30)
  A = 1
  !$omp target data map( A(1:4) )
    ! Cannot map distinct parts of the same array
    !$omp target map( A(8:27) )
      A(3) = 0
    !$omp end target map
  !$omp end target data
end subroutine
```

Fortran

This example shows the invalid usage of two separate sections of the same array inside of a **target** construct.



*Example 54.2c*

```

void foo ()
{
    int A[30], *p;
    #pragma omp target data map( A[0:4] )
    {
        p = &A[0];
        /* invalid because p[3] and A[3] are the same
         * location on the host but the array section
         * specified via p[...] is not a subset of A[0:4] */
        #pragma omp target map( p[3:20] )
        {
            A[2] = 0;
            p[8] = 0;
        }
    }
}

```

*Example 54.2f*

```

subroutine foo()
integer,target :: A(30)
integer,pointer :: p(:)
A=1
!$omp target data map( A(1:4) )
p=>A
! invalid because p(4) and A(4) are the same
! location on the host but the array section
! specified via p(...) is not a subset of A(1:4)
!$omp target map( p(4:23) )
A(3) = 0
p(9) = 0
!$omp end target
!$omp end target data
end subroutine

```

This example shows the valid usage of two separate sections of the same array inside of a **target** construct.

*Example 54.3c*

```

void foo ()
{
    int A[30], *p;
    #pragma omp target data map( A[0:4] )
    {
        p = &A[0];
        #pragma omp target map( p[7:20] )
        {
            A[2] = 0;
            p[8] = 0;
        }
    }
}

```

*Example 54.3f*

```

subroutine foo()
integer,target :: A(30)
integer,pointer :: p(:)
!$omp target data map( A(1:4) )
p=>A
!$omp target map( p(8:27) )
A(3) = 0
p(9) = 0
!$omp end target map
!$omp end target data
end subroutine

```

This example shows the valid usage of a wholly contained array section of an already mapped array section inside of a **target** construct.

*Example 54.4c*

```

void foo ()
{
    int A[30];
    #pragma omp target data map( A[0:10] )
    {
        p = &A[0];
        #pragma omp target map( p[3:7] )
    }
}

```

```

    {
        A[2] = 0;
        p[8] = 0;
        A[8] = 1;
    }
}

```

C/C++

Fortran

*Example 54.4f*

```

subroutine foo()
integer,target :: A(30)
integer,pointer :: p(:)
!$omp target data map( A(1:10) )
  p=>A
!$omp target map( p(4:10) )
  A(3) = 0
  p(9) = 0
  A(9) = 1
!$omp end target
!$omp end target data
end subroutine

```

Fortran

## omp\_is\_initial\_device Routine

The following example shows how the `omp_is_initial_device` runtime library routine can be used to query if a code is executing on the initial host device or on a target device. The example then sets the number of threads in the **parallel** region based on where the code is executing.

C/C++

*Example 55.1c*

```
#include <stdio.h>
#include <omp.h>
#pragma omp declare target
void vec_mult(float *p, float *v1, float *v2, int N);
extern float *p, *v1, *v2;
extern int N;
#pragma omp end declare target
extern void init_vars(float *, float *, int);
extern void output(float *, int);
void foo()
{
    N = init_vars(&p, &v1, &v2);
    #pragma omp target device(42) map(p[:N], v1[:N], v2[:N])
    {
        vec_mult(p, v1, v2, N);
    }
    output(p, N);
}
void vec_mult(float *p, float *v1, float *v2, int N)
{
    int i;
    int nthreads = omp_is_initial_device() ? 8 : 1024;
    if (!omp_is_initial_device())
    {
        printf("1024 threads on target device\n");
        nthreads = 1024;
    }
    else
    {
        printf("8 threads on initial device\n");
        nthreads = 8;
    }
    #pragma omp parallel for private(i) num_threads(nthreads);
```

```

    for (i=0; i<N; i++)
        p[i] = v1[i] * v2[i];
    }

```

C/C++

Fortran

### *Example 55.1f*

