

# OpenMP Application Program Interface

Version 4.0 - RC 2 - March 2013

Public Review Release Candidate 2

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This is a public draft release candidate, based on version 3.1, incorporating the following internal change tickets:

RC1: 71, 77, 90, 88, 111, 125, 126, 136, 137, 138, 144, 146, 147, 148, 149, 150, 151, 152, 154, 155, 156, 157, 164, 166, 168, 169, 170, 172, 173, 174, 175, 176

RC2: 114, 116, 117, 133, 161, 165, 167, 171, 177, 179, 181, 182, 183, 184, 185, 187, 188, 192, 193, 197, 198, 200, 201

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#### 1 CHAPTER 1

## Introduction

3	The collection of compiler directives, library routines, and environment variables
4	described in this document collectively define the specification of the OpenMP
5	Application Program Interface (OpenMP API) for shared-memory parallelism in C, C++
6	and Fortran programs.
7	This specification provides a model for parallel programming that is portable across
8	shared memory architectures from different vendors. Compilers from numerous vendors
9	support the OpenMP API. More information about the OpenMP API can be found at the
10	following web site
11	http://www.openmp.org
12	The directives, library routines, and environment variables defined in this document
13	allow users to create and manage parallel programs while permitting portability. The
14	directives extend the C, C++ and Fortran base languages with single program multiple
15	data (SPMD) constructs, tasking constructs, worksharing constructs, and
16	synchronization constructs, and they provide support for sharing and privatizing data.
17	The functionality to control the runtime environment is provided by library routines and
18	environment variables. Compilers that support the OpenMP API often include a
19	command line option to the compiler that activates and allows interpretation of all
20	OpenMP directives.

## 1.1 Scope

The OpenMP API covers only user-directed parallelization, wherein the programmer explicitly specifies the actions to be taken by the compiler and runtime system in order to execute the program in parallel. OpenMP-compliant implementations are not required to check for data dependencies, data conflicts, race conditions, or deadlocks, any of which may occur in conforming programs. In addition, compliant implementations are not required to check for code sequences that cause a program to be classified as non-

## 4 1.2 Glossary

1

3

## 1.2.1 Threading Concepts

О		
7 8	thread	An execution entity with a stack and associated static memory, called <i>threadprivate memory</i> .
9	OpenMP thread	A thread that is managed by the OpenMP runtime system.
10 11	thread-safe routine	A routine that performs the intended function even when executed concurrently (by more than one <i>thread</i> ).
12 13	processor	Implementation defined hardware unit on which one or more <i>OpenMP threads</i> can execute
14	device	An implementation defined logical execution engine.
15		COMMENT: A device could have one or more processors.
16	league	The set of thread teams created by a teams construct.
17	contention group	The threads of a team in a <i>league</i> and their descendent threads.

## 1.2.2 OpenMP Language Terminology

19 20 21	base language	A programming language that serves as the foundation of the OpenMP specification.
22 23		COMMENT: See Section 1.6 on page 20 for a listing of current <i>base languages</i> for the OpenMP API.
24	base program	A program written in a base language.

1 2	structured block	For C/C++, an executable statement, possibly compound, with a single entry at the top and a single exit at the bottom, or an OpenMP construct.
3 4		For Fortran, a block of executable statements with a single entry at the top and a single exit at the bottom, or an OpenMP construct.
5		COMMENTS:
6		For all base languages,
7		• Access to the <i>structured block</i> must not be the result of a branch.
8 9		• The point of exit cannot be a branch out of the <i>structured block</i> .
10		For C/C++:
11		• The point of entry must not be a call to setjmp().
12		• longjmp() and throw() must not violate the entry/exit criteria.
13		• Calls to exit() are allowed in a structured block.
14 15 16 17 18		<ul> <li>An expression statement, iteration statement, selection statement, or try block is considered to be a <i>structured block</i> if the corresponding compound statement obtained by enclosing it in { and } would be a <i>structured block</i>.</li> </ul>
19		For Fortran:
20		• STOP statements are allowed in a structured block.
21 22	enclosing context	In C/C++, the innermost scope enclosing an OpenMP directive.
23		In Fortran, the innermost scoping unit enclosing an OpenMP directive.
24 25	directive	In C/C++, a <b>#pragma</b> , and in Fortran, a comment, that specifies <i>OpenMP</i> program behavior.
26 27		COMMENT: See Section 2.1 on page 24 for a description of OpenMP <i>directive</i> syntax.
28	white space	A non-empty sequence of space and/or horizontal tab characters.
29 30	OpenMP program	A program that consists of a <i>base program</i> , annotated with OpenMP <i>directives</i> and runtime library routines.
31 32	conforming program	An <i>OpenMP program</i> that follows all the rules and restrictions of the OpenMP specification.

1 2 3	declarative directive	An OpenMP <i>directive</i> that may only be placed in a declarative context. A <i>declarative directive</i> has no associated executable user code, but instead has one or more associated user declarations.
4 5	executable directive	An OpenMP <i>directive</i> that is not declarative. That is, it may be placed in an executable context.
6	stand-alone directive	An OpenMP executable directive that has no associated executable user code.
7 8	loop directive	An OpenMP <i>executable directive</i> whose associated user code must be a loop nest that is a <i>structured block</i> .
9	$associated \ loop(s) \\$	The loop(s) controlled by a loop directive.
10 11		COMMENT: If the <i>loop directive</i> contains a <b>collapse</b> clause then there may be more than one <i>associated loop</i> .
12 13 14 15	construct	An OpenMP <i>executable directive</i> (and for Fortran, the paired <b>end</b> <i>directive</i> , if any) and the associated statement, loop or <i>structured block</i> , if any, not including the code in any called routines. That is, in the lexical extent of an <i>executable directive</i> .
16 17 18 19 20 21	region	All code encountered during a specific instance of the execution of a given <i>construct</i> or of an OpenMP library routine. A <i>region</i> includes any code in called routines as well as any implicit code introduced by the OpenMP implementation. The generation of a <i>task</i> at the point where a <b>task</b> directive is encountered is a part of the <i>region</i> of the <i>encountering thread</i> , but the <i>explicit task region</i> associated with the <b>task</b> directive is not.
22		COMMENTS:
23 24		A <i>region</i> may also be thought of as the dynamic or runtime extent of a <i>construct</i> or of an OpenMP library routine.
25 26		During the execution of an <i>OpenMP program</i> , a <i>construct</i> may give rise to many <i>regions</i> .
27		
28 29	active parallel region	A parallel <i>region</i> that is executed by a <i>team</i> consisting of more than one <i>thread</i> .
30	inactive parallel region	A parallel region that is executed by a team of only one thread.

1 2 3	sequential part	All code encountered during the execution of an <i>OpenMP program</i> that is not part of a parallel <i>region</i> corresponding to a parallel <i>construct</i> or a task <i>region</i> corresponding to a task <i>construct</i> .
4		COMMENTS:
5 6		The sequential part executes as if it were enclosed by an inactive parallel region.
7 8 9		Executable statements in called routines may be in both the <i>sequential</i> part and any number of explicit <b>parallel</b> regions at different points in the program execution.
10 11	master thread	The <i>thread</i> that encounters a <b>parallel</b> <i>construct</i> , creates a <i>team</i> , generates a set of <i>tasks</i> , then executes one of those <i>tasks</i> as <i>thread</i> number 0.
12 13 14 15 16	parent thread	The thread that encountered the parallel construct and generated a parallel region is the parent thread of each of the threads in the team of that parallel region. The master thread of a parallel region is the same thread as its parent thread with respect to any resources associated with an OpenMP thread.
17	ancestor thread	For a given thread, its parent thread or one of its parent thread's ancestor threads.
18 19	team	A set of one or more <i>threads</i> participating in the execution of a <b>parallel</b> <i>region</i> .
20		COMMENTS:
21 22		For an active parallel region, the team comprises the master thread and at least one additional thread.
23 24		For an <i>inactive parallel region</i> , the <i>team</i> comprises only the <i>master thread</i> .
25	initial thread	The thread that executes the sequential part.
26 27	implicit parallel region	The inactive parallel region that encloses the sequential part of an OpenMP program.
28	nested construct	A construct (lexically) enclosed by another construct.
29 30	nested region	A <i>region</i> (dynamically) enclosed by another <i>region</i> . That is, a <i>region</i> encountered during the execution of another <i>region</i> .
31 32		COMMENT: Some nestings are <i>conforming</i> and some are not. See Section 2.16 on page 158 for the restrictions on nesting.
33 34	closely nested region	A region nested inside another region with no parallel region nested between them.

1	all threads	All OpenMP threads participating in the OpenMP program.
2	current team	All threads in the team executing the innermost enclosing parallel region.
3	encountering thread	For a given region, the thread that encounters the corresponding construct.
4	all tasks	All tasks participating in the OpenMP program.
5 6 7	current team tasks	All tasks encountered by the corresponding team. Note that the implicit tasks constituting the parallel region and any descendant tasks encountered during the execution of these implicit tasks are included in this binding task set.
8	generating task	For a given region the task whose execution by a thread generated the region.
9 10	binding thread set	The set of <i>threads</i> that are affected by, or provide the context for, the execution of a <i>region</i> .
11 12		The binding thread set for a given region can be all threads, the current team, or the encountering thread.
13 14		COMMENT: The <i>binding thread set</i> for a particular <i>region</i> is described in its corresponding subsection of this specification.
15 16	binding task set	The set of <i>tasks</i> that are affected by, or provide the context for, the execution of a <i>region</i> .
17 18		The binding task set for a given region can be all tasks, the current team tasks, or the generating task.
19 20		COMMENT: The <i>binding task set</i> for a particular <i>region</i> (if applicable) is described in its corresponding subsection of this specification.
21 22	binding region	The enclosing <i>region</i> that determines the execution context and limits the scope of the effects of the bound <i>region</i> is called the <i>binding region</i> .
23 24 25		Binding region is not defined for regions whose binding thread set is all threads or the encountering thread, nor is it defined for regions whose binding task set is all tasks.
26		COMMENTS:
27 28		The <i>binding region</i> for an <b>ordered</b> <i>region</i> is the innermost enclosing <i>loop region</i> .
29 30		The binding region for a taskwait region is the innermost enclosing task region.
31 32 33		For all other regions for which the binding thread set is the current team or the binding task set is the current team tasks, the binding region is the innermost enclosing parallel region.

1 2		For regions for which the binding task set is the generating task, the binding region is the region of the generating task.
3 4		A parallel region need not be active nor explicit to be a binding region.
5		A task region need not be explicit to be a binding region.
6 7		A <i>region</i> never binds to any <i>region</i> outside of the innermost enclosing parallel <i>region</i> .
8 9	orphaned construct	A construct that gives rise to a region whose binding thread set is the current team, but is not nested within another construct giving rise to the binding region.
10 11	worksharing construct	A <i>construct</i> that defines units of work, each of which is executed exactly once by one of the <i>threads</i> in the <i>team</i> executing the <i>construct</i> .
12		For C/C++, worksharing constructs are for, sections, and single.
13 14		For Fortran, worksharing constructs are do, sections, single and workshare.
15	sequential loop	A loop that is not associated with any OpenMP loop directive.
15 16 17	sequential loop place	A loop that is not associated with any OpenMP <i>loop directive</i> .  Unordered set of <i>processors</i> that is treated by the execution environment as a location unit when dealing with OpenMP thread affinity.
16		Unordered set of <i>processors</i> that is treated by the execution environment as a
16 17 18	place	Unordered set of <i>processors</i> that is treated by the execution environment as a location unit when dealing with OpenMP thread affinity.  The ordered list that describes all OpenMP <i>places</i> available to the execution
16 17 18 19 20 21	place place list	Unordered set of <i>processors</i> that is treated by the execution environment as a location unit when dealing with OpenMP thread affinity.  The ordered list that describes all OpenMP <i>places</i> available to the execution environment.  An ordered list that corresponds to a contiguous interval in the OpenMP <i>place list</i> . It describes the places currently available to the execution environment for a given
16 17 18 19 20 21 22	place place list place partition	Unordered set of <i>processors</i> that is treated by the execution environment as a location unit when dealing with OpenMP thread affinity.  The ordered list that describes all OpenMP <i>places</i> available to the execution environment.  An ordered list that corresponds to a contiguous interval in the OpenMP <i>place list</i> . It describes the places currently available to the execution environment for a given parallel region.
16 17 18 19 20 21 22 23	place place list place partition SIMD instruction	Unordered set of <i>processors</i> that is treated by the execution environment as a location unit when dealing with OpenMP thread affinity.  The ordered list that describes all OpenMP <i>places</i> available to the execution environment.  An ordered list that corresponds to a contiguous interval in the OpenMP <i>place list</i> . It describes the places currently available to the execution environment for a given parallel region.  A single machine instruction that can compute multiple data.  A software or hardware mechanism capable of processing one data element from a

1	1.2.3 Sync	chronization Terminology
2	barrier	A point in the execution of a program encountered by a team of threads, beyond
3 4		which no <i>thread</i> in the team may execute until all <i>threads</i> in the <i>team</i> have reached the barrier and all <i>explicit tasks</i> generated by the <i>team</i> have executed to
5		completion. If cancellation has been requested, threads may proceed to the end of
6		the canceled region even if some threads in the team have not reached the barrier.
7 8	cancellation	An action that aborts an OpenMP region and the executing implicit and explicit task jump to at the end of the canceled region.
9 10	cancellation point	A point at which implicit and explicit tasks check if cancellation has been requested. If cancellation has been observed, they perform the <i>cancellation</i> .
11		COMMENT: Only the following are cancellation points:
12		<ul> <li>encountered implicit barriers</li> </ul>
13		• encountered barrier regions
14		• encountered cancel regions
15		<ul> <li>cancellation point regions</li> </ul>

16	1.2.4	Tasking Terminology	
17 18		task	A specific instance of executable code and its data environment, generated when a <i>thread</i> encounters a <b>task</b> <i>construct</i> or a <b>parallel</b> <i>construct</i> .
19		task region	A region consisting of all code encountered during the execution of a task.
20			COMMENT: A parallel region consists of one or more implicit task regions.
21		explicit task	A task generated when a task construct is encountered during execution.
22 23		implicit task	A <i>task</i> generated by the <i>implicit parallel region</i> or generated when a <b>parallel</b> <i>construct</i> is encountered during execution.
24		initial task	The implicit task associated with the implicit parallel region.
25 26		current task	For a given <i>thread</i> , the <i>task</i> corresponding to the <i>task region</i> in which it is executing.
27 28		child task	A task is a child task of the region of its generating task. A child task region is not part of its generating task region.
29		sibling tasks	Tasks that are child tasks of the same task region.
30 31	de	scendant task	A task that is the child task of a task region or of one of its descendant task regions.

1 2	task completion	Task completion occurs when the end of the structured block associated with the construct that generated the task is reached.
3		COMMENT: Completion of the initial task occurs at program exit.
4 5 6	task scheduling point	A point during the execution of the current <i>task region</i> at which it can be suspended to be resumed later; or the point of <i>task completion</i> , after which the executing thread may switch to a different <i>task region</i> .
7		COMMENT: For a list of task scheduling points, see Section 2.11.3 on page 94.
8	task switching	The act of a thread switching from the execution of one task to another task.
9 10	tied task	A <i>task</i> that, when its <i>task region</i> is suspended, can be resumed only by the same <i>thread</i> that suspended it. That is, the <i>task</i> is tied to that <i>thread</i> .
11 12	untied task	A <i>task</i> that, when its <i>task region</i> is suspended, can be resumed by any <i>thread</i> in the team. That is, the <i>task</i> is not tied to any <i>thread</i> .
13 14 15	undeferred task	A <i>task</i> for which execution is not deferred with respect to its generating task region. That is, its generating <i>task region</i> is suspended until execution of the <i>undeferred task</i> is completed.
16 17	included task	A <i>task</i> for which execution is sequentially included in the generating <i>task region</i> . That is, it is <i>undeferred</i> and executed immediately by the encountering thread.
18 19	merged task	A <i>task</i> whose data environment, inclusive of ICVs, is the same as that of its generating <i>task region</i> .
20	final task	A task that forces all of its child tasks to become final and included tasks.
21 22 23	task dependence	An ordering relation between two <i>sibling tasks</i> : the <i>dependent task</i> and a previously generated <i>predecessor task</i> . The <i>task dependence</i> is fulfilled when the <i>predecessor task</i> has completed.
24 25	dependent task	A <i>task</i> that because of a <i>task dependence</i> cannot be executed until its <i>predecessor tasks</i> have completed.
26	predecessor task	A task that must complete before its dependent tasks can be executed.
27	task synchronization construct	A taskwait, taskgroup, or a barrier construct.

# 28 1.2.5 Data Terminology

29 30	variable	A named data storage block, whose value can be defined and redefined during the execution of a program.
31	array section	An array section designates a subset of the elements of an array

1 2 3	private variable	With respect to a given set of <i>task regions</i> that bind to the same <b>parallel</b> <i>region</i> , a <i>variable</i> whose name provides access to a different block of storage for each <i>task region</i> .
4 5		A <i>variable</i> that is part of another variable (as an array or structure element) cannot be made private independently of other components.
6 7 8	shared variable	With respect to a given set of <i>task regions</i> that bind to the same <b>parallel</b> <i>region</i> , a <i>variable</i> whose name provides access to the same block of storage for each <i>task region</i> .
9 10 11		A <i>variable</i> that is part of another variable (as an array or structure element) cannot be <i>shared</i> independently of the other components, except for static data members of C++ classes.
12 13 14	threadprivate variable	A <i>variable</i> that is replicated, one instance per <i>thread</i> , by the OpenMP implementation. Its name then provides access to a different block of storage for each <i>thread</i> .
15 16 17		A <i>variable</i> that is part of another variable (as an array or structure element) cannot be made <i>threadprivate</i> independently of the other components, except for static data members of C++ classes.
18	threadprivate memory	The set of threadprivate variables associated with each thread.
19	data environment	The variables associated with the execution of a given region.
20	device data environment	A data environment defined by a target data or target construct.
21		
22 23	mapped variable	An original <i>variable</i> in a data environment with a corresponding <i>variable</i> in a device data environment.
24 25		COMMENT: The original and corresponding variable may share storage.

1	mappable type	A type that is valid for a mapped variable.
2		For C:
3		The type must be a complete type.
4		For C++:
5		The type must be a complete type.
6		For class types:
7 8		<ul> <li>All members functions accessed in any target region must appear in a declare target directive.</li> </ul>
9		• All data members must be non-static.
10 11		• It cannot contain virtual members.
12		For Fortran:
13		The type must be definable.
14 15		COMMENT: Pointer types are mappable but the memory block to which the pointer refers is not mapped.
16	defined	For variables, the property of having a valid value.
17		For C:
18		For the contents of variables, the property of having a valid value.
19		For C++:
20 21		For the contents of <i>variables</i> of POD (plain old data) type, the property of having a valid value.
22 23		For <i>variables</i> of non-POD class type, the property of having been constructed but not subsequently destructed.
24		For Fortran:
25 26		For the contents of <i>variables</i> , the property of having a valid value. For the allocation or association status of <i>variables</i> , the property of having a valid status.
27 28		COMMENT: Programs that rely upon <i>variables</i> that are not <i>defined</i> are <i>non-conforming programs</i> .
29	class type	For C++: Variables declared with one of the <b>class</b> , <b>struct</b> , or <b>union</b> keywords.
30	sequentially consistent atomic construct	An atomic construct for which the seq_cst clause is specified.
	non-sequentially consistent atomic	
31	construct	An atomic construct for which the seq_cst clause is not specified.

# 1 1.2.6 Implementation Terminology

2	supporting n levels of parallelism	Implies allowing an active parallel region to be enclosed by $n-1$ active parallel regions.
4	supporting the OpenMP API	Supporting at least one level of parallelism.
5	supporting nested parallelism	Supporting more than one level of parallelism.
6 7	internal control variable	A conceptual variable that specifies run-time behavior of a set of <i>threads</i> or <i>tasks</i> in an <i>OpenMP program</i> .
8 9		COMMENT: The acronym ICV is used interchangeably with the term <i>internal control variable</i> in the remainder of this specification.
10 11	compliant implementation	An implementation of the OpenMP specification that compiles and executes any <i>conforming program</i> as defined by the specification.
12 13		COMMENT: A compliant implementation may exhibit unspecified behavior when compiling or executing a non-conforming program.
14 15	unspecified behavior	A behavior or result that is not specified by the OpenMP specification or not known prior to the compilation or execution of an OpenMP program.
16		Such unspecified behavior may result from:
17 18		• Issues documented by the OpenMP specification as having <i>unspecified behavior</i> .
19		• A non-conforming program.
20		• A conforming program exhibiting an implementation defined behavior.
21	immlementation	
22 23 24	implementation defined	Behavior that must be documented by the implementation, and is allowed to vary among different <i>compliant implementations</i> . An implementation is allowed to define this behavior as <i>unspecified</i> .
25 26		COMMENT: All features that have <i>implementation defined</i> behavior are documented in Appendix E.

#### 1.3 Execution Model

The OpenMP API uses the fork-join model of parallel execution. Multiple threads of execution perform tasks defined implicitly or explicitly by OpenMP directives. The OpenMP API is intended to support programs that will execute correctly both as parallel programs (multiple threads of execution and a full OpenMP support library) and as sequential programs (directives ignored and a simple OpenMP stubs library). However, it is possible and permitted to develop a program that executes correctly as a parallel program but not as a sequential program, or that produces different results when executed as a parallel program compared to when it is executed as a sequential program. Furthermore, using different numbers of threads may result in different numeric results because of changes in the association of numeric operations. For example, a serial addition reduction may have a different pattern of addition associations than a parallel reduction. These different associations may change the results of floating-point addition.

An OpenMP program begins as a single thread of execution, called the initial thread. The initial thread executes sequentially, as if enclosed in an implicit task region, called the initial task region, that is defined by an implicit inactive **parallel** region surrounding the whole program.

The thread that executes the initial OpenMP program executes on the *host device*. An implementation may support other *target devices*. If supported, one or more devices are available to the host device for offloading code and data. Each device has its own threads that are distinct from threads that execute on another device. Threads cannot migrate from one device to another device. The execution model is host-centric such that the host device offloads target regions to target devices.

A target region begins as a single thread of execution, called the initial device thread. The initial device thread executes sequentially, as if enclosed in an implicit task region, called the initial device task region, that is defined by an implicit inactive parallel region that surrounds the entire target region.

When a target construct is encountered, the target region is executed by the implicit device task. The task that encounters the target construct waits at the end of the construct until execution of the region completes. If a target device does not exist, or the target device is not supported by the implementation, or the target device cannot execute the target construct then the target region is executed by the host device.

If a construct creates a data environment, the data environment is created at the time the construct is encountered. Whether a construct creates a data environment is defined in the description of the construct.

When any thread encounters a **parallel** construct, the thread creates a team of itself and zero or more additional threads and becomes the master of the new team. A set of implicit tasks, one per thread, is generated. The code for each task is defined by the code

inside the parallel construct. Each task is assigned to a different thread in the team and becomes tied; that is, it is always executed by the thread to which it is initially assigned. The task region of the task being executed by the encountering thread is suspended, and each member of the new team executes its implicit task. There is an implicit barrier at the end of the parallel construct. Only the master thread resumes execution beyond the end of the parallel construct, resuming the task region that was suspended upon encountering the parallel construct. Any number of parallel constructs can be specified in a single program.

parallel regions may be arbitrarily nested inside each other. If nested parallelism is disabled, or is not supported by the OpenMP implementation, then the new team that is created by a thread encountering a parallel construct inside a parallel region will consist only of the encountering thread. However, if nested parallelism is supported and enabled, then the new team can consist of more than one thread. A parallel construct may include a proc\_bind clause to specify the places to use for the threads in the team within the parallel region.

When any team encounters a worksharing construct, the work inside the construct is divided among the members of the team, and executed cooperatively instead of being executed by every thread. There is a default barrier at the end of each worksharing construct unless the **nowait** clause is present. Redundant execution of code by every thread in the team resumes after the end of the worksharing construct.

When any thread encounters a task construct, a new explicit task is generated. Execution of explicitly generated tasks is assigned to one of the threads in the current team, subject to the thread's availability to execute work. Thus, execution of the new task could be immediate, or deferred until later according to task scheduling constraints and thread availability. Threads are allowed to suspend the current task region at a task scheduling point in order to execute a different task. If the suspended task region is for a tied task, the initially assigned thread later resumes execution of the suspended task region. If the suspended task region is for an untied task, then any thread may resume its execution. Completion of all explicit tasks bound to a given parallel region is guaranteed before the master thread leaves the implicit barrier at the end of the region. Completion of a subset of all explicit tasks bound to a given parallel region may be specified through the use of task synchronization constructs. Completion of all explicit tasks bound to the implicit parallel region is guaranteed by the time the program exits.

When any thread encounters a **simd** construct, the iterations of the loop associated with the construct may be executed concurrently using the SIMD lanes that are available to the thread.

Synchronization constructs and library routines are available in the OpenMP API to coordinate tasks and data access in parallel regions. In addition, library routines and environment variables are available to control or to query the runtime environment of OpenMP programs.

The OpenMP specification makes no guarantee that input or output to the same file is synchronous when executed in parallel. In this case, the programmer is responsible for synchronizing input and output statements (or routines) using the provided synchronization constructs or library routines. For the case where each thread accesses a different file, no synchronization by the programmer is necessary.

## 1.4 Memory Model

#### 1.4.1 Structure of the OpenMP Memory Model

The OpenMP API provides a relaxed-consistency, shared-memory model. All OpenMP threads have access to a place to store and to retrieve variables, called the *memory*. In addition, each thread is allowed to have its own *temporary view* of the memory. The temporary view of memory for each thread is not a required part of the OpenMP memory model, but can represent any kind of intervening structure, such as machine registers, cache, or other local storage, between the thread and the memory. The temporary view of memory allows the thread to cache variables and thereby to avoid going to memory for every reference to a variable. Each thread also has access to another type of memory that must not be accessed by other threads, called *threadprivate memory*.

A directive that accepts data-sharing attribute clauses determines two kinds of access to variables used in the directive's associated structured block: shared and private. Each variable referenced in the structured block has an original variable, which is the variable by the same name that exists in the program immediately outside the construct. Each reference to a shared variable in the structured block becomes a reference to the original variable. For each private variable referenced in the structured block, a new version of the original variable (of the same type and size) is created in memory for each task that contains code associated with the directive. Creation of the new version does not alter the value of the original variable. However, the impact of attempts to access the original variable during the region associated with the directive is unspecified; see Section 2.14.3.3 on page 134 for additional details. References to a private variable in the structured block refer to the current task's private version of the original variable. The relationship between the value of the original variable and the initial or final value of the private version depends on the exact clause that specifies it. Details of this issue, as well as other issues with privatization, are provided in Section 2.14 on page 122.

The minimum size at which a memory update may also read and write back adjacent variables that are part of another variable (as array or structure elements) is implementation defined but is no larger than required by the base language.

A single access to a variable may be implemented with multiple load or store instructions, and hence is not guaranteed to be atomic with respect to other accesses to the same variable. Accesses to variables smaller than the implementation defined minimum size or to C or C++ bit-fields may be implemented by reading, modifying, and rewriting a larger unit of memory, and may thus interfere with updates of variables or fields in the same unit of memory.

If multiple threads write without synchronization to the same memory unit, including cases due to atomicity considerations as described above, then a data race occurs. Similarly, if at least one thread reads from a memory unit and at least one thread writes without synchronization to that same memory unit, including cases due to atomicity considerations as described above, then a data race occurs. If a data race occurs then the result of the program is unspecified.

A private variable in a task region that eventually generates an inner nested **parallel** region is permitted to be made shared by implicit tasks in the inner **parallel** region. A private variable in a task region can be shared by an explicit **task** region generated during its execution. However, it is the programmer's responsibility to ensure through synchronization that the lifetime of the variable does not end before completion of the explicit **task** region sharing it. Any other access by one task to the private variables of another task results in unspecified behavior.

#### 1.4.2 Device Data Environments

When an OpenMP program begins, each device has an initial device data environment. The initial device data environment for the host device is the data environment associated with the initial task region. Directives that accept data-mapping attribute clauses determine how an original variable is mapped to a corresponding variable in a device data environment. The original variable is the variable with the same name that exists in the data environment of the task that encounters the directive.

If a corresponding variable is present in the enclosing device data environment, the new device data environment inherits the corresponding variable from the enclosing device data environment. If a corresponding variable is not present in the enclosing device data environment, a new corresponding variable (of the same type and size) is created in the new device data environment. In the latter case, the initial value of the new corresponding variable is determined from the clauses and the data environment of the encountering thread.

The corresponding variable in the device data environment may share storage with the original variable. Writes to the corresponding variable may alter the value of the original variable. The impact of this on memory consistency is discussed in Section 1.4.4 on page 18. When a task executes in the context of a device data environment, references to the original variable refer to the corresponding variable in the device data environment.

The relationship between the value of the original variable and the initial or final value of the corresponding variable depends on the *map-type*. Details of this issue, as well as other issues with mapping a variable, are provided in Section 2.14.5 on page 151.

The original variable in a data environment and the corresponding variable(s) in one or more device data environments may share storage. Without intervening synchronization data races can occur.

#### 1.4.3 The Flush Operation

The memory model has relaxed-consistency because a thread's temporary view of memory is not required to be consistent with memory at all times. A value written to a variable can remain in the thread's temporary view until it is forced to memory at a later time. Likewise, a read from a variable may retrieve the value from the thread's temporary view, unless it is forced to read from memory. The OpenMP flush operation enforces consistency between the temporary view and memory.

The flush operation is applied to a set of variables called the *flush-set*. The flush operation restricts reordering of memory operations that an implementation might otherwise do. Implementations must not reorder the code for a memory operation for a given variable, or the code for a flush operation for the variable, with respect to a flush operation that refers to the same variable.

If a thread has performed a write to its temporary view of a shared variable since its last flush of that variable, then when it executes another flush of the variable, the flush does not complete until the value of the variable has been written to the variable in memory. If a thread performs multiple writes to the same variable between two flushes of that variable, the flush ensures that the value of the last write is written to the variable in memory. A flush of a variable executed by a thread also causes its temporary view of the variable to be discarded, so that if its next memory operation for that variable is a read, then the thread will read from memory when it may again capture the value in the temporary view. When a thread executes a flush, no later memory operation by that thread for a variable involved in that flush is allowed to start until the flush completes. The completion of a flush of a set of variables executed by a thread is defined as the point at which all writes to those variables performed by the thread before the flush are visible in memory to all other threads and that thread's temporary view of all variables involved is discarded.

The flush operation provides a guarantee of consistency between a thread's temporary view and memory. Therefore, the flush operation can be used to guarantee that a value written to a variable by one thread may be read by a second thread. To accomplish this, the programmer must ensure that the second thread has not written to the variable since its last flush of the variable, and that the following sequence of events happens in the specified order:

- 1. The value is written to the variable by the first thread.
- 2. The variable is flushed by the first thread.
- 3. The variable is flushed by the second thread.
- 4. The value is read from the variable by the second thread.

**Note** – OpenMP synchronization operations, described in Section 2.12 on page 95 and in Section 3.3 on page 196, are recommended for enforcing this order. Synchronization through variables is possible but is not recommended because the proper timing of flushes is difficult as shown in Section A.2 on page 222.

### 1.4.4 OpenMP Memory Consistency

The restrictions in Section 1.4.3 on page 17 on reordering with respect to flush operations guarantee the following:

- If the intersection of the flush-sets of two flushes performed by two different threads
  is non-empty, then the two flushes must be completed as if in some sequential order,
  seen by all threads.
- If two operations performed by the same thread either access, modify, or flush the same variable, then they must be completed as if in that thread's program order, as seen by all threads.
- If the intersection of the flush-sets of two flushes is empty, the threads can observe these flushes in any order.

The flush operation can be specified using the **flush** directive, and is also implied at various locations in an OpenMP program: see Section 2.12.7 on page 110 for details. For an example illustrating the memory model, see Section A.2 on page 222.

**Note** – Since flush operations by themselves cannot prevent data races, explicit flush operations are only useful in combination with non-sequentially consistent atomic directives.

1	OpenMP programs that:
2	• do not use non-seque
3 4	<ul> <li>do not rely on the accomp_test_nest_</li> </ul>
5	<ul> <li>correctly avoid data i</li> </ul>
6	behave as though operat
7	consistent with the orde

do not use non-sequentially consistent atomic directives,

do not rely on the accuracy of a *false* result from omp\_test\_lock and omp test nest lock, and

correctly avoid data races as required in Section 1.4.1 on page 15

behave as though operations on shared variables were simply interleaved in an order consistent with the order in which they are performed by each thread. The relaxed consistency model is invisible for such programs, and any explicit flush operations in such programs are redundant.

Implementations are allowed to relax the ordering imposed by implicit flush operations when the result is only visible to programs using non-sequentially consistent atomic directives.

## 1.5 OpenMP Compliance

An implementation of the OpenMP API is compliant if and only if it compiles and executes all conforming programs according to the syntax and semantics laid out in Chapters 1, 2, 3 and 4. Appendices A, B, C, D, E and F and sections designated as Notes (see Section 1.7 on page 21) are for information purposes only and are not part of the specification.

The OpenMP API defines constructs that operate in the context of the base language that is supported by an implementation. If the base language does not support a language construct that appears in this document, a compliant OpenMP implementation is not required to support it, with the exception that for Fortran, the implementation must allow case insensitivity for directive and API routines names, and must allow identifiers of more than six characters.

All library, intrinsic and built-in routines provided by the base language must be thread-safe in a compliant implementation. In addition, the implementation of the base language must also be thread-safe. For example, **ALLOCATE** and **DEALLOCATE** statements must be thread-safe in Fortran. Unsynchronized concurrent use of such routines by different threads must produce correct results (although not necessarily the same as serial execution results, as in the case of random number generation routines).

In both Fortran 90 and Fortran 95, variables with explicit initialization have the **SAVE** attribute implicitly. This is not the case in Fortran 77. However, a compliant OpenMP Fortran implementation must give such a variable the **SAVE** attribute, regardless of the underlying base language version.

#### 1.6 Normative References

ISO/IEC 9899:1990, Information Technology - Programming Languages - C.
 This OpenMP API specification refers to ISO/IEC 9899:1990 as C90.

ISO/IEC 9899:1999, Information Technology - Programming Languages - C.
 This OpenMP API specification refers to ISO/IEC 9899:1999 as C99.

• ISO/IEC 14882:1998, *Information Technology - Programming Languages - C++*.

This OpenMP API specification refers to ISO/IEC 14882:1998 as C++.

ISO/IEC 1539:1980, Information Technology - Programming Languages - Fortran.
 This OpenMP API specification refers to ISO/IEC 1539:1980 as Fortran 77.

• ISO/IEC 1539:1991, *Information Technology - Programming Languages - Fortran*.

This OpenMP API specification refers to ISO/IEC 1539:1991 as Fortran 90.

ISO/IEC 1539-1:1997, Information Technology - Programming Languages - Fortran.
 This OpenMP API specification refers to ISO/IEC 1539-1:1997 as Fortran 95.

• ISO/IEC 1539-1:2004, *Information Technology - Programming Languages - Fortran*.

This OpenMP API specification refers to ISO/IEC 1539-1:2004 as Fortran 2003. The

following features are not supported:

• IEEE Arithmetic issues covered in Fortran 2003 Section 14

• Parameterized derived types

Allocatable enhancement

1	<ul> <li>Finalization</li> </ul>
2	<ul> <li>Procedures bound by name to a type</li> </ul>
3	• The PASS attribute
4	<ul> <li>Procedures bound to a type as operators</li> </ul>
5	Type extension
6	Overriding a type-bound procedure
7	<ul> <li>Polymorphic entities</li> </ul>
8	• SELECT TYPE construct
9	<ul> <li>Deferred bindings and abstract types</li> </ul>
10	Controlling IEEE underflow
11	Another IEEE class value
12	Where this OpenMP API specification refers to C, C++ or Fortran, reference is made to
13	the base language supported by the implementation.

## 14 1.7 Organization of this document

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15	The remainder of this document is structured as follows:
16	• Chapter 2: Directives
17	Chapter 3: Runtime Library Routines
18	Chapter 4: Environment Variables
19	Appendix A: Examples
20	Appendix B: Stubs for Runtime Library Routines
21	<ul> <li>Appendix C: OpenMP C and C++ Grammar</li> </ul>
22	Appendix D: Interface Declarations
23	Appendix E: OpenMP Implementation Defined Behaviors
24	Appendix F: Features History
25 26 27	Some sections of this document only apply to programs written in a certain base language. Text that applies only to programs whose base language is C or C++ is shown as follows:
28	C/C++ specific text  C/C++

Text that applies only to programs whose base language is C only is shown as follows:

4	Constitute to
1	C specific text
2	Text that applies only to programs whose base language is C++ only is shown as
3	follows:
	C++
4	C++ specific text
	C++
5	Text that applies only to programs whose base language is Fortran is shown as follows
	Fortran
6	Fortran specific text
	Fortran —
7	Where an entire page consists of, for example, Fortran specific text, a marker is shown
8	at the top of the page like this:
	Fortran (cont.)
9	Some text is for information only, and is not part of the normative specification. Such
0	text is designated as a note, like this:
1	▼ Note - Non-normative text
•	A

## 1 CHAPTER **2**

# **Directives**

3 4	This chapter describes the syntax and behavior of OpenMP directives, and is divided into the following sections:
5	• The language-specific directive format (Section 2.1 on page 24)
6	<ul> <li>Mechanisms to control conditional compilation (Section 2.2 on page 28)</li> </ul>
7	• Control of OpenMP API ICVs (Section 2.3 on page 30)
8	<ul> <li>Details of each OpenMP directive (Section 2.5 on page 37 to Section 2.16 on page 158)</li> </ul>
	C/C++
10 11	In C/C++, OpenMP directives are specified by using the <b>#pragma</b> mechanism provided by the C and C++ standards.
	C/C++
	Fortran —
12 13 14	In Fortran, OpenMP directives are specified by using special comments that are identified by unique sentinels. Also, a special comment form is available for conditiona compilation.
	Fortran —
15 16 17 18 19 20	Compilers can therefore ignore OpenMP directives and conditionally compiled code if support of the OpenMP API is not provided or enabled. A compliant implementation must provide an option or interface that ensures that underlying support of all OpenMI directives and OpenMP conditional compilation mechanisms is enabled. In the remainder of this document, the phrase <i>OpenMP compilation</i> is used to mean a compilation with these OpenMP features enabled.
	•

#### Restrictions

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20 21 The following restriction applies to all OpenMP directives:

• OpenMP directives may not appear in **PURE** or **ELEMENTAL** procedures.

——— Fortran —

Fortran -

#### 4 2.1 Directive Format

#### - C/C++ -

OpenMP directives for C/C++ are specified with the **pragma** preprocessing directive. The syntax of an OpenMP directive is formally specified by the grammar in Appendix C, and informally as follows:

#pragma omp directive-name [clause[ [,] clause]...] new-line

Each directive starts with **#pragma omp**. The remainder of the directive follows the conventions of the C and C++ standards for compiler directives. In particular, white space can be used before and after the **#**, and sometimes white space must be used to separate the words in a directive. Preprocessing tokens following the **#pragma omp** are subject to macro replacement.

Some OpenMP directives may be composed of consecutive OpenMP directives if specified in their syntax.

Directives are case-sensitive.

An OpenMP executable directive applies to at most one succeeding statement, which must be a structured block.

- C/C++ -

----- Fortran

OpenMP directives for Fortran are specified as follows:

sentinel directive-name [clause[[,] clause]...]

All OpenMP compiler directives must begin with a directive *sentinel*. The format of a sentinel differs between fixed and free-form source files, as described in Section 2.1.1 on page 25 and Section 2.1.2 on page 26.

Directives are case-insensitive. Directives cannot be embedded within continued 1 2 statements, and statements cannot be embedded within directives. 3 In order to simplify the presentation, free form is used for the syntax of OpenMP directives for Fortran in the remainder of this document, except as noted. 4 Fortran -Only one directive-name can be specified per directive (note that this includes combined 5 directives, see Section 2.10 on page 82). The order in which clauses appear on directives 6 is not significant. Clauses on directives may be repeated as needed, subject to the 7 restrictions listed in the description of each clause. 8 9 Some data-sharing attribute clauses (Section 2.14.3 on page 130), data copying clauses (Section 2.14.4 on page 147), the threadprivate directive (Section 2.14.2 on page 10 126) and the **flush** directive (Section 2.12.7 on page 110) accept a list. A list consists 11 12 of a comma-separated collection of one or more *list items*. \_\_\_\_ C/C++ \_\_\_\_ A list item is a variable name, subject to the restrictions specified in each of the sections 13 14 describing clauses and directives for which a *list* appears. C/C++ ----- Fortran A list item is a variable name or a common block name (enclosed in slashes), subject to 15 16 the restrictions specified in each of the sections describing clauses and directives for which a *list* appears. 17 ——— Fortran ———— 18 ----- Fortran -**Fixed Source Form Directives** 2.1.1 20 The following sentinels are recognized in fixed form source files: qmo\$! c\$omp | \*\$omp 21 Sentinels must start in column 1 and appear as a single word with no intervening characters. Fortran fixed form line length, white space, continuation, and column rules 22 apply to the directive line. Initial directive lines must have a space or zero in column 6, 23 24 and continuation directive lines must have a character other than a space or a zero in

25

column 6.

Comments may appear on the same line as a directive. The exclamation point initiates a comment when it appears after column 6. The comment extends to the end of the source line and is ignored. If the first non-blank character after the directive sentinel of an initial or continuation directive line is an exclamation point, the line is ignored.

Note – in the following example, the three formats for specifying the directive are equivalent (the first line represents the position of the first 9 columns):

8 c23456789

!\$omp parallel do shared(a,b,c)

c\$omp parallel do
c\$omp+shared(a,b,c)

c\$omp paralleldoshared(a,b,c)

#### 15 2.1.2 Free Source Form Directives

The following sentinel is recognized in free form source files:

!\$omp

The sentinel can appear in any column as long as it is preceded only by white space (spaces and tab characters). It must appear as a single word with no intervening character. Fortran free form line length, white space, and continuation rules apply to the directive line. Initial directive lines must have a space after the sentinel. Continued directive lines must have an ampersand (&) as the last nonblank character on the line, prior to any comment placed inside the directive. Continuation directive lines can have an ampersand after the directive sentinel with optional white space before and after the ampersand.

Comments may appear on the same line as a directive. The exclamation point (1) initiates a comment. The comment extends to the end of the source line and is ignored. If the first nonblank character after the directive sentinel is an exclamation point, the line is ignored.

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One or more blanks or horizontal tabs must be used to separate adjacent keywords in directives in free source form, except in the following cases, where white space is optional between the given pair of keywords:

end atomic end critical end do end do simd end master end ordered end parallel end parallel do end parallel do simd end sections end simd end single end task end taskgroup end workshare parallel do parallel sections

parallel workshare

```
Note — in the following example the three formats for specifying the directive are equivalent (the first line represents the position of the first 9 columns):

123456789

150mp parallel do &

150mp shared(a,b,c)

161

172

185mp parallel &

185mp shared(a,b,c)
```

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Fortran -

# 2.2 Conditional Compilation

In implementations that support a preprocessor, the \_OPENMP macro name is defined to have the decimal value *yyyymm* where *yyyy* and *mm* are the year and month designations of the version of the OpenMP API that the implementation supports.

If this macro is the subject of a **#define** or a **#undef** preprocessing directive, the behavior is unspecified.

For examples of conditional compilation, see Section A.3 on page 229.

#### - Fortran -

The OpenMP API requires Fortran lines to be compiled conditionally, as described in the following sections.

# 2.2.1 Fixed Source Form Conditional Compilation Sentinels

The following conditional compilation sentinels are recognized in fixed form source files:

```
!$ | *$ | c$
```

To enable conditional compilation, a line with a conditional compilation sentinel must satisfy the following criteria:

- The sentinel must start in column 1 and appear as a single word with no intervening white space.
- After the sentinel is replaced with two spaces, initial lines must have a space or zero in column 6 and only white space and numbers in columns 1 through 5.
- After the sentinel is replaced with two spaces, continuation lines must have a character other than a space or zero in column 6 and only white space in columns 1 through 5.

1 If these criteria are met, the sentinel is replaced by two spaces. If these criteria are not 2 met, the line is left unchanged. 3 4 5 ▼------Fortran (cont.) --------6 7 **Note** – in the following example, the two forms for specifying conditional compilation in fixed source form are equivalent (the first line represents the position of the first 9 8 9 columns): 10 c23456789 11 !\$ 10 iam = omp get thread num() + 12 index 13 14 15 #ifdef OPENMP 16 10 iam = omp get thread num() + 17 & index 18 #endif

# 19 2.2.2 Free Source Form Conditional Compilation Sentinel

The following conditional compilation sentinel is recognized in free form source files:

```
!$
```

To enable conditional compilation, a line with a conditional compilation sentinel must satisfy the following criteria:

- The sentinel can appear in any column but must be preceded only by white space.
- The sentinel must appear as a single word with no intervening white space.
- Initial lines must have a space after the sentinel.

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 Continued lines must have an ampersand as the last nonblank character on the line, prior to any comment appearing on the conditionally compiled line. Continued lines can have an ampersand after the sentinel, with optional white space before and after the ampersand.

If these criteria are met, the sentinel is replaced by two spaces. If these criteria are not met, the line is left unchanged.

**Note** – in the following example, the two forms for specifying conditional compilation in free source form are equivalent (the first line represents the position of the first 9 columns):

```
c23456789
!$ iam = omp_get_thread_num() + &
!$& index

#ifdef _OPENMP
    iam = omp_get_thread_num() + &
        index

#endif
```

Fortran

# 2.3 Internal Control Variables

An OpenMP implementation must act as if there were internal control variables (ICVs) that control the behavior of an OpenMP program. These ICVs store information such as the number of threads to use for future <code>parallel</code> regions, the schedule to use for worksharing loops and whether nested parallelism is enabled or not. The ICVs are given values at various times (described below) during the execution of the program. They are initialized by the implementation itself and may be given values through OpenMP environment variables and through calls to OpenMP API routines. The program can retrieve the values of these ICVs only through OpenMP API routines.

For purposes of exposition, this document refers to the ICVs by certain names, but an implementation is not required to use these names or to offer any way to access the variables other than through the ways shown in Section 2.3.2 on page 32.

# 1 2.3.1 ICV Descriptions

data environment.

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4 5 6	• <i>dyn-var</i> - controls whether dynamic adjustment of the number of threads is enabled for encountered <b>parallel</b> regions. There is one copy of this ICV per data environment.
7 8	• <i>nest-var</i> - controls whether nested parallelism is enabled for encountered <b>parallel</b> regions. There is one copy of this ICV per data environment.
9 10	• <i>nthreads-var</i> - controls the number of threads requested for encountered <b>parallel</b> regions. There is one copy of this ICV per data environment.
11 12	• <i>thread-limit-var</i> - controls the maximum number of threads participating in the OpenMP program. There is one copy of this ICV per device.
13 14	• <i>max-active-levels-var</i> - controls the maximum number of nested active <b>parallel</b> regions. There is one copy of this ICV per device.
15 16 17	• place-partition-var — controls the place partition available to the execution environment for encountered parallel regions. There is one copy of this ICV per implicit task.
18	The following ICVs store values that affect the operation of loop regions.
19 20	• run-sched-var - controls the schedule that the runtime schedule clause uses for loop regions. There is one copy of this ICV per data environment.
21 22	• <i>def-sched-var</i> - controls the implementation defined default scheduling of loop regions. There is one copy of this ICV per device.
23	The following ICVs store values that affect the program execution.
24 25 26 27	• <i>bind-var</i> - controls the binding of OpenMP threads to places. When binding is requested, the variable indicates that the execution environment is advised not to move threads between places. The variable can also provide default thread affinity policies. There is one copy of this ICV per data environment.
28 29	• <i>stacksize-var</i> - controls the stack size for threads that the OpenMP implementation creates. There is one copy this ICV per device.
30 31	• wait-policy-var - controls the desired behavior of waiting threads. There is one copy of this ICV per device.
32 33	• <i>cancel-var</i> - controls the desired behavior of the <b>cancel</b> construct and cancellation points. There is one copy of the ICV for the whole program.

• default-device-var - controls the default device. There is one copy of this ICV per

The following ICVs store values that affect the operation of parallel regions.

# Modifying and Retrieving ICV Values

The following table shows the methods for retrieving the values of the ICVs as well as their initial values:

ICV	Scope	Ways to modify value	Way to retrieve value	Initial value
dyn-var	data environment	OMP_DYNAMIC omp_set_dynamic()	omp_get_dynamic()	See comments below
nest-var	data environment	OMP_NESTED omp_set_nested()	<pre>omp_get_nested()</pre>	false
nthreads-var	data environment	OMP_NUM_THREADS omp_set_num_threads()	<pre>omp_get_max_threads()</pre>	Implementation defined
run-sched-var	data environment	OMP_SCHEDULE omp_set_schedule()	<pre>omp_get_schedule()</pre>	Implementation defined
def-sched-var	device	(none)	(none)	Implementation defined
bind-var	data environment	OMP_PROC_BIND	<pre>omp_get_proc_bind()</pre>	Implementation defined
stacksize-var	device	OMP_STACKSIZE	(none)	Implementation defined
wait-policy-var	device	OMP_WAIT_POLICY	(none)	Implementation defined
thread-limit-var	device	OMP_THREAD_LIMIT	<pre>omp_get_thread_limit()</pre>	Implementation defined
max-active-levels-var	device	OMP_MAX_ACTIVE_LEVELS omp_set_max_active_ levels()	<pre>omp_get_max_active_ levels()</pre>	See comments below
place-partition-var	implicit task	OMP_PLACES	(none)	Implementation defined
cancel-var	global	OMP_CANCELLATION	<pre>omp_get_cancellation()</pre>	false
default-device-var	data environment	OMP_DEFAULT-DEVICE omp_set_default_ device()	<pre>omp_get_default_ device()</pre>	Implementation defined

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#### Comments:

- There is one copy per device of each ICV with device scope.
- The value of the nthreads-var ICV is a list. The runtime call omp\_set\_num\_threads() sets the value of the first element of this list, and omp\_get\_max\_threads() retrieves the value of the first element of this list.
- The value of the *bind-var* ICV is a list. The runtime call <code>omp\_get\_proc\_bind()</code> retrieves the value of the first element of this list.
- The initial value of *dyn-var* is implementation defined if the implementation supports dynamic adjustment of the number of threads; otherwise, the initial value is *false*.

The initial value of *max-active-levels-var* is the number of levels of parallelism that the implementation supports. See the definition of *supporting n levels of parallelism* in Section 1.2.6 on page 12 for further details.

After the initial values are assigned, but before any OpenMP construct or OpenMP API routine executes, the values of any OpenMP environment variables that were set by the user are read and the associated ICVs are modified accordingly. After this point, no changes to any OpenMP environment variables will affect the ICVs.

Clauses on OpenMP constructs do not modify the values of any of the ICVs.

# 2.3.3 How the ICVs Work in target Regions

Each device has its own copy of the ICVs. When a **teams**, **target** or **target data** construct is encountered, the device data environment inherits the values of *dyn-var*, *nest-var*, *run-sched-var*, *nthreads-var* and *default-device-var* from the generating task's ICV values.

## 14 2.3.4 How the Per-Data Environment ICVs Work

Each data environment has its own copies of internal variables *bind-var*, *dyn-var*, *nest-var*, *nthreads-var*, *run-sched-var*. In addition, each implicit task has its own copy of the internal variable *place-partition-var*.

Calls to omp\_set\_num\_threads(), omp\_set\_dynamic(), omp\_set\_nested(), and omp\_set\_schedule() modify only the ICVs in the data environment of their binding task.

When a **task** construct or **parallel** construct is encountered, the generated task(s) inherit the values of *dyn-var*, *nest-var*, and *run-sched-var* from the generating task's ICV values.

When a task construct is encountered, the generated task inherits the value of *nthreads-var* from the generating task's *nthreads-var* value. When a parallel construct is encountered, and the generating task's *nthreads-var* list contains a single element, the generated task(s) inherit that list as the value of *nthreads-var*. When a parallel construct is encountered, and the generating task's *nthreads-var* list contains multiple elements, the generated task(s) inherit the value of *nthreads-var* as the list obtained by deletion of the first element from the generating task's *nthreads-var* value. The *bind-var* ICV is handled in the same way as the *nthreads-var* ICV.

When a **teams**, **target**, or **target data** construct is encountered, the generated data environment(s) inherit the values of *dyn-var*, *nest-var*, *run-sched-var*, *nthreads-var* and *default-device-var* from the data environment of the generating task's ICV values.

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# 2.3.5 ICV Override Relationships

The override relationships among various construct clauses, OpenMP API routines, environment variables, and the initial values of ICVs are shown in the following table:

construct clause, if used	overrides call to API routine	overrides setting of environment variable	overrides initial value of
(none)	<pre>omp_set_dynamic()</pre>	OMP_DYNAMIC	dyn-var
(none)	<pre>omp_set_nested()</pre>	OMP_NESTED	nest-var
${\tt num\_threads}$	<pre>omp_set_num_threads()</pre>	OMP_NUM_THREADS	nthreads-var*
schedule	omp_set_schedule()	OMP_SCHEDULE	run-sched-var
(none)	(none)	OMP_PROC_BIND	bind-var
schedule	(none)	(none)	def-sched-var
(none)	(none)	OMP_STACKSIZE	stacksize-var
(none)	(none)	OMP_WAIT_POLICY	wait-policy-var
(none)	(none)	OMP_THREAD_LIMIT	thread-limit-var
(none)	<pre>omp_set_max_active_levels()</pre>	OMP_MAX_ACTIVE_LEVELS	max-active-levels-var
proc_bind	(none)	OMP_PROC_BIND	bind-var**
(none)	(none)	OMP_PLACES	place-partition-var
(none)	(none)	OMP CANCELLATION	cancel-var

- \* The num\_threads clause and omp\_set\_num\_threads() override the value of the OMP\_NUM\_THREADS environment variable and the initial value of the first element of the nthreads-var ICV.
- \*\* The proc\_bind clause overrides the value of the OMP\_PROC\_BIND environment variable and the initial value of the first element of the *bind-var* ICV.

#### **Cross References:**

- parallel construct, see Section 2.5 on page 37.
- num threads clause, see Section 2.5.1 on page 40.
- proc bind clause, Section 2.5 on page 37

1 • Loop construct, see Section 2.7.1 on page 47. 2 **schedule** clause, see Section 2.7.1.1 on page 52. 3 omp set num threads routine, see Section 3.2.1 on page 162. 4 omp get max threads routine, see Section 3.2.3 on page 165. 5 omp set dynamic routine, see Section 3.2.7 on page 169. 6 • omp get dynamic routine, see Section 3.2.8 on page 171. • omp set nested routine, see Section 3.2.10 on page 173. 7 8 • omp get nested routine, see Section 3.2.11 on page 174. 9 omp set schedule routine, see Section 3.2.12 on page 176. 10 • omp get schedule routine, see Section 3.2.13 on page 178. 11 omp get thread limit routine, see Section 3.2.14 on page 179. 12 omp set max active levels routine, see Section 3.2.15 on page 180. 13 omp get max active levels routine, see Section 3.2.16 on page 182. 14 **OMP SCHEDULE** environment variable, see Section 4.1 on page 210. 15 **OMP NUM THREADS** environment variable, see Section 4.2 on page 211. **OMP DYNAMIC** environment variable, see Section 4.3 on page 212. 16 17 **OMP PROC BIND** environment variable, see Section 4.4 on page 213 18 **OMP PLACES** environment variable, see Section 4.5 on page 213 19 **OMP NESTED** environment variable, see Section 4.6 on page 215. 20 **OMP STACKSIZE** environment variable, see Section 4.7 on page 216. 21 • OMP WAIT POLICY environment variable, see Section 4.8 on page 217. 22 • OMP MAX ACTIVE LEVELS environment variable, see Section 4.9 on page 217. 23 • OMP THREAD LIMIT environment variable, see Section 4.10 on page 218.

• OMP CANCELLATION environment variable, see Section 4.11 on page 218

# 2.4 Array Sections

An array section designates a subset of the elements in an array. An array section can appear only in clauses where it is explicitly allowed.

—— C/C++ —

To specify an array section in an OpenMP construct, array subscript notation is extended with the following syntax:

[ lower-bound : length ] or

[ lower-bound : ] or

[ : *length* ] or

[:]

The *lower-bound* and *length* are integral type expressions. When evaluated they represent a set of integer values as follows:

 $\{ lower-bound, lower-bound + 1, lower-bound + 2, ..., lower-bound + length - 1 \}$ 

The lower-bound and length must evaluate to non-negative integers.

When the *lower-bound* is absent it defaults to 0.

When the size of the array dimension is not known, the *length* must be specified explicitly.

When the *length* is absent, it defaults to the size of the array dimension minus the *lower-bound*.

The array section must be a subset of the original array.

Array sections are allowed on multidimensional arrays.

C/C++

Fortran

Fortran has built-in support for array sections but the following restrictions apply for OpenMP constructs:

- Only unit stride is supported.
- The upper bound for the last dimension of an assumed-size dummy array must be specified.

Fortran ———

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# 1 2.5 parallel Construct

Summary 2 3 This fundamental construct starts parallel execution. See Section 1.3 on page 13 for a general description of the OpenMP execution model. 4 **Syntax** 5 6 The syntax of the parallel construct is as follows: #pragma omp parallel [clause[[, ]clause]...] new-line structured-block 7 where *clause* is one of the following: if (scalar-expression) num threads (integer-expression) default(shared | none) private(list) firstprivate(list) shared(list) copyin (list) reduction(operator: list) proc bind(master | close | spread) C/C++ 8 Fortran 9 The syntax of the **parallel** construct is as follows: !\$omp parallel [clause[[,] clause]...] structured-block !\$omp end parallel

where *clause* is one of the following:

```
1
```

```
if (scalar-logical-expression)
num_threads (scalar-integer-expression)
default(private | firstprivate | shared | none)
private (list)
firstprivate (list)
shared (list)
copyin (list)
reduction ({operator | intrinsic_procedure_name} : list)
proc_bind(master | close | spread)
```

The end parallel directive denotes the end of the parallel construct.

Fortran -

## Binding

The binding thread set for a **parallel** region is the encountering thread. The encountering thread becomes the master thread of the new team.

## Description

When a thread encounters a **parallel** construct, a team of threads is created to execute the **parallel** region (see Section 2.5.1 on page 40 for more information about how the number of threads in the team is determined, including the evaluation of the **if** and **num\_threads** clauses). The thread that encountered the **parallel** construct becomes the master thread of the new team, with a thread number of zero for the duration of the new **parallel** region. All threads in the new team, including the master thread, execute the region. Once the team is created, the number of threads in the team remains constant for the duration of that **parallel** region.

The optional proc\_bind clause, described in Section 2.5.2 on page 42, specifies the mapping of OpenMP threads to places within the current place partition, that is, within the places listed in the *place-partition-var* ICV for the current thread.

Within a parallel region, thread numbers uniquely identify each thread. Thread numbers are consecutive whole numbers ranging from zero for the master thread up to one less than the number of threads in the team. A thread may obtain its own thread number by a call to the omp get thread num library routine.

1 A set of implicit tasks, equal in number to the number of threads in the team, is 2 generated by the encountering thread. The structured block of the parallel construct 3 determines the code that will be executed in each implicit task. Each task is assigned to 4 a different thread in the team and becomes tied. The task region of the task being 5 executed by the encountering thread is suspended and each thread in the team executes 6 its implicit task. Each thread can execute a path of statements that is different from that 7 of the other threads. 8 The implementation may cause any thread to suspend execution of its implicit task at a 9 task scheduling point, and switch to execute any explicit task generated by any of the 10 threads in the team, before eventually resuming execution of the implicit task (for more details see Section 2.11 on page 88). 11 12 There is an implied barrier at the end of a parallel region. After the end of a 13 parallel region, only the master thread of the team resumes execution of the 14 enclosing task region. 15 If a thread in a team executing a parallel region encounters another parallel 16 directive, it creates a new team, according to the rules in Section 2.5.1 on page 40, and it becomes the master of that new team. 17 18 If execution of a thread terminates while inside a parallel region, execution of all 19 threads in all teams terminates. The order of termination of threads is unspecified. All work done by a team prior to any barrier that the team has passed in the program is 20 21 guaranteed to be complete. The amount of work done by each thread after the last 22 barrier that it passed and before it terminates is unspecified. 23 For an example of the parallel construct, see Section A.5 on page 232. For an example of the **num threads** clause, see Section A.7 on page 237. 24 Restrictions 25 26 Restrictions to the parallel construct are as follows:

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- A program that branches into or out of a parallel region is non-conforming.
- A program must not depend on any ordering of the evaluations of the clauses of the parallel directive, or on any side effects of the evaluations of the clauses.
- At most one if clause can appear on the directive.
- At most one **proc** bind clause can appear on the directive.
- At most one num threads clause can appear on the directive. The num threads expression must evaluate to a positive integer value.

3 4 5 6 7 8 9 9 9 9 1 1 1 2 2	When execution encounters a parallel directive, the value of the if clause or num_threads clause (if any) on the directive, the current parallel context, and the values of the nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs are used to determine the number of threads to use in the region.  Note that using a variable in an if or num_threads clause expression of a parallel construct causes an implicit reference to the variable in all enclosing constructs. The if clause expression and the num_threads clause expression are evaluated in the context outside of the parallel construct, and no ordering of those evaluations is specified. It is also unspecified whether, in what order, or how many times any side-effects of the evaluation of the num_threads or if clause expressions occur.  When a thread encounters a parallel construct, the number of threads is determined according to Algorithm 2.1.  Algorithm 2.1
4 5 6 7 8 9 20 21 22	num_threads clause (if any) on the directive, the current parallel context, and the values of the nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs are used to determine the number of threads to use in the region. Note that using a variable in an if or num_threads clause expression of a parallel construct causes an implicit reference to the variable in all enclosing constructs. The if clause expression and the num_threads clause expression are evaluated in the context outside of the parallel construct, and no ordering of those evaluations is specified. It is also unspecified whether, in what order, or how many times any side-effects of the evaluation of the num_threads or if clause expressions occur. When a thread encounters a parallel construct, the number of threads is determined
4 5 6 7 8 9 20	num_threads clause (if any) on the directive, the current parallel context, and the values of the nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs are used to determine the number of threads to use in the region. Note that using a variable in an if or num_threads clause expression of a parallel construct causes an implicit reference to the variable in all enclosing constructs. The if clause expression and the num_threads clause expression are evaluated in the context outside of the parallel construct, and no ordering of those evaluations is specified. It is also unspecified whether, in what order, or how many times
4 5 6 7 8 9	num_threads clause (if any) on the directive, the current parallel context, and the values of the nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs are used to determine the number of threads to use in the region. Note that using a variable in an if or num_threads clause expression of a parallel construct causes an implicit reference to the variable in all enclosing constructs. The if clause expression and the num_threads clause expression are evaluated in the context outside of the parallel construct, and no ordering of those
4 5 6 7 8	<ul> <li>num_threads clause (if any) on the directive, the current parallel context, and the values of the nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs are used to determine the number of threads to use in the region.</li> <li>Note that using a variable in an if or num_threads clause expression of a parallel construct causes an implicit reference to the variable in all enclosing</li> </ul>
4 5 6	<ul> <li>num_threads clause (if any) on the directive, the current parallel context, and the values of the nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs are used to determine the number of threads to use in the region.</li> <li>Note that using a variable in an if or num_threads clause expression of a</li> </ul>
4 5	num_threads clause (if any) on the directive, the current parallel context, and the values of the <i>nthreads-var</i> , <i>dyn-var</i> , <i>thread-limit-var</i> , <i>max-active-level-var</i> , and <i>nest-var</i>
4	num_threads clause (if any) on the directive, the current parallel context, and the
1 <b>2.5.1</b>	Determining the Number of Threads for a parallel Region
0	• omp_get_thread_num routine, see Section 3.2.4 on page 166.
9	• copyin clause, see Section 2.14.4 on page 147.
7 8	• default, shared, private, firstprivate, and reduction clauses, see Section 2.14.3 on page 130.
6	Cross References
	Fortran —
5	has unspecified behavior.
4	<ul> <li>Unsynchronized use of Fortran I/O statements by multiple threads on the same unit</li> </ul>
	C/C++
3	must catch it.  C/C++
1 2 3	

let ActiveParRegions be the number of enclosing active parallel regions;

if an if clause exists

```
then let IfClauseValue = true;
else let IfClauseValue = true;
if a num_threads clause exists
then let ThreadsRequested be the value of the num_threads clause
expression;
else let ThreadsRequested = value of the first element of nthreads-var;
let ThreadsAvailable = (thread-limit-var - ThreadsBusy + 1);
if (IfClauseValue = false)
then number of threads = 1;
else if (ActiveParRegions >= 1) and (nest-var = false)
then number of threads = 1;
else if (ActiveParRegions = max-active-levels-var)
then number of threads = 1;
else if (dyn-var = true) and (ThreadsRequested <= ThreadsAvailable)
then number of threads = [ 1 : ThreadsRequested ];</pre>
```

**else if** (dyn-var = true) **and** (ThreadsRequested > ThreadsAvailable)

**else if** (dyn-var = false) **and** (ThreadsRequested <= ThreadsAvailable)

**else if** (dyn-var = false) **and** (ThreadsRequested > ThreadsAvailable)

**then** number of threads = [1: ThreadsAvailable];

**then** number of threads = *ThreadsRequested*;

then behavior is implementation defined;

**Note** – Since the initial value of the *dyn-var* ICV is implementation defined, programs that depend on a specific number of threads for correct execution should explicitly disable dynamic adjustment of the number of threads.

#### **Cross References**

• nthreads-var, dyn-var, thread-limit-var, max-active-level-var, and nest-var ICVs, see Section 2.3 on page 30.

# 2.5.2 Controlling OpenMP Thread Affinity

When creating a team for a parallel region, the **proc\_bind** clause specifies a policy for assigning OpenMP threads to places within the current place partition, that is, the places listed in the *place-partition-var* ICV for the current thread. Once a thread is assigned to a place, the OpenMP implementation should not move it to another place.

The **master** thread affinity policy instructs the execution environment to assign every thread in the team to the same place as the master thread. The place partition is not changed by this policy, and each implicit task inherits the *place-partition-var* ICV of the parent implicit task.

The **close** thread affinity policy instructs the execution environment to assign the threads to places close to the place of the parent thread. The master thread executes on the parent's place and the remaining threads in the team execute on places from the place list consecutive from the parent's position in the list, with wrap around with respect to the encountering thread's place partition. The place partition is not changed by this policy, and each implicit task inherits the *place-partition-var* ICV of the parent implicit task.

The purpose of the **spread** thread affinity policy is to create a sparse distribution for a team of T threads among the P places of the parent's place partition. It accomplishes this by first subdividing the parent partition into T subpartitions if T is less than or equal to P, or P subpartitions if T is greater than P. Then it assigns 1 (T <= P) or a set of threads (T > P) to each subpartition. The place-partition-var ICV of each thread is set to its subpartition. The subpartitioning is not only a mechanism for achieving a sparse distribution, it is also a subset of places for a thread to use when creating a nested parallel region.

- *T*<=*P*. The parent's partition is split into *T* subpartitions, where each subpartition contains at least *S*=*floor*(*P*/*T*) consecutive places. A single thread is assigned to each subpartition. The master thread executes on the place of the parent thread and is assigned to the subpartition that includes that place. For the other threads, assignment is to the first place in the corresponding subpartition. When *T* does not divide *P* evenly, the assignment of the remaining *P*-*T*\**S* places into subpartitions is implementation defined.
- T>P. The parent's partition is split into P unit-sized subpartitions. Each place is assigned S=floor(T/P) threads. When P does not divide T evenly, the assignment of the remaining T-P\*S threads into places is implementation defined.

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For the close and spread thread affinity policies, the threads with the smallest thread numbers execute on the place of the master thread, then the threads with the next smaller thread numbers execute on the next place in the partition; and so on, with wrap around with respect to the encountering thread's place partition.

The determination of whether the affinity request can be fulfilled is implementation defined. If not, the number of threads in the team and their mapping to places become implementation defined.

# 8 2.6 Canonical Loop Form

C/C++

A loop has *canonical loop form* if it conforms to the following:

```
for (init-expr; test-expr; incr-expr) structured-block
init-expr
                  One of the following:
                     var = lb
                     integer-type\ var = lb
                     random-access-iterator-type var = lb
                     pointer-type var = lb
test-expr
                  One of the following:
                    var relational-op b
                    b relational-op var
incr-expr
                  One of the following:
                     ++var
                     var++
                     - - var
                     var--
                     var += incr
                     var -= incr
                     var = var + incr
                     var = incr + var
                     var = var - incr
```

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15 16 lb and b

Loop invariant expressions of a type compatible with the type of *yar*.

incr

A loop invariant integer expression.

The canonical form allows the iteration count of all associated loops to be computed before executing the outermost loop. The computation is performed for each loop in an integer type. This type is derived from the type of *var* as follows:

- If var is of an integer type, then the type is the type of var.
- For C++, if *var* is of a random access iterator type, then the type is the type that would be used by *std::distance* applied to variables of the type of *var*.
- For C, if var is of a pointer type, then the type is ptrdiff\_t.

The behavior is unspecified if any intermediate result required to compute the iteration count cannot be represented in the type determined above.

There is no implied synchronization during the evaluation of the *lb*, *b*, or *incr* expressions. It is unspecified whether, in what order, or how many times any side effects within the *lb*, *b*, or *incr* expressions occur.

**Note** – Random access iterators are required to support random access to elements in constant time. Other iterators are precluded by the restrictions since they can take linear time or offer limited functionality. It is therefore advisable to use tasks to parallelize those cases.

#### Restrictions 1 2 The following restrictions also apply: 3 • If test-expr is of the form var relational-op b and relational-op is < or <= then 4 incr-expr must cause var to increase on each iteration of the loop. If test-expr is of 5 the form var relational-op b and relational-op is > or >= then incr-expr must cause 6 var to decrease on each iteration of the loop. 7 • If test-expr is of the form b relational-op var and relational-op is < or <= then incr-expr must cause var to decrease on each iteration of the loop. If test-expr is of 8 the form b relational-op var and relational-op is > or >= then incr-expr must cause 9 10 var to increase on each iteration of the loop. 11 • For C++, in the **simd** construct the only random access iterator type that are allowed 12

for var are pointer types.

C/C++

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# **14 2.7 Worksharing Constructs**

A worksharing construct distributes the execution of the associated region among the members of the team that encounters it. Threads execute portions of the region in the context of the implicit tasks each one is executing. If the team consists of only one thread then the worksharing region is not executed in parallel.

A worksharing region has no barrier on entry; however, an implied barrier exists at the end of the worksharing region, unless a **nowait** clause is specified. If a **nowait** clause is present, an implementation may omit the barrier at the end of the worksharing region. In this case, threads that finish early may proceed straight to the instructions following the worksharing region without waiting for the other members of the team to finish the worksharing region, and without performing a flush operation (see Section A.10 on page 242 for an example).

The OpenMP API defines the following worksharing constructs, and these are described in the sections that follow:

- loop construct
- sections construct
- single construct
- workshare construct

#### Restrictions

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6 7 The following restrictions apply to worksharing constructs:

- Each worksharing region must be encountered by all threads in a team or by none at all, unless cancellation has been requested for the innermost enclosing parallel region.
- The sequence of worksharing regions and **barrier** regions encountered must be the same for every thread in a team.

# 1 2.7.1 Loop Construct

2 Summary

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The loop construct specifies that the iterations of one or more associated loops will be executed in parallel by threads in the team in the context of their implicit tasks. The iterations are distributed across threads that already exist in the team executing the parallel region to which the loop region binds.

Syntax

The syntax of the loop construct is as follows:

#pragma omp for [clause[[,] clause] ... ] new-line
 for-loops

9 where *clause* is one of the following:

```
private (list)
firstprivate (list)
lastprivate (list)
reduction (operator: list)
schedule (kind[, chunk_size])
collapse (n)
ordered
nowait
```

The **for** directive places restrictions on the structure of all associated *for-loops*. Specifically, all associated *for-loops* must have *canonical loop form* (see Section 2.6 on page 43).

C/C++ -

The syntax of the loop construct is as follows:

```
!$omp do [clause], | clause] ... ]
     do-loops
/!$omp end do /nowait//
```

2 where *clause* is one of the following:

```
private(list)
firstprivate(list)
lastprivate(list)
reduction({operator/intrinsic_procedure_name}:list)
schedule(kind[, chunk_size])
collapse(n)
ordered
```

If an end do directive is not specified, an end do directive is assumed at the end of the do-loop.

All associated do-loops must be do-constructs as defined by the Fortran standard. If an end do directive follows a do-construct in which several loop statements share a DO termination statement, then the directive can only be specified for the outermost of these **DO** statements. See Section A.8 on page 239 for examples.

If any of the loop iteration variables would otherwise be shared, they are implicitly made private on the loop construct. See Section A.9 on page 241 for examples. Unless the loop iteration variables are specified lastprivate on the loop construct, their values after the loop are unspecified.

Fortran -

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14 The binding thread set for a loop region is the current team. A loop region binds to the 15

Binding

innermost enclosing parallel region. Only the threads of the team executing the binding parallel region participate in the execution of the loop iterations and the

implied barrier of the loop region if the barrier is not eliminated by a nowait clause.

## Description

The loop construct is associated with a loop nest consisting of one or more loops that follow the directive.

There is an implicit barrier at the end of a loop construct unless a **nowait** clause is specified.

The **collapse** clause may be used to specify how many loops are associated with the loop construct. The parameter of the **collapse** clause must be a constant positive integer expression. If no **collapse** clause is present, the only loop that is associated with the loop construct is the one that immediately follows the loop directive.

If more than one loop is associated with the loop construct, then the iterations of all associated loops are collapsed into one larger iteration space that is then divided according to the **schedule** clause. The sequential execution of the iterations in all associated loops determines the order of the iterations in the collapsed iteration space.

The iteration count for each associated loop is computed before entry to the outermost loop. If execution of any associated loop changes any of the values used to compute any of the iteration counts, then the behavior is unspecified.

The integer type (or kind, for Fortran) used to compute the iteration count for the collapsed loop is implementation defined.

A worksharing loop has logical iterations numbered 0,1,...,N-1 where N is the number of loop iterations, and the logical numbering denotes the sequence in which the iterations would be executed if the associated loop(s) were executed by a single thread. The **schedule** clause specifies how iterations of the associated loops are divided into contiguous non-empty subsets, called chunks, and how these chunks are distributed among threads of the team. Each thread executes its assigned chunk(s) in the context of its implicit task. The *chunk\_size* expression is evaluated using the original list items of any variables that are made private in the loop construct. It is unspecified whether, in what order, or how many times, any side-effects of the evaluation of this expression occur. The use of a variable in a **schedule** clause expression of a loop construct causes an implicit reference to the variable in all enclosing constructs.

Different loop regions with the same schedule and iteration count, even if they occur in the same parallel region, can distribute iterations among threads differently. The only exception is for the **static** schedule as specified in Table 2-1. Programs that depend on which thread executes a particular iteration under any other circumstances are non-conforming.

See Section 2.7.1.1 on page 52 for details of how the schedule for a worksharing loop is determined.

The schedule *kind* can be one of those specified in Table 2-1.

#### static

When **schedule**(**static**, *chunk\_size*) is specified, iterations are divided into chunks of size *chunk\_size*, and the chunks are assigned to the threads in the team in a round-robin fashion in the order of the thread number.

When no *chunk\_size* is specified, the iteration space is divided into chunks that are approximately equal in size, and at most one chunk is distributed to each thread. Note that the size of the chunks is unspecified in this case.

A compliant implementation of the **static** schedule must ensure that the same assignment of logical iteration numbers to threads will be used in two loop regions if the following conditions are satisfied: 1) both loop regions have the same number of loop iterations, 2) both loop regions have the same value of *chunk\_size* specified, or both loop regions have no *chunk\_size* specified, 3) both loop regions bind to the same parallel region, and 4) neither loop is associated with a SIMD construct. A data dependence between the same logical iterations in two such loops is guaranteed to be satisfied allowing safe use of the **nowait** clause (see Section A.10 on page 242 for examples).

#### dynamic

When schedule (dynamic, chunk\_size) is specified, the iterations are distributed to threads in the team in chunks as the threads request them. Each thread executes a chunk of iterations, then requests another chunk, until no chunks remain to be distributed.

Each chunk contains *chunk\_size* iterations, except for the last chunk to be distributed, which may have fewer iterations.

When no *chunk\_size* is specified, it defaults to 1.

#### guided

When **schedule**(**guided**, *chunk\_size*) is specified, the iterations are assigned to threads in the team in chunks as the executing threads request them. Each thread executes a chunk of iterations, then requests another chunk, until no chunks remain to be assigned.