```

module params
    integer,parameter :: N=1024
end module params
module vmult
contains
    subroutine vec_mult(p, v1, v2, N)
        use omp_lib, ONLY : omp_is_initial_device
        !$omp declare target
        real    :: p(N), v1(N), v2(N)
        integer :: i, nthreads, N
        if (.not. omp_is_initial_device()) then
            print*, "1024 threads on target device"
            nthreads = 1024
        else
            print*, "8 threads on initial device"
            nthreads = 8
        endif
        !$omp parallel do private(i) num_threads(nthreads)
        do i = 1,N
            p(i) = v1(i) * v2(i)
        end do
    end subroutine vec_mult
end module vmult
program prog_vec_mult
    use params
    use vmult
    real :: p(N), v1(N), v2(N)
    call init(v1,v2,N)
    !$omp target device(42) map(p, v1, v2)
        call vec_mult(p, v1, v2, N)
    !$omp end target
    call output(p, N)
end program

```

Fortran

## omp\_get\_num\_devices Routine

The following example shows how the `omp_get_num_devices` runtime library routine can be used to determine the number of devices.

### C/C++

#### *Example 55.2c*

```
#include <omp.h>
extern void init(float *, float *, int);
extern void output(float *, int);
void vec_mult(float *p, float *v1, float *v2, int N)
{
    int i;
    init(v1, v2, N);
    int ndev = omp_get_num_devices();
    int do_offload = (ndev>0 && N>1000000);
    #pragma omp target if(do_offload) map(to: v1[0:N], v2[:N]) map(from: p[0:N])
    #pragma omp parallel for if(N>1000) private(i)
    for (i=0; i<N; i++)
        p[i] = v1[i] * v2[i];
    output(p, N);
}
```

### C/C++

### Fortran

#### *Example 55.2f*

```
subroutine vec_mult(p, v1, v2, N)
use omp_lib, ONLY : omp_get_num_devices
real    :: p(N), v1(N), v2(N)
integer :: N, i, ndev
logical :: do_offload
    call init(v1, v2, N)
    ndev = omp_get_num_devices()
    do_offload = (ndev>0) .and. (N>1000000)
    !$omp target if(do_offload) map(to: v1, v2) map(from: p)
    !$omp parallel do if(N>1000)
        do i=1,N
            p(i) = v1(i) * v2(i)
        end do
    !$omp end target
    call output(p, N)
end subroutine
```

### Fortran

## omp\_set\_default\_device and omp\_get\_default\_device Routines

The following example shows how the `omp_set_default_device` and `omp_get_default_device` runtime library routines can be used to set the default device and determine the default device respectively.

C/C++

*Example 55.3c*

```
#include <omp.h>
#include <stdio.h>
void foo(void)
{
    int default_device = omp_get_default_device();
    printf("Default device = %d\n", default_device);
    omp_set_default_device(default_device+1);
    if (omp_get_default_device() != default_device+1)
        printf("Default device is still = %d\n", default_device);
}
```

C/C++

Fortran

*Example 55.3f*

```
program foo
use omp_lib, ONLY : omp_get_default_device, omp_set_default_device
integer :: old_default_device, new_default_device
old_default_device = omp_get_default_device()
print*, "Default device = ", old_default_device
new_default_device = old_default_device + 1
call omp_set_default_device(new_default_device)
if (omp_get_default_device() == old_default_device) &
    print*, "Default device is STILL = ", old_default_device
end program
```

Fortran

*Example 56.1f*

This is an invalid example of specifying an associate name on a data-sharing attribute clause. The constraint in the Data Sharing Attribute Rules section in the OpenMP 4.0 API Specifications states that an associate name preserves the association with the selector established at the **ASSOCIATE** statement. The associate name `b` is associated with the shared variable `a`. With the predetermined data-sharing attribute rule, the associate name `b` is not allowed to be specified on the **private** clause.

```
program example
  real :: a, c
  associate (b => a)
!$omp parallel private(b, c)      ! invalid to privatize b
  c = 2.0*b
!$omp end parallel
  end associate
end program
```

*Example 56.2f*

In this example, within the parallel construct, the association name `thread_id` is associated with the private copy of `i`. The print statement should output the unique thread number.

```
program example
  use omp_lib
  integer i
!$omp parallel private(i)
  i = omp_get_thread_num()
  associate(thread_id => i)
    print *, thread_id      ! print private i value
  end associate
!$omp end parallel
end program
```



### Example 56.3f

This example illustrates the effect of specifying a selector name on a data-sharing attribute clause. The associate name `u` is associated with `v` and the variable `v` is specified on the **private** clause of the **parallel** construct. The construct association is established prior to the **parallel** region. The association between `u` and the original `v` is retained (see the Data Sharing Attribute Rules section in the OpenMP 4.0 API Specifications). Inside the **parallel** region, `v` has the value of -1 and `u` has the value of the original `v`.

```
program example
  integer :: v
  v = 15
  associate(u => v)
  !$omp parallel private(v)
    v = -1
    print *, v           ! private v=-1
    print *, u           ! original v=15
  !$omp end parallel
end associate
end program
```

Fortran