For a *chunk\_size* of 1, the size of each chunk is proportional to the number of unassigned iterations divided by the number of threads in the team, decreasing to 1. For a *chunk\_size* with value k (greater than 1), the size of each chunk is determined in the same way, with the restriction that the chunks do not contain fewer than k iterations (except for the last chunk to be assigned, which may have fewer than k iterations).

When no *chunk\_size* is specified, it defaults to 1.

#### auto

When **schedule(auto)** is specified, the decision regarding scheduling is delegated to the compiler and/or runtime system. The programmer gives the implementation the freedom to choose any possible mapping of iterations to threads in the team.

runtime

When schedule (runtime) is specified, the decision regarding scheduling is deferred until run time, and the schedule and chunk size are taken from the run-sched-var ICV. If the ICV is set to auto, the schedule is implementation defined.

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**Note** – For a team of p threads and a loop of n iterations, let  $\lceil n/p \rceil$  be the integer q that satisfies n = p\*q - r, with  $0 \le r < p$ . One compliant implementation of the **static** schedule (with no specified chunk size) would behave as though chunk size had been specified with value q. Another compliant implementation would assign q iterations to the first p-r threads, and q-1 iterations to the remaining r threads. This illustrates why a conforming program must not rely on the details of a particular implementation.

A compliant implementation of the **guided** schedule with a *chunk\_size* value of kwould assign  $q = \lceil n/p \rceil$  iterations to the first available thread and set n to the larger of n-q and p\*k. It would then repeat this process until q is greater than or equal to the number of remaining iterations, at which time the remaining iterations form the final chunk. Another compliant implementation could use the same method, except with  $q = \lceil n/(2p) \rceil$ , and set n to the larger of n-q and 2\*p\*k.

#### Restrictions

Restrictions to the loop construct are as follows:

- All loops associated with the loop construct must be perfectly nested; that is, there must be no intervening code nor any OpenMP directive between any two loops.
- The values of the loop control expressions of the loops associated with the loop construct must be the same for all the threads in the team.
- Only one **schedule** clause can appear on a loop directive.
- Only one collapse clause can appear on a loop directive.
- *chunk\_size* must be a loop invariant integer expression with a positive value.
- The value of the *chunk size* expression must be the same for all threads in the team.
- The value of the run-sched-var ICV must be the same for all threads in the team.
- When schedule (runtime) or schedule (auto) is specified, chunk size must not be specified.
- Only one **ordered** clause can appear on a loop directive.
- The ordered clause must be present on the loop construct if any ordered region ever binds to a loop region arising from the loop construct.
- The loop iteration variable may not appear in a **threadprivate** directive.

<ul> <li>Only an iteration of the innermost associated loop may be curtailed by a continue</li> </ul>
statement.
<ul> <li>No statement can branch to any associated for statement.</li> </ul>
<ul> <li>Only one nowait clause can appear on a for directive.</li> </ul>
<ul> <li>A throw executed inside a loop region must cause execution to resume within the same iteration of the loop region, and the same thread that threw the exception must catch it.</li> </ul>
C/C++
Fortran
• The associated do-loops must be structured blocks.
<ul> <li>Only an iteration of the innermost associated loop may be curtailed by a CYCLE statement.</li> </ul>
<ul> <li>No statement in the associated loops other than the DO statements can cause a branch out of the loops.</li> </ul>
• The <i>do-loop</i> iteration variable must be of type integer.
• The do-loop cannot be a DO WHILE or a DO loop without loop control.
Fortran —
Cross References
• private, firstprivate, lastprivate, and reduction clauses, see Section 2.14.3 on page 130.
• OMP_SCHEDULE environment variable, see Section 4.1 on page 210.
• ordered construct, see Section 2.12.8 on page 114.
Determining the Schedule of a Worksharing Loop
When execution encounters a loop directive, the <b>schedule</b> clause (if any) on the directive, and the <i>run-sched-var</i> and <i>def-sched-var</i> ICVs are used to determine how loop iterations are assigned to threads. See Section 2.3 on page 30 for details of how the values of the ICVs are determined. If the loop directive does not have a <b>schedule</b> clause then the current value of the <i>def-sched-var</i> ICV determines the schedule. If the loop directive has a <b>schedule</b> clause that specifies the <b>runtime</b> schedule kind then

the current value of the run-sched-var ICV determines the schedule. Otherwise, the

value of the schedule clause determines the schedule. Figure 2-1 describes how the

C/C++ -

schedule for a worksharing loop is determined.

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#### **Cross References**

• ICVs, see Section 2.3 on page 30.

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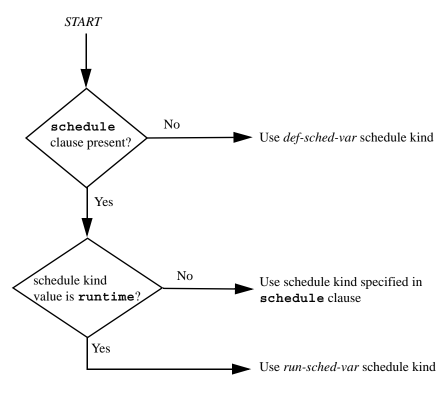
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**FIGURE 2-1** Determining the schedule for a worksharing loop.

## 5 2.7.2 sections Construct

## 6 Summary

The **sections** construct is a noniterative worksharing construct that contains a set of structured blocks that are to be distributed among and executed by the threads in a team. Each structured block is executed once by one of the threads in the team in the context of its implicit task.

```
Syntax
```

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C/C++

The syntax of the **sections** construct is as follows:

where *clause* is one of the following:

```
private (list)
firstprivate (list)
lastprivate (list)
reduction (operator: list)
nowait
```

C/C++ =

Fortran -

The syntax of the **sections** construct is as follows:

```
!$omp sections [clause[[,] clause]...]
  [!$omp section]
    structured-block
  [!$omp section
    structured-block]
...
!$omp end sections [nowait]
```

where *clause* is one of the following:

```
private(list)
```

	firstprivate(list)
	lastprivate(list)
	<pre>reduction({operator intrinsic_procedure_name}: list)</pre>
1	Fortran
2	Binding
3 4 5 6 7	The binding thread set for a <b>sections</b> region is the current team. A <b>sections</b> region binds to the innermost enclosing <b>parallel</b> region. Only the threads of the team executing the binding <b>parallel</b> region participate in the execution of the structured blocks and the implied barrier of the <b>sections</b> region if the barrier is not eliminated by a <b>nowait</b> clause.
8	Description
9 10	Each structured block in the <b>sections</b> construct is preceded by a <b>section</b> directive except possibly the first block, for which a preceding <b>section</b> directive is optional.
11 12	The method of scheduling the structured blocks among the threads in the team is implementation defined.
13 14	There is an implicit barrier at the end of a <b>sections</b> construct unless a <b>nowait</b> clause is specified.
15	Restrictions
16	Restrictions to the sections construct are as follows:
17 18 19	<ul> <li>Orphaned section directives are prohibited. That is, the section directives must appear within the sections construct and must not be encountered elsewhere in the sections region.</li> </ul>
20	• The code enclosed in a <b>sections</b> construct must be a structured block.
21	<ul> <li>Only a single nowait clause can appear on a sections directive.</li> </ul>
	C/C++
22 23 24	<ul> <li>A throw executed inside a sections region must cause execution to resume within the same section of the sections region, and the same thread that threw the exception must catch it.</li> </ul>

C/C++

#### Cross References

• private, firstprivate, lastprivate, and reduction clauses, see Section 2.14.3 on page 130.

# 4 2.7.3 single Construct

## 5 Summary

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The **single** construct specifies that the associated structured block is executed by only one of the threads in the team (not necessarily the master thread), in the context of its implicit task. The other threads in the team, which do not execute the block, wait at an implicit barrier at the end of the **single** construct unless a **nowait** clause is specified.

## Syntax

C/C++ -

The syntax of the **single** construct is as follows:

```
#pragma omp single [clause[[,] clause]...] new-line
structured-block
```

where *clause* is one of the following:

```
private (list)
firstprivate (list)
copyprivate (list)
nowait
```

C/C++ ----

The syntax of the **single** construct is as follows:

```
!$omp single [clause[[,] clause] ...]

structured-block
!$omp end single [end_clause[[,] end_clause] ...]
```

Fortran

1 where *clause* is one of the following: private(list) firstprivate(list) 2 and end clause is one of the following: copyprivate(list) nowait 3 Fortran **Binding** 4 5 The binding thread set for a single region is the current team. A single region binds to the innermost enclosing parallel region. Only the threads of the team 6 7 executing the binding parallel region participate in the execution of the structured block and the implied barrier of the single region if the barrier is not eliminated by a 8 9 nowait clause. Description 10 11 The method of choosing a thread to execute the structured block is implementation defined. There is an implicit barrier at the end of the **single** construct unless a 12 13 nowait clause is specified. For an example of the **single** construct, see Section A.14 on page 252. 14 Restrictions 15 16 Restrictions to the **single** construct are as follows: • The copyprivate clause must not be used with the nowait clause. 17 18 • At most one **nowait** clause can appear on a **single** construct. C/C++• A throw executed inside a single region must cause execution to resume within the 19 20 same single region, and the same thread that threw the exception must catch it.

– C/C++ -

#### Cross References

- private and firstprivate clauses, see Section 2.14.3 on page 130.
- **copyprivate** clause, see Section 2.14.4.2 on page 149.

#### Fortran

## 4 2.7.4 workshare Construct

## 5 Summary

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The workshare construct divides the execution of the enclosed structured block into separate units of work, and causes the threads of the team to share the work such that each unit is executed only once by one thread, in the context of its implicit task.

### Syntax

The syntax of the **workshare** construct is as follows:

```
!$omp workshare
    structured-block
!$omp end workshare [nowait]
```

The enclosed structured block must consist of only the following:

- · array assignments
- scalar assignments
- FORALL statements
- FORALL constructs
- WHERE statements
- WHERE constructs
- atomic constructs
- critical constructs
- parallel constructs
- Statements contained in any enclosed **critical** construct are also subject to these restrictions. Statements in any enclosed **parallel** construct are not restricted.

## Binding

The binding thread set for a **workshare** region is the current team. A **workshare** region binds to the innermost enclosing **parallel** region. Only the threads of the team executing the binding **parallel** region participate in the execution of the units of work and the implied barrier of the **workshare** region if the barrier is not eliminated by a **nowait** clause.

### **Description**

There is an implicit barrier at the end of a **workshare** construct unless a **nowait** clause is specified.

An implementation of the **workshare** construct must insert any synchronization that is required to maintain standard Fortran semantics. For example, the effects of one statement within the structured block must appear to occur before the execution of succeeding statements, and the evaluation of the right hand side of an assignment must appear to complete prior to the effects of assigning to the left hand side.

The statements in the workshare construct are divided into units of work as follows:

- For array expressions within each statement, including transformational array intrinsic functions that compute scalar values from arrays:
  - Evaluation of each element of the array expression, including any references to **ELEMENTAL** functions, is a unit of work.
  - Evaluation of transformational array intrinsic functions may be freely subdivided into any number of units of work.
- For an array assignment statement, the assignment of each element is a unit of work.
- For a scalar assignment statement, the assignment operation is a unit of work.
- For a **WHERE** statement or construct, the evaluation of the mask expression and the masked assignments are each a unit of work.
- For a **FORALL** statement or construct, the evaluation of the mask expression, expressions occurring in the specification of the iteration space, and the masked assignments are each a unit of work.
- For an atomic construct, the atomic operation on the storage location designated as x is the unit of work.
- For a **critical** construct, the construct is a single unit of work.
- For a parallel construct, the construct is a unit of work with respect to the workshare construct. The statements contained in the parallel construct are executed by a new thread team.
- If none of the rules above apply to a portion of a statement in the structured block, then that portion is a unit of work.

1 2 3	The transformational array intrinsic functions are MATMUL, DOT_PRODUCT, SUM, PRODUCT, MAXVAL, MINVAL, COUNT, ANY, ALL, SPREAD, PACK, UNPACK, RESHAPE, TRANSPOSE, EOSHIFT, CSHIFT, MINLOC, and MAXLOC.
4 5	It is unspecified how the units of work are assigned to the threads executing a <b>workshare</b> region.
6 7 8	If an array expression in the block references the value, association status, or allocation status of private variables, the value of the expression is undefined, unless the same value would be computed by every thread.
9 10	If an array assignment, a scalar assignment, a masked array assignment, or a <b>FORALL</b> assignment assigns to a private variable in the block, the result is unspecified.
11 12	The workshare directive causes the sharing of work to occur only in the workshare construct, and not in the remainder of the workshare region.
13	For examples of the workshare construct, see Section A.17 on page 273.
14	Restrictions
15	The following restrictions apply to the workshare construct:
16 17	<ul> <li>All array assignments, scalar assignments, and masked array assignments must be intrinsic assignments.</li> </ul>
18 19	• The construct must not contain any user defined function calls unless the function is <b>ELEMENTAL</b> .
	Fortran —

# 20 2.8 SIMD Constructs

## 21 2.8.1 simd construct

## Summary

The **simd** construct can be applied to a loop to indicate that the loop can be transformed into a SIMD loop (that is, multiple iterations of the loop can be executed concurrently using SIMD instructions).

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```
Syntax
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                    The syntax of the simd construct is as follows:
                                                           C/C++
3
                       #pragma omp simd [clause[[,] clause] ...] new-line
                            for-loops
                    where clause is one of the following:
4
                                safelen(length)
                                linear(list[:linear-step])
                                aligned(list[:alignment])
                               private(list)
                                lastprivate(list)
                                reduction(operator:list)
                                collapse(n)
5
                    The simd directive places restrictions on the structure of the associated for-loops.
6
                    Specifically, all associated for-loops must have canonical loop form (Section 2.6 on
7
                    page 43).
                                                           Fortran -
8
                       !$omp simd [clause[[,] clause ...]
                            do-loops
                       [!$omp end simd]
9
                    where clause is one of the following:
                                safelen(length)
                                linear(list[:linear-step])
                                aligned(list[:alignment])
```

private(list) lastprivate(list) reduction (operator:list) collapse(n)

1 2

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If an end simd directive is not specified, an end simd directive is assumed at the end of the do-loops.

All associated do-loops must be do-constructs as defined by the Fortran standard. If an

end simd directive follows a do-construct in which several loop statements share a

**DO** termination statement, then the directive can only be specified for the outermost of these **DO** statements.

- Fortran -

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Description 11

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25 26 Binding

A simd region binds to the current task region. The binding thread set of the simd region is the current team.

The **simd** construct enables the execution of multiple iterations of the associated loops concurrently by means of SIMD instructions.

The collapse clause may be used to specify how many loops are associated with the construct. The parameter of the collapse clause must be a constant positive integer expression. If no collapse clause is present, the only loop that is associated with the loop construct is the one that immediately follows the directive.

If more than one loop is associated with the simd construct, then the iterations of all associated loops are collapsed into one larger iteration space that is then executed with SIMD instructions. The sequential execution of the iterations in all associated loops determines the order of the iterations in the collapsed iteration space.

The iteration count for each associated loop is computed before entry to the outermost loop. If execution of any associated loop changes any of the values used to compute any of the iteration counts, then the behavior is unspecified.

The integer type (or kind, for Fortran) used to compute the iteration count for the collapsed loop is implementation defined.

1 A SIMD loop has logical iterations numbered 0,1,...,N-1 where N is the number of loop 2 iterations, and the logical numbering denotes the sequence in which the iterations would be executed if the associated loop(s) were executed with no SIMD instructions. If the 3 4 safelen clause is used then no two iterations executed concurrently with SIMD 5 instructions can have a greater distance in the logical iteration space than its value. The 6 parameter of the safelen clause must be a constant positive integer expression. The 7 number of iterations that are executed concurrently at any given time is implementation defined. Each concurrent iteration will be executed by a different SIMD lane. Each set 8 of concurrent iterations is a SIMD chunk. 9 10 The aligned clause declares one or more list items to be aligned to the number of bytes expressed in the optional parameter of the aligned clause. The optional 11 parameter of the aligned clause, alignment, must be a constant positive integer 12 13 expression. If no optional parameter is specified, the default alignment that SIMD 14 instructions in the target platforms use is assumed. Restrictions 15 16 • All loops associated with the construct must be perfectly nested; that is, there must be no intervening code nor any OpenMP directive between any two loops. 17 • The associated loops must be structured blocks. 18 • Only one collapse clause can appear on a simd directive. 19 20 • A list-item cannot appear in more than one aligned clause. • Only one **safelen** clause can appear on a **simd** directive. 21 22 • No OpenMP construct can appear in the simd region. - C/C++ -• The simd region cannot contain calls to the longjmp or setjmp functions. 23 C/C++ ----C++24 • No exception can be raised in the **simd** region. Fortran • The *do-loop* iteration variable must be of type **integer**. 25 26 • The do-loop cannot be a **DO WHILE** or a **DO** loop without loop control. — Fortran ————

#### Cross References

• private, lastprivate, linear and reduction clauses, see Section 2.14.3 on page 130.

#### 4 2.8.2 declare simd construct

#### Summary

The **declare simd** construct can be applied to a function (C, C++ and Fortran) or a subroutine (Fortran) to enable the creation of one or more versions that can process multiple arguments using SIMD instructions from a single invocation from a SIMD loop. The **declare simd** directive is a declarative directive. There may be multiple **declare simd** directives for a function (C, C++, Fortran) or subroutine (Fortran).

## Syntax

The syntax of the **declare** simd construct is as follows:

C/C++

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```
#pragma omp declare simd [clause[[,] clause] ...] new-line
[#pragma omp declare simd [clause[[,] clause] ...] new-line]
[...]
function definition or declaration
```

14

where *clause* is one of the following:

```
simdlen(length)
linear(argument-list[:linear-step])
aligned(argument-list[:alignment])
uniform(argument-list)
reduction(operator:list)
inbranch
notinbranch
```

C/C++ 1 Fortran !\$omp declare simd(proc-name) [clause[[,]clause]...] 2 3 where *clause* is one of the following:: simdlen(length) linear(argument-list[:linear-step]) aligned(argument-list[:alignment]) uniform(argument-list) reduction(operator:list) inbranch notinbranch **Description** 4 C/C++ -The use of a declare simd construct on a function enables the creation of SIMD 5 6 versions of the associated function that can be used to process multiple arguments from 7 a single invocation from a SIMD loop concurrently. - C/C++ -Fortran -8 9

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The use of a **declare simd** construct enables the creation of SIMD versions of the specified subroutine or function that can be used to process multiple arguments from a single invocation from a SIMD loop concurrently. If a **declare simd** directive contains multiple declarations, then for each declaration one or more SIMD versions will be created.

- Fortran -

If a SIMD version is created, the number of concurrent arguments for the function is determined by the **simdlen** clause. If the **simdlen** is used its value corresponds to the number of concurrent arguments of the function. The parameter of the **simdlen** clause must be a constant positive integer expression. Otherwise, the number of concurrent arguments for the function is implementation defined.

The uniform clause declares one or more arguments to have an invariant value for all 1 2 concurrent invocations of the function in the execution of a single SIMD loop. 3 The **aligned** clause declares one or more list items to be aligned to the number of 4 bytes expressed in the optional parameter of the aligned clause. The optional 5 parameter of the aligned clause, alignment, must be a constant positive integer expression. If no optional parameter is specified, the default alignment that SIMD 6 instructions in the target platforms use is assumed. 7 The **inbranch** clause specifies that the function will always be called from inside a 8 9 conditional statement of a SIMD loop. The **notinbranch** clause specifies that the 10 function will never be called from inside a conditional statement of a SIMD loop. If neither clause is specified, then the function may or may not be called from inside a 11 conditional statement of a SIMD loop. 12 Restrictions 13 14 • Each argument can appear in at most one uniform or linear clause. • At most one **simdlen** clause can appear in a declare **simd** directive. 15 • When inbranch is specified, notinbranch must not be specified. 16 17 • The function or subroutine body must be a structured block. 18 • The execution of the function or subroutine, when called from a SIMD loop, cannot result in the execution of an OpenMP construct. 19 • The execution of the function or subroutine cannot have any side-effects that would 20 alter its execution for concurrent iterations of a SIMD chunk. 21 22 • A program that branches into or out of the function region is non-conforming. \_\_\_\_\_ C/C++ \_\_\_\_ • If the function has any declarations, then the declare simd construct for any 23 declaration that has one must be equivalent to the one specified for the definition. 24 25 Otherwise, the result is unspecified. • The function cannot contain calls to the *longjmp* or *setjmp* functions. 26 • The function declaration or definition shall not have an exception specification. 27 • The execution of the function cannot raise any exception. 28 Fortran

• proc-name must not be a generic name, procedure pointer or entry name.

• Any declare simd directive must appear in a specification part of a subroutine 1 2 subprogram, function subprogram or interface body to which it applies. 3 • If a declare simd directive is specified in an interface block for a procedure, it must match a **declare simd** directive in the definition of the procedure. 4 5 • If a procedure is declared via a procedure declaration statement, the procedure 6 proc-name should appear in the same specification. 7 • If a declare simd directive is specified for a procedure name with explicit interface and the definition of the procedure then they must match. Otherwise, the 8 9 result is unspecified. 10 Procedure pointers may not be used to access versions created by the declare 11 simd directive. Fortran -12 2.8.3 Loop SIMD construct Summary 13 14 The loop SIMD construct specifies a loop that can be executed concurrently using SIMD 15 instructions and that those iterations will also be executed in parallel by threads in the 16 team. Syntax 17 C/C++ #pragma omp for simd[clause[[,] clause] ...] new-line for-loops 18 where clause can be any of the clauses accepted by the for or simd directives with 19 identical meanings and restrictions. C/C++ Fortran -20 !\$omp do simd[clause[[,]clause]...] do-loops !\$omp end do simd /nowait/

where *clause* can be any of the clauses accepted by the **simd** or **do** directives, with identical meanings and restrictions.

If an **end do simd** directive is not specified, an **end do simd** directive is assumed at the end of the do-loop.

Fortran -

#### Description

The loop SIMD construct will first convert the associated loop(s) to a SIMD loop in a manner consistent with any clauses that apply to the <code>simd</code> construct. The resulting SIMD chunks and any remaining iterations will be distributed across the implicit tasks of the parallel region in a manner consistent with any clauses that apply to the loop construct.

#### Restrictions

The restrictions for the loop construct and the **simd** construct apply.

#### **Cross References**

- loop construct, see Section 2.7.1 on page 47.
- simd construct, see Section 2.8.1 on page 60.
- Data attribute clauses, see Section 2.14.3 on page 130.

# 17 2.9 target Constructs

# 18 2.9.1 target data Construct

#### Summary

Create a device data environment for the extent of the region.

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**Syntax** 1 C/C++ The syntax of the target data construct is as follows: 2 #pragma omp target data [clause[[,] clause],...] new-line structured-block 3 where *clause* is one of the following: device(integer-expression) 5 map ( [map-type : ] list ) 6 if( scalar-expression ) C/C++ -7 Fortran 8 The syntax of the target data construct is as follows: 9 !\$omp target data [clause[[,] clause],...] structured-block !\$omp end target data where *clause* is one of the following: 10 11 device ( scalar-integer-expression ) 12 map([map-type:]list)13 if( scalar-logical-expression ) The end target data directive denotes the end of the target data construct. 14 Fortran -**Binding** 15 The binding task region for a target data construct is the encountering task. The 16 target region binds to the enclosing parallel or task region. 17

## Description

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When a target data construct is encountered, a new device data environment is created, and the encountering task executes the target data region. If there is no device clause, the default device is determined by the default-device-var ICV. The new device data environment is constructed from the enclosing device data environment, the data environment of the encountering task and any data-mapping clauses on the construct. When an if clause is present and the if clause expression evaluates to false, the device is the host.

#### Restrictions

- A program must not depend on any ordering of the evaluations of the clauses of the target data directive, or on any side effects of the evaluations of the clauses.
- At most one **device** clause may appear on the directive. The **device** expression must evaluate to a non-negative integer value.
- At most one **if** clause can appear on the directive.

#### **Cross References**

- map clause, see Section 2.14.5 on page 151.
- default-device-var, see Section 2.3 on page 30.

# 18 2.9.2 target Construct

#### Summary

Create a device data environment and execute the construct on the same device.

# Syntax

C/C++

The syntax of the target construct is as follows:

#pragma omp target[clause[[,] clause],...] new-line
structured-block

23 where *clause* is one of the following:

```
1
                        device(integer-expression)
 2
                        map([map-type:]list)
 3
                        if( scalar-expression )
                                                           C/C++
                                                          Fortran
 4
                     The syntax of the target construct is as follows:
                        !$omp target [clause[[,] clause],...]
                       structured-block
                        !$omp end target
                     where clause is one of the following:
 5
 6
                        device ( scalar-integer-expression )
 7
                        map([map-type:]list)
 8
                        if( scalar-logical-expression )
 9
                     The end target directive denotes the end of the target construct.
                                                        - Fortran -
                     Binding
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                     The binding task for a target construct is the encountering task. The target region
                     binds to the enclosing parallel or task region.
12
                     Description
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                     The target construct provides a superset of the functionality and restrictions provided
                     by the target data construct. In addition, the target construct specifies that the
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                     region is executed by a device. The encountering task waits for the device to complete
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17
                     the target region. When an if clause is present and the if clause expression evaluates
```

to *false*, the target region is executed by the host device.

#### Restrictions

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• If a target, target update, or target data construct appears within a target region then the behavior is unspecified.

 The result of an omp set default device, omp get default device, 1 2 or omp get num devices routine called within a target region is unspecified. 3 • The effect of an access to a **threadprivate** variable in a target region is 4 unspecified. 5 • A variable referenced in a target construct that is not declared in the construct is implicitly treated as if it had appeared in a map clause with a map-type of 6 7 tofrom. 8 A variable referenced in a target region but not the target construct that is not declared in the target region must appear in a target declare directive. 9 - C/C++ 10 • A throw executed inside a target region must cause execution to resume within the same target region, and the same thread that threw the exception must catch it. 11 - C/C++ -Cross References 12 • target data construct, see Section 2.9.1 on page 68. 13 • map clause, see Section 2.14.5 on page 151. 14 2.9.3 target update Construct **Summary** 16 The target update directive makes the corresponding list items in the device data 17 18 environment consistent with their original list items, according to the specified motion 19 clauses. **Syntax** 20 C/C++The syntax of the target update construct is as follows: 21 #pragma omp target update motion-clause[, clause[,] clause],...] new-line 22 where *motion-clause* is one of the following: 23 to ( list )

```
from ( list )
 1
 2
                     and where clause is one of the following:
                       device(integer-expression)
 3
 4
                        if( scalar-expression )
                                                          C/C++ -
                                                         Fortran ————
 5
                     The syntax of the target update construct is as follows:
                       !$omp target update motion-clause[, clause[,] clause],...]
 6
                     where motion-clause is one of the following:
 7
                       to ( list )
 8
                       from ( list )
 9
                     and where clause is one of the following:
                       device ( scalar-integer-expression )
10
11
                       if( scalar-logical-expression )
                                                        - Fortran -
                     Binding
12
                     The binding task for a target update construct is the encountering task.
13
                     Description
14
                     The target update directive is a stand-alone directive.
15
16
                     For each list item in a to or from clause there is a corresponding list item and an
17
                     original list item. If the corresponding list item is not present in the device data
                     environment, the behavior is unspecified. Otherwise, each corresponding list item in the
18
                     device data environment has an original list item in the current task's data environment.
19
```

For each list item in a **from** clause the value of the corresponding list item is assigned

For each list item in a to clause the value of the original list item is assigned to the

to the original list item.

corresponding list item.

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The device is specified in the **device** clause. If there is no **device** clause, the device 1 2 is determined by the default-device-var ICV. When an if clause is present and the if clause expression evaluates to false then no assignments occur. Restrictions 5 • A list item may only appear in a to or from clause, but not both. · At most one device clause may appear on the directive. The device expression must evaluate to a non-negative integer value. • At most one if clause can appear on the directive. Cross References 9 10 • target data, see Section 2.9.1 on page 68. 11 2.9.4 declare target Directive **Summary** 12 13 The declare target directive specifies that variables, functions (C, C++ and Fortran), and subroutines (Fortran) are mapped to a device. The declare target 14 directive is a declarative directive. 15 **Syntax** 16 C/C++ -17 The syntax of the **declare target** directive is as follows: #pragma omp declare target new-line declarations-definition-seq #pragma omp end declare target new-line C/C++ -18 Fortran -19 The syntax of the **declare target** directive is as follows:

For variables, functions and subroutines: 1 !\$omp declare target( list ) 2 3 where *list* is a comma-separated list of named variables, procedure names and named 4 common blocks. Common block names must appear between slashes. 5 For functions and subroutines: !\$omp declare target **Description** 6 —— C/C++ ——— 7 Variable and routine declarations that appear between the declare target and end declare target directives form an implicit list where each list item is the variable 8 9 or function name. Fortran ———— 10 If a declare target does not have an explicit list, then an implicit list of one item is formed from the name of the enclosing subroutine subprogram, function subprogram or 11 12 interface body to which it applies. - Fortran — If a list item is a function (C, C++, Fortran) or subroutine (Fortran) then a 13 device-specific version of the routine is created that can be called from a target region. 14 15 If a list item is a variable then the original variable is mapped to a corresponding variable in the initial device data environment for all devices. If the original variable is 16 initialized, the corresponding variable in the device data environment is initialized with 17 the same value. 18 Restrictions 19 20 • A threadprivate variable cannot appear in a **declare target** directive. 21 • A variable declared in a **declare target** directive must have a mappable type. \_\_\_\_ C/C++ \_\_\_\_ 22 • A variable declared in a **declare target** directive must be at file or namespace 23 scope.

- A function declared in a **declare target** directive must be at file, namespace, or class scope.
- All declarations and definitions for a function must have a **declare target** directive if one is specified for any of them. Otherwise, the result is unspecified.



- If a list item is a procedure name, it must not be a generic name, procedure pointer or entry name.
- Any **declare target** directive with a list can only appear in a specification part of a subroutine subprogram, function subprogram, program or module.
- Any declare target directive without a list can only appear in a specification
  part of a subroutine subprogram, function subprogram or interface body to which it
  applies.
- If a declare target directive is specified in an interface block for a procedure, it must match a declare target directive in the definition of the procedure.
- If any procedure is declared via a procedure declaration statement, any **declare** target directive with the procedure name must appear in the same specification part.
- A variable that is part of another variable (as an array or structure element) cannot appear in a **declare target** directive.
- The declare target directive must appear in the declaration section of a scoping unit in which the common block or variable is declared. Although variables in common blocks can be accessed by use association or host association, common block names cannot. This means that a common block name specified in a declare target directive must be declared to be a common block in the same scoping unit in which the declare target directive appears.
- If a declare target directive specifying a common block name appears in one program unit, then such a directive must also appear in every other program unit that contains a COMMON statement specifying the same name. It must appear after the last such COMMON statement in the program unit.
- If a declare target variable or a declare target common block is declared with the BIND attribute, the corresponding C entities must also be specified in a declare target directive in the C program.
- A blank common block cannot appear in a declare target directive.
- A variable can only appear in a declare target directive in the scope in which it
  is declared. It must not be an element of a common block or appear in an
  EQUIVALENCE statement.

• A variable that appears in a **declare target** directive must be declared in the Fortran scope of a module or have the **SAVE** attribute, either explicitly or implicitly.

Fortran —

## 3 2.9.5 teams Construct

## 4 Summary

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The **teams** construct creates a league of thread teams where the master thread of each team executes the region.

## 7 Syntax

C/C++ =

The syntax of the **teams** construct is as follows:

```
#pragma omp teams [clause[[,] clause],...] new-line
structured-block
```

9 where *clause* is one of the following:

```
10 num_teams ( integer-expression )
```

11 num threads (integer-expression)

12 default (shared | none)

13 private(list)

14 firstprivate(list)

15 shared(list)

16 reduction ( operator : list )

Fortran

C/C++ -

The syntax of the **teams** construct is as follows:

```
!$omp teams [clause[[,] clause],...]
structured-block
!$omp end teams
```

1	where clause is one of the following:
2	<pre>num_teams ( scalar-integer-expression )</pre>
3	<pre>num_threads( scalar-integer-expression )</pre>
4	<pre>default(shared firstprivate private none)</pre>
5	<pre>private( list )</pre>
6	<pre>firstprivate( list )</pre>
7	shared( list )
8	<pre>reduction({operator/intrinsic_procedure_name} : list)</pre>
a	The and teams directive denotes the end of the teams construct

- Fortran -

#### Binding

The binding thread set for a **teams** region is the encountering thread.

#### **Description**

When a thread encounters a **teams** construct, a league of thread teams is created and the master thread of each thread team executes the **teams** region.

The number of teams is determined by the **num\_teams** clause or a default value that is implementation defined.

The number of threads in each team is determined by the **num\_threads** clause or a default value that is implementation defined.

Once the teams are created, the number of teams remains constant for the duration of the **teams** region.

Within a **teams** region, team numbers uniquely identify each team. Team numbers are consecutive whole numbers ranging from zero to one less than the number of teams. A thread may obtain its own team number by a call to the **omp\_get\_team\_num** library routine.

The **teams** region is executed by the master thread of each team.

The threads other than the master thread do not begin execution until the master thread encounters a parallel region.

1 2		After the teams have completed execution of the teams region, the encountering thread resumes execution of the enclosing target region.
3		Restrictions
4		Restrictions to the teams construct are as follows:
5		• A program that branches into or out of a teams region is non-conforming.
6 7		• A program must not depend on any ordering of the evaluations of the clauses of the <b>teams</b> directive, or on any side effects of the evaluation of the clauses.
8 9		<ul> <li>At most one num_threads clause can appear on the directive. The num_threads expression must evaluate to a positive integer value.</li> </ul>
10 11		<ul> <li>At most one num_teams clause can appear on the directive. The num_teams expression must evaluate to a positive integer value.</li> </ul>
12 13 14		<ul> <li>If specified, a teams construct must be contained within a target construct. That target construct must contain no statements or directives outside of the teams construct.</li> </ul>
15 16 17		<ul> <li>distribute, parallel, , parallel sections, parallel workshare, parallel loop, and parallel loop simd are the only OpenMP constructs that can be closely nested in the teams region.</li> </ul>
18		Cross References:
19 20		• default, shared, private, firstprivate, and reduction clauses, see Section 2.14.3 on page 130.
21		• omp_get_team_num routine, see Section 3.2.27 on page 195.
22	2.9.6	distribute Construct
23		Summary
24 25 26 27		The <b>distribute</b> construct specifies that the iterations of one or more loops will be executed by the thread teams in the context of their implicit tasks. The iterations are distributed across the master threads of all teams that execute the <b>teams</b> region to which the <b>distribute</b> region binds.

# **Syntax** 1 C/C++ 2 The syntax of the **distribute** construct is as follows: #pragma omp distribute [clause[[,] clause],...] new-line for-loops 3 Where *clause* is one of the following: private( list ) firstprivate( list ) collapse(n) dist schedule(kind[, chunk\_size]) All associated for-loops must have the canonical form described in Section 2.6 on page 10 C/C++ -Fortran -11 The syntax of the **distribute** construct is as follows: !\$omp distribute [clause][,] clause],...] do-loops [ !\$omp end distribute ] 12 Where *clause* is one of the following: 13 private( list ) 14 firstprivate( list ) 15 collapse(n) 16 dist schedule(kind[, chunk\_size])

If an end distribute directive is not specified, an end distribute directive

is assumed at the end of the do-loop.

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1 All associated do-loops must be do-constructs as defined by the Fortran standard. If an 2 end do directive follows a do-construct in which several loop statements share a DO 3 termination statement, then the directive can only be specified for the outermost of these 4 **DO** statements. - Fortran -**Binding** 5 The binding thread set for a **distribute** region is the set of master threads created by 6 7 a teams construct. A distribute region binds to the innermost enclosing teams 8 region. Only the threads executing the binding teams region participate in the 9 execution of the loop iterations. Description 10 The distribute construct is associated with a loop nest consisting of one or more 11 12 loops that follow the directive. There is no implicit barrier at the end of a **distribute** construct. 13 14 The collapse clause may be used to specify how many loops are associated with the 15 distribute construct. The parameter of the collapse clause must be a constant 16 positive integer expression. If no collapse clause is present, the only loop that is 17 associated with the distribute construct is the one that immediately follows the distribute construct. 18 19 If more than one loop is associated with the **distribute** construct, then the iteration 20 of all associated loops are collapsed into one larger iteration space. The sequential 21 execution of the iterations in all associated loops determines the order of the iterations in 22 the collapsed iteration space. 23 If dist schedule is specified, kind must be static. If specified, iterations are 24 divided into chunks of size chunk\_size, chunks are assigned to the teams of the league in 25 a round-robin fashion in the order of the team number. When no *chunk\_size* is specified, the iteration space is divided into chunks that are approximately equal in size, and at 26 27 most one chunk is distributed to each team of the league. Note that the size of the 28 chunks is unspecified in this case. Restrictions 29

Restrictions to the **distribute** construct are as follows:

• The distribute construct inherits the restrictions of the loop construct.

• A distribute construct must be closely nested in a teams region.

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#### Cross References:

- loop construct, see Section 2.7.1 on page 47.
- **teams** construct, see Section 2.9.5 on page 77.

# 2.10 Combined Constructs

Combined constructs are shortcuts for specifying one construct immediately nested inside another construct. The semantics of the combined constructs are identical to that of explicitly specifying the second construct containing one instance of the first construct and no other statements.

Some combined constructs have clauses that are permitted on both constructs that were combined. If applying the clause to one construct would result in different program behavior than applying the clause to the other construct then the program's behavior is unspecified.

# **2.10.1** Parallel Loop Construct

## Summary

The parallel loop construct is a shortcut for specifying a **parallel** construct containing one or more associated loops and no other statements.

## Syntax

The syntax of the parallel loop construct is as follows:

#pragma omp parallel for [clause[[,] clause] ...] new-line
 for-loop

where *clause* can be any of the clauses accepted by the **parallel** or **for** directives, except the **nowait** clause, with identical meanings and restrictions.

C/C++

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- Fortran -1 The syntax of the parallel loop construct is as follows: !\$omp parallel do [clause[[,] clause]...] do-loop /!\$omp end parallel do/ where *clause* can be any of the clauses accepted by the parallel or do directives, 2 with identical meanings and restrictions. 3 If an end parallel do directive is not specified, an end parallel do directive is 4 assumed at the end of the do-loop. nowait may not be specified on an end 5 parallel do directive. 6 — Fortran – **Description** 7 — C/C++ —— The semantics are identical to explicitly specifying a parallel directive immediately 8 followed by a for directive. 9 - Fortran -----10 The semantics are identical to explicitly specifying a parallel directive immediately 11 followed by a do directive, and an end do directive immediately followed by an end parallel directive. 12 Fortran Restrictions 13 14 The restrictions for the **parallel** construct and the loop construct apply. Cross References 15 • parallel construct, see Section 2.5 on page 37. 16 • loop construct, see Section 2.7.1 on page 47. 17 • Data attribute clauses, see Section 2.14.3 on page 130. 18

# 1 2.10.2 parallel sections Construct

## 2 Summary

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The parallel sections construct is a shortcut for specifying a parallel construct containing one sections construct and no other statements.

#### **Syntax**

— C/C++ —

The syntax of the **parallel sections** construct is as follows:

where *clause* can be any of the clauses accepted by the **parallel** or **sections** directives, except the **nowait** clause, with identical meanings and restrictions.

\_\_\_\_\_ C/C++

Fortran -

The syntax of the **parallel sections** construct is as follows:

```
!$omp parallel sections [clause[[,] clause]...]
    [!$omp section]
    structured-block
    [!$omp section
        structured-block]
...
!$omp end parallel sections
```

where *clause* can be any of the clauses accepted by the **parallel** or **sections** directives, with identical meanings and restrictions.

1 The last section ends at the end parallel sections directive. nowait cannot be 2 specified on an end parallel sections directive. Fortran **Description** 3 - C/C++ -----The semantics are identical to explicitly specifying a parallel directive immediately 4 followed by a sections directive. 5 \_\_\_\_ C/C++ \_\_\_\_\_\_\_ Fortran — The semantics are identical to explicitly specifying a parallel directive immediately 6 7 followed by a sections directive, and an end sections directive immediately followed by an end parallel directive. 8 Fortran — 9 For an example of the parallel sections construct, see Section A.12 on page 249. Restrictions 10 11 The restrictions for the parallel construct and the sections construct apply. **Cross References:** 12 • parallel construct, see Section 2.5 on page 37. 13 • sections construct, see Section 2.7.2 on page 53. 14 15 • Data attribute clauses, see Section 2.14.3 on page 130. ——— Fortran ——— 16 **2.10.3** parallel workshare Construct Summary 17 18 The parallel workshare construct is a shortcut for specifying a parallel 19 construct containing one workshare construct and no other statements.

#### Syntax

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The syntax of the parallel workshare construct is as follows:

!\$omp parallel workshare [clause[[,] clause]...]
 structured-block
!\$omp end parallel workshare

where *clause* can be any of the clauses accepted by the **parallel** directive, with identical meanings and restrictions. **nowait** may not be specified on an **end parallel workshare** directive.

#### Description

The semantics are identical to explicitly specifying a **parallel** directive immediately followed by a **workshare** directive, and an **end workshare** directive immediately followed by an **end parallel** directive.

#### Restrictions

The restrictions for the parallel construct and the workshare construct apply.

#### **Cross References**

- parallel construct, see Section 2.5 on page 37.
- workshare construct, see Section 2.7.4 on page 58.
- Data attribute clauses, see Section 2.14.3 on page 130.

Fortran -

# **2.10.4 Parallel Loop SIMD Construct**

## Summary

The parallel loop SIMD construct is a shortcut for specifying a **parallel** construct containing one SIMD loop construct and no other statement.

**Syntax** 1 C/C++ #pragma omp parallel for simd[clause[],] clause]...] new-line for-loops 2 where clause can be any of the clauses accepted by the parallel, for or simd 3 directives, except the **nowait** clause, with identical meanings and restrictions. C/C++ Fortran • 4 !\$omp parallel do simd[clause[[,]clause]...] do-loops !\$omp end parallel do simd 5 where clause can be any of the clauses accepted by the parallel, do or simd 6 directives, with identical meanings and restrictions. 7 If an end parallel do simd directive is not specified, an end parallel do 8 simd directive is assumed at the end of the do-loop. nowait may not be specified on 9 an end parallel do simd directive. Fortran -**Description** 10 The semantics of the parallel loop SIMD construct are identical to explicitly specifying 11 12 a parallel directive immediately followed by a loop SIMD directive. Restrictions 13 The restrictions for the **parallel** construct and the loop SIMD construct apply. 14 Cross References 15 16 parallel construct, see Section 2.5 on page 37. loop SIMD construct, see Section 2.8.3 on page 67. 17 18 Data attribute clauses, see Section 2.14.3 on page 130.

# **1 2.11 Tasking Constructs**

#### 2 2.11.1 task Construct

3 Summary

The **task** construct defines an explicit task.

Syntax

C/C++

The syntax of the **task** construct is as follows:

#pragma omp task [clause[[,] clause]...] new-line
 structured-block

where *clause* is one of the following:

if (scalar-expression)

final (scalar-expression)

untied

default(shared | none)

mergeable

private(list)

firstprivate(list)

shared (list)

depend (dependence-type : list)

C/C++

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Fortran

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The syntax of the task construct is as follows:

```
!$omp task [clause[[,] clause] ...]

structured-block
!$omp end task
```

2 where *clause* is one of the following:

```
if (scalar-logical-expression)
final (scalar-logical-expression)
untied
default (private | firstprivate | shared | none)
mergeable
private (list)
firstprivate (list)
shared (list)
depend (dependence-type : list)
```

Fortran -

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#### Binding

TEN 1: 1:

The binding thread set of the task region is the current team. A task region binds to the innermost enclosing parallel region.

## Description

When a thread encounters a **task** construct, a task is generated from the code for the associated structured block. The data environment of the task is created according to the data-sharing attribute clauses on the **task** construct, per-data environment ICVs, and any defaults that apply.

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The encountering thread may immediately execute the task, or defer its execution. In the latter case, any thread in the team may be assigned the task. Completion of the task can be guaranteed using task synchronization constructs. A task construct may be nested inside an outer task, but the task region of the inner task is not a part of the task region of the outer task.

When an if clause is present on a task construct, and the if clause expression evaluates to false, an undeferred task is generated, and the encountering thread must suspend the current task region, for which execution cannot be resumed until the generated task is completed. Note that the use of a variable in an if clause expression of a task construct causes an implicit reference to the variable in all enclosing constructs.

When a final clause is present on a task construct and the final clause expression evaluates to true, the generated task will be a final task. All task constructs encountered during execution of a final task will generate final and included tasks. Note that the use of a variable in a final clause expression of a task construct causes an implicit reference to the variable in all enclosing constructs.

The if clause expression and the final clause expression are evaluated in the context outside of the task construct, and no ordering of those evaluations is specified.

A thread that encounters a task scheduling point within the task region may temporarily suspend the task region. By default, a task is tied and its suspended task region can only be resumed by the thread that started its execution. If the untied clause is present on a task construct, any thread in the team can resume the task region after a suspension. The untied clause is ignored if a final clause is present on the same task construct and the final clause expression evaluates to true, or if a task is an included task.

The task construct includes a task scheduling point in the task region of its generating task, immediately following the generation of the explicit task. Each explicit task region includes a task scheduling point at its point of completion.

When a mergeable clause is present on a task construct, and the generated task is an undeferred task or an included task, the implementation might generate a merged task instead.

**Note** – When storage is shared by an explicit **task** region, it is the programmer's responsibility to ensure, by adding proper synchronization, that the storage does not reach the end of its lifetime before the explicit task region completes its execution.

1	Restrictions
2	Restrictions to the task construct are as follows:
3	<ul> <li>A program that branches into or out of a task region is non-conforming.</li> </ul>
4 5	<ul> <li>A program must not depend on any ordering of the evaluations of the clauses of the task directive, or on any side effects of the evaluations of the clauses.</li> </ul>
6	<ul> <li>At most one if clause can appear on the directive.</li> </ul>
7	<ul> <li>At most one final clause can appear on the directive.</li> </ul>
8	• A throw executed inside a task region must cause execution to resume within the same task region, and the same thread that threw the exception must catch it.  C/C++
	Fortran
10 11	<ul> <li>Unsynchronized use of Fortran I/O statements by multiple tasks on the same unit has unspecified behavior.</li> </ul>
	Fortran
12 <b>2.</b> 13	.1 depend Clause Summary
14	The depend clause enforces additional constraints on the scheduling of tasks. These
15	constraints establish dependences only between sibling tasks. The clause consists of a
16	dependence-type with one or more list items.
17	Syntax
18	The syntax of the depend clause is as follows:
	depend( dependence-type : list )
19	Description
20 21	Task dependences are derived from the <i>dependence-type</i> of a <b>depend</b> clause and its list items, where <i>dependence-type</i> is one of the following:

The **in** dependence-type. The generated task will be a dependent task of all previously 1 2 generated sibling tasks that reference at least one of the list items in an out or inout clause. The **out** and **inout** dependence-types. The generated task will be a dependent task of all previously generated sibling tasks that reference at least one of the list items in an 5 in, out, or inout clause. The list items that appear in the **depend** clause may include array sections. 8 **Note** – The enforced task dependence establishes a synchronization of accesses to the list item performed by a dependent task with respect to accesses to the same list item 10 performed by the predecessor tasks. However, it is the responsibility of the programmer to properly synchronize with respect to other concurrent accesses to the list item. 11 Restrictions 12 13 List items used in depend clauses of the same task or sibling tasks must indicate identical storage or disjoint storage. 14 Cross References 15 16 • Array sections, Section 2.4 on page 36. 17 • Task scheduling constraints, Section 2.11.3 on page 94. 18 **2.11.2** taskyield Construct

# Summary

The taskyield construct specifies that the current task can be suspended in favor of execution of a different task.

#### Syntax

C/C++

The syntax of the taskyield construct is as follows:

#pragma omp taskyield new-line

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1 Because the taskyield construct is a stand-alone directive, there are some 2 restrictions on its placement within a program. The taskyield directive may be 3 placed only at a point where a base language statement is allowed. The taskyield 4 directive may not be used in place of the statement following an if, while, do, 5 switch, or label. See Appendix C for the formal grammar. The examples in 6 Section A.25 on page 296 illustrate these restrictions. Fortran -7 The syntax of the taskyield construct is as follows: !\$omp taskyield Because the taskyield construct is a stand-alone directive, there are some 8 9 restrictions on its placement within a program. The taskyield directive may be 10 placed only at a point where a Fortran executable statement is allowed. The taskyield directive may not be used as the action statement in an if statement or as 11 12 the executable statement following a label if the label is referenced in the program. The examples in Section A.25 on page 296 illustrate these restrictions. 13 Fortran -**Binding** 14 A taskyield region binds to the current task region. The binding thread set of the 15 16 taskyield region is the current team. **Description** 17 18 The taskyield region includes an explicit task scheduling point in the current task region. 19 Cross References 20 21 • Task scheduling, see Section 2.11.3 on page 94.

2.11.3 Task Scheduling Whenever a thread reaches a task scheduling point, the implementation may cause it to 2 3 perform a task switch, beginning or resuming execution of a different task bound to the current team. Task scheduling points are implied at the following locations: 5 • the point immediately following the generation of an explicit task • after the point of completion of a task region 6 7 • in taskyield regions 8 • in taskwait regions • at the end of a taskgroup region 9 in implicit and explicit barrier regions. 10 11 • the point immediately following the generation of a target region 12 • at the beginning and end of a target data region 13 • in a target update region 14 When a thread encounters a task scheduling point it may do one of the following, subject to the Task Scheduling Constraints (below): 15 16 begin execution of a tied task bound to the current team 17 resume any suspended task region, bound to the current team, to which it is tied 18 begin execution of an untied task bound to the current team 19 resume any suspended untied task region bound to the current team. 20 If more than one of the above choices is available, it is unspecified as to which will be 21 chosen. 22

Task Scheduling Constraints are as follows:

- 1. An included task is executed immediately after generation of the task.
- 2. Scheduling of new tied tasks is constrained by the set of task regions that are currently tied to the thread, and that are not suspended in a barrier region. If this set is empty, any new tied task may be scheduled. Otherwise, a new tied task may be scheduled only if it is a descendant of every task in the set.
- 3. A dependent task shall not be scheduled until its task dependences are fulfilled.
- 4. When an explicit task is generated by a construct containing an **if** clause for which the expression evaluated to false, and the previous constraints are already met, the task is executed immediately after generation of the task.

A program relying on any other assumption about task scheduling is non-conforming.

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**Note** – Task scheduling points dynamically divide task regions into parts. Each part is executed uninterrupted from start to end. Different parts of the same task region are executed in the order in which they are encountered. In the absence of task synchronization constructs, the order in which a thread executes parts of different schedulable tasks is unspecified.

A correct program must behave correctly and consistently with all conceivable scheduling sequences that are compatible with the rules above.

For example, if **threadprivate** storage is accessed (explicitly in the source code or implicitly in calls to library routines) in one part of a task region, its value cannot be assumed to be preserved into the next part of the same task region if another schedulable task exists that modifies it (see Example A.15.7c on page 262, Example A.15.7f on page 262, Example A.15.8c on page 263 and Example A.15.8f on page 263).

As another example, if a lock acquire and release happen in different parts of a task region, no attempt should be made to acquire the same lock in any part of another task that the executing thread may schedule. Otherwise, a deadlock is possible. A similar situation can occur when a critical region spans multiple parts of a task and another schedulable task contains a critical region with the same name (see Example A.15.9c on page 264, Example A.15.9f on page 265, Example A.15.10c on page 266 and Example A.15.10f on page 267).

The use of threadprivate variables and the use of locks or critical sections in an explicit task with an **if** clause must take into account that when the **if** clause evaluates to *false*, the task is executed immediately, without regard to *Task Scheduling Constraint* 2.

# 2.12 Master and Synchronization Constructs

OpenMP provides the following synchronization constructs:

• the master construct.

- the critical construct.
- the barrier construct.
- the taskwait construct.
- the atomic construct.
- the flush construct.
- the ordered construct.

#### 1 Restrictions

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2 OpenMP synchronization constructs have the following restrictions:

• The use of synchronization constructs between threads in different contention groups results in unspecified behavior.

#### 2.12.1 master Construct

#### 6 Summary

The master construct specifies a structured block that is executed by the master thread of the team.

Fortran -

----- C/C++ -

## Syntax

The syntax of the master construct is as follows:

#pragma omp master new-line
 structured-block

C/C++ ----

The syntax of the **master** construct is as follows:

!\$omp master structured-block

!\$omp end master

Fortran —

#### **Binding** 1 The binding thread set for a master region is the current team. A master region 2 3 binds to the innermost enclosing parallel region. Only the master thread of the team 4 executing the binding parallel region participates in the execution of the structured 5 block of the master region. **Description** 6 7 Other threads in the team do not execute the associated structured block. There is no implied barrier either on entry to, or exit from, the master construct. 8 9 For an example of the **master** construct, see Section A.18 on page 277. Restrictions 10 C/C++ 11 A throw executed inside a master region must cause execution to resume within the 12 same master region, and the same thread that threw the exception must catch it. - C/C++ -13 2.12.2 critical Construct **Summary** 14 The critical construct restricts execution of the associated structured block to a 15 single thread at a time. 16 **Syntax** 17 C/C++ 18 The syntax of the **critical** construct is as follows: #pragma omp critical [(name)] new-line

C/C++ -

structured-block

#### - Fortran -

The syntax of the **critical** construct is as follows:

```
!$omp critical [(name)]
    structured-block
!$omp end critical [(name)]
```

---- Fortran -----

# Binding

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The binding thread set for a **critical** region is all threads. Region execution is restricted to a single thread at a time among all the threads in the program, without regard to the team(s) to which the threads belong.

## **Description**

An optional *name* may be used to identify the critical construct. All critical constructs without a name are considered to have the same unspecified name. A thread waits at the beginning of a critical region until no thread is executing a critical region with the same name. The critical construct enforces exclusive access with respect to all critical constructs with the same name in all threads, not just those threads in the current team.

#### \_\_\_\_\_ C/C++ \_\_\_\_\_

Identifiers used to identify a **critical** construct have external linkage and are in a name space that is separate from the name spaces used by labels, tags, members, and ordinary identifiers.

\_\_\_\_\_ C/C++ \_\_\_\_\_

#### — Fortran ——

The names of **critical** constructs are global entities of the program. If a name conflicts with any other entity, the behavior of the program is unspecified.

Fortran

For an example of the **critical** construct, see Section A.19 on page 279.

#### Restrictions 1 C/C++ -· A throw executed inside a critical region must cause execution to resume within 2 3 the same critical region, and the same thread that threw the exception must catch – C/C++ – Fortran ——— The following restrictions apply to the **critical** construct: 5 6 • If a *name* is specified on a **critical** directive, the same *name* must also be specified on the end critical directive. 7 8 • If no *name* appears on the **critical** directive, no *name* can appear on the **end** 9 critical directive. Fortran — 2.12.3 barrier Construct **Summary** 11 12 The barrier construct specifies an explicit barrier at the point at which the construct 13 appears. **Syntax** 14 \_\_\_\_\_ C/C++ -15 The syntax of the barrier construct is as follows: #pragma omp barrier new-line 16 Because the **barrier** construct is a stand-alone directive, there are some restrictions on its placement within a program. The barrier directive may be placed only at a 17

point where a base language statement is allowed. The barrier directive may not be

used in place of the statement following an if, while, do, switch, or label. See

C/C++ -

Appendix C for the formal grammar. The examples in Section A.25 on page 296

illustrate these restrictions.

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#### Fortran

The syntax of the **barrier** construct is as follows:

#### !\$omp barrier

Because the **barrier** construct is a stand-alone directive, there are some restrictions on its placement within a program. The **barrier** directive may be placed only at a point where a Fortran executable statement is allowed. The **barrier** directive may not be used as the action statement in an **if** statement or as the executable statement following a label if the label is referenced in the program. The examples in Section A.25 on page 296 illustrate these restrictions.

#### Fortran -

#### **Binding**

The binding thread set for a **barrier** region is the current team. A **barrier** region binds to the innermost enclosing **parallel** region. See Section A.21 on page 282 for examples.

#### **Description**

All threads of the team executing the binding **parallel** region must execute the **barrier** region and must not continue execution beyond the barrier until they complete execution of all explicit tasks bound to this **parallel** region.

The **barrier** region includes an implicit task scheduling point in the current task region.

#### Restrictions

The following restrictions apply to the **barrier** construct:

- Each **barrier** region must be encountered by all threads in a team or by none at all, unless cancellation has been requested for the innermost enclosing parallel region.
- The sequence of worksharing regions and **barrier** regions encountered must be the same for every thread in a team.

# 1 2.12.4 taskwait Construct

2	Summary
3 4	The taskwait construct specifies a wait on the completion of child tasks of the current task.
5	Syntax
6	The syntax of the taskwait construct is as follows:
	#pragma omp taskwait newline
7 8 9 10 11	Because the taskwait construct is a stand-alone directive, there are some restrictions on its placement within a program. The taskwait directive may be placed only at a point where a base language statement is allowed. The taskwait directive may not be used in place of the statement following an if, while, do, switch, or label. See Appendix C for the formal grammar. The examples in Section A.25 on page 296 illustrate these restrictions.
	C/C++
13	The syntax of the taskwait construct is as follows:
	!\$omp taskwait
14 15 16 17 18	Because the taskwait construct is a stand-alone directive, there are some restrictions on its placement within a program. The taskwait directive may be placed only at a point where a Fortran executable statement is allowed. The taskwait directive may not be used as the action statement in an if statement or as the executable statement following a label if the label is referenced in the program. The examples in Section A.25 on page 296 illustrate these restrictions.
	Fortran —
20	Binding
21 22	A taskwait region binds to the current task region. The binding thread set of the taskwait region is the current team.

#### 1 Description

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The taskwait region includes an implicit task scheduling point in the current task region. The current task region is suspended at the task scheduling point until all child tasks that it generated before the taskwait region complete execution.

# 5 2.12.5 taskgroup Construct

#### 6 Summary

The **taskgroup** construct specifies a wait on completion of child tasks of the current task and their descendant tasks.

C/C++ -

#### Syntax

The syntax of the **taskgroup** construct is as follows:

#pragma omp taskgroup new-line
 structured-block

C/C++ -

Fortran —

The syntax of the taskgroup construct is as follows:

structured-block
!\$omp end taskgroup

!\$omp taskgroup

Binding

A **taskgroup** region binds to the current task region. The binding thread set of the **taskgoup** region is the current team.

Fortran -

#### 1 Description

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When a thread encounters a **taskgroup** construct, it starts executing the region. There is an implicit task scheduling point at the end of the **taskgroup** region. The current task is suspended at the task scheduling point until all child tasks that it generated in the **taskgroup** region and all of their descendant tasks complete execution.

#### Cross References

• Task scheduling, see Section 2.11.3 on page 94

## 8 2.12.6 atomic Construct

## 9 Summary

The atomic construct ensures that a specific storage location is accessed atomically, rather than exposing it to the possibility of multiple, simultaneous reading and writing threads that may result in indeterminate values.

## 13 Syntax

C/C++

The syntax of the **atomic** construct takes either of the following forms:

```
#pragma omp atomic [read | write | update |
  capture][seq_cst] new-line
  expression-stmt
```

15 or:

```
#pragma omp atomic capture [seq_cst] new-line
structured-block
```

where *expression-stmt* is an expression statement with one of the following forms:

If clause is read:

v = x;

If clause is write:

20 x = expr;

```
1
                     • If clause is update or not present:
 2
                        x++;
 3
                        x - - ;
 4
                        ++x;
 5
                        --x;
 6
                        x \ binop = expr;
 7
                        x = x \ binop \ expr;
 8
                        x = expr binop x;
 9
                     • If clause is capture:
10
                        v = x + + ;
11
                          = x - - 
12
                           = ++x;
13
                        v = --x;
14
                        v = x \ binop = expr;
15
                        v = x = x \ binop \ expr;
                        v = x = expr \ binop \ x;
16
17
                     and where structured-block is a structured block with one of the following forms:
                        \{v = x; x \ binop = expr;\}
```

 $\{v = x; x = x \text{ binop } expr; \}$  $\{v = x; x = expr \ binop \ x; \}$  $\{x = x \ binop \ expr; \ v = x;\}$  $\{x = expr \ binop \ x; \ v = x; \}$  $\{v = x; x = expr;\}$  $\{v = x; x++;\}$  $\{v = x; ++x;\}$ 

 $\{x \ binop = expr; \ v = x; \}$ 

 $\{++x; v = x;\}$  $\{x++; v = x;\}$  $\{v = x; x--;\}$  $\{v = x; --x;\}$ 

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 $\{--x; v = x;\}$  $\{x - -; v = x;\}$ 

In the preceding expressions:

- x and y (as applicable) are both *l-value* expressions with scalar type.
- During the execution of an atomic region, multiple syntactic occurrences of x must designate the same storage location.
- Neither of v and expr (as applicable) may access the storage location designated by x.
- Neither of x and expr (as applicable) may access the storage location designated by v.
- expr is an expression with scalar type.
- binop is one of +, \*, -, /, &, ^, |, <<, or >>.

```
1
                    • binop, binop=, ++, and -- are not overloaded operators.
 2
                    • The expression x binop expr must be mathematically equivalent to x binop (expr).
 3
                       This requirement is satisfied if the operators in expr have precedence greater than
                       binop, or by using parentheses around expr or subexpressions of expr.
 4
 5
                    • The expression expr binop x must be mathematically equivalent to (expr) binop x.
 6
                       This requirement is satisfied if the operators in expr have precedence equal to or
 7
                       greater than binop, or by using parentheses around expr or subexpressions of expr.
                    • For forms that allow multiple occurrences of x, the number of times that x is
 8
 9
                       evaluated is unspecified.
                                                         C/C++
                                                        Fortran
10
                    The syntax of the atomic construct takes any of the following forms:
                       !$omp atomic read [seq cst]
                           capture-statement
                       /!$omp end atomic/
11
                    or
                       !$omp atomic write /seq cst/
                           write-statement
                       /!$omp end atomic/
12
                    or
                       !$omp atomic [update] [seq cst]
                           update-statement
                       /!$omp end atomic/
13
                    or
                       !$omp atomic capture [seq cst]
                           update-statement
                           capture-statement
                       !$omp end atomic
```

```
----- Fortran (cont.) -------
 1
                    or
                       !$omp atomic capture /seq cst/
                           capture-statement
                           update-statement
                       !$omp end atomic
 2
                    or
                       !$omp atomic capture /seq cst/
                           capture-statement
                           write-statement
                       !$omp end atomic
 3
                    where write-statement has the following form (if clause is write):
                       x = expr
                     where capture-statement has the following form (if clause is capture or read):
                       v = x
                    and where update-statement has one of the following forms (if clause is update,
                     capture, or not present):
                       x = x operator expr
10
                       x = expr \ operator \ x
11
                       x = intrinsic procedure name (x, expr list)
                       x = intrinsic\_procedure\_name (expr_list, x)
12
13
                    In the preceding statements:
                    • x and v (as applicable) are both scalar variables of intrinsic type.
14

    x must not be an allocatable variable.

15
                    • During the execution of an atomic region, multiple syntactic occurrences of x must
16
17
                       designate the same storage location.
18

    None of v, expr and expr_list (as applicable) may access the same storage location as

19
20
                    • None of x, expr and expr_list (as applicable) may access the same storage location as
21
                       ν.
```

1	• expr is a scalar expression.
2 3 4	<ul> <li>expr_list is a comma-separated, non-empty list of scalar expressions. If     intrinsic_procedure_name refers to IAND, IOR, or IEOR, exactly one expression     must appear in expr_list.</li> </ul>
5	• intrinsic_procedure_name is one of MAX, MIN, IAND, IOR, or IEOR.
6	• operator is one of +, *, -, /, .AND., .OR., .EQV., or .NEQV
7 8 9	• The expression <i>x operator expr</i> must be mathematically equivalent to <i>x operator</i> ( <i>expr</i> ). This requirement is satisfied if the operators in <i>expr</i> have precedence greater than <i>operator</i> , or by using parentheses around <i>expr</i> or subexpressions of <i>expr</i> .
10 11 12	• The expression <i>expr operator x</i> must be mathematically equivalent to <i>(expr) operator x</i> . This requirement is satisfied if the operators in <i>expr</i> have precedence equal to or greater than <i>operator</i> , or by using parentheses around <i>expr</i> or subexpressions of <i>exp</i>
13 14	<ul> <li>intrinsic_procedure_name must refer to the intrinsic procedure name and not to other program entities.</li> </ul>
15	• operator must refer to the intrinsic operator and not to a user-defined operator.
16	All assignments must be intrinsic assignments.
17 18	• For forms that allow multiple occurrences of x, the number of times that x is evaluated is unspecified.
	Fortran
19 20 21	<ul> <li>In all atomic construct forms, the seq_cst clause and the clause that denotes the type of the atomic construct can appear in any order. In addition, an optional comma may be used to separate the clauses.</li> </ul>
22	Binding
23 24 25 26	The binding thread set for an atomic region is all threads. <b>atomic</b> regions enforce exclusive access with respect to other <b>atomic</b> regions that access the same storage location x among all the threads in the program without regard to the teams to which the threads belong.
27	Description
28 29	The <b>atomic</b> construct with the <b>read</b> clause forces an atomic read of the location designated by $x$ regardless of the native machine word size.
30	The atomic construct with the write clause forces an atomic write of the location

designated by x regardless of the native machine word size.

The **atomic** construct with the **update** clause forces an atomic update of the location designated by x using the designated operator or intrinsic. Note that when no clause is present, the semantics are equivalent to atomic update. Only the read and write of the location designated by x are performed mutually atomically. The evaluation of expr or  $expr\_list$  need not be atomic with respect to the read or write of the location designated by x. No task scheduling points are allowed between the read and the write of the location designated by x.

The atomic construct with the capture clause forces an atomic update of the location designated by x using the designated operator or intrinsic while also capturing the original or final value of the location designated by x with respect to the atomic update. The original or final value of the location designated by x is written in the location designated by y depending on the form of the atomic construct structured block or statements following the usual language semantics. Only the read and write of the location designated by x are performed mutually atomically. Neither the evaluation of expr or  $expr_list$ , nor the write to the location designated by y need be atomic with respect to the read or write of the location designated by x. No task scheduling points are allowed between the read and the write of the location designated by x.

Any atomic construct with a seq\_cst clause forces the atomically performed operation to include an implicit flush operation without a list.

**Note** – As with other implicit flush regions, Section 1.4.4 on page 18 reduces the ordering that must be enforced. The intent is that, when the analogous operation exists in C++11 or C11, a sequentially consistent atomic construct has the same semantics as a memory\_order\_seq\_cst atomic operation in C++11/C11. Similarly, a non-sequentially consistent atomic construct has the same semantics as a memory order relaxed atomic operation in C++11/C11.

Unlike non-sequentially consistent atomic constructs, sequentially consistent atomic constructs preserve the interleaving (sequentially consistent) behavior of correct, data-race-free programs. However, they are not designed to replace the flush directive as a mechanism to enforce ordering for non-sequentially consistent atomic constructs, and attempts to do so require extreme caution. For example, a sequentially consistent atomic write construct may appear to be reordered with a subsequent non-sequentially consistent atomic write construct, since such reordering would not be observable by a correct program if the second write were outside an atomic directive.

For all forms of the **atomic** construct, any combination of two or more of these **atomic** constructs enforces mutually exclusive access to the locations designated by x. To avoid race conditions, all accesses of the locations designated by x that could potentially occur in parallel must be protected with an **atomic** construct.

1 atomic regions do not guarantee exclusive access with respect to any accesses outside 2 of atomic regions to the same storage location x even if those accesses occur during a 3 critical or ordered region, while an OpenMP lock is owned by the executing task, or during the execution of a reduction clause. 4 5 However, other OpenMP synchronization can ensure the desired exclusive access. For example, a barrier following a series of atomic updates to x guarantees that subsequent 6 accesses do not form a race with the atomic accesses. 7 A compliant implementation may enforce exclusive access between atomic regions 8 9 that update different storage locations. The circumstances under which this occurs are 10 implementation defined. 11 If the storage location designated by x is not size-aligned (that is, if the byte alignment 12 of x is not a multiple of the size of x), then the behavior of the **atomic** region is implementation defined. 13 14 For an example of the **atomic** construct, see Section A.22 on page 284. Restrictions 15 \_\_\_\_\_ C/C++ -The following restriction applies to the **atomic** construct: 16 • All atomic accesses to the storage locations designated by x throughout the program 17 are required to have a compatible type. See Section A.23 on page 290 for examples. 18 — C/C++ —— Fortran ———— 19 The following restriction applies to the **atomic** construct: 20 • All atomic accesses to the storage locations designated by x throughout the program 21 are required to have the same type and type parameters. See Section A.23 on page 290 for examples. 22 Fortran Cross References 23 24 • critical construct, see Section 2.12.2 on page 97. 25 • barrier construct, see Section 2.12.3 on page 99. 26 • **flush** construct, see Section 2.12.7 on page 110. 27 • ordered construct, see Section 2.12.8 on page 114. • reduction clause, see Section 2.14.3.6 on page 142. 28

• lock routines, see Section 3.3 on page 196.

## 2 2.12.7 flush Construct

## з Summary

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The **flush** construct executes the OpenMP flush operation. This operation makes a thread's temporary view of memory consistent with memory, and enforces an order on the memory operations of the variables explicitly specified or implied. See the memory model description in Section 1.4 on page 15 for more details.

#### **Syntax**

C/C++

The syntax of the **flush** construct is as follows:

#pragma omp flush [(list)] new-line

Because the **flush** construct is a stand-alone directive, there are some restrictions on its placement within a program. The **flush** directive may be placed only at a point where a base language statement is allowed. The **flush** directive may not be used in place of the statement following an **if**, **while**, **do**, **switch**, or **label**. See Appendix C for the formal grammar. See Section A.25 on page 296 for an example that illustrates these placement restrictions.

C/C++

Fortran

The syntax of the **flush** construct is as follows:

!\$omp flush [(list)]

Because the **flush** construct is a stand-alone directive, there are some restrictions on its placement within a program. The **flush** directive may be placed only at a point where a Fortran executable statement is allowed. The **flush** directive may not be used as the action statement in an **if** statement or as the executable statement following a label if the label is referenced in the program. The examples in Section A.25 on page 296 illustrate these restrictions.

Fortran -

## Binding

The binding thread set for a **flush** region is the encountering thread. Execution of a **flush** region affects the memory and the temporary view of memory of only the thread that executes the region. It does not affect the temporary view of other threads. Other threads must themselves execute a flush operation in order to be guaranteed to observe the effects of the encountering thread's flush operation.

#### **Description**

A flush construct without a list, executed on a given thread, operates as if the whole thread-visible data state of the program, as defined by the base language, is flushed. A flush construct with a list applies the flush operation to the items in the list, and does not return until the operation is complete for all specified list items. Use of a flush construct with a list is extremely error prone and users are strongly discouraged from attempting it. An implementation may implement a flush with a list by ignoring the list, and treating it the same as a flush without a list.

– C/C++ —

If a pointer is present in the list, the pointer itself is flushed, not the memory block to which the pointer refers.

C/C++

Fortran

If the list item or a subobject of the list item has the **POINTER** attribute, the allocation or association status of the **POINTER** item is flushed, but the pointer target is not. If the list item is a Cray pointer, the pointer is flushed, but the object to which it points is not. If the list item is of type **C\_PTR**, the variable is flushed, but the storage that corresponds to that address is not flushed. If the list item or the subobject of the list item has the **ALLOCATABLE** attribute and has an allocation status of currently allocated, the allocated variable is flushed; otherwise the allocation status is flushed.

——— Fortran ———

For examples of the **flush** construct, see Section A.25 on page 296.

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**Note** – the following examples illustrate the ordering properties of the flush operation. In the following incorrect pseudocode example, the programmer intends to prevent simultaneous execution of the protected section by the two threads, but the program does not work properly because it does not enforce the proper ordering of the operations on variables a and b. Any shared data accessed in the protected section is not guaranteed to be current or consistent during or after the protected section. The atomic notation in the pseudocode in the following two examples indicates that the accesses to a and b are ATOMIC writes and captures. Otherwise both examples would contain data races and automatically result in unspecified behavior.

```
Incorrect example:
                            a = b = 0
            thread 1
                                                    thread 2
      atomic(b = 1)
                                              atomic(a = 1)
     flush (b)
                                              flush (a)
     flush (a)
                                              flush (b)
     atomic(tmp = a)
                                              atomic(tmp = b)
      if (tmp == 0) then
                                               if (tmp == 0) then
                                                  protected section
        protected section
      end if
                                              end if
```

The problem with this example is that operations on variables a and b are not ordered with respect to each other. For instance, nothing prevents the compiler from moving the flush of **b** on thread 1 or the flush of **a** on thread 2 to a position completely after the protected section (assuming that the protected section on thread 1 does not reference b and the protected section on thread 2 does not reference a). If either re-ordering happens, both threads can simultaneously execute the protected section.

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13 14 The following pseudocode example correctly ensures that the protected section is executed by not more than one of the two threads at any one time. Notice that execution of the protected section by neither thread is considered correct in this example. This occurs if both flushes complete prior to either thread executing its if statement.

```
Correct example:
                      a = b = 0
       thread 1
                                              thread 2
 atomic(b = 1)
                                        atomic(a = 1)
 flush (a,b)
                                        flush (a,b)
 atomic(tmp = a)
                                        atomic(tmp = b)
 if (tmp == 0) then
                                        if (tmp == 0) then
   protected section
                                            protected section
 end if
                                        end if
```

The compiler is prohibited from moving the flush at all for either thread, ensuring that the respective assignment is complete and the data is flushed before the **if** statement is executed.

A **flush** region without a list is implied at the following locations:

- During a barrier region.
- At entry to and exit from parallel, critical, and ordered regions.
- At exit from worksharing regions unless a **nowait** is present.
- At entry to and exit from combined parallel worksharing regions.
- At entry to and exit from the **atomic** operation (read, write, update, or capture) performed in a sequentially consistent atomic region.
- During omp\_set\_lock and omp\_unset\_lock regions.
- During omp\_test\_lock, omp\_set\_nest\_lock, omp\_unset\_nest\_lock and omp\_test\_nest\_lock regions, if the region causes the lock to be set or unset.
- Immediately before and immediately after every task scheduling point.

A **flush** region with a list is implied at the following locations:

• At entry to and exit from the **atomic** operation (read, write, update, or capture) performed in a non-sequentially consistent **atomic** region, where the list contains only the storage location designated as x according to the description of the syntax of the **atomic** construct in Section 2.12.6 on page 103.

- 1 **Note** A **flush** region is not implied at the following locations:
  - At entry to worksharing regions.
  - At entry to or exit from a master region.

## 4 2.12.8 ordered Construct

## 5 Summary

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The **ordered** construct specifies a structured block in a loop region that will be executed in the order of the loop iterations. This sequentializes and orders the code within an **ordered** region while allowing code outside the region to run in parallel.

C/C++ -

C/C++ -

## 9 Syntax

The syntax of the **ordered** construct is as follows:

#pragma omp ordered new-line
 structured-block

Fortran -

The syntax of the **ordered** construct is as follows:

!\$omp ordered
 structured-block
!\$omp end ordered

----- Fortran -

1	Binding
2 3 4	The binding thread set for an <b>ordered</b> region is the current team. An <b>ordered</b> region binds to the innermost enclosing loop region. <b>ordered</b> regions that bind to different loop regions execute independently of each other.
5	Description
6 7 8 9 10	The threads in the team executing the loop region execute <b>ordered</b> regions sequentially in the order of the loop iterations. When the thread executing the first iteration of the loop encounters an <b>ordered</b> construct, it can enter the <b>ordered</b> region without waiting. When a thread executing any subsequent iteration encounters an <b>ordered</b> region, it waits at the beginning of that <b>ordered</b> region until execution of all the <b>ordered</b> regions belonging to all previous iterations have completed.
12	For examples of the <b>ordered</b> construct, see Section A.26 on page 299.
13	Restrictions
14	Restrictions to the ordered construct are as follows:
15 16	<ul> <li>The loop region to which an ordered region binds must have an ordered clause specified on the corresponding loop (or parallel loop) construct.</li> </ul>
17 18 19	<ul> <li>During execution of an iteration of a loop or a loop nest within a loop region, a thread must not execute more than one ordered region that binds to the same loop region.</li> </ul>
20 21 22	<ul> <li>A throw executed inside a ordered region must cause execution to resume within the same ordered region, and the same thread that threw the exception must catch it.</li> </ul>
	C/C++
23	Cross References
24	• loop construct, see Section 2.7.1 on page 47.
25	• parallel loop construct, see Section 2.10.1 on page 82.

# **2.13 Cancellation Constructs**

## 2 2.13.1 cancel Construct

4	
5	The cancel construct requests cancellation of the innermost enclosing region of the type specified.
6	Syntax
7	C/C++ The syntax of the cancel construct is as follows:
	<pre>#pragma omp cancel clause [[,] clause ] new-line</pre>
8	where <i>clause</i> is one of the following
9	parallel
10	sections
11	for
12	taskgroup
13	if (scalar-expression)
14 15 16 17	Because the <b>cancel</b> construct is a stand-alone directive, there are some restrictions on its placement within a program. The <b>cancel</b> directive may be placed only at a point where a base language statement is allowed. The <b>cancel</b> directive may not be used in place of the statement following an <b>if</b> , <b>while</b> , <b>do</b> , <b>switch</b> , or label. See Appendix C for the formal grammar.
	Fortran —

1	where <i>clause</i> is one of the following
2	parallel
3	sections
4	do
5	taskgroup
6	if (scalar-logical-expression)
7 8	Because the <b>cancel</b> construct is a stand-alone directive, there are some restrictions on its placement within a program. The <b>cancel</b> directive may be placed only at a point
9	where a Fortran executable statement is allowed. The cancel directive may not be
10	used as the action statement in an <b>if</b> statement or as the executable statement following
11	a label if the label is referenced in the program.
	Fortran —
12	Binding
13	The binding thread set of the cancel region is the current team. The cancel region
14	binds to the innermost enclosing region of the type specified in the clause.
15	Description
15	Description
16	The cancel construct requests cancellation of the innermost enclosing region of the
17	type specified and causes the encountering implicit and explicit task to proceed to the
18	end of the canceled construct.
19	Cancellation points are implied at the following locations:
20	<ul> <li>implicit barriers</li> </ul>
21	• barrier regions
22	• cancel regions
23	• cancellation point regions
24	When a thread reaches one of the above cancellation points and if the cancellation point
25	is active, the thread immediately checks for cancellation and performs the cancellation if
26	observed.
27	When cancellation of tasks is requested through the cancel taskgroup construct,
28	the innermost enclosing taskgroup will be canceled. The task that encountered the
29	cancel taskgroup construct jumps to the end of its task region. Any task that

belongs to the innermost enclosing taskgroup and has already begun execution must

run to completion or until a cancellation point is reached. Upon reaching a cancellation

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point, if a cancellation request has been observed, the task must jump to the end of its 2 task region and is considered completed. Any task that belongs to the innermost enclosing taskgroup and has not begun execution may be discarded, in which case it 3 is considered completed. When cancellation is observed for a parallel, sections, for, or do region, each thread of the binding thread set resumes execution at the end of the canceled region. If 6 the cancelled region is a parallel region, any tasks that have been created by a task 7 construct and their descendants are cancelled according to the above taskgroup 8 9 cancellation semantics. If the cancelled region is a sections, for, or do region, no 10 task cancellation occurs. C++ The usual C++ rules for object destruction are followed when cancellation is performed. 11 ----- Fortran All private objects or subobjects with ALLOCATABLE attribute that are allocated inside 12 the canceled construct are deallocated. 13 Fortran Note – The user is responsible for releasing locks and similar data structures that might 14 cause a deadlock when a cancel construct is encountered and blocked threads cannot 15 be canceled. 16 If the canceled construct contains a reduction or lastprivate clause, the final 17 value of the reduction or lastprivate variable is undefined. 18 19 When an if clause is present on a cancel construct and the if expression evaluates to false, the cancel construct does not request cancellation. The cancellation 20 point associated with the cancel construct is always encountered regardless of the 21 value of the if expression. 22 Restrictions 23 24 The restrictions to the cancel construct are as follows:

• Only one construct type may be specified in the clause list. If more than one

• The behavior for concurrent cancellation of a region and a region nested within it is

construct is specified, then the behavior is unspecified.

• At most one if clause can appear on the directive.

unspecified.

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1 2 3 4		• If the clause is taskgroup, the cancel construct must be closely nested inside a task construct. Otherwise, the cancel construct must be closely nested inside an OpenMP construct that matches the type specified in the clause of the cancel construct.
5		• A worksharing construct that is cancelled must not have a nowait clause.
6		<ul> <li>A loop construct that is cancelled must not have an ordered clause.</li> </ul>
7 8 9		• A construct that may be subject to cancellation must not encounter an orphaned cancellation point. That is, a cancellation point must only be encountered within that construct and must not be encountered elsewhere in its region.
10		• The cancel construct is a stand-alone directive.
11 12 13 14		• The cancel directive may be placed only at a point where a base language statement is allowed. The cancel directive may not be used in place of the statement following an if, while, do, switch, or label. See Appendix C for the formal grammar.
		C/C++
		Fortran —
15 16 17 18		• The cancel directive may be placed only at a point where a Fortran executable statement is allowed. The cancel directive may not be used as the action statement in an if statement or as the executable statement following a label if the label is referenced in the program.
		Fortran —
19 20		Cross References:  • cancel-var, see Section 2.3.1 on page 31
21		• cancellation point construct, see Section 2.13.2 on page 119
22		• omp_get_cancellation routine, see Section 3.2.9 on page 172
23	2.13.2	cancellation point Construct
24		Summary
25 26 27		The cancellation point construct introduces a user-defined cancellation point at which implicit or explicit tasks check if cancellation of the innermost enclosing region of the type specified has been requested

#### Syntax 1 \_\_\_\_\_ C/C++ -2 The syntax of the cancellation point construct is as follows: #pragma omp cancellation point clause new-line where *clause* is one of the following parallel sections for taskgroup Because the cancellation point construct is a stand-alone directive, there are 9 some restrictions on its placement within a program. The cancellation point directive may be placed only at a point where a base language statement is allowed. The 10 cancellation point directive may not be used in place of the statement following 11 an if, while, do, switch, or label. See Appendix C for the formal grammar. 12 C/C++ ----Fortran The syntax of the cancellation point construct is as follows: 13 !\$omp cancellation point clause where *clause* is one of the following 14 15 parallel 16 sections 17 do 18 taskgroup 19 Because the **cancellation point** construct is a stand-alone directive, there are 20 some restrictions on its placement within a program. The cancellation point directive may be placed only at a point where a Fortran executable statement is allowed. 21 The cancellation point directive may not be used as the action statement in an 22 23 if statement or as the executable statement following a label if the label is referenced 24 in the program.

- Fortran –

#### **Binding** 1 2 A cancellation point region binds to the current task region. **Description** 3 This directive introduces a user-defined cancellation point at which an implicit or 4 5 explicit task must check if cancellation of the innermost enclosing region of the type 6 specified in the clause has been requested. This construct does not implement a 7 synchronization between threads or tasks. When an implicit or explicit task reaches a user-defined cancellation point, the task 8 immediately checks for cancellation of the region specified in the clause and performs 9 cancellation of this region if cancellation is observed. If the clause specified is 10 taskgroup then the current task region is canceled. 11 Restrictions 12 • If the clause is taskgroup, the cancellation point construct must be nested 13 inside a task construct. Otherwise, the cancellation point construct must be 14 15 closely nested inside an OpenMP construct that matches the type specified in the 16 clause of the cancellation point construct. 17 • An OpenMP program with orphaned cancellation point constructs is non-conforming. 18 19 • The cancellation point construct is a stand-alone directive. - C/C++ ----• The cancellation point directive may be placed only at a point where a base 20 language statement is allowed. The cancellation point directive may not be 21 used in place of the statement following an if, while, do, switch, or label. 22 See Appendix C for the formal grammar. 23 - C/C++ ----- Fortran -----• The cancellation point directive may be placed only at a point where a 24 Fortran executable statement is allowed. The cancellation point directive may 25 not be used as the action statement in an if statement or as the executable statement 26 27 following a label if the label is referenced in the program.

Fortran —

#### Cross References:

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- cancel-var, see Section 2.3.1 on page 31
- cancellation point construct, see Section 2.13.2 on page 119
- omp\_get\_cancellation routine, see Section 3.2.9 on page 172

## 2.14 Data Environment

This section presents a directive and several clauses for controlling the data environment during the execution of parallel, task, simd, and worksharing regions.

- Section 2.14.1 on page 122 describes how the data-sharing attributes of variables referenced in parallel, task, simd, and worksharing regions are determined.
- The **threadprivate** directive, which is provided to create threadprivate memory, is described in Section 2.14.2 on page 126.
- Clauses that may be specified on directives to control the data-sharing attributes of variables referenced in parallel, task, simd or worksharing constructs are described in Section 2.14.3 on page 130.
- Clauses that may be specified on directives to copy data values from private or threadprivate variables on one thread to the corresponding variables on other threads in the team are described in Section 2.14.4 on page 147.

# 2.14.1 Data-sharing Attribute Rules

This section describes how the data-sharing attributes of variables referenced in **parallel**, **task**, **simd**, and worksharing regions are determined. The following two cases are described separately:

- Section 2.14.1.1 on page 122 describes the data-sharing attribute rules for variables referenced in a construct.
- Section 2.14.1.2 on page 125 describes the data-sharing attribute rules for variables referenced in a region, but outside any construct.

# 2.14.1.1 Data-sharing Attribute Rules for Variables Referenced in a Construct

The data-sharing attributes of variables that are referenced in a construct can be *predetermined*, *explicitly determined*, or *implicitly determined*, according to the rules outlined in this section.

1	Specifying a variable on a firstprivate, lastprivate, linear, or
2	reduction clause of an enclosed construct causes an implicit reference to the variable
3 4	in the enclosing construct. Such implicit references are also subject to the data-sharing attribute rules outlined in this section.
5	Certain variables and objects have predetermined data-sharing attributes as follows:
5	
6	• Variables appearing in <b>threadprivate</b> directives are threadprivate.
7	<ul> <li>Variables with automatic storage duration that are declared in a scope inside the</li> </ul>
8	construct are private.
9	<ul> <li>Objects with dynamic storage duration are shared.</li> </ul>
10	• Static data members are shared.
11 12	<ul> <li>The loop iteration variable(s) in the associated for-loop(s) of a for or parallel for construct is (are) private.</li> </ul>
13 14	• The loop iteration variable(s) in the associated for-loop(s) of a simd construct is (are) linear with a <i>linear-step</i> that is the increment of the associated loop.
15	<ul> <li>Variables with const-qualified type having no mutable member are shared.</li> </ul>
16 17	<ul> <li>Variables with static storage duration that are declared in a scope inside the construct are shared.</li> </ul>
18	<ul> <li>Variables appearing in depend clauses are shared.</li> </ul>
	C/C++
	Fortran —
19 20	<ul> <li>Variables and common blocks appearing in threadprivate directives are threadprivate.</li> </ul>
21 22	• The loop iteration variable(s) in the associated <i>do-loop(s)</i> of a <b>do</b> or <b>parallel do</b> construct is (are) private.
23 24	• The loop iteration variable(s) in the associated <i>do-loop(s)</i> of a <b>simd</b> construct is (are) linear with a <i>linear-step</i> that is the increment of the associated loop
25 26	• A loop iteration variable for a sequential loop in a parallel or task construct is private in the innermost such construct that encloses the loop.
27	<ul> <li>Implied-do indices and forall indices are private.</li> </ul>
28 29	<ul> <li>Cray pointees inherit the data-sharing attribute of the storage with which their Cray pointers are associated.</li> </ul>
30	<ul> <li>Assumed-size arrays are shared.</li> </ul>
31 32	<ul> <li>An associate name preserves the association with the selector established at the ASSOCIATE statement.</li> </ul>

• Variables appearing in **depend** clauses are shared. 1 - Fortran -2 Variables with predetermined data-sharing attributes may not be listed in data-sharing 3 attribute clauses, except for the cases listed below. For these exceptions only, listing a predetermined variable in a data-sharing attribute clause is allowed and overrides the 5 variable's predetermined data-sharing attributes. ----- C/C++ - The loop iteration variable(s) in the associated for-loop(s) of a for or parallel for construct may be listed in a private or lastprivate clause. 7 • Variables with const-qualified type having no mutable member may be listed in a 8 9 firstprivate clause, even if they are static data members. \_\_\_\_\_\_ C/C++ \_\_\_\_\_ - Fortran ----• The loop iteration variable(s) in the associated do-loop(s) of a do or parallel do 10 construct may be listed in a private or lastprivate clause. 11 12 Variables used as loop iteration variables in sequential loops in a parallel or task construct may be listed in data-sharing clauses on the construct itself, and on 13 enclosed constructs, subject to other restrictions. 14 15 • Assumed-size arrays may be listed in a **shared** clause. ------ Fortran -----Additional restrictions on the variables that may appear in individual clauses are 16 described with each clause in Section 2.14.3 on page 130. 17 18 Variables with explicitly determined data-sharing attributes are those that are referenced 19 in a given construct and are listed in a data-sharing attribute clause on the construct. 20 Variables with *implicitly determined* data-sharing attributes are those that are referenced in a given construct, do not have predetermined data-sharing attributes, and are not 21 listed in a data-sharing attribute clause on the construct. 22 23 Rules for variables with *implicitly determined* data-sharing attributes are as follows: 24 • In a parallel or task construct, the data-sharing attributes of these variables are determined by the **default** clause, if present (see Section 2.14.3.1 on page 131). 25 26 • In a parallel construct, if no default clause is present, these variables are shared. 27 28 • For constructs other than task, if no default clause is present, these variables inherit their data-sharing attributes from the enclosing context. 29 30 • In a task construct, if no default clause is present, a variable that in the 31 enclosing context is determined to be shared by all implicit tasks bound to the current

team is shared.

		Fortran —
1 2		• In an orphaned task construct, if no default clause is present, dummy arguments are firstprivate.
		Fortran —
3 4		• In a task construct, if no default clause is present, a variable whose data-sharing attribute is not determined by the rules above is firstprivate.
5 6 7		Additional restrictions on the variables for which data-sharing attributes cannot be implicitly determined in a task construct are described in Section 2.14.3.4 on page 137.
8 9	2.14.1.2	Data-sharing Attribute Rules for Variables Referenced in a Region but not in a Construct
10		The data-sharing attributes of variables that are referenced in a region, but not in a
11		construct, are determined as follows:
		C/C++
12 13		<ul> <li>Variables with static storage duration that are declared in called routines in the region are shared.</li> </ul>
14 15		<ul> <li>Variables with const-qualified type having no mutable member, and that are declared in called routines, are shared.</li> </ul>
16 17		• File-scope or namespace-scope variables referenced in called routines in the region are shared unless they appear in a <b>threadprivate</b> directive.
18		Objects with dynamic storage duration are shared.
19		• Static data members are shared unless they appear in a threadprivate directive.
20 21		• Formal arguments of called routines in the region that are passed by reference inherit the data-sharing attributes of the associated actual argument.
22		• Other variables declared in called routines in the region are private.  C/C++
		Fortran —
23 24 25		• Local variables declared in called routines in the region and that have the save attribute, or that are data initialized, are shared unless they appear in a threadprivate directive.
26 27 28		<ul> <li>Variables belonging to common blocks, or declared in modules, and referenced in called routines in the region are shared unless they appear in a threadprivate directive.</li> </ul>
29 30		• Dummy arguments of called routines in the region that are passed by reference inherit the data-sharing attributes of the associated actual argument.

 Cray pointees inherit the data-sharing attribute of the storage with which their Cray 1 2 pointers are associated. • Implied-do indices, forall indices, and other local variables declared in called 3 routines in the region are private. Fortran -5 **2.14.2** threadprivate Directive **Summary** The threadprivate directive specifies that variables are replicated, with each thread having its own copy. The **threadprivate** directive is a declarative directive. **Syntax** 9 ———— C/C++ — 10 The syntax of the **threadprivate** directive is as follows: #pragma omp threadprivate(list) new-line 11 where *list* is a comma-separated list of file-scope, namespace-scope, or static block-scope variables that do not have incomplete types. 12 C/C++ ----- Fortran -----13 The syntax of the **threadprivate** directive is as follows:

where *list* is a comma-separated list of named variables and named common blocks.

!\$omp threadprivate(list)

Common block names must appear between slashes.

Fortran

## 

#### Description

Each copy of a threadprivate variable is initialized once, in the manner specified by the program, but at an unspecified point in the program prior to the first reference to that copy. The storage of all copies of a threadprivate variable is freed according to how static variables are handled in the base language, but at an unspecified point in the program.

A program in which a thread references another thread's copy of a threadprivate variable is non-conforming.

The content of a threadprivate variable can change across a task scheduling point if the executing thread switches to another task that modifies the variable. For more details on task scheduling, see Section 1.3 on page 13 and Section 2.11 on page 88.

In **parallel** regions, references by the master thread will be to the copy of the variable in the thread that encountered the **parallel** region.

During the sequential part references will be to the initial thread's copy of the variable. The values of data in the initial thread's copy of a threadprivate variable are guaranteed to persist between any two consecutive references to the variable in the program.

The values of data in the threadprivate variables of non-initial threads are guaranteed to persist between two consecutive active **parallel** regions only if all the following conditions hold:

- Neither parallel region is nested inside another explicit parallel region.
- The number of threads used to execute both **parallel** regions is the same.
- The thread affinity policies used to execute both **parallel** regions are the same.
- The value of the *dyn-var* internal control variable in the enclosing task region is *false* at entry to both **parallel** regions.

If these conditions all hold, and if a threadprivate variable is referenced in both regions, then threads with the same thread number in their respective regions will reference the same copy of that variable.

#### C/C++

If the above conditions hold, the storage duration, lifetime, and value of a thread's copy of a threadprivate variable that does not appear in any **copyin** clause on the second region will be retained. Otherwise, the storage duration, lifetime, and value of a thread's copy of the variable in the second region is unspecified.

If the value of a variable referenced in an explicit initializer of a threadprivate variable is modified prior to the first reference to any instance of the threadprivate variable, then the behavior is unspecified.

The order in which any constructors for different threadprivate variables of class type 1 2 are called is unspecified. The order in which any destructors for different threadprivate 3 variables of class type are called is unspecified. C/C++ -Fortran -A variable is affected by a **copyin** clause if the variable appears in the **copyin** clause 5 or it is in a common block that appears in the **copyin** clause. 6 If the above conditions hold, the definition, association, or allocation status of a thread's copy of a threadprivate variable or a variable in a threadprivate common 8 block, that is not affected by any copyin clause that appears on the second region, will 9 10 be retained. Otherwise, the definition and association status of a thread's copy of the variable in the second region is undefined, and the allocation status of an allocatable 11 variable will be implementation defined. 12 13 If a threadprivate variable or a variable in a threadprivate common block is not affected by any copyin clause that appears on the first parallel region in which 14 it is referenced, the variable or any subobject of the variable is initially defined or 15 undefined according to the following rules: 16 17 If it has the ALLOCATABLE attribute, each copy created will have an initial 18 allocation status of not currently allocated. 19 • If it has the **POINTER** attribute: 20 if it has an initial association status of disassociated, either through explicit initialization or default initialization, each copy created will have an association 21 status of disassociated; 22 23 • otherwise, each copy created will have an association status of undefined. 24 • If it does not have either the **POINTER** or the **ALLOCATABLE** attribute: 25 • if it is initially defined, either through explicit initialization or default 26 initialization, each copy created is so defined; 27 • otherwise, each copy created is undefined. Fortran

For examples of the **threadprivate** directive, see Section A.27 on page 304.

#### Restrictions

The restrictions to the **threadprivate** directive are as follows:

 A threadprivate variable must not appear in any clause except the copyin, copyprivate, schedule, num\_threads, and if clauses.

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1	A program in which an unfield task accesses threadprivate storage is non-conforming
2 3 4	• A variable that is part of another variable (as an array or structure element) cannot appear in a threadprivate clause unless it is a static data member of a C++ class.
5 6 7	<ul> <li>A threadprivate directive for file-scope variables must appear outside any definition or declaration, and must lexically precede all references to any of the variables in its list.</li> </ul>
8 9 10	• A threadprivate directive for static class member variables must appear in the class definition, in the same scope in which the member variables are declared, and must lexically precede all references to any of the variables in its list.
11 12 13	<ul> <li>A threadprivate directive for namespace-scope variables must appear outside any definition or declaration other than the namespace definition itself, and must lexically precede all references to any of the variables in its list.</li> </ul>
14 15 16	<ul> <li>Each variable in the list of a threadprivate directive at file, namespace, or class scope must refer to a variable declaration at file, namespace, or class scope that lexically precedes the directive.</li> </ul>
17 18 19	<ul> <li>A threadprivate directive for static block-scope variables must appear in the scope of the variable and not in a nested scope. The directive must lexically precede all references to any of the variables in its list.</li> </ul>
20 21 22	<ul> <li>Each variable in the list of a threadprivate directive in block scope must refer to a variable declaration in the same scope that lexically precedes the directive. The variable declaration must use the static storage-class specifier.</li> </ul>
23 24 25	<ul> <li>If a variable is specified in a threadprivate directive in one translation unit, it must be specified in a threadprivate directive in every translation unit in which it is declared.</li> </ul>
26	• The address of a threadprivate variable is not an address constant.
27	• A threadprivate variable must not have an incomplete type or a reference type.
28	• A threadprivate variable with class type must have:
29 30	<ul> <li>an accessible, unambiguous default constructor in case of default initialization without a given initializer;</li> </ul>
31 32	<ul> <li>an accessible, unambiguous constructor accepting the given argument in case of direct initialization;</li> </ul>
33 34	<ul> <li>an accessible, unambiguous copy constructor in case of copy initialization with an explicit initializer.</li> </ul>
25	C/C++
35	Fortran —

• A variable that is part of another variable (as an array or structure element) cannot 1 2 appear in a threadprivate clause. 3 • The **threadprivate** directive must appear in the declaration section of a scoping unit in which the common block or variable is declared. Although variables in 4 5 common blocks can be accessed by use association or host association, common 6 block names cannot. This means that a common block name specified in a 7 threadprivate directive must be declared to be a common block in the same 8 scoping unit in which the threadprivate directive appears. • If a threadprivate directive specifying a common block name appears in one 9 10 program unit, then such a directive must also appear in every other program unit that contains a **COMMON** statement specifying the same name. It must appear after the last 11 12 such **COMMON** statement in the program unit. 13 • If a threadprivate variable or a threadprivate common block is declared 14 with the BIND attribute, the corresponding C entities must also be specified in a threadprivate directive in the C program. 15 • A blank common block cannot appear in a threadprivate directive. 16 17 • A variable can only appear in a **threadprivate** directive in the scope in which it is declared. It must not be an element of a common block or appear in an 18 19 **EQUIVALENCE** statement. 20 A variable that appears in a threadprivate directive must be declared in the 21 scope of a module or have the **SAVE** attribute, either explicitly or implicitly. —— Fortran – Cross References: 22 23 • dyn-var ICV, see Section 2.3 on page 30. 24 • number of threads used to execute a **parallel** region, see Section 2.5.1 on page 40. 25 • copyin clause, see Section 2.14.4.1 on page 148. 2.14.3 **Data-Sharing Attribute Clauses** 27 Several constructs accept clauses that allow a user to control the data-sharing attributes

of variables referenced in the construct. Data-sharing attribute clauses apply only to

variables for which the names are visible in the construct on which the clause appears.

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Most of the clauses accept a comma-separated list of list items (see Section 2.1 on page 2 24). All list items appearing in a clause must be visible, according to the scoping rules 3 of the base language. With the exception of the **default** clause, clauses may be 4 repeated as needed. A list item that specifies a given variable may not appear in more 5 than one clause on the same directive, except that a variable may be specified in both 6 firstprivate and lastprivate clauses. - C/C++ ----If a variable referenced in a data-sharing attribute clause has a type derived from a 7 8 template, and there are no other references to that variable in the program, then any 9 behavior related to that variable is unspecified. \_\_\_\_\_ C/C++ \_\_\_\_\_ - Fortran ———— 10 A named common block may be specified in a list by enclosing the name in slashes. When a named common block appears in a list, it has the same meaning as if every 11 12 explicit member of the common block appeared in the list. An explicit member of a common block is a variable that is named in a COMMON statement that specifies the 13 common block name and is declared in the same scoping unit in which the clause 14 15 appears. Although variables in common blocks can be accessed by use association or host 16 17 association, common block names cannot. As a result, a common block name specified 18 in a data-sharing attribute clause must be declared to be a common block in the same scoping unit in which the data-sharing attribute clause appears. 19 20 When a named common block appears in a private, firstprivate, 21 lastprivate, or shared clause of a directive, none of its members may be declared 22 in another data-sharing attribute clause in that directive (see Section A.29 on page 311 23 for examples). When individual members of a common block appear in a private, firstprivate, lastprivate, or reduction clause of a directive, the storage of 24 25 the specified variables is no longer associated with the storage of the common block 26 itself (see Section A.33 on page 320 for examples). ——— Fortran — 27 **2.14.3.1** default clause

## Summary

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The default clause explicitly determines the data-sharing attributes of variables that are referenced in a parallel or task construct and would otherwise be implicitly determined (see Section 2.14.1.1 on page 122).

## **Syntax** 1 C/C++ — 2 The syntax of the **default** clause is as follows: default(shared | none) C/C++ - Fortran -5 The syntax of the **default** clause is as follows: default(private | firstprivate | shared | none) - Fortran – 6 Description The default (shared) clause causes all variables referenced in the construct that have implicitly determined data-sharing attributes to be shared. ——— Fortran — The default (firstprivate) clause causes all variables in the construct that have 10 implicitly determined data-sharing attributes to be firstprivate. 11 12 The default (private) clause causes all variables referenced in the construct that have implicitly determined data-sharing attributes to be private. 13 Fortran — The default (none) clause requires that each variable that is referenced in the 14 construct, and that does not have a predetermined data-sharing attribute, must have its 15 data-sharing attribute explicitly determined by being listed in a data-sharing attribute 16 17 clause. See Section A.30 on page 313 for examples. Restrictions 18 19 The restrictions to the **default** clause are as follows: 20 Only a single default clause may be specified on a parallel or task directive.

#### 2.14.3.2 shared clause

#### 2 Summary

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The **shared** clause declares one or more list items to be shared by tasks generated by a **parallel** or **task** construct.

#### Syntax

The syntax of the **shared** clause is as follows:

shared(list)

#### Description

All references to a list item within a task refer to the storage area of the original variable at the point the directive was encountered.

It is the programmer's responsibility to ensure, by adding proper synchronization, that storage shared by an explicit task region does not reach the end of its lifetime before the explicit task region completes its execution.

#### · Fortran -

The association status of a shared pointer becomes undefined upon entry to and on exit from the **parallel** or **task** construct if it is associated with a target or a subobject of a target that is in a **private**, **firstprivate**, **lastprivate**, or **reduction** clause inside the construct.

Under certain conditions, passing a shared variable to a non-intrinsic procedure may result in the value of the shared variable being copied into temporary storage before the procedure reference, and back out of the temporary storage into the actual argument storage after the procedure reference. It is implementation defined when this situation occurs. See Section A.31 on page 315 for an example of this behavior.

**Note** – Use of intervening temporary storage may occur when the following three conditions hold regarding an actual argument in a reference to a non-intrinsic procedure:

- a. The actual argument is one of the following:
- A shared variable.
- A subobject of a shared variable.

 An object associated with a shared variable. 1 • An object associated with a subobject of a shared variable. b. The actual argument is also one of the following: 3 · An array section. • An array section with a vector subscript. · An assumed-shape array. A pointer array. c. The associated dummy argument for this actual argument is an explicit-shape array 8 9 or an assumed-size array. 10 These conditions effectively result in references to, and definitions of, the temporary 11 storage during the procedure reference. Any references to (or definitions of) the shared 12 storage that is associated with the dummy argument by any other task must be synchronized with the procedure reference to avoid possible race conditions. 13 14 Fortran -2.14.3.3 15 private clause 16 Summary 17 The private clause declares one or more list items to be private to a task or a SIMD 18 lane. Syntax 1 4 1 19 20 The syntax of the **private** clause is as follows: private(list) **Description** 21 22 Each task that references a list item that appears in a **private** clause in any statement 23 in the construct receives a new list item. Each SIMD lane used in a simd construct that 24 references a list item that appears in a private clause in any statement in the construct 25 receives a new list item. Language-specific attributes for new list items are derived from

the corresponding original list items. In the rest of the region, it is unspecified whether 1 2 references are to the new list item or the original list item. Therefore, if an attempt is 3 made to reference the original item, its value after the region is also unspecified. If a 4 SIMD construct or a task does not reference a list item that appears in a private clause, it is unspecified whether SIMD lanes or the task receive a new list item. 5 6 The value and/or allocation status of the original list item will change only: 7 • if accessed and modified via pointer, 8 • if possibly accessed in the region but outside of the construct, 9 • as a side effect of directives or clauses, or - Fortran -• if accessed and modified via construct association. 10 - Fortran —— 11 List items that appear in a private, firstprivate, or reduction clause in a parallel construct may also appear in a private clause in an enclosed parallel, 12 task, or worksharing construct or simd construct. List items that appear in a 13 private or firstprivate clause in a task construct may also appear in a 14 private clause in an enclosed parallel or task construct. List items that appear 15 in a private, firstprivate, lastprivate, or reduction clause in a 16 worksharing construct may also appear in a private clause in an enclosed parallel 17 or task construct. See Section A.32 on page 316 for an example. 18 \_\_\_\_\_ C/C++ \_\_\_\_ 19 A new list item of the same type, with automatic storage duration, is allocated for the construct. The storage and thus lifetime of these list items lasts until the block in which 20 they are created exits. The size and alignment of the new list item are determined by the 21 type of the variable. This allocation occurs once for each task generated by the construct 22 and/or once for each SIMD lane used by the construct. 23 24 The new list item is initialized, or has an undefined initial value, as if it had been locally 25 declared without an initializer. The order in which any default constructors for different private variables of class type are called is unspecified. The order in which any 26 destructors for different private variables of class type are called is unspecified. 27 C/C++ Fortran ———— 28 If any statement of the construct references a list item, a new list item of the same type and type parameters is allocated: once for each implicit task in the parallel 29 construct; once for each task generated by a task construct; and once for each SIMD 30 lane used by a SIMD construct. The initial value of the new list item is undefined. 31 Within a parallel, worksharing, task region, or simd region, the initial status of a 32 33 private pointer is undefined.

For a list item or the subobject of a list item with the **ALLOCATABLE** attribute: 1 • if the allocation status is "not currently allocated", the new list item or the subobject 3 of the new list item will have an initial allocation status of "not currently allocated"; • if the allocation status is "currently allocated", the new list item or the subobject of 4 the new list item will have an initial allocation status of "currently allocated". If it is 5 an array, the bounds will be the same as that of the original list item or the subobject 6 7 of the original list item. 8 A list item that appears in a **private** clause may be storage-associated with other variables when the **private** clause is encountered. Storage association may exist 9 because of constructs such as **EQUIVALENCE** or **COMMON**. If A is a variable appearing 10 in a **private** clause and B is a variable that is storage-associated with A, then: 11 12 • The contents, allocation, and association status of B are undefined on entry to the 13 parallel, task, or simd region. 14 A list item that appears in a private clause may be a selector of an ASSOCIATE construct. If the construct association is established prior to a parallel region, the 15 association between the associate name and the original list item will be retained in 16 17 the region. • Any definition of A, or of its allocation or association status, causes the contents, 18 allocation, and association status of B to become undefined. 19 20 • Any definition of B, or of its allocation or association status, causes the contents, allocation, and association status of A to become undefined. 21 22 For examples, see Section A.33 on page 320. — Fortran – 23 For examples of the **private** clause, see Section A.32 on page 316. Restrictions 24 25 The restrictions to the **private** clause are as follows: 26 • A variable that is part of another variable (as an array or structure element) cannot 27 appear in a **private** clause. \_\_\_\_\_ C/C++ \_\_\_\_ 28 • A variable of class type (or array thereof) that appears in a **private** clause requires an accessible, unambiguous default constructor for the class type. 29 30 A variable that appears in a private clause must not have a const-qualified type

unless it is of class type with a **mutable** member. This restriction does not apply to

the firstprivate clause.

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C/C++ Fortran
— Fortran
private clause must either be definable, or an criction does not apply to the firstprivate clause.
elist statements, in variable format expressions, and in ction definitions, may not appear in a <b>private</b> clause.
N) attribute may not appear in the private clause.  y to the firstprivate clause.
Fortran —
clares one or more list items to be private to a task, and value that the corresponding original item has when the
ate clause is as follows:
ovides a superset of the functionality provided by the
estprivate clause is subject to the private clause 2.14.3.3 on page 134, except as noted. In addition, the the original list item existing before the construct. The is done once for each task that references the list item to the initialization is done prior to the execution of the

For a **firstprivate** clause on a **parallel** or **task** construct, the initial value of the new list item is the value of the original list item that exists immediately prior to the construct in the task region where the construct is encountered. For a **firstprivate** clause on a worksharing construct, the initial value of the new list item for each implicit task of the threads that execute the worksharing construct is the value of the original list item that exists in the implicit task immediately prior to the point in time that the worksharing construct is encountered.

To avoid race conditions, concurrent updates of the original list item must be synchronized with the read of the original list item that occurs as a result of the **firstprivate** clause.

If a list item appears in both firstprivate and lastprivate clauses, the update required for lastprivate occurs after all the initializations for firstprivate.

### \_\_\_\_\_ C/C++ \_\_\_\_\_

For variables of non-array type, the initialization occurs by copy assignment. For an array of elements of non-array type, each element is initialized as if by assignment from an element of the original array to the corresponding element of the new array. For variables of class type, a copy constructor is invoked to perform the initialization. The order in which copy constructors for different variables of class type are called is unspecified.

#### C/C++

## ------ Fortran

If the original list item does not have the **POINTER** attribute, initialization of the new list items occurs as if by intrinsic assignment, unless the original list item has the allocation status of not currently allocated, in which case the new list items will have the same status.

If the original list item has the **POINTER** attribute, the new list items receive the same association status of the original list item as if by pointer assignment.

## ------ Fortran

#### Restrictions

The restrictions to the **firstprivate** clause are as follows:

- A variable that is part of another variable (as an array or structure element) cannot appear in a firstprivate clause.
- A list item that is private within a parallel region must not appear in a firstprivate clause on a worksharing construct if any of the worksharing regions arising from the worksharing construct ever bind to any of the parallel regions arising from the parallel construct.

1 2 3 4		• A list item that appears in a reduction clause of a parallel construct must not appear in a firstprivate clause on a worksharing or task construct if any of the worksharing or task regions arising from the worksharing or task construct ever bind to any of the parallel regions arising from the parallel construct.
5 6 7		• A list item that appears in a <b>reduction</b> clause in a worksharing construct must not appear in a <b>firstprivate</b> clause in a task construct encountered during execution of any of the worksharing regions arising from the worksharing construct.
8		• A variable of class type (or array thereof) that appears in a <b>firstprivate</b> clause requires an accessible, unambiguous copy constructor for the class type.
10 11		• A variable that appears in a <b>firstprivate</b> clause must not have an incomplete type or a reference type.
		C/C++
12		Fortran —
13 14 15		<ul> <li>Variables that appear in namelist statements, in variable format expressions, and in expressions for statement function definitions, may not appear in a firstprivate clause.</li> </ul>
		Fortran —
16	2.14.3.5	lastprivate <b>clause</b>
17		Summary
18 19 20		The lastprivate clause declares one or more list items to be private to an implicit task or to a SIMD lane, and causes the corresponding original list item to be updated after the end of the region.
21		Syntax
22		The syntax of the lastprivate clause is as follows:
		lastprivate(list)

#### **Description** 1 The lastprivate clause provides a superset of the functionality provided by the 2 private clause. A list item that appears in a lastprivate clause is subject to the private clause 4 5 semantics described in Section 2.14.3.3 on page 134. In addition, when a 6 lastprivate clause appears on the directive that identifies a worksharing construct or a SIMD construct, the value of each new list item from the sequentially last iteration 7 8 of the associated loops, or the lexically last section construct, is assigned to the original list item. 9 \_\_\_\_\_ C/C++ \_\_\_\_ For an array of elements of non-array type, each element is assigned to the 10 corresponding element of the original array. 11 C/C++ Fortran If the original list item does not have the **POINTER** attribute, its update occurs as if by 12 intrinsic assignment. 13 If the original list item has the **POINTER** attribute, its update occurs as if by pointer 14 assignment. 15 Fortran — 16 List items that are not assigned a value by the sequentially last iteration of the loops, or by the lexically last **section** construct, have unspecified values after the construct. 17 18 Unassigned subcomponents also have unspecified values after the construct. 19 The original list item becomes defined at the end of the construct if there is an implicit 20 barrier at that point. To avoid race conditions, concurrent reads or updates of the original list item must be synchronized with the update of the original list item that occurs as a 21 result of the lastprivate clause. 22 23 If the lastprivate clause is used on a construct to which nowait is applied, accesses to the original list item may create a data race. To avoid this, synchronization 24 25 must be inserted to ensure that the sequentially last iteration or lexically last section construct has stored and flushed that list item. 26

If a list item appears in both firstprivate and lastprivate clauses, the update required for lastprivate occurs after all initializations for firstprivate.

For an example of the lastprivate clause, see Section A.35 on page 324.

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1	Restrictions	
2	The restrictions to the lastprivate clause are as follows:	
3	<ul> <li>A variable that is part of another variable (as an array or structure element) cannot</li></ul>	
4	appear in a lastprivate clause.	
5	<ul> <li>A list item that is private within a parallel region, or that appears in the</li></ul>	
6	reduction clause of a parallel construct, must not appear in a lastprivate	
7	clause on a worksharing construct if any of the corresponding worksharing regions	
8	ever binds to any of the corresponding parallel regions.	
9 10 11	<ul> <li>C/C++</li> <li>A variable of class type (or array thereof) that appears in a lastprivate clause requires an accessible, unambiguous default constructor for the class type, unless the list item is also specified in a firstprivate clause.</li> </ul>	
12	<ul> <li>A variable of class type (or array thereof) that appears in a lastprivate clause</li></ul>	
13	requires an accessible, unambiguous copy assignment operator for the class type. The	
14	order in which copy assignment operators for different variables of class type are	
15	called is unspecified.	
16	<ul> <li>A variable that appears in a lastprivate clause must not have a const-qualified</li></ul>	
17	type unless it is of class type with a mutable member.	
18	<ul> <li>A variable that appears in a lastprivate clause must not have an incomplete type</li></ul>	
19	or a reference type.	
	C/C++	
	Fortran	
20	• A variable that appears in a lastprivate clause must be definable.	
21	<ul> <li>An original list item with the ALLOCATABLE attribute in the sequentially last</li></ul>	
22	iteration or lexically last section must have an allocation status of allocated upon exit	
23	from that iteration or section.	
24	<ul> <li>Variables that appear in namelist statements, in variable format expressions, and in</li></ul>	
25	expressions for statement function definitions, may not appear in a lastprivate	
26	clause.	
	Fortran	

#### 1 2.14.3.6 reduction clause

### 2 Summary

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The **reduction** clause specifies a *reduction-identifier* and one or more list items. For each list item, a private copy is created in each implicit task or SIMD lane, and is initialized with the initializer value of the *reduction-identifier*. After the end of the region, the original list item is updated with the values of the private copies using the combiner associated with the *reduction-identifier*.

- C/C++ -

### **Syntax**

The syntax of the **reduction** clause is as follows:

reduction (reduction-identifier: list)

where:

reduction-identifier is either an identifier or one of the following operators: +, -, \*,

&, |, ^, && and ||

C++

reduction-identifier is either an id-expression or one of the following operators: +, -,

\*, &, |, ^, && and ||

The following table lists each *reduction-identifier* that is implicitly declared at every scope for arithmetic types and its semantic initializer value. The actual initializer value is that value as expressed in the data type of the reduction list item.

Identifier	Initializer	Combiner
+	omp_priv = 0	omp_out += omp_in
*	omp_priv = 1	omp_out *= omp_in
-	omp_priv = 0	omp_out += omp_in
&	omp_priv = ~0	omp_out &= omp_in
1	omp_priv = 0	omp_out  = omp_in

omp out ^= omp in

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8 9 10 where omp\_in and omp\_out correspond to two identifiers that refer to storage of the type of the list item. omp\_out holds the final value of the combiner operation.

C/C++

Fortran

The syntax of the **reduction** clause is as follows:

reduction (reduction-identifier: list)

omp priv = 0

where reduction-identifier is either a base language identifier, or a user-defined operator, or one of the following operators: +, -, \*, .and., .or., .eqv., .neqv., or one of the following intrinsic procedure names: max, min, iand, ior, ieor.

The following table lists each *reduction-identifier* that is implicitly declared for numeric and logical types and its semantic initializer value. The actual initializer value is that value as expressed in the data type of the reduction list item..

Identifier	Initializer	Combiner
+	omp_priv = 0	omp_out = omp_in + omp_out
*	omp_priv = 1	<pre>omp_out = omp_in * omp_out</pre>
-	omp_priv = 0	<pre>omp_out = omp_in + omp_out</pre>
.and.	<pre>omp_priv = .true.</pre>	<pre>omp_out = omp_in .and. omp_out</pre>
.or.	<pre>omp_priv = .false.</pre>	<pre>omp_out = omp_in .or. omp_out</pre>
.eqv.	<pre>omp_priv = .true.</pre>	<pre>omp_out = omp_in .eqv. omp_out</pre>
.neqv.	<pre>omp_priv = .false.</pre>	<pre>omp_out = omp_in .neqv. omp_out</pre>
max	<pre>omp_priv = Least representable number in the reduction list item type</pre>	<pre>omp_out = max(omp_in, omp_out)</pre>

min	<pre>omp_priv = Largest representable number in the reduction list item type</pre>	<pre>omp_out = min(omp_in, omp_out)</pre>
iand	<pre>omp_priv = All bits on</pre>	<pre>omp_out = iand(omp_in, omp_out)</pre>
ior	omp_priv = 0	<pre>omp_out = ior(omp_in, omp_out)</pre>
ieor	omp_priv = 0	<pre>omp_out = ieor(omp_in, omp_out)</pre>

– Fortran –

 Any reduction-identifier that is defined with the **declare reduction** directive is also valid. In that case, the initializer and combiner of the reduction-identifier are specified by the initializer-clause and the combiner in the **declare reduction** directive.

## **Description**

The reduction clause can be used to perform some forms of recurrence calculations (involving mathematically associative and commutative operators) in parallel.

For parallel and worksharing constructs, a private copy of each list item is created, one for each implicit task, as if the private clause had been used. For the simd construct, a private copy of each list item is created, one for each SIMD lane as if the private clause had been used. The private copy is then initialized as specified above. At the end of the region for which the reduction clause was specified, the original list item is updated by combining its original value with the final value of each of the private copies, using the combiner of the specified reduction-identifier.

The *reduction-identifier* specified in the **reduction** clause must match a previously declared *reduction-identifier* of the same name and type for each of the list items. This match is done by means of a name lookup in the base language.

C++

If the type of a list item is a reference to a type T then the type will considered to be T for all purposes of this clause.

If the type is a derived class, then any *reduction-identifier* that matches its base classes are also a match, if there is no specific match for the type.

If the *reduction-identifier* is not an *id-expression* then it is implicitly converted to one by prepending the keyword operator (for example, + becomes *operator*+).

If the *reduction-identifier* is qualified then a qualified name lookup is used to find the declaration.

2 3 4 5 6 7 8 9 ensured through a barrier synchronization. 10 11 12 13 14 the same location in the OpenMP program. 15 16 17 18 reduction computation. Restrictions 19 20 The restrictions to the **reduction** clause are as follows: 21 22 from the worksharing construct bind. 23 24 25 26 27 28 29 C/C++ -30 31 32 33

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If the reduction-identifier is unqualified then an argument-dependent name lookup must be performed using the type of each list item. If nowait is not used, the reduction computation will be complete at the end of the construct; however, if the reduction clause is used on a construct to which nowait is also applied, accesses to the original list item will create a race and, thus, have unspecified effect unless synchronization ensures that they occur after all threads have executed all of their iterations or **section** constructs, and the reduction computation has completed and stored the computed value of that list item. This can most simply be

The location in the OpenMP program at which the values are combined and the order in which the values are combined are unspecified. Therefore, when comparing sequential and parallel runs, or when comparing one parallel run to another (even if the number of threads used is the same), there is no guarantee that bit-identical results will be obtained or that side effects (such as floating point exceptions) will be identical or take place at

To avoid race conditions, concurrent reads or updates of the original list item must be synchronized with the update of the original list item that occurs as a result of the

- A list item that appears in a **reduction** clause of a worksharing construct must be shared in the parallel regions to which any of the worksharing regions arising
- A list item that appears in a **reduction** clause of the innermost enclosing worksharing or parallel construct may not be accessed in an explicit task.
- Any number of **reduction** clauses can be specified on the directive, but a list item can appear only once in the **reduction** clauses for that directive.
- For a reduction-identifier declared with the declare reduction construct, the directive must appear before its use in a reduction clause.

• The type of a list item that appears in a **reduction** clause must be valid for the reduction-identifier. For a max or min reduction in C, the type of the list item must be an allowed arithmetic data type: char, int, float, double, or Bool, possibly modified with long, short, signed, or unsigned. For a max or min

1 2 3	reduction in C++, the type of the list item must be an allowed arithmetic data type: char, wchar_t, int, float, double, or bool, possibly modified with long, short, signed, or unsigned.
4	• Arrays may not appear in a reduction clause.
5	• A list item that appears in a reduction clause must not be const-qualified.
6 7	<ul> <li>If a list-item is a reference type then it must bind to the same object for all threads of the team.</li> </ul>
8	• The <i>reduction-identifier</i> for any list item must be unambiguous and accessible.  C/C++
	Fortran
9 10	• The type of a list item that appears in a <b>reduction</b> clause must be valid for the reduction operator or intrinsic.
11	• A list item that appears in a reduction clause must be definable.
12	• A procedure pointer may not appear in a reduction clause.
13 14	<ul> <li>A pointer with the INTENT(IN) attribute may not appear in the reduction clause.</li> </ul>
15 16 17	<ul> <li>An original list item with the ALLOCATABLE attribute must be in the allocated state at entry to the construct containing the reduction clause. Additionally, the list item must not be deallocated and/or allocated within the region.</li> </ul>
18 19 20	<ul> <li>If the reduction-identifier is defined in a declare reduction directive, the declare reduction directive must be in the same subprogram, or accessible by host or use association.</li> </ul>
21 22	• If the <i>reduction-identifier</i> is a user-defined operator, the same explicit interface for that operator must be accessible as at the <b>declare reduction</b> directive.
23 24 25 26	• If the <i>reduction-identifier</i> is defined in a <b>declare reduction</b> directive, any subroutine or function referenced in the initializer clause or combiner expression must be an intrinsic function, or must have an explicit interface where the same explicit interface is accessible as at the <b>declare reduction</b> directive.
	Fortran —

## **2.14.3.7** linear clause

## Summary

The linear clause declares one or more list items to be private to a SIMD lane and to have a linear relationship with respect to the iteration space of a loop.

## Syntax

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The syntax of the linear clause is as follows:

linear(list[:linear-step])

### Description

The linear clause provides a superset of the functionality provided by the private clause.

A list item that appears in a linear clause is subject to the private clause semantics described in Section 2.14.3.3 on page 134 except as noted. In addition, the value of the new list item on each iteration of the associated loop(s) corresponds to the value of the original list item before entering the construct plus the logical number of the iteration times *linear-step*. If *linear-step* is not specified it is assumed to be 1. The value corresponding to the sequentially last iteration of the associated loops is assigned to the original list item.

#### Restrictions

- The *linear-step* expression must be invariant during the execution of the region associated with the construct. Otherwise, the execution results in unspecified behavior.
- A *list-item* cannot appear in more than one linear clause.
- A list-item that appears in a linear clause cannot appear in any other data-sharing attribute clause.

C/C++

• A *list-item* that appears in a **linear** clause must be of integral or pointer type.

---- Fortran -

• A *list-item* that appears in a **linear** clause must be of type **integer**.

- Fortran -

## 22 2.14.4 Data Copying Clauses

This section describes the **copyin** clause (allowed on the **parallel** directive and combined parallel worksharing directives) and the **copyprivate** clause (allowed on the **single** directive).

These clauses support the copying of data values from private or threadprivate variables on one implicit task or thread to the corresponding variables on other implicit tasks or threads in the team.

The clauses accept a comma-separated list of list items (see Section 2.1 on page 24). All list items appearing in a clause must be visible, according to the scoping rules of the base language. Clauses may be repeated as needed, but a list item that specifies a given variable may not appear in more than one clause on the same directive.

### 2.14.4.1 copyin clause

### Summary

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23 24 The **copyin** clause provides a mechanism to copy the value of the master thread's threadprivate variable to the threadprivate variable of each other member of the team executing the **parallel** region.

### Syntax

The syntax of the **copyin** clause is as follows:

copyin(list)

## Description

is unspecified.

The copy is done after the team is formed and prior to the start of execution of the associated structured block. For variables of non-array type, the copy occurs by copy assignment. For an array of elements of non-array type, each element is copied as if by assignment from an element of the master thread's array to the corresponding element of the other thread's array. For class types, the copy assignment operator is invoked. The order in which copy assignment operators for different variables of class type are called

C/C++

C/C++

Fortran —

The copy is done, as if by assignment, after the team is formed and prior to the start of execution of the associated structured block.

1 2 3		On entry to any parallel region, each thread's copy of a variable that is affected by a copyin clause for the parallel region will acquire the allocation, association, and definition status of the master thread's copy, according to the following rules:
4 5		• If the original list item has the <b>POINTER</b> attribute, each copy receives the same association status of the master thread's copy as if by pointer assignment.
6 7 8 9		• If the original list item does not have the <b>POINTER</b> attribute, each copy becomes defined with the value of the master thread's copy as if by intrinsic assignment, unless it has the allocation status of not currently allocated, in which case each copy will have the same status.  Fortran
10		For an example of the <b>copyin</b> clause, see Section A.37 on page 331.
11		Restrictions
12		The restrictions to the <b>copyin</b> clause are as follows:
		C/C++
13		• A list item that appears in a copyin clause must be threadprivate.
14 15		<ul> <li>A variable of class type (or array thereof) that appears in a copyin clause requires an accessible, unambiguous copy assignment operator for the class type.</li> </ul>
		Fortran —
16 17 18		<ul> <li>A list item that appears in a copyin clause must be threadprivate. Named variables appearing in a threadprivate common block may be specified: it is not necessary to specify the whole common block.</li> </ul>
19 20		<ul> <li>A common block name that appears in a copyin clause must be declared to be a common block in the same scoping unit in which the copyin clause appears.</li> </ul>
		Fortran —
21	2.14.4.2	copyprivate <b>clause</b>
22		Summary
23		The <b>copyprivate</b> clause provides a mechanism to use a private variable to broadcast
24		a value from the data environment of one implicit task to the data environments of the
25		other implicit tasks belonging to the parallel region.

To avoid race conditions, concurrent reads or updates of the list item must be synchronized with the update of the list item that occurs as a result of the **copyprivate** clause.

### **Syntax**

The syntax of the **copyprivate** clause is as follows:

copyprivate(list)

### **Description**

The effect of the **copyprivate** clause on the specified list items occurs after the execution of the structured block associated with the **single** construct (see Section 2.7.3 on page 56), and before any of the threads in the team have left the barrier at the end of the construct.

#### C/C++ —

In all other implicit tasks belonging to the <code>parallel</code> region, each specified list item becomes defined with the value of the corresponding list item in the implicit task whose thread executed the structured block. For variables of non-array type, the definition occurs by copy assignment. For an array of elements of non-array type, each element is copied by copy assignment from an element of the array in the data environment of the implicit task associated with the thread that executed the structured block to the corresponding element of the array in the data environment of the other implicit tasks. For class types, a copy assignment operator is invoked. The order in which copy assignment operators for different variables of class type are called is unspecified.

#### \_\_\_\_\_ C/C++ -

## Fortran ———

If a list item does not have the **POINTER** attribute, then in all other implicit tasks belonging to the **parallel** region, the list item becomes defined as if by intrinsic assignment with the value of the corresponding list item in the implicit task associated with the thread that executed the structured block.

If the list item has the **POINTER** attribute, then, in all other implicit tasks belonging to the **parallel** region, the list item receives, as if by pointer assignment, the same association status of the corresponding list item in the implicit task associated with the thread that executed the structured block.

## - Fortran -----

For examples of the **copyprivate** clause, see Section A.38 on page 333.

1		<b>Note</b> – The <b>copyprivate</b> clause is an alternative to using a shared variable for the
2		value when providing such a shared variable would be difficult (for example, in a
3		recursion requiring a different variable at each level).
		<b>A</b>
4		Restrictions
5		The restrictions to the <b>copyprivate</b> clause are as follows:
6 7		• All list items that appear in the <b>copyprivate</b> clause must be either threadprivate or private in the enclosing context.
8 9		<ul> <li>A list item that appears in a copyprivate clause may not appear in a private or firstprivate clause on the single construct.</li> </ul>
10 11		• A variable of class type (or array thereof) that appears in a <b>copyprivate</b> clause requires an accessible unambiguous copy assignment operator for the class type.  C/C++
		Fortran —
12		• A common block that appears in a <b>copyprivate</b> clause must be threadprivate.
13 14		<ul> <li>Pointers with the INTENT (IN) attribute may not appear in the copyprivate clause.</li> </ul>
		Fortran
15	2.14.5	map Clause
16		Summary
17 18		Map a variable from the current task's data environment to the device data environment associated with the construct.

**Syntax** 

The syntax of the map clause is as follows:

map ( [map-type : ] list )

## **Description**

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The list items that appear in a map clause may include array sections.

For list items that appear in a map clause, corresponding new list items are created in the device data environment associated with the construct.

The original and corresponding list items may share storage such that writes to either item by one task followed by a read or write of the other item by another task without intervening synchronization can result in data races.

If a corresponding list item of the original list item is in the enclosing device data environment, the new device data environment uses the corresponding list item from the enclosing device data environment. No additional storage is allocated in the new device data environment and neither initialization nor assignment is performed, regardless of the *map-type* that is specified.

If a corresponding list item is not in the enclosing device data environment, a new list item with language-specific attributes is derived from the original list item and created in the new device data environment. This new list item becomes the corresponding list item to the original list item in the new device data environment. Initialization and assignment are performed if specified by the *map-type*.

#### — C/C++ —

If a new list item is created then a new list item of the same type, with automatic storage duration, is allocated for the construct. The storage and thus lifetime of this list item lasts until the block in which it is created exits. The size and alignment of the new list item are determined by the type of the variable. This allocation occurs if the region references the list item in any statement. \_\_\_\_\_ C/C++ \_\_\_\_\_

## ----- Fortran -----

If a new list item is created then a new list item of the same type, type parameter, and rank is allocated.

## - Fortran ----

The *map-type* determines how the new list item is initialized.

The alloc map-type declares that on entry to the region each new corresponding list item has an undefined initial value.

The to map-type declares that on entry to the region each new corresponding list item is initialized with the original list item's value.

The **from** map-type declares that on exit from the region the corresponding list item's value is assigned to each original list item.

1 The tofrom map-type declares that on entry to the region each new corresponding list 2 item is initialized with the original list item's value and that on exit from the region the corresponding list item's value is assigned to each original list item. 3 4 If a *map-type* is not specified, the *map-type* defaults to tofrom. Restrictions 5 • If a list item is an array section it must specify contiguous storage. 6 7 • If variables that share storage are mapped, the behavior is unspecified. • threadprivate variables cannot appear in a map clause. 8 - C/C++ -· Initialization and assignment are through bitwise copy. 9 \_\_\_\_\_ C/C++ -Fortran — 10 The map initialization and assignment are done as if by intrinsic assignment. Fortran —

#### 11 **2.15** declare reduction Directive

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where:

## **Summary** The following section describes the directive for declaring user-defined reductions. The 13 14 declare reduction directive declares a reduction-identifier that can be used in a reduction clause. The declare reduction directive is a declarative directive. 15 **Syntax** 16 17 #pragma omp declare reduction(reduction-identifier: typename-list: combiner ) [initializer-clause] new-line

1 2	• reduction-identifier is either a base language identifier or one of the following operators: +, -, *, &,  , ^, && and	
3	• typename-list is list of type names	
4	• combiner is an expression	
5	<ul> <li>initializer-clause is initializer ( omp_priv = initializer   function-name ( argument-list ))</li> </ul>	
7	C ++	
	<pre>#pragma omp declare reduction( reduction-identifier : typename-list :    combiner) [initializer-clause] new-line</pre>	
8	where:	
9 10	<ul> <li>reduction-identifier is either a base language identifier, an operator-function-id, or one of the following operators: +, -, *, &amp;,  , ^, &amp;&amp; and    </li> </ul>	
11	• typename-list is list of type names	
12	• combiner is an expression	
13 14	<ul> <li>initializer-clause is initializer ( omp_priv initializer   function-name ( argument-list ))</li> </ul>	
	C++	
15	Fortran —	
	!\$omp declare reduction( reduction-identifier: type-list: combiner) [initializer-clause]	
16	where:	
17 18 19	<ul> <li>reduction-identifier is either a base language identifier, or a user-defined operator, or one of the following operators: +, -, *, .and., .or., .eqv., .neqv., or one of the following intrinsic procedure names: max, min, iand, ior, ieor.</li> </ul>	
20	• type-list is a list of type specifiers	
21 22	<ul> <li>combiner is either an assignment statement or a subroutine name followed by an argument list</li> </ul>	
23 24	<ul> <li>initializer-clause is initializer ( omp_priv = initializer-expr   subroutine-name ( argument-list ) )</li> </ul>	
	Fortran	

## **Description**

Custom reductions can be defined using the **declare reduction** directive; the *reduction-identifier* and the type identify the **declare reduction** directive. The *reduction-identifier* can later be used in a **reduction** clause using variables of the type or types specified in the **declare reduction** directive. If the directive applies to several types then it is considered as if there were multiple **declare reduction** directives, one for each type.

The visibility and accessibility of this declaration are the same as those of a variable declared at the same point in the program. The enclosing context of the *combiner* and of the *initializer* will be that of the **declare reduction** directive. The *combiner* and the *initializer* must be correct in the base language as if they were the body of a function defined at the same point in the program.

The *combiner* specifies how partial results can be combined into a single value. The *combiner* can use the special variable identifiers **omp\_in** and **omp\_out** that are of the type of the variables being reduced with this *reduction-identifier*. Each of them will denote one of the values to be combined before executing the *combiner*. It is assumed that the special **omp\_out** identifier will refer to the storage that holds the resulting combined value after executing the *combiner*.

The number of times the *combiner* is executed, and the order of these executions, for any **reduction** clause is unspecified.

#### Fortran -

If the *combiner* is a subroutine name with an argument list, the *combiner* is evaluated by calling the subroutine with the specified argument list.

If the *combiner* is an assignment statement, the *combiner* is evaluated by executing the assignment statement.

#### · Fortran -

As the *initializer* value of a user-defined reduction is not known a *priori* the *initializer-clause* can be used to specify one. Then the contents of the *initializer-clause* will be used as the initializer for private copies of reduction list items where the **omp\_priv** identifier will refer to the storage to be initialized. The special identifier **omp\_orig** can also appear in the initializer-clause and it will refer to the storage of the original variable to be reduced.

The number of times that the *initializer* is evaluated, and the order of these evaluations, is unspecified.

	C/C++
1 2	If the <i>initializer</i> is a function name with an argument list, the <i>initializer</i> is evaluated by calling the function with the specified argument list. Otherwise, the <i>initializer</i> specifies
3	how omp_priv is declared and initialized.  C/C++
	C/C++
	C
4	If no <i>initializer-clause</i> is specified, the private variables will be initialized following the
5	rules for initialization of objects with static storage duration.
	C++
6 7	If no <i>initializer-clause</i> is specified, the private variables will be initialized following the rules for <i>default-initialization</i> .
	C++
	Fortran
8 9	If the <i>initializer</i> is a subroutine name with an argument list, the <i>initializer</i> is evaluated by calling the subroutine with the specified argument list.
10 11	If the <i>initializer</i> is an assignment statement, the <i>initializer</i> is evaluated by executing the assignment statement.
12	If no initializer-clause is specified, the private variables will be initialized as follows:
13	• For complex, real, or integer types, the value 0 will be used.
14	• For logical types, the value .false. will be used.
15 16	<ul> <li>For derived types for which default initialization is specified, default initialization will be used.</li> </ul>
17	• Otherwise, not specifying an initializer-clause results in unspecified behavior.
	Fortran
18	Restrictions
19	<ul> <li>Only the variables omp in and omp out are allowed in the combiner.</li> </ul>
20	<ul> <li>Only the variables omp priv and omp orig are allowed in the <i>initializer-clause</i>.</li> </ul>
21	<ul> <li>If the variable omp_orig is modified in the <i>initializer-clause</i>, the behavior is</li> </ul>
22	unspecified.
23	• If execution of the <i>combiner</i> or the <i>initializer</i> results in the execution of an OpenMP

construct or an OpenMP API call, then the behavior is unspecified.

• A user-defined reduction may not be re-declared in the current scope.

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1	• At most one <i>initializer-clause</i> can be specified.
	C/C++
2	• A type name in a <b>declare reduction</b> directive cannot be a function type, an
3 4	array type, a reference type, or a type qualified with const, volatile or restrict.
4	C/C++
	C
5	• If the initializer-clause is a function name with an argument list, then one of the
6	arguments must be the address of omp_priv.
	C
	C++
7 8	<ul> <li>If the <i>initializer-clause</i> is a function name with an argument list, then one of the arguments must be omp_priv or the address of omp_priv.</li> </ul>
	C++
	Fortran —
9	• If the <i>initializer-clause</i> is a subroutine name with an argument list, then one of the
10	arguments must be omp_priv.
11	• If the declare reduction directive appears in a module and the corresponding
12	reduction clause does not appear in the same module, the reduction-identifier must be
13	a user-defined operator, one of the allowed operators or one of the allowed intrinsic
14	procedures.
15	• If the <i>reduction-identifier</i> is a user-defined operator or an extended operator, the
16 17	interface for that operator must be defined in the same subprogram, or must be accessible by host or use association.
18	<ul> <li>If the declare reduction directive appears in a module, any user-defined</li> </ul>
19	operators used in the combiner must be defined in the same subprogram, or must be
20	accessible by host or use association. The user-defined operators must also be
21	accessible by host or use association in the subprogram in which the corresponding
22	reduction clause appears.
23	• Any subroutine or function used in the initializer clause or <i>combiner</i>
24 25	expression must be an intrinsic function, or must have an explicit interface in the
25	same subprogram or must be accessible by host or use association.
	Fortran
26	Cross References
27	• reduction clause Section 2.14.3.6 on page 142

## 2.16 Nesting of Regions

This section describes a set of restrictions on the nesting of regions. The restrictions on nesting are as follows:

- A worksharing region may not be closely nested inside a worksharing, explicit task,
   critical, ordered, atomic, or master region.
- A barrier region may not be closely nested inside a worksharing, explicit task, critical, ordered, atomic, or master region.
- A master region may not be closely nested inside a worksharing, atomic, or explicit task region.
- An ordered region may not be closely nested inside a critical, atomic, or explicit task region.
- An **ordered** region must be closely nested inside a loop region (or parallel loop region) with an **ordered** clause.
- A critical region may not be nested (closely or otherwise) inside a critical region with the same name. Note that this restriction is not sufficient to prevent deadlock.
- parallel, flush, critical, atomic, taskyield, and explicit task regions may not be closely nested inside an atomic region.
- OpenMP constructs may not be nested inside a **simd** region.

For examples illustrating these rules, see Section A.20 on page 281, Section A.39 on page 338, Section A.40 on page 341, and Section A.15 on page 253.

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# **Runtime Library Routines**

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3 4	This chapter describes the OpenMP API runtime library routines and is divided into the following sections:
5	• Runtime library definitions (Section 3.1 on page 160).
6 7	• Execution environment routines that can be used to control and to query the parallel execution environment (Section 3.2 on page 161).
8 9	<ul> <li>Lock routines that can be used to synchronize access to data (Section 3.3 on page 196).</li> </ul>
10	• Portable timer routines (Section 3.4 on page 203).
11	
12 13	Throughout this chapter, <i>true</i> and <i>false</i> are used as generic terms to simplify the description of the routines.
	C/C++
14	true means a nonzero integer value and false means an integer value of zero.
	C/C++
	Fortran
15	true means a logical value of .TRUE. and false means a logical value of .FALSE
	Fortran
	Fortran
16	Restrictions
17	The following restriction applies to all OpenMP runtime library routines:
18 19	<ul> <li>OpenMP runtime library routines may not be called from PURE or ELEMENTAL procedures.</li> </ul>
	Fortran

## 3.1 Runtime Library Definitions

For each base language, a compliant implementation must supply a set of definitions for the OpenMP API runtime library routines and the special data types of their parameters. The set of definitions must contain a declaration for each OpenMP API runtime library routine and a declaration for the *simple lock*, *nestable lock*, *schedule*, and *thread affinity policy* data types. In addition, each set of definitions may specify other implementation specific values.

### C/C++ -

The library routines are external functions with "C" linkage.

Prototypes for the C/C++ runtime library routines described in this chapter shall be provided in a header file named omp.h. This file defines the following:

- The prototypes of all the routines in the chapter.
- The type omp lock t.
- The type omp nest lock t.
- The type omp sched t.
- The type omp proc bind t.

See Section D.1 on page 388 for an example of this file.

## C/C++

#### - Fortran ---

The OpenMP Fortran API runtime library routines are external procedures. The return values of these routines are of default kind, unless otherwise specified.

Interface declarations for the OpenMP Fortran runtime library routines described in this chapter shall be provided in the form of a Fortran include file named omp\_lib.h or a Fortran 90 module named omp\_lib. It is implementation defined whether the include file or the module file (or both) is provided.

These files define the following:

- The interfaces of all of the routines in this chapter.
- The integer parameter omp lock kind.
- The integer parameter omp nest lock kind.
- · The integer parameter omp sched kind.
- The integer parameter omp proc bind kind.

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1	• The integer parameter openmp version with a value yyyymm where yyyy
2	and mm are the year and month designations of the version of the OpenMP Fortran
3	API that the implementation supports. This value matches that of the C preprocessor
1	macro <b>_OPENMP</b> , when a macro preprocessor is supported (see Section 2.2 on page
5	28).
6	See Section D.2 on page 390 and Section D.3 on page 392 for examples of these files.

It is implementation defined whether any of the OpenMP runtime library routines that take an argument are extended with a generic interface so arguments of different **KIND** type can be accommodated. See Appendix D.4 for an example of such an extension.

Fortran -

## 10 3.2 Execution Environment Routines

The routines described in this section affect and monitor threads, processors, and the parallel environment.

- the omp set num threads routine.
- the omp get num threads routine.
- the omp get max threads routine.
- the omp get thread num routine.
- the omp get num procs routine.
- the omp in parallel routine.
  - the omp set dynamic routine.
- the omp get dynamic routine.
- the omp get cancellation routine
- the omp set nested routine.

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- the omp get nested routine.
- the omp set schedule routine.
  - the omp get schedule routine.
- the omp get thread limit routine.
  - the omp set max active levels routine.
- the omp get max active levels routine.
- the omp get level routine.
- the omp get ancestor thread num routine.
- the omp get team size routine.

• the omp get active level routine. 1 • the omp in final routine. • the omp get proc bind routine. 3 • the omp set default device routine. • the omp get default device routine. 5 • the omp get num devices routine. • the omp get num teams routine. the omp get team num routine. 9 3.2.1 omp set num threads Summary 10 11 The omp set num threads routine affects the number of threads to be used for 12 subsequent parallel regions that do not specify a num threads clause, by setting the value of the first element of the nthreads-var ICV of the current task. 13 **Format** 14 C/C++ void omp set num threads(int num\_threads); C/C++ ----15 Fortran -

subroutine omp set num threads(num\_threads)

- 17 Constraints on Arguments
  - The value of the argument passed to this routine must evaluate to a positive integer, or else the behavior of this routine is implementation defined.

- Fortran -----

integer num\_threads

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1	Binding
2	The binding task set for an omp_set_num_threads region is the generating task.
3	Effect
4 5	The effect of this routine is to set the value of the first element of the <i>nthreads-var</i> ICV of the current task to the value specified in the argument.
6 7	See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a parallel region.
8 9	For an example of the omp_set_num_threads routine, see Section A.41 on page 348.
10	Cross References
11	• nthreads-var ICV, see Section 2.3 on page 30.
12	• OMP_NUM_THREADS environment variable, see Section 4.2 on page 211.
13	• omp_get_max_threads routine, see Section 3.2.3 on page 165.
14	• parallel construct, see Section 2.5 on page 37.
15	• num_threads clause, see Section 2.5 on page 37.
16 3.2.2	omp_get_num_threads
17	Summary
18	The omp_get_num_threads routine returns the number of threads in the current

19

team.

1	Format
	C/C++
	<pre>int omp_get_num_threads(void);</pre>
2	C/C++
2	Fortran —
	<pre>integer function omp_get_num_threads()</pre>
3	Fortran —
4	Binding
5 6	The binding region for an omp_get_num_threads region is the innermost enclosing parallel region.
7	Effect
8 9 10 11	The omp_get_num_threads routine returns the number of threads in the team executing the parallel region to which the routine region binds. If called from the sequential part of a program, this routine returns 1. For examples, see Section A.42 on page 349.
12 13	See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a parallel region.

### **Cross References**

- parallel construct, see Section 2.5 on page 37.
- omp\_set\_num\_threads routine, see Section 3.2.1 on page 162.
- OMP NUM THREADS environment variable, see Section 4.2 on page 211.

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## 1 3.2.3 omp get max threads

## Summary 2 3 The omp get max threads routine returns an upper bound on the number of 4 threads that could be used to form a new team if a parallel region without a 5 num threads clause were encountered after execution returns from this routine. **Format** 6 C/C++ int omp get max threads(void); C/C++ -7 Fortran integer function omp get max threads() - Fortran -8 **Binding** 9 10 The binding task set for an omp get max threads region is the generating task. Effect 11 12 The value returned by omp get max threads is the value of the first element of the nthreads-var ICV of the current task. This value is also an upper bound on the 13 14 number of threads that could be used to form a new team if a parallel region without a 15 num threads clause were encountered after execution returns from this routine. 16 See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a parallel region. 17

1	<b>Note</b> - The return value of the omp_get_max_threads routine can be used to
2	dynamically allocate sufficient storage for all threads in the team formed at the
3	subsequent active parallel region.
	^
4	Cross References
5	• nthreads-var ICV, see Section 2.3 on page 30.
6	• parallel construct, see Section 2.5 on page 37.
7	• num threads clause, see Section 2.5 on page 37.
8	• omp set num threads routine, see Section 3.2.1 on page 162.
9	• OMP_NUM_THREADS environment variable, see Section 4.2 on page 211.
<b>3.2.4</b>	<pre>omp_get_thread_num</pre>
11	Summary
12 13	The omp_get_thread_num routine returns the thread number, within the current team, of the calling thread.
14	Format
	C/C++
	<pre>int omp_get_thread_num(void);</pre>
	C/C++
15	Fortran —
	▼ Tottian -
	<pre>integer function omp_get_thread_num()</pre>
10	Fortran
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## 1 Binding

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The binding thread set for an omp\_get\_thread\_num region is the current team. The binding region for an omp\_get\_thread\_num region is the innermost enclosing parallel region.

#### Effect

The omp\_get\_thread\_num routine returns the thread number of the calling thread, within the team executing the parallel region to which the routine region binds. The thread number is an integer between 0 and one less than the value returned by omp\_get\_num\_threads, inclusive. The thread number of the master thread of the team is 0. The routine returns 0 if it is called from the sequential part of a program.

**Note** — The thread number may change during the execution of an untied task. The value returned by <code>omp\_get\_thread\_num</code> is not generally useful during the execution of such a task region.

### **Cross References**

• omp get num threads routine, see Section 3.2.2 on page 163.

## omp get num procs

## 17 Summary

The omp\_get\_num\_procs routine returns the number of processors available to the program.

## 1 Format

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C/C++

int omp get num procs(void);

C/C++

Fortran —

integer function omp get num procs()

Fortran —

## Binding

The binding thread set for an omp\_get\_num\_procs region is all threads. The effect of executing this routine is not related to any specific region corresponding to any construct or API routine.

#### Effect

The omp\_get\_num\_procs routine returns the number of processors that are available to the program at the time the routine is called. Note that this value may change between the time that it is determined by the omp\_get\_num\_procs routine and the time that it is read in the calling context due to system actions outside the control of the OpenMP implementation.

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## Summary

The omp\_in\_parallel routine returns *true* if the call to the routine is enclosed by an active parallel region; otherwise, it returns *false*.

**Format** 1 C/C++ int omp in parallel(void); C/C++ -2 Fortran logical function omp in parallel() Fortran -3 **Binding** 4 5 The binding thread set for an omp in parallel region is all threads. The effect of 6 executing this routine is not related to any specific parallel region but instead depends on the state of all enclosing parallel regions. 7 **Effect** 8 9 omp in parallel returns true if any enclosing parallel region is active. If the routine call is enclosed by only inactive parallel regions (including the implicit 10 11 parallel region), then it returns false. 12 **3.2.7** omp set dynamic **Summary** 13 14 The omp set dynamic routine enables or disables dynamic adjustment of the 15 number of threads available for the execution of subsequent parallel regions by 16 setting the value of the dyn-var ICV.

## **Format** 1 C/C++ void omp set dynamic(int dynamic\_threads); C/C++ -2 Fortran subroutine omp set dynamic (dynamic\_threads) logical dynamic threads - Fortran -3 **Binding** The binding task set for an **omp set dynamic** region is the generating task. Effect For implementations that support dynamic adjustment of the number of threads, if the argument to omp set dynamic evaluates to true, dynamic adjustment is enabled for the current task; otherwise, dynamic adjustment is disabled for the current task. For 10 implementations that do not support dynamic adjustment of the number of threads this

For an example of the omp set dynamic routine, see Section A.41 on page 348.

See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a **parallel** region.

#### **Cross References:**

- dyn-var ICV, see Section 2.3 on page 30.
- omp get num threads routine, see Section 3.2.2 on page 163.
- omp get dynamic routine, see Section 3.2.8 on page 171.

routine has no effect: the value of dyn-var remains false.

• **OMP\_DYNAMIC** environment variable, see Section 4.3 on page 212.

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# 1 3.2.8 omp\_get\_dynamic

2	Summary	
3 4	The omp_get_dynamic routine returns the value of the <i>dyn-var</i> ICV, which determines whether dynamic adjustment of the number of threads is enabled or disabled	
5	Format C/C++	
	<pre>int omp_get_dynamic(void);</pre>	
6	C/C++	
	Fortran —	
	logical function omp_get_dynamic()	
7	Fortran	
8	Binding	
9	The binding task set for an omp_get_dynamic region is the generating task.	
0	Effect	
1 2 3	This routine returns <i>true</i> if dynamic adjustment of the number of threads is enabled for the current task; it returns <i>false</i> , otherwise. If an implementation does not support dynamic adjustment of the number of threads, then this routine always returns <i>false</i> .	
4 5	See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a parallel region.	
6	Cross References	
7	• dyn-var ICV, see Section 2.3 on page 30.	
8	• omp_set_dynamic routine, see Section 3.2.7 on page 169.	

• OMP DYNAMIC environment variable, see Section 4.3 on page 212.

# 2 3.2.9 omp get cancellation

# **Summary** 3 The omp get cancellation routine returns the value of the cancel-var ICV, which 5 controls the behavior of the cancel construct and cancellation points. **Format** 6 — C/C++ — 7 int omp get cancellation(void); 8 C/C++ ----Fortran -9 logical function omp get cancellation() 10 Fortran **Binding** 11 The binding task set for an omp get cancellation region is the generating task. 12 **Effect** 13 14 This routine returns *true* if cancellation is activated for the current task set. It returns 15 false otherwise.

# **Cross References:** 1 2 • cancel-var ICV, see Section 2.3.1 on page 31. 3 • OMP CANCELLATION environment variable, see Section 4.11 on page 218. 4 3.2.10 omp set nested **Summary** 5 6 The omp set nested routine enables or disables nested parallelism, by setting the nest-var ICV. 7 **Format** 8 C/C++ void omp set nested(int nested); – C/C++ – 9 Fortran subroutine omp set nested (nested)

Fortran -

logical nested

1	Binding
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The binding task set for an **omp set nested** region is the generating task.

#### Effect

For implementations that support nested parallelism, if the argument to omp\_set\_nested evaluates to true, nested parallelism is enabled for the current task; otherwise, nested parallelism is disabled for the current task. For implementations that do not support nested parallelism, this routine has no effect: the value of nest-var remains false.

See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a parallel region.

#### Cross References

- *nest-var* ICV, see Section 2.3 on page 30.
- omp set max active levels routine, see Section 3.2.15 on page 180.
- omp get max active levels routine, see Section 3.2.16 on page 182.
- omp get nested routine, see Section 3.2.11 on page 174.
- OMP NESTED environment variable, see Section on page 213.

# 17 3.2.11 omp get nested

# Summary

The omp\_get\_nested routine returns the value of the *nest-var* ICV, which determines if nested parallelism is enabled or disabled.

**Format** 1 C/C++ int omp get nested(void); C/C++ -2 Fortran logical function omp get nested() - Fortran -3 **Binding** 4 The binding task set for an omp get nested region is the generating task. 5 **Effect** 6 This routine returns true if nested parallelism is enabled for the current task; it returns 7 false, otherwise. If an implementation does not support nested parallelism, this routine 8 9 always returns false. 10 See Section 2.5.1 on page 40 for the rules governing the number of threads used to execute a parallel region. 11 **Cross References** 12 13 • nest-var ICV, see Section 2.3 on page 30. • omp set nested routine, see Section 3.2.10 on page 173. 14 • OMP NESTED environment variable, see Section on page 213. 15

# 1 3.2.12 omp set schedule

## 2 Summary

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14 15 The omp\_set\_schedule routine affects the schedule that is applied when runtime is used as schedule kind, by setting the value of the *run-sched-var* ICV.

#### Format

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void omp\_set\_schedule(omp\_sched\_t kind, int modifier);

C/C++

C/C++ -

Fortran —

subroutine omp\_set\_schedule(kind, modifier)
integer (kind=omp\_sched\_kind) kind
integer modifier

Fortran

## **Constraints on Arguments**

The first argument passed to this routine can be one of the valid OpenMP schedule kinds (except for runtime) or any implementation specific schedule. The C/C++ header file (omp.h) and the Fortran include file (omp\_lib.h) and/or Fortran 90 module file (omp\_lib) define the valid constants. The valid constants must include the following, which can be extended with implementation specific values:

C/C++

1

```
typedef enum omp_sched_t {
   omp_sched_static = 1,
   omp_sched_dynamic = 2,
   omp_sched_guided = 3,
   omp_sched_auto = 4
} omp_sched_t;
```

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```
Fortran —
```

C/C++

```
integer(kind=omp_sched_kind), parameter :: omp_sched_static = 1
integer(kind=omp_sched_kind), parameter :: omp_sched_dynamic = 2
integer(kind=omp_sched_kind), parameter :: omp_sched_guided = 3
integer(kind=omp_sched_kind), parameter :: omp_sched_auto = 4
```

Fortran

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## **Binding**

The binding task set for an omp set schedule region is the generating task.

## Effect

The effect of this routine is to set the value of the *run-sched-var* ICV of the current task to the values specified in the two arguments. The schedule is set to the schedule type specified by the first argument **kind**. It can be any of the standard schedule types or any other implementation specific one. For the schedule types **static**, **dynamic**, and **guided** the *chunk\_size* is set to the value of the second argument, or to the default *chunk\_size* if the value of the second argument is less than 1; for the schedule type **auto** the second argument has no meaning; for implementation specific schedule types, the values and associated meanings of the second argument are implementation defined.

# **Cross References** 1 • run-sched-var ICV, see Section 2.3 on page 30. • omp get schedule routine, see Section 3.2.13 on page 178. 3 • **OMP SCHEDULE** environment variable, see Section 4.1 on page 210. 5 • Determining the schedule of a worksharing loop, see Section 2.7.1.1 on page 52. 3.2.13 omp get schedule **Summary** 7 The omp get schedule routine returns the schedule that is applied when the 9 runtime schedule is used. **Format** 10 11 C/C++ void omp get schedule(omp sched t \* kind, int \* modifier ); C/C++ \_\_\_\_ 12 Fortran subroutine omp\_get\_schedule(kind, modifier) integer (kind=omp sched kind) kind integer modifier Fortran — 13 **Binding** 14

The binding task set for an omp\_get\_schedule region is the generating task.

# **Effect** 1 This routine returns the run-sched-var ICV in the task to which the routine binds. The 2 3 first argument kind returns the schedule to be used. It can be any of the standard 4 schedule types as defined in Section 3.2.12 on page 176, or any implementation specific schedule type. The second argument is interpreted as in the omp set schedule call. 5 6 defined in Section 3.2.12 on page 176. Cross References 7 8 • run-sched-var ICV, see Section 2.3 on page 30. • omp set schedule routine, see Section 3.2.12 on page 176. 9 • OMP SCHEDULE environment variable, see Section 4.1 on page 210. 10 11 • Determining the schedule of a worksharing loop, see Section 2.7.1.1 on page 52. 12 **3.2.14** omp get thread limit Summary 13 14 The omp get thread limit routine returns the maximum number of OpenMP threads available to the program. 15 **Format** 16 17 C/C++ int omp get thread limit(void); C/C++18 Fortran integer function omp get thread limit() Fortran -

# **Binding** 1 2 The binding thread set for an **omp get thread limit** region is all threads. The effect of executing this routine is not related to any specific region corresponding to any 4 construct or API routine. Effect 5 The omp get thread limit routine returns the maximum number of OpenMP 7 threads available to the program as stored in the ICV thread-limit-var. Cross References 8 • thread-limit-var ICV, see Section 2.3 on page 30. 9 • OMP THREAD LIMIT environment variable, see Section 4.10 on page 218. 10 11 **3.2.15** omp set max active levels **Summary** 12 13 The omp set max active levels routine limits the number of nested active 14 parallel regions, by setting the max-active-levels-var ICV. **Format** 15 16 C/C++ void omp set max active levels (int max levels); C/C++ -

	•	1

#### Fortran

subroutine omp\_set\_max\_active\_levels (max\_levels)
integer max\_levels

Fortran

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## **Constraints on Arguments**

The value of the argument passed to this routine must evaluate to a non-negative integer, otherwise the behavior of this routine is implementation defined.

## 6 Binding

When called from the sequential part of the program, the binding thread set for an omp\_set\_max\_active\_levels region is the encountering thread. When called from within any explicit parallel region, the binding thread set (and binding region, if required) for the omp set max active levels region is implementation defined.

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#### Effect

The effect of this routine is to set the value of the *max-active-levels-var* ICV to the value specified in the argument.

If the number of parallel levels requested exceeds the number of levels of parallelism supported by the implementation, the value of the *max-active-levels-var* ICV will be set to the number of parallel levels supported by the implementation.

This routine has the described effect only when called from the sequential part of the program. When called from within an explicit **parallel** region, the effect of this routine is implementation defined.

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#### Cross References

- max-active-levels-var ICV, see Section 2.3 on page 30.
- omp get max active levels routine, see Section 3.2.16 on page 182.
- OMP MAX ACTIVE LEVELS environment variable, see Section 4.9 on page 217.

# 1 3.2.16 omp get max active levels

# 2 Summary

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The omp\_get\_max\_active\_levels routine returns the value of the *max-active-levels-var* ICV, which determines the maximum number of nested active parallel regions.

#### **Format**

c/C++
int omp\_get\_max\_active\_levels(void);

c/C++

Fortran

integer function omp\_get\_max\_active\_levels()

# **Binding**

When called from the sequential part of the program, the binding thread set for an omp\_get\_max\_active\_levels region is the encountering thread. When called from within any explicit parallel region, the binding thread set (and binding region, if required) for the omp\_get\_max\_active\_levels region is implementation defined.

#### Effect

The omp\_get\_max\_active\_levels routine returns the value of the *max-active-levels-var* ICV, which determines the maximum number of nested active parallel regions.

# **Cross References** 1 • max-active-levels-var ICV, see Section 2.3 on page 30. 2 3 • omp set max active levels routine, see Section 3.2.15 on page 180. • OMP MAX ACTIVE LEVELS environment variable, see Section 4.9 on page 217. 4 5 **3.2.17** omp get level Summary 6 The omp get level routine returns the number of nested parallel regions 7 8 enclosing the task that contains the call. **Format** 9 10 – C/C++ int omp get level(void); - C/C++ -11 - Fortran integer function omp get level() Fortran -12 **Binding** 13 14 The binding task set for an **omp get level** region is the generating task. The 15 binding region for an omp get level region is the innermost enclosing parallel region. 16

# Effect 1 2 The omp get level routine returns the number of nested parallel regions 3 (whether active or inactive) enclosing the task that contains the call, not including the 4 implicit parallel region. The routine always returns a non-negative integer, and returns 0 if it is called from the sequential part of the program. **Cross References** 6 • omp get active level routine, see Section 3.2.20 on page 187. • OMP MAX ACTIVE LEVELS environment variable, see Section 4.9 on page 217. 9 3.2.18 omp get ancestor thread num **Summary** 10 11 The omp get ancestor thread num routine returns, for a given nested level of 12 the current thread, the thread number of the ancestor or the current thread. **Format** 13 14 C/C++ int omp get ancestor thread num(int level); C/C++ \_\_\_\_\_ 15 Fortran -

integer function omp get ancestor thread num(level)

- Fortran —

integer level

Binding

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The binding thread set for an omp\_get\_ancestor\_thread\_num region is the encountering thread. The binding region for an omp\_get\_ancestor\_thread\_num region is the innermost enclosing parallel region.

#### Effect

The omp\_get\_ancestor\_thread\_num routine returns the thread number of the ancestor at a given nest level of the current thread or the thread number of the current thread. If the requested nest level is outside the range of 0 and the nest level of the current thread, as returned by the omp\_get level routine, the routine returns -1.

**Note** - When the omp\_get\_ancestor\_thread\_num routine is called with a value of level=0, the routine always returns 0. If level=omp\_get\_level(), the routine has the same effect as the omp\_get\_thread\_num routine.

#### Cross References

- omp get level routine, see Section 3.2.17 on page 183.
- omp get thread num routine, see Section 3.2.4 on page 166.
  - omp get team size routine, see Section 3.2.19 on page 185.

# 17 3.2.19 omp get team size

## 18 Summary

The omp\_get\_team\_size routine returns, for a given nested level of the current thread, the size of the thread team to which the ancestor or the current thread belongs.

#### Format

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C/C++

int omp\_get\_team\_size(int level);

C/C++ -

Fortran —

integer function omp\_get\_team\_size(level)
integer level

---- Fortran -----

# **Binding**

The binding thread set for an omp\_get\_team\_size region is the encountering thread. The binding region for an omp\_get\_team\_size region is the innermost enclosing parallel region.

#### **Effect**

The <code>omp\_get\_team\_size</code> routine returns the size of the thread team to which the ancestor or the current thread belongs. If the requested nested level is outside the range of 0 and the nested level of the current thread, as returned by the <code>omp\_get\_level</code> routine, the routine returns -1. Inactive parallel regions are regarded like active parallel regions executed with one thread.

**Note** — When the <code>omp\_get\_team\_size</code> routine is called with a value of <code>level=0</code>, the routine always returns 1. If <code>level=omp\_get\_level()</code>, the routine has the same effect as the <code>omp\_get\_num\_threads</code> routine.

# **Cross References** 1 2 • omp get num threads routine, see Section 3.2.2 on page 163. • omp get level routine, see Section 3.2.17 on page 183. 3 4 • omp get ancestor thread num routine, see Section 3.2.18 on page 184. 3.2.20 omp get active level Summary 6 7 The omp get active level routine returns the number of nested, active 8 parallel regions enclosing the task that contains the call. **Format** 9 10 C/C++ --int omp get active level(void); C/C++ -11 Fortran integer function omp get active level() - Fortran -12 **Binding** 13 The binding task set for the an omp get active level region is the generating 14 task. The binding region for an omp get active level region is the innermost 15 enclosing parallel region. 16

# **Effect** 1 2 The omp get active level routine returns the number of nested, active parallel regions enclosing the task that contains the call. The routine always returns a non-4 negative integer, and returns 0 if it is called from the sequential part of the program. **Cross References** 5 • omp get level routine, see Section 3.2.17 on page 183. 3.2.21 omp in final **Summary** The omp in final routine returns true if the routine is executed in a final task 10 region; otherwise, it returns false. **Format** 11 12 C/C++ int omp in final(void); \_\_\_\_\_ C/C++ \_\_\_\_\_ 13 Fortran logical function omp in final() Fortran -14

# Binding

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The binding task set for an **omp** in **final** region is the generating task.

**Effect** 1 2 omp in final returns true if the enclosing task region is final. Otherwise, it returns 3 false. 3.2.22 omp get proc bind Summary 5 6 The omp get proc bind routine returns the thread affinity policy to be used for the 7 next most closely nested parallel region. **Format** 8 9 C/C++ omp proc bind t omp get proc bind(void); C/C++ 10 Fortran -11 integer (kind=omp proc bind kind) function omp get proc bind() Fortran -12 **Constraints on Arguments** 13 The value returned by this routine must be one of the valid affinity policy kinds. The C/ 14 C++ header file (omp.h) and the Fortran include file (omp lib.h) and/or Fortran 90 15 module file (omp lib) define the valid constants. The valid constants must include the 16 17 following:

```
1
                                                 C/C++
 2
                 typedef enum omp proc bind t {
 3
                   omp proc bind false = 0,
 4
                   omp proc bind true = 1,
5
                   omp proc bind master = 2,
6
                   omp proc bind close = 3,
 7
                   omp proc bind spread = 4
 8
                 } omp proc bind t;
                                                 C/C++ -
 9
                                                Fortran -
10
                 integer (kind=omp_proc_bind_kind), &
11
                                  parameter :: omp proc bind false = 0
                 integer (kind=omp proc bind kind), &
13
                                  parameter :: omp proc bind true = 1
14
                 integer (kind=omp_proc_bind_kind), &
15
                                  parameter :: omp proc bind master = 2
16
                 integer (kind=omp proc bind kind), &
17
                                  parameter :: omp proc bind close = 3
18
                 integer (kind=omp_proc_bind_kind), &
19
                                  parameter :: omp_proc_bind_spread = 4
                                             — Fortran ————
```

## **Binding**

The binding task set for an **omp get proc bind** region is the generating task.

#### Effect

The effect of this routine is to return the value of the first element of the *bind-var* ICV of the current task. See Section 2.5.2 on page 42 for the rules governing the thread affinity policy.

#### **Cross References**

- bind-var ICV, see Section 2.3 on page 30.
- OMP PROC BIND environment variable, see Section 4.4 on page 213.

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page 42. 3.2.23 omp set default device **Summary** 4 The omp\_set\_default\_device routine assigns the value of the default-device-var 5 6 ICV, which determines the default device number. **Format** 7 8 C/C++ void omp set default device(int device\_num ); C/C++ -9 Fortran subroutine omp set default device( device\_num ) integer device\_num Fortran – **Binding** 10 The binding task set for an omp set default device region is the generating 11 12 task. Effect 13 14 The effect of this routine is to set the value of the default-device-var ICV of the current 15 task to the value specified in the argument. **Cross References:** 16 • default-device-var, see Section 2.3 on page 30. 17

• Determining the thread affinity policy of a parallel construct, see Section 2.5.2 on

• omp get default device, see Section 3.2.24 on page 192. 1 2 • OMP DEFAULT DEVICE environment variable, see Section 4.13 on page 220 3.2.24 omp get default device **Summary** 4 The omp get default device routine returns the value of the default-device-var ICV, which determines the default device number. 6 **Format** 7 C/C++ int omp\_get\_default\_device(void); \_\_\_\_\_ C/C++ \_\_\_\_ 9 - Fortran integer function omp get default device() Fortran – 10 **Binding** 11 12 The binding task set for an *omp\_get\_default\_device* region is the generating task. **Effect** 13 14 The omp get default device routine returns the value of the default-device-var

ICV of the current task.

# **Cross References** 1 • default-device-var, see Section 2.3 on page 30. 2 • omp set default device, see Section 3.2.23 on page 191. 3 • OMP DEFAULT DEVICE environment variable, see Section 4.13 on page 220. 4 3.2.25 omp get num devices **Summary** 6 The omp get num devices routine returns the number of target devices. 7 **Format** 8 C/C++ -9 int omp get num devices(void); C/C++ 10 - Fortran integer function omp get num devices() - Fortran -----11 **Binding** 12 The binding task set for an omp get num devices region is the generating task. 13 Effect 14 The omp get num devices routine returns the number of available target devices. 15

2 None. **3.2.26** omp\_get num teams **Summary** The omp get num teams routine returns the number of teams in the current teams region. **Format** 7 C/C++ -8 int omp get num teams(void); - C/C++ ----9 Fortran integer function omp\_get\_num\_teams() — Fortran — 10 **Binding** 11 The binding task set for an omp get num teams region is the generating task. 12 **Effect** 13 14 The effect of this routine is to return the number of teams in the current team region. 15 The routine returns 1 if it is called from outside of a team region.

**Cross References:** 

# **Cross References:** 1 2 • **teams** construct, see Section 2.9.5 on page 77. **3 3.2.27** omp get team num Summary 4 The omp get team num routine returns the team number of the calling thread. 5 **Format** 6 C/C++ — 7 int omp get team num(void); - C/C++ ----8 Fortran integer function omp get team num() Fortran — 9 **Binding** 10 The binding task set for an omp get team num region is the generating task. 11 **Effect** 12 13 The omp get team num routine returns the team number of the calling thread. The team number is an integer between 0 and one less than the value returned by 14 omp get num teams, inclusive. The routine returns 0 if it is called outside of a 15 teams region. 16

#### **Cross References:**

- **teams** construct, see Section 2.9.5 on page 77.
- omp get num teams routine, see Section 3.2.26 on page 194.

# 3.3 Lock Routines

The OpenMP runtime library includes a set of general-purpose lock routines that can be used for synchronization. These general-purpose lock routines operate on OpenMP locks that are represented by OpenMP lock variables. OpenMP lock variables must be accessed only through the routines described in this section; programs that otherwise access OpenMP lock variables are non-conforming.

An OpenMP lock can be in one of the following states: *uninitialized*, *unlocked*, or *locked*. If a lock is in the unlocked state, a task can *set* the lock, which changes its state to *locked*. The task that sets the lock is then said to *own* the lock. A task that owns a lock can *unset* that lock, returning it to the *unlocked* state. A program in which a task unsets a lock that is owned by another task is non-conforming.

Two types of locks are supported: *simple locks* and *nestable locks*. A nestable lock can be set multiple times by the same task before being unset; a *simple lock* cannot be set if it is already owned by the task trying to set it. *Simple lock* variables are associated with *simple locks* and can only be passed to *simple lock* routines. *Nestable lock* variables are associated with *nestable locks* and can only be passed to *nestable lock* routines.

Constraints on the state and ownership of the lock accessed by each of the lock routines are described with the routine. If these constraints are not met, the behavior of the routine is unspecified.

The OpenMP lock routines access a lock variable in such a way that they always read and update the most current value of the lock variable. It is not necessary for an OpenMP program to include explicit **flush** directives to ensure that the lock variable's value is consistent among different tasks.

See Section A.45 on page 354 and Section A.46 on page 357, for examples of using the simple and the nestable lock routines, respectively.

## Binding

The binding thread set for all lock routine regions is all threads. As a consequence, for each OpenMP lock, the lock routine effects relate to all tasks that call the routines, without regard to which teams the threads executing the tasks belong.

#### Simple Lock Routines 1 The type omp lock t is a data type capable of representing a simple lock. For the 2 following routines, a simple lock variable must be of omp lock t type. All simple 3 lock routines require an argument that is a pointer to a variable of type omp lock t. 4 \_\_\_\_\_ C/C++ -------Fortran 5 For the following routines, a simple lock variable must be an integer variable of kind=omp lock kind. 6 ——— Fortran — The simple lock routines are as follows: 7 • The omp init lock routine initializes a simple lock. 8 • The omp destroy lock routine uninitializes a simple lock. 9 • The omp set lock routine waits until a simple lock is available, and then sets it. 10 • The omp unset lock routine unsets a simple lock. 11 12 • The omp test lock routine tests a simple lock, and sets it if it is available. 13 14 Nestable Lock Routines: — C/C++ ——— The type omp nest lock t is a data type capable of representing a nestable lock. 15 For the following routines, a nested lock variable must be of omp nest lock t type. 16 All nestable lock routines require an argument that is a pointer to a variable of type 17 18 omp nest lock t. \_\_\_\_\_ C/C++ \_\_\_\_\_ - Fortran -----For the following routines, a nested lock variable must be an integer variable of 19 20 kind=omp nest lock kind. Fortran — 21 The nestable lock routines are as follows: 22 • The omp init nest lock routine initializes a nestable lock. 23 • The omp destroy nest lock routine uninitializes a nestable lock.

1 • The omp set nest lock routine waits until a nestable lock is available, and then 2 sets it. 3 • The omp unset nest lock routine unsets a nestable lock. • The omp test nest lock routine tests a nestable lock, and sets it if it is available. Restrictions 7 OpenMP lock routines have the following restrictions: • The use of the same OpenMP lock in different contention groups results in 9 unspecified behavior. 10 3.3.1 omp init lock and omp init nest lock Summary 11 12 These routines provide the only means of initializing an OpenMP lock. Format 13 C/C++ void omp init lock(omp lock t \*lock); void omp init nest lock(omp nest lock t \*lock); C/C++ -14 Fortran subroutine omp init lock(svar) integer (kind=omp lock kind) svar subroutine omp init nest lock(nvar) integer (kind=omp nest lock kind) nvar

Fortran –

## 1 Constraints on Arguments

A program that accesses a lock that is not in the uninitialized state through either routine is non-conforming.

#### 4 Effect

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The effect of these routines is to initialize the lock to the unlocked state; that is, no task owns the lock. In addition, the nesting count for a nestable lock is set to zero.

For an example of the **omp init lock** routine, see Section A.43 on page 352.

# 8 3.3.2 omp\_destroy\_lock and 9 omp\_destroy\_nest\_lock

## 10 Summary

These routines ensure that the OpenMP lock is uninitialized.

#### Format

void omp\_destroy\_lock(omp\_lock\_t \*lock);

C/C++

void omp\_destroy\_lock(omp\_lock\_t \*lock);
void omp\_destroy\_nest\_lock(omp\_nest\_lock\_t \*lock);

C/C++ Fortran

```
subroutine omp_destroy_lock(svar)
integer (kind=omp_lock_kind) svar

subroutine omp_destroy_nest_lock(nvar)
integer (kind=omp_nest_lock kind) nvar
```

Fortran —

## 1 Constraints on Arguments

A program that accesses a lock that is not in the unlocked state through either routine is non-conforming.

#### Effect

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The effect of these routines is to change the state of the lock to uninitialized.

# 6 3.3.3 omp set lock and omp set nest lock

# Summary

These routines provide a means of setting an OpenMP lock. The calling task region is suspended until the lock is set.

#### Format

```
void omp_set_lock(omp_lock_t *lock);
void omp set nest lock(omp nest lock t *lock);
```

— C/C++ —

C/C++

Fortran —

```
subroutine omp_set_lock(svar)
integer (kind=omp_lock_kind) svar

subroutine omp_set_nest_lock(nvar)
integer (kind=omp_nest_lock_kind) nvar
```

12 Fortran

## Constraints on Arguments

A program that accesses a lock that is in the uninitialized state through either routine is non-conforming. A simple lock accessed by **omp\_set\_lock** that is in the locked state must not be owned by the task that contains the call or deadlock will result.

### 5 Effect

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Each of these routines causes suspension of the task executing the routine until the specified lock is available and then sets the lock.

A simple lock is available if it is unlocked. Ownership of the lock is granted to the task executing the routine.

A nestable lock is available if it is unlocked or if it is already owned by the task executing the routine. The task executing the routine is granted, or retains, ownership of the lock, and the nesting count for the lock is incremented.

# 13 3.3.4 omp unset lock and omp unset nest lock

### Summary

These routines provide the means of unsetting an OpenMP lock.

#### 16 Format

C/C++

```
void omp_unset_lock(omp_lock_t *lock);
void omp_unset_nest_lock(omp_nest_lock_t *lock);
```

C/C++
Fortran

```
subroutine omp_unset_lock(svar)
integer (kind=omp_lock_kind) svar

subroutine omp_unset_nest_lock(nvar)
integer (kind=omp_nest_lock_kind) nvar
```

1		Fortran —
2		Constraints on Arguments
3 4		A program that accesses a lock that is not in the locked state or that is not owned by the task that contains the call through either routine is non-conforming.
5		Effect
6		For a simple lock, the omp_unset_lock routine causes the lock to become unlocked.
7 8		For a nestable lock, the <code>omp_unset_nest_lock</code> routine decrements the nesting count, and causes the lock to become unlocked if the resulting nesting count is zero.
9 10 11		For either routine, if the lock becomes unlocked, and if one or more task regions were suspended because the lock was unavailable, the effect is that one task is chosen and given ownership of the lock.
12	3.3.5	<pre>omp_test_lock and omp_test_nest_lock</pre>
13		Summary

These routines attempt to set an OpenMP lock but do not suspend execution of the task

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executing the routine.

# **Format** 1 C/C++ int omp test lock(omp lock t \*lock); int omp test nest lock(omp nest lock t \*lock); 2 Fortran logical function omp test lock(svar) integer (kind=omp lock kind) svar integer function omp test\_nest\_lock(nvar) integer (kind=omp nest lock kind) nvar Fortran -3 **Constraints on Arguments** 4 5 A program that accesses a lock that is in the uninitialized state through either routine is non-conforming. The behavior is unspecified if a simple lock accessed by 6 7 omp test lock is in the locked state and is owned by the task that contains the call. Effect 8 9 These routines attempt to set a lock in the same manner as omp set lock and 10 omp set nest lock, except that they do not suspend execution of the task 11 executing the routine. 12 For a simple lock, the omp test lock routine returns true if the lock is successfully

# 16 3.4 Timing Routines

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set; otherwise, it returns false.

The routines described in this section support a portable wall clock timer.

if the lock is successfully set; otherwise, it returns zero.

For a nestable lock, the omp test nest lock routine returns the new nesting count

- the omp get wtime routine.
  - the omp get wtick routine.

# 3.4.1 omp get wtime

## u Summary

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16 17 The omp get wtime routine returns elapsed wall clock time in seconds.

#### Format

C/C++

double omp\_get\_wtime(void);

C/C++

Fortran

double precision function omp\_get\_wtime()

Fortran

## Binding

The binding thread set for an **omp\_get\_wtime** region is the encountering thread. The routine's return value is not guaranteed to be consistent across any set of threads.

#### Effect

The omp\_get\_wtime routine returns a value equal to the elapsed wall clock time in seconds since some "time in the past". The actual "time in the past" is arbitrary, but it is guaranteed not to change during the execution of the application program. The time returned is a "per-thread time", so it is not required to be globally consistent across all the threads participating in an application.

**Note** – It is anticipated that the routine will be used to measure elapsed times as shown 1 2 in the following example: C/C++ double start; double end; start = omp get wtime(); ... work to be timed ... end = omp get wtime(); printf("Work took %f seconds\n", end - start); C/C++ -3 Fortran -4 DOUBLE PRECISION START, END START = omp get wtime() ... work to be timed ... END = omp get wtime() PRINT \*, "Work took", END - START, "seconds" Fortran -5 6 **3.4.2** omp\_get\_wtick **Summary** 7 The omp get wtick routine returns the precision of the timer used by 8 9 omp\_get\_wtime.

C/C++

double omp\_get\_wtick(void);

C/C++

Fortran

double precision function omp\_get\_wtick()

Fortran

## Binding

The binding thread set for an **omp\_get\_wtick** region is the encountering thread. The routine's return value is not guaranteed to be consistent across any set of threads.

#### Effect

The omp\_get\_wtick routine returns a value equal to the number of seconds between successive clock ticks of the timer used by omp\_get\_wtime.

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# **Environment Variables**

3 4 5 6 7 8 9	This chapter describes the OpenMP environment variables that specify the settings of the ICVs that affect the execution of OpenMP programs (see Section 2.3 on page 30). The names of the environment variables must be upper case. The values assigned to the environment variables are case insensitive and may have leading and trailing white space. Modifications to the environment variables after the program has started, even if modified by the program itself, are ignored by the OpenMP implementation. However, the settings of some of the ICVs can be modified during the execution of the OpenMP program by the use of the appropriate directive clauses or OpenMP API routines.
11	The environment variables are as follows:
12 13	• <b>OMP_SCHEDULE</b> sets the <i>run-sched-var</i> ICV that specifies the runtime schedule type and chunk size. It can be set to any of the valid OpenMP schedule types.
14 15	• OMP_NUM_THREADS sets the <i>nthreads-var</i> ICV that specifies the number of threads to use for parallel regions.
16 17	<ul> <li>OMP_DYNAMIC sets the dyn-var ICV that specifies the dynamic adjustment of threads to use for parallel regions.</li> </ul>
18 19 20	<ul> <li>OMP_PROC_BIND sets the bind-var ICV that controls the OpenMP thread affinity policy. If the OMP_PROC_BIND variable is not defined, the initial value of this ICV is implementation defined.</li> </ul>
21	• OMP_NESTED sets the <i>nest-var</i> ICV that enables or disables nested parallelism.
22 23	• <b>OMP_STACKSIZE</b> sets the <i>stacksize-var</i> ICV that specifies the size of the stack for threads created by the OpenMP implementation.
24 25	• OMP_WAIT_POLICY sets the wait-policy-var ICV that controls the desired behavior of waiting threads.
26 27	• OMP_MAX_ACTIVE_LEVELS sets the <i>max-active-levels-var</i> ICV that controls the maximum number of nested active parallel regions.
28 29	• <b>OMP_THREAD_LIMIT</b> sets the <i>thread-limit-var</i> ICV that controls the maximum number of threads participating in the OpenMP program.

• OMP PLACES sets the place-partition-var ICV that defines the OpenMP places that 1 2 are available to the execution environment. 3 • OMP DISPLAY ENV instructs the runtime to display the OpenMP version number and the initial values of the ICVs, once, during initialization of the runtime. 4 5 • OMP DEFAULT DEVICE sets the default-device-var ICV that controls the default device number. 6 The examples in this chapter only demonstrate how these variables might be set in Unix 8 C shell (csh) environments. In Korn shell (ksh) and DOS environments the actions are similar, as follows: 10 csh: setenv OMP SCHEDULE "dynamic" 11 ksh: export OMP SCHEDULE="dynamic" 12 • DOS: set OMP SCHEDULE=dynamic 4.1 OMP SCHEDULE 14 The **OMP SCHEDULE** environment variable controls the schedule type and chunk size 15 of all loop directives that have the schedule type runtime, by setting the value of the run-sched-var ICV. 16 The value of this environment variable takes the form: 17 18 *type*[,*chunk*] 19 where 20 type is one of static, dynamic, guided, or auto 21 • chunk is an optional positive integer that specifies the chunk size 22 If chunk is present, there may be white space on either side of the ",". See Section 2.7.1 on page 47 for a detailed description of the schedule types. 23

The behavior of the program is implementation defined if the value of **OMP SCHEDULE** 

does not conform to the above format.

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Implementation specific schedules cannot be specified in **OMP\_SCHEDULE**. They can only be specified by calling **omp\_set\_schedule**, described in Section 3.2.12 on page 176.

Example:

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24 25 setenv OMP\_SCHEDULE "guided,4"
setenv OMP\_SCHEDULE "dynamic"

#### Cross References

- run-sched-var ICV, see Section 2.3 on page 30.
- Loop construct, see Section 2.7.1 on page 47.
  - Parallel loop construct, see Section 2.10.1 on page 82.
  - omp set schedule routine, see Section 3.2.12 on page 176.
    - omp get schedule routine, see Section 3.2.13 on page 178.

## 4.2 OMP NUM THREADS

The OMP\_NUM\_THREADS environment variable sets the number of threads to use for parallel regions by setting the initial value of the *nthreads-var* ICV. See Section 2.3 on page 30 for a comprehensive set of rules about the interaction between the OMP\_NUM\_THREADS environment variable, the num\_threads clause, the omp\_set\_num\_threads library routine and dynamic adjustment of threads, and Section 2.5.1 on page 40 for a complete algorithm that describes how the number of threads for a parallel region is determined.

The value of this environment variable must be a list of positive integer values. The values of the list set the number of threads to use for **parallel** regions at the corresponding nested level.

The behavior of the program is implementation defined if any value of the list specified in the **OMP\_NUM\_THREADS** environment variable leads to a number of threads which is greater than an implementation can support, or if any value is not a positive integer.

Example:

setenv OMP NUM THREADS 4,3,2

Cross	References:
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- nthreads-var ICV, see Section 2.3 on page 30.
- num threads clause, Section 2.5 on page 37.
- omp set num threads routine, see Section 3.2.1 on page 162.
- omp get num threads routine, see Section 3.2.2 on page 163.
- omp get max threads routine, see Section 3.2.3 on page 165.
- omp get team size routine, see Section 3.2.19 on page 185.

## 4.3 OMP DYNAMIC

The OMP\_DYNAMIC environment variable controls dynamic adjustment of the number of threads to use for executing parallel regions by setting the initial value of the dyn-var ICV. The value of this environment variable must be true or false. If the environment variable is set to true, the OpenMP implementation may adjust the number of threads to use for executing parallel regions in order to optimize the use of system resources. If the environment variable is set to false, the dynamic adjustment of the number of threads is disabled. The behavior of the program is implementation defined if the value of OMP\_DYNAMIC is neither true nor false.

Example:

setenv OMP DYNAMIC true

#### Cross References:

- dyn-var ICV, see Section 2.3 on page 30.
- omp set dynamic routine, see Section 3.2.7 on page 169.
- omp get dynamic routine, see Section 3.2.8 on page 171.

## 4.4 OMP PROC BIND

The OMP\_PROC\_BIND environment variable sets the initial value of bind-var ICV. The value of this environment variable is either true, false, or a comma separated list of master, close, or spread. The values of the list set the thread affinity policy to be used for parallel regions at the corresponding nested level.

If the environment variable is set to **false**, the execution environment may move OpenMP threads between OpenMP places, thread affinity is disabled, and **proc\_bind** clauses on **parallel** constructs are ignored.

Otherwise, the execution environment should not move OpenMP threads between OpenMP places, thread affinity is enabled, and the initial thread is bound to the first place in the OpenMP place list.

The behavior of the program is implementation defined if any of the values in the OMP\_PROC\_BIND environment variable is not true, false, or a comma separated list of master, close, or spread. The behavior is also implementation defined if the initial thread cannot be bound to the first place in the OpenMP place list.

Example:

```
setenv OMP_PROC_BIND false
setenv OMP_PROC_BIND "spread, spread, close"
```

#### Cross References:

- bind-var ICV, see Section 2.3 on page 30.
  - proc bind clause, see Section 2.5.2 on page 42
- omp get proc bind routine, see Section 3.2.22 on page 189

## 4.5 OMP PLACES

A list of places can be specified in the **OMP\_PLACES** environment variable. The *place-partition-var* ICV obtains its initial value from the **OMP\_PLACES** value, and makes the list available to the execution environment. The value of **OMP\_PLACES** can be one of two types of values: either an abstract name describing a set of places or an explicit list of places described by nonnegative numbers.

The **OMP\_PLACES** environment variable can be defined using an explicit ordered list of places. A place is defined by an unordered set of nonnegative numbers enclosed by braces and separated by commas. The meaning of the numbers and how the numbering is done are implementation defined. Generally, the numbers represent the smallest unit of execution exposed by the execution environment, typically a hardware thread.

Intervals can be specified using the *<lower-bound>*: *<length>*: *<stride>* notation to represent the following list of numbers: "*<lower-bound>*, *<lower-bound>* + *<stride>*, ..., *<lower-bound>* + (*<length>*-1)\**<stride>*." When *<stride>* is omitted, a unit stride is assumed. Intervals can specify numbers within a place as well as sequences of places.

An exclusion operator "!" can also be used to exclude the number or place immediately following the operator.

Alternatively, the abstract names listed in TABLE 4-1 should be understood by the execution and runtime environment. The precise definitions of the abstract names are implementation defined. An implementation may also add abstract names as appropriate for the target platform.

The abstract name may be appended by a positive number in parentheses to denote the length of the place list to be created, that is <code>abstract\_name(num-places)</code>. When requesting fewer places than available on the system, the determination of which resources of type <code>abstract\_name</code> are to be included in the place list is implementation defined. When requesting more resources than available, the length of the place list is implementation defined.

TABLE 4-1 List of defined abstract names for OMP PLACES

Abstract Name	Meaning
threads	Each place corresponds to a single hardware thread on the target machine.
cores	Each place corresponds to a single core (having one or more hardware threads) on the target machine.
sockets	Each place corresponds to a single socket (consisting of one or more cores) on the target machine.

The behavior of the program is implementation defined when the execution environment cannot map a numerical value (either explicitly defined or implicitly derived from an interval) within the **OMP\_PLACES** list to a processor on the target platform, or if it maps to an unavailable processor. The behavior is also implementation defined when the **OMP\_PLACES** environment variable is defined using an abstract name.

1 Example:

```
setenv OMP_PLACES threads
setenv OMP_PLACES "threads(4)"
setenv OMP_PLACES "{0,1,2,3},{4,5,6,7},{8,9,10,11},{12,13,14,15}"
setenv OMP_PLACES "{0:4},{4:4},{8:4},{12:4}"
setenv OMP_PLACES "{0:4}:4:4"
```

where each of the last three definitions corresponds to the same 4 places including the smallest units of execution exposed by the execution environment numbered, in turn, 0 to 3, 4 to 7, 8 to 11, and 12 to 15.

#### **Cross References**

- place-partition-var, Section 2.3 on page 30
- thread affinity, Section 2.5.2 on page 42.

## 4.6 OMP NESTED

The OMP\_NESTED environment variable controls nested parallelism by setting the initial value of the *nest-var* ICV. The value of this environment variable must be true or false. If the environment variable is set to true, nested parallelism is enabled; if set to false, nested parallelism is disabled. The behavior of the program is implementation defined if the value of OMP NESTED is neither true nor false.

Example:

```
setenv OMP_NESTED false
```

#### Cross References

- nest-var ICV, see Section 2.3 on page 30.
- omp set nested routine, see Section 3.2.10 on page 173.
- omp\_get\_nested routine, see Section 3.2.19 on page 185.

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## 4.7 OMP STACKSIZE

The **OMP\_STACKSIZE** environment variable controls the size of the stack for threads created by the OpenMP implementation, by setting the value of the *stacksize-var* ICV. The environment variable does not control the size of the stack for the initial thread.

The value of this environment variable takes the form:

```
size | sizeB | sizeK | sizeM | sizeG
```

#### where:

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- *size* is a positive integer that specifies the size of the stack for threads that are created by the OpenMP implementation.
- B, K, M, and G are letters that specify whether the given size is in Bytes, Kilobytes (1024 Bytes), Megabytes (1024 Kilobytes), or Gigabytes (1024 Megabytes), respectively. If one of these letters is present, there may be white space between *size* and the letter.

If only size is specified and none of **B**, **K**, **M**, or **G** is specified, then size is assumed to be in Kilobytes.

The behavior of the program is implementation defined if **OMP\_STACKSIZE** does not conform to the above format, or if the implementation cannot provide a stack with the requested size.

#### Examples:

```
setenv OMP_STACKSIZE 2000500B
setenv OMP_STACKSIZE "3000 k "
setenv OMP_STACKSIZE 10M
setenv OMP_STACKSIZE " 10 M "
setenv OMP_STACKSIZE "20 m "
setenv OMP_STACKSIZE "1G"
setenv OMP_STACKSIZE 20000
```

#### Cross References

• stacksize-var ICV, see Section 2.3 on page 30.

## 4.8 OMP WAIT POLICY

The **OMP\_WAIT\_POLICY** environment variable provides a hint to an OpenMP implementation about the desired behavior of waiting threads by setting the *wait-policy-var* ICV. A compliant OpenMP implementation may or may not abide by the setting of the environment variable.

The value of this environment variable takes the form:

#### ACTIVE | PASSIVE

The **ACTIVE** value specifies that waiting threads should mostly be active, consuming processor cycles, while waiting. An OpenMP implementation may, for example, make waiting threads spin.

The **PASSIVE** value specifies that waiting threads should mostly be passive, not consuming processor cycles, while waiting. For example, an OpenMP implementation may make waiting threads yield the processor to other threads or go to sleep.

The details of the **ACTIVE** and **PASSIVE** behaviors are implementation defined.

15 Examples:

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```
setenv OMP_WAIT_POLICY ACTIVE
setenv OMP_WAIT_POLICY active
setenv OMP_WAIT_POLICY PASSIVE
setenv OMP_WAIT_POLICY passive
```

#### Cross References

• wait-policy-var ICV, see Section 2.3 on page 24.

## 4.9 OMP MAX ACTIVE LEVELS

The **OMP\_MAX\_ACTIVE\_LEVELS** environment variable controls the maximum number of nested active parallel regions by setting the initial value of the *max-active-levels-var* ICV.

The value of this environment variable must be a non-negative integer. The behavior of the program is implementation defined if the requested value of

OMP\_MAX\_ACTIVE\_LEVELS is greater than the maximum number of nested active parallel levels an implementation can support, or if the value is not a non-negative integer.

#### **Cross References**

- max-active-levels-var ICV, see Section 2.3 on page 30.
- omp\_set\_max\_active\_levels routine, see Section 3.2.15 on page 180.
- omp get max active levels routine, see Section 3.2.16 on page 182.

## 4.10 OMP THREAD LIMIT

The **OMP\_THREAD\_LIMIT** environment variable sets the number of OpenMP threads to use for the whole OpenMP program by setting the *thread-limit-var* ICV.

The value of this environment variable must be a positive integer. The behavior of the program is implementation defined if the requested value of **OMP\_THREAD\_LIMIT** is greater than the number of threads an implementation can support, or if the value is not a positive integer.

#### Cross References

- thread-limit-var ICV, see Section 2.3 on page 30.
- omp get thread limit routine

## 20 4.11 OMP CANCELLATION

The **OMP\_CANCELLATION** environment variable sets the initial value of the *cancel-var* ICV.

The value of this environment variable must be **true** or **false**. If set to **true**, the effects of the **cancel construct** and of **cancellation points** are enabled and cancellation is activated. If set to **false**, cancellation is disabled and the **cancel** construct and cancellation points are effectively ignored.

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## 1 Cross References:

- *cancel-var*, see Section 2.3.1 on page 31.
- cancel construct, see Section 2.13.1 on page 116.
- cancellation point construct, see Section 2.13.2 on page 119
  - omp get cancellation routine, see Section 3.2.9 on page 172

## 4.12 OMP DISPLAY ENV

The OMP\_DISPLAY\_ENV environment variable instructs the runtime to display the OpenMP version number and the value of the ICVs associated with the environment variables described in Chapter 4, as name=value pairs. The runtime displays this information once, after processing the environment variables and before any user calls to change the ICV values by runtime routines defined in Chapter 3.

The value of the **OMP\_DISPLAY\_ENV** environment variable may be set to one of these values:

#### TRUE | FALSE | VERBOSE

The TRUE value instructs the runtime to display the OpenMP version number defined by the \_OPENMP macro and the initial ICV values for the environment variables listed in Chapter 4. The VERBOSE value indicates that the runtime may also display the values of runtime variables that may be modified by vendor-specific environment variables. The runtime does not display any information when the OMP\_DISPLAY\_ENV environment is FALSE, undefined, or any other value than TRUE or VERBOSE.

The display begins with "OPENMP DISPLAY ENVIRONMENT BEGIN", followed by the \_OPENMP version macro value and ICV values, in the format NAME ' = ' VALUE. NAME corresponds to the macro or environment variable name, optionally prepended by a bracketed-device type. VALUE corresponds to the value of the macro or ICV associated with this environment variable. Values should be enclosed in single quotes. The display is terminated with "OPENMP DISPLAY ENVIRONMENT END".

#### Example:

1

```
OPENMP DISPLAY ENVIRONMENT BEGIN

_OPENMP='201301'

[host] OMP_SCHEDULE='GUIDED,4'

[host] OMP_NUM_THREADS='4,3,2'

[device] OMP_NUM_THREADS='2'

[host,device] OMP_DYNAMIC='TRUE'

[host] OMP_PLACES='{0:4},{4:4},{8:4},{12:4}'

...

OPENMP DISPLAY ENVIRONMENT END
```

## 4.13 OMP DEFAULT DEVICE

The **OMP\_DEFAULT\_DEVICE** environment variable sets the device number to use in **target** constructs by setting the initial value of the *default-device-var* ICV.

The value of this environment variable must be a non-negative integer value.

#### **Cross References:**

- default-device-var ICV, see Section 2.3 on page 30.
- target constructs, Section 2.9 on page 68

### 1 APPENDIX A

7

8

9

## **Examples**

The following are examples of the constructs and routines defined in this document.

C/C++

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

A.1 A Simple Parallel Loop

The following example demonstrates how to parallelize a simple loop using the parallel loop construct (Section 2.10.1 on page 82). The loop iteration variable is private by default, so it is not necessary to specify it explicitly in a **private** clause.

C/C++

```
C/C++
                   Example A.1.1c
10
11
                   void simple(int n, float *a, float *b)
12
13
                       int i;
14
15
                   #pragma omp parallel for
16
                       for (i=1; i<n; i++) /* i is private by default */
17
                           b[i] = (a[i] + a[i-1]) / 2.0;
18
                   }
                                                      C/C++
```

```
1
2
3
4
5
6
7
8
9
10
11
12
```

#### **Fortran**

#### Example A.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
```

Fortran

## A.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.

4

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```
C/C++
```

#### Example A.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++

27 28

#### Example A.2.1f

```
PROGRAM MEMMODEL
 INCLUDE "omp lib.h"
                          ! or USE OMP LIB
 INTEGER X
 X = 2
!$OMP PARALLEL NUM_THREADS(2) SHARED(X)
    IF (OMP GET THREAD NUM() .EQ. 0) THEN
      X = 5
    ! PRINT 1: The following read of x has a race
     PRINT *,"1: THREAD# ", OMP GET THREAD NUM(), "X = ", X
    ENDIF
 !SOMP BARRIER
   IF (OMP GET THREAD NUM() .EQ. 0) THEN
      PRINT *,"2: THREAD# ", OMP GET THREAD NUM(), "X = ", X
   ELSE
    ! PRINT 3
      PRINT *, "3: THREAD# ", OMP GET THREAD NUM(), "X = ", X
    ENDIF
!$OMP END PARALLEL
END PROGRAM MEMMODEL
```

Fortran •

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 A.2.2c

```
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35
36
37
38
39
40
41
```

42

43

```
#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;
}
                                  C/C++
```

#### 2 3 4 5 6 7 8 9

#### 9 10 11 12 13 14 15 16 17

18

19

32

33

34

26

35 36

37

38

```
Example A.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
```

#### Fortran

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 A.2.3c

```
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39
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41
42
43
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```

45

1

```
#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
                       /* 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;
}
                                   C/C++
```

Appendix A Examples

```
1
```

14

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30 31 32

33 34 35

36

37

```
Example A.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
```

Fortran –

END

## **A.3** Conditional Compilation

	C/C++
2	The following example illustrates the use of conditional compilation using the OpenMP
3	macro <b>OPENMP</b> (Section 2.2 on page 28). With OpenMP compilation, the <b>OPENMP</b>
4	macro becomes defined.
5	Example A.3.1c
6 7	<pre>#include <stdio.h></stdio.h></pre>
8	<pre>int main()</pre>
9	{
10	
11 12	# ifdef _OPENMP
13	<pre>printf("Compiled by an OpenMP-compliant implementation.\n"); # endif</pre>
14	"
15	return 0;
16	}
	C/C++
	Fortran
17	The following example illustrates the use of the conditional compilation sentinel (see
18	Section 2.2 on page 28). With OpenMP compilation, the conditional compilation
19	sentinel !\$ is recognized and treated as two spaces. In fixed form source, statements
20	guarded by the sentinel must start after column 6.
21	Example A.3.1f
22	PROGRAM EXAMPLE
23	
24 25	C234567890 !\$ PRINT *, "Compiled by an OpenMP-compliant implementation."
26	., IXIXI , compiled by an opening-compilant implementation.
27	END PROGRAM EXAMPLE
	Fortran

## A.4 Internal Control Variables (ICVs)

According to Section 2.3 on page 30, 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 <code>omp\_set\_nested</code>, <code>omp\_set\_max\_active\_levels</code>, <code>omp\_set\_dynamic</code>, and <code>omp\_set\_num\_threads</code> respectively. These ICVs affect the operation of <code>parallel</code> regions. Each implicit task generated by a <code>parallel</code> 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 *maxactive-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 A.4.1c

}

return 0;

}

```
1
 2
 3
 4
 5
 6
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8
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10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
```

29

30 31

32

33

34 35

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37

38

39

40

41

42 43

44

```
#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
           */
```

C/C++

omp get max threads());

printf ("Outer: max act lev=%d, num thds=%d, max thds=%d\n",

omp get max active levels(), omp get num threads(),

**Fortran** 

#### 1 Example A.4.1f

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37

38

```
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
```

Fortran

## A.5 The parallel Construct

The **parallel** construct (Section 2.5 on page 37) 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:

```
1
```

Example A.5.1c

```
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 4
 5
 6
 7
 8
 9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
```

34

35

```
#include <omp.h>
void subdomain(float *x, int istart, int ipoints)
 int i;
  for (i = 0; i < ipoints; i++)</pre>
      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++

34 35

36

37

38

#### Example A.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
!$OMP PARALLEL DEFAULT(PRIVATE) SHARED(X, NPOINTS)
          IAM = OMP GET THREAD NUM()
         NT = OMP_GET_NUM_THREADS()
          IPOINTS = NPOINTS/NT
          ISTART = IAM * IPOINTS
          IF (IAM .EQ. NT-1) THEN
              IPOINTS = NPOINTS - ISTART
          CALL SUBDOMAIN(X, ISTART, IPOINTS)
!$OMP END PARALLEL
     END SUBROUTINE SUB
      PROGRAM PAREXAMPLE
         REAL ARRAY(10000)
         CALL SUB (ARRAY, 10000)
     END PROGRAM PAREXAMPLE
```

Fortran -

# A.6 Controlling the Number of Threads on Multiple Nesting Levels

The following examples demonstrate how to use the **OMP\_NUM\_THREADS** environment variable (Section 2.3.2 on page 32) to control the number of threads on multiple nesting levels:

C/C++

#### Example A.6.1c

```
6
 7
 8
 9
10
11
12
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18
19
20
21
22
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25
26
27
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29
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33
34
35
36
37
38
39
```

40

41

42

43

2

3

4

```
#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());
```

```
1
                                   }
2
3
                               #pragma omp barrier
4
                               #pragma omp single
5
6
7
                                   * If OMP NUM THREADS=2,3 was set, the following should print:
8
                                   * Outer: num thds=2
9
10
                                   printf ("Outer: num thds=%d\n", omp get num threads());
                               }
11
12
                           }
13
                           return 0;
14
                    }
                                                       C/C++ -

    Fortran

                    Example A.6.1f
15
16
                           program icv
17
                             use omp lib
18
                             call omp_set_nested(.true.)
19
                             call omp set dynamic(.false.)
20
                    !$omp parallel
21
                    !$omp parallel
22
                    !$omp single
23
                             ! If OMP_NUM_THREADS=2,3 was set, the following should print:
24
                             ! Inner: num thds= 3
25
                             ! Inner: num thds= 3
26
                             ! If nesting is not supported, the following should print:
27
                             ! Inner: num thds= 1
28
                             ! Inner: num thds= 1
29
                             print *, "Inner: num_thds=", omp_get_num_threads()
30
                    !$omp end single
31
                    !$omp end parallel
32
                    !$omp barrier
33
                             call omp set nested(.false.)
34
                    !$omp parallel
35
                    !$omp single
36
                             ! Even if OMP NUM THREADS=2,3 was set, the following should print,
37
                             ! because nesting is disabled:
38
                             ! Inner: num thds= 1
39
                             ! Inner: num_thds= 1
                             print *, "Inner: num_thds=", omp_get_num_threads()
40
41
                    !$omp end single
42
                    !$omp end parallel
43
                    !$omp barrier
44
                    !$omp single
45
                             ! If OMP_NUM_THREADS=2,3 was set, the following should print:
46
                              ! Outer: num thds= 2
47
                             print *, "Outer: num thds=", omp get num threads()
```

1 !\$omp end single
2 !\$omp end parallel
3 end
Fortran

# A.7 Interaction Between the num\_threads Clause and omp set dynamic

The following example demonstrates the num\_threads clause (Section 2.5 on page 37) and the effect of the omp\_set\_dynamic routine (Section 3.2.7 on page 169) 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.

C/C++ Example A.7.1c 17 18 #include <omp.h> 19 int main() 20 21 omp set dynamic(0); 22 #pragma omp parallel num\_threads(10) 23 24 /\* do work here \*/ 25 26 return 0; 27 } C/C++

Fortran —

#### Example A.7.1f

5

7

8 9

10

11

12

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PROGRAM EXAMPLE
INCLUDE "omp\_lib.h" ! or USE OMP\_LIB

```
1
                            CALL OMP SET DYNAMIC (.FALSE.)
                    !$OMP
                              PARALLEL NUM THREADS (10)
 3
                                 ! do work here
 4
                              END PARALLEL
                    ! SOMP
                          END PROGRAM EXAMPLE
                                                       Fortran
                    The call to the omp set dynamic routine with a non-zero argument in C/C++, or
 6
                    .TRUE. in Fortran, allows the OpenMP implementation to choose any number of
                    threads between 1 and 10 (see also Algorithm 2.1 in Section 2.5.1 on page 40).
                                                        C/C++
                    Example A.7.2c
 9
10
                    #include <omp.h>
11
                    int main()
13
                      omp_set_dynamic(1);
                      #pragma omp parallel num threads(10)
15
                         /* do work here */
16
                      }
17
18
                      return 0;
19
                                                        C/C++
                                                        Fortran
                    Example A.7.2f
20
21
                          PROGRAM EXAMPLE
22
                            INCLUDE "omp lib.h"
                                                       ! or USE OMP LIB
23
                            CALL OMP SET DYNAMIC (.TRUE.)
                              PARALLEL NUM THREADS (10)
24
                    !$OMP
25
                                 ! do work here
26
                              END PARALLEL
                    !$OMP
27
                          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.

28

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6

## A.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. For more information, see Section 2.7.1 on page 47. The following example contains correct usages of loop constructs:

```
Example A.8.1f
```

2

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20 21

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42

43

44

45

```
SUBROUTINE WORK(I, J)
      INTEGER I,J
      END SUBROUTINE WORK
      SUBROUTINE DO GOOD ()
        INTEGER I, J
        REAL A(1000)
        DO 100 I = 1,10
!$OMP
          DO 100 J = 1,10
            CALL WORK (I, J)
100
        CONTINUE ! !$OMP ENDDO implied here
! SOMP
       DO
        DO 200 J = 1,10
        A(I) = I + 1
200
!$OMP
        ENDDO
! SOMP
       DΩ
        DO 300 I = 1,10
          DO 300 J = 1,10
            CALL WORK (I, J)
300
        CONTINUE
!$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 A.8.2f

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

SUBROUTINE DO_WRONG
INTEGER I, J

DO 100 I = 1,10

PO 100 J = 1,10

CALL WORK(I, J)

CONTINUE

SOMP ENDDO
END SUBROUTINE DO WRONG
```

Fortran

## A.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 Section 2.7.1 on page 47 and Section 2.14.1 on page 122). In the following example of a sequential loop in a **parallel** construct the loop iteration variable *I* will be private.

#### Example A.9.1f

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

#### Example A.9.2f

1 2

3

4

5

6

7 8

9

10

11

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24 25

26

27

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30

31

```
SUBROUTINE PLOOP 2(A,B,N,I1,I2)
REAL A(*), B(*)
INTEGER I1, I2, N
!$OMP PARALLEL SHARED(A,B,I1,I2)
!SOMP SECTIONS
!$OMP SECTION
    DO I1 = I1, N
       IF (A(I1).NE.0.0) EXIT
     ENDDO
!SOMP SECTION
    DO I2 = I2, 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.'
!SOMP END SINGLE
!$OMP END PARALLEL
```

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

Fortran -

## 28 A.10 The nowait clause

END SUBROUTINE PLOOP 2

If there are multiple independent loops within a **parallel** region, you can use the **nowait** clause (see Section 2.7.1 on page 47) to avoid the implied barrier at the end of the loop construct, as follows:

```
C/C++
                    Example A.10.1c
 1
 2
                    #include <math.h>
 3
4
                    void nowait example(int n, int m, float *a, float *b, float *y, float *z)
5
6
                      int i;
7
                      #pragma omp parallel
8
9
                        #pragma omp for nowait
10
                          for (i=1; i<n; i++)
11
                            b[i] = (a[i] + a[i-1]) / 2.0;
12
13
                        #pragma omp for nowait
14
                          for (i=0; i<m; i++)
15
                            y[i] = sqrt(z[i]);
16
                      }
17
                    }
                                                       C/C++ -
                                                     Fortran
                    Example A.10.1f
18
19
                            SUBROUTINE NOWAIT EXAMPLE(N, M, A, B, Y, Z)
20
21
                            INTEGER N, M
22
                            REAL A(*), B(*), Y(*), Z(*)
23
24
                            INTEGER I
25
26
                    !$OMP PARALLEL
27
28
                    !$OMP DO
29
                            DO I=2,N
30
                              B(I) = (A(I) + A(I-1)) / 2.0
31
32
                    !$OMP END DO NOWAIT
33
34
                    !$OMP DO
35
                            DO I=1,M
36
                              Y(I) = SQRT(Z(I))
37
                            ENDDO
38
                    !$OMP END DO NOWAIT
39
40
                    !$OMP END PARALLEL
41
42
                            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 A.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 A.10.2f
   SUBROUTINE NOWAIT EXAMPLE2(N, A, B, C, Y, Z)
   INTEGER N
  REAL A(*), B(*), C(*), Y(*), Z(*)
```

```
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
```

## A.11 The collapse clause

For the following three examples, see Section 2.7.1 on page 47 for a description of the **collapse** clause, Section 2.12.8 on page 114 for a description of the **ordered** construct, and Section 2.14.3.5 on page 139 for a description of the **lastprivate** 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.

C/C++

#### Example A.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);
}</pre>
```

Appendix A Examples

#### Example A.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.

```
C/C++
                   Example A.11.2c
1
2
                    #include <stdio.h>
3
                    void test()
4
5
                       int j, k, jlast, klast;
6
                       #pragma omp parallel
7
8
                          #pragma omp for collapse(2) lastprivate(jlast, klast)
9
                          for (k=1; k<=2; k++)
10
                             for (j=1; j<=3; j++)
11
12
                                jlast=j;
13
                                klast=k;
14
15
                          #pragma omp single
16
                          printf("%d %d\n", klast, jlast);
17
                       }
18
                    }
                                                       C/C++ -
                                                     Fortran -
                   Example A.11.2f
19
20
                          program test
21
                    !$omp parallel
22
                    !$omp do private(j,k) collapse(2) lastprivate(jlast, klast)
23
                          do k = 1,2
24
                            do j = 1,3
25
                              jlast=j
26
                              klast=k
27
                            enddo
28
                          enddo
29
                    !$omp end do
30
                    !$omp single
31
                                    print *, klast, jlast
32
                    !$omp end single
33
                    !$omp end parallel
34
                          end program test

    Fortran -
```

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

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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 Section 2.12.8 on page 114, 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).

C/C++

#### The code prints

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

#### Example A.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++

```
Fortran
                    Example A.11.3f
 1
2
                          program test
3
                          include 'omp lib.h'
4
                    !$omp parallel num threads(2)
5
                    !$omp do collapse(2) ordered private(j,k) schedule(static,3)
6
                          do k = 1,3
7
                            do j = 1,2
8
                    !$omp ordered
9
                              print *, omp get thread num(), k, j
10
                    !$omp end ordered
11
                              call work(a,j,k)
12
                            enddo
13
                          enddo
14
                    !$omp end do
15
                    !$omp end parallel
16
                          end program test
                                                       Fortran
```

## A.12 The parallel sections Construct

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In the following example (for Section 2.10.2 on page 84) routines **XAXIS**, 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 A.12.1c
22
23
                    void XAXIS();
24
                    void YAXIS();
25
                   void ZAXIS();
26
27
                   void sect example()
28
29
                      #pragma omp parallel sections
30
31
                        #pragma omp section
32
                          XAXIS();
33
34
                        #pragma omp section
35
                          YAXIS();
36
```

```
1
                        #pragma omp section
 2
                           ZAXIS();
 3
                      }
                    }
                                                        C/C++
                                                        Fortran
                    Example A.12.1f
 5
6
                           SUBROUTINE SECT EXAMPLE()
7
                    !$OMP PARALLEL SECTIONS
9
10
                    !SOMP SECTION
11
                             CALL XAXIS()
12
13
                    !SOMP SECTION
                             CALL YAXIS()
15
16
                    !$OMP SECTION
17
                             CALL ZAXIS()
18
19
                    !$OMP END PARALLEL SECTIONS
20
                          END SUBROUTINE SECT EXAMPLE
                                                       Fortran •
```

# A.13 The firstprivate Clause and the sections Construct

In the following example of the **sections** construct (Section 2.7.2 on page 53) 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 A.13.1c
```

#include <omp.h>

22

23

24

25

26

27

28

29

30

31

```
#include <stdio.h>
1
2
                    #define NT 4
3
                    int main() {
4
                        int section count = 0;
5
                        omp set dynamic(0);
6
                        omp set num threads(NT);
7
                    #pragma omp parallel
8
                    #pragma omp sections firstprivate( section count )
9
10
                    #pragma omp section
11
                        {
12
                            section count++;
13
                            /* may print the number one or two */
14
                            printf( "section count %d\n", section count );
15
16
                    #pragma omp section
17
18
                            section count++;
19
                            /* may print the number one or two */
20
                            printf( "section_count %d\n", section_count );
21
22
                    }
23
                        return 1;
24
                                                       C/C++
                                                       Fortran
                    Example A.13.1f
25
26
                   program section
27
                        use omp_lib
28
                        integer :: section count = 0
29
                        integer, parameter :: NT = 4
30
                        call omp set dynamic (.false.)
                        call omp set num threads(NT)
31
32
                    !$omp parallel
33
                    !$omp sections firstprivate ( section count )
34
                    !$omp section
35
                        section count = section count + 1
36
                    ! may print the number one or two
37
                        print *, 'section count', section count
38
                    !$omp section
39
                        section count = section count + 1
                    ! may print the number one or two
40
41
                        print *, 'section count', section count
42
                    !$omp end sections
43
                    !$omp end parallel
44
                    end program section
                                                       Fortran
```

The following example demonstrates the **single** construct (Section 2.7.3 on page 56). 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 A.14.1c

#### Example A.14.1f

1

31

32

33

34

35

36

```
2
                           SUBROUTINE WORK1()
3
                           END SUBROUTINE WORK1
4
5
                           SUBROUTINE WORK2()
6
                          END SUBROUTINE WORK2
7
8
                          PROGRAM SINGLE EXAMPLE
9
                    !$OMP PARALLEL
10
11
                    !$OMP SINGLE
12
                            print *, "Beginning work1."
13
                    !$OMP END SINGLE
14
15
                             CALL WORK1()
16
17
                    !$OMP SINGLE
18
                             print *, "Finishing work1."
19
                    !$OMP END SINGLE
20
21
                    !SOMP SINGLE
22
                            print *, "Finished work1 and beginning work2."
23
                    !$OMP END SINGLE NOWAIT
24
25
                             CALL WORK2()
26
27
                    !$OMP END PARALLEL
28
29
                           END PROGRAM SINGLE_EXAMPLE
```

Fortran -

## A.15 Tasking Constructs

The following example shows how to traverse a tree-like structure using explicit tasks (see Section 2.11 on page 88). 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 A.15.1c
 1
2
                   struct node {
3
                     struct node *left;
4
                     struct node *right;
5
6
7
                   extern void process(struct node *);
                   void traverse( struct node *p ) {
                     if (p->left)
9
                   #pragma omp task // p is firstprivate by default
10
                         traverse(p->left);
11
                     if (p->right)
12
                   #pragma omp task
                                     // p is firstprivate by default
13
                         traverse (p->right);
14
                     process(p);
15
                   }
                                                      C/C++ -
                                                    - Fortran -
                   Example A.15.1f
16
17
                          RECURSIVE SUBROUTINE traverse ( P )
18
                             TYPE Node
19
                               TYPE(Node), POINTER :: left, right
20
                             END TYPE Node
21
                             TYPE(Node) :: P
22
                             IF (associated(P%left)) THEN
23
                                      !$OMP TASK ! P is firstprivate by default
24
                                          call traverse(P%left)
25
                                      !$OMP END TASK
26
                             ENDIF
27
                             IF (associated(P%right)) THEN
28
                                      !$OMP TASK ! P is firstprivate by default
29
                                          call traverse(P%right)
30
                                      !$OMP END TASK
31
                             ENDIF
32
                             CALL process ( P )
33
                           END SUBROUTINE
                                                    - Fortran –
```

1 In the next example, we force a postorder traversal of the tree by adding a taskwait 2 directive (see Section 2.12.4 on page 101). Now, we can safely assume that the left and 3 right sons have been executed before we process the current node. C/C++ Example A.15.2c 4 5 struct node { 6 struct node \*left; 7 struct node \*right; 8 **}**; 9 extern void process(struct node \*); 10 void postorder traverse( struct node \*p ) { 11 if (p->left) 12 #pragma omp task // p is firstprivate by default 13 postorder traverse(p->left); 14 if (p->right) 15 #pragma omp task // p is firstprivate by default 16 postorder traverse(p->right); 17 #pragma omp taskwait 18 process(p); 19 } C/C++ - Fortran · Example A.15.2f 20 21 RECURSIVE SUBROUTINE traverse ( P ) 22 TYPE Node 23 TYPE(Node), POINTER :: left, right 24 END TYPE Node 25 TYPE(Node) :: P 26 IF (associated(P%left)) THEN 27 !\$OMP TASK ! P is firstprivate by default 28 call traverse(P%left) 29 !SOMP END TASK 30 ENDIF 31 IF (associated(P%right)) THEN 32 !\$OMP TASK ! P is firstprivate by default 33 call traverse (P%right) 34 !\$OMP END TASK 35 ENDIF 36 !\$OMP TASKWAIT CALL process ( P ) 37 38 END SUBROUTINE Fortran –

31

32

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 (see page 125).

#### \_\_\_\_\_ C/C++

```
Example A.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 A.15.3f

1

```
2
                          MODULE LIST
3
                             TYPE NODE
4
                                 INTEGER :: PAYLOAD
5
                                 TYPE (NODE), POINTER :: NEXT
6
                             END TYPE NODE
7
                          CONTAINS
8
                              SUBROUTINE PROCESS(p)
9
                                 TYPE (NODE), POINTER :: P
10
                                      ! do work here
11
                              END SUBROUTINE
12
                              SUBROUTINE INCREMENT LIST ITEMS (HEAD)
13
                                  TYPE (NODE), POINTER :: HEAD
14
                                   TYPE (NODE), POINTER :: P
15
                                   !$OMP PARALLEL PRIVATE(P)
16
                                      !$OMP SINGLE
17
                                           P => HEAD
18
                                           DO
19
                                              !$OMP TASK
20
                                                  ! P is firstprivate by default
21
                                                  CALL PROCESS (P)
22
                                              !$OMP END TASK
23
                                              P => P%NEXT
24
                                              IF ( .NOT. ASSOCIATED (P) ) EXIT
25
                                           END DO
26
                                     !$OMP END SINGLE
27
                                 !$OMP END PARALLEL
28
                              END SUBROUTINE
29
                           END MODULE
```

- Fortran -

The fib() function should be called from within a parallel region for the different 1 2 specified tasks to be executed in parallel. Also, only one thread of the parallel 3 region should call fib () unless multiple concurrent Fibonacci computations are desired. 4 C/C++ Example A.15.4c 5 int fib(int n) { 7 int i, j; 8 if (n<2) 9 return n; 10 else { 11 #pragma omp task shared(i) 12 i=fib(n-1); 13 #pragma omp task shared(j) 14 j=fib(n-2); 15 #pragma omp taskwait 16 return i+j; 17 } 18 } C/C++ Fortran Example A.15.4f 19 20 RECURSIVE INTEGER FUNCTION fib(n) RESULT(res) 21 INTEGER n, i, j 22 IF ( n .LT. 2) THEN 23 res = n 24 ELSE 25 !\$OMP TASK SHARED(i) 26 i = fib(n-1)27 !\$OMP END TASK 28 !\$OMP TASK SHARED(j) 29 i = fib(n-2)30 !\$OMP END TASK 31 !\$OMP TASKWAIT 32 res = i+j33 END IF 34 END FUNCTION

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

Fortran

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1 The following example demonstrates a way to generate a large number of tasks with one 2 thread and execute them with the threads in the team (see Section 2.11.3 on page 94). 3 While generating these tasks, the implementation may reach its limit on unassigned 4 tasks. If it does, the implementation is allowed to cause the thread executing the task 5 generating loop to suspend its task at the task scheduling point in the task directive, 6 and start executing unassigned tasks. Once the number of unassigned tasks is 7 sufficiently low, the thread may resume execution of the task generating loop. C/C++ Example A.15.5c 8 9 #define LARGE NUMBER 10000000 10 double item[LARGE NUMBER]; 11 extern void process(double); 12 13 int main() { 14 #pragma omp parallel 15 16 #pragma omp single 17 18 int i; 19 for (i=0; i<LARGE NUMBER; i++) 20 #pragma omp task // i is firstprivate, item is shared 21 process(item[i]); 22 23 } 24 } C/C++ Fortran Example A.15.5f 25 26 real\*8 item(10000000) 27 integer i 28 29 !\$omp parallel 30 !\$omp single ! loop iteration variable i is private 31 do i=1,1000000032 !\$omp task 33 ! i is firstprivate, item is shared 34 call process(item(i)) 35 !\$omp end task 36 end do 37 !\$omp end single 38 !\$omp end parallel 39 end Fortran

The following example is the same as the previous one, except that the tasks are generated in an untied task (see Section 2.11 on page 88). 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++

C/C++

#### Example A.15.6c

#### Example A.15.6f

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#### Fortran

The following two examples demonstrate how the scheduling rules illustrated in Section 2.11.3 on page 94 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.

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.

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42
43
44
45
```

```
C/C++ -
Example A.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 A.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 -

end subroutine

1

In this example, scheduling constraints (see Section 2.11.3 on page 94) 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.

```
C/C++
Example A.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
        }
    }
}
                                   C/C++ -
                                 - Fortran -
Example A.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
```

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- 1		
•		

end module

The following two examples demonstrate how the scheduling rules illustrated in Section 2.11.3 on page 94 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.

Fortran

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.

#### C/C++

C/C++

#### Example A.15.9c

## 1 Example A.15.9f

```
2
                           module example
3
                           contains
4
                           subroutine work
5
                    !$omp task
                           ! Task 1
7
                    !$omp task
8
                           ! Task 2
9
                    !$omp critical
10
                           ! Critical region 1
11
                           ! do work here
12
                    !$omp end critical
13
                    !$omp end task
14
                    !$omp critical
15
                           ! Critical region 2
16
                           ! Capture data for the following task
17
                    !$omp task
18
                           !Task 3
19
                           ! do work here
20
                    !$omp end task
21
                    !$omp end critical
22
                    !$omp end task
23
                          end subroutine
24
                          end module
```

Fortran -

28

In the following example, **lock** is held across a task scheduling point. However, according to the scheduling restrictions outlined in Section 2.11.3 on page 94, 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 A.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++

#### Example A.15.10f

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```
module example
      include 'omp lib.h'
      integer (kind=omp lock kind) lock
      integer i
      contains
      subroutine work
      call omp init lock(lock)
!$omp parallel
     !somp do
      do i=1,100
         !$omp task
              ! Outer task
              call omp set lock(lock)
                                          ! lock is shared by
                                          ! default in the task
                     ! Capture data for the following task
                                     ! Task Scheduling Point 1
                     !$omp task
                               ! do work here
                     !$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 (see Section 2.11.1 on page 88). 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 (see Section 1.2.4 on page 8). 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 A.15.11c

```
#include <stdio.h>
void foo ( )
{
   int x = 2;
```

```
1
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 3
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```

```
#pragma omp task shared(x) mergeable
      x++;
   #pragma omp taskwait
   printf("%d\n",x); // prints 3
                                  C/C++ -
                                  Fortran
Example A.15.11f
subroutine foo()
 integer :: x
!$omp task shared(x) mergeable
 x = x + 1
!$omp end task
!$omp taskwait
 print *, x
                 ! prints 3
end subroutine
```

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.

Fortran

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

#### Example A.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 (see Section 2.11.1 on page 88) and the **omp\_in\_final** API call (see Section 3.2.21 on page 188) 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 <code>omp\_in\_final</code> 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 <code>final</code> 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 <code>new\_state</code> in the stack could also be avoided but it would make this example less clear. The <code>final</code> clause is most effective when used in conjunction with the <code>mergeable</code> clause since all tasks created in a <code>final</code> task region are included tasks that can be merged if the <code>mergeable</code> clause is present.

#### C/C++

#### Example A.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() ) {
```

```
1
                            memcpy(new state, state, pos);
2
                            state = new state;
3
4
                          state[pos] = 0;
5
                          bin search(pos+1, n, state );
6
7
                       #pragma omp task final( pos > LIMIT ) mergeable
8
9
                          char new state[n];
10
                          if (! omp in final() ) {
11
                            memcpy (new state, state, pos );
12
                            state = new state;
14
                          state[pos] = 1;
15
                          bin search(pos+1, n, state );
16
17
                       #pragma omp taskwait
18
                    }
                                                       C/C++
                                                       Fortran
                    Example A.15.13f
19
20
                    recursive subroutine bin search(pos, n, state)
21
                      use omp_lib
22
                      integer :: pos, n
23
                      character, pointer :: state(:)
24
                      character, target, dimension(n) :: new_state1, new_state2
25
                      integer, parameter :: LIMIT = 3
26
                      if (pos .eq. n) then
27
                        call check solution(state)
28
                        return
29
                      endif
30
                    !$omp task final(pos > LIMIT) mergeable
31
                      if (.not. omp in final()) then
32
                        new state1(1:pos) = state(1:pos)
33
                        state => new state1
34
                      endif
35
                      state(pos+1) = 'z'
36
                      call bin search(pos+1, n, state)
37
                    !$omp end task
38
                    !$omp task final(pos > LIMIT) mergeable
39
                      if (.not. omp in final()) then
40
                        new state2(1:pos) = state(1:pos)
41
                        state => new state2
42
                      endif
43
                      state(pos+1) = 'y'
44
                      call bin search(pos+1, n, state)
45
                    !$omp end task
```

!\$omp taskwait

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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.

C/C++

Fortran

## Example A.15.14c

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

C/C++

#### Fortran

#### Example A.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
```

```
1
 2
 3
 4
 5
 6
 7
 8
9
10
11
12
13
15
16
17
```

```
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
```

## 14 A.16 The taskyield Directive

Example A.16.1c

The following example illustrates the use of the taskyield directive (see Section 2.11.2 on page 92). 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.

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```

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31
32
```

```
C/C++
```

Fortran Example A.16.1f 1 2 subroutine foo ( lock, n ) 3 use omp lib 4 integer (kind=omp lock kind) :: lock 5 integer n 6 integer i 7 8 do i = 1, n 9 !\$omp task shared(lock) 10 call something useful() 11 do while ( .not. omp test lock(lock) ) 12 !\$omp taskyield 13 end do 14 call something critical() 15 call omp\_unset\_lock(lock) 16 !\$omp end task 17 end do 18 19 end subroutine Fortran 20 21 22 Fortran 23

### A.17 The workshare Construct

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The following are examples of the **workshare** construct (see Section 2.7.4 on page 58).

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 A.17.1f 1 2 SUBROUTINE WSHARE1 (AA, BB, CC, DD, EE, FF, N) 3 REAL AA(N,N), BB(N,N), CC(N,N), DD(N,N), EE(N,N), FF(N,N) 5 6 !\$OMP PARALLEL 7 !\$OMP WORKSHARE 8 AA = BB9 CC = DD10 EE = FF11 !\$OMP END WORKSHARE 12 ! SOMP END PARALLEL 13 14 END SUBROUTINE WSHARE1 15 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 16 EE = FF when they are done with CC = DD. 17 Example A.17.2f 18 19 SUBROUTINE WSHARE2 (AA, BB, CC, DD, EE, FF, N) 20 INTEGER N 21 REAL AA(N,N), BB(N,N), CC(N,N) 22 REAL DD(N,N), EE(N,N), FF(N,N) 23 24 !\$OMP PARALLEL 25 !\$OMP WORKSHARE 26 AA = BB27 CC = DD28 !\$OMP END WORKSHARE NOWAIT 29 !\$OMP WORKSHARE 30 EE = FF31 !\$OMP END WORKSHARE 32 !\$OMP END PARALLEL 33 END SUBROUTINE WSHARE2 34 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 A.17.3f
 1
 2
                         SUBROUTINE WSHARE3 (AA, BB, CC, DD, N)
 3
                         INTEGER N
 4
                         REAL AA(N,N), BB(N,N), CC(N,N), DD(N,N)
 5
                         REAL R
 6
 7
                          R=0
 8
                   !$OMP
                          PARALLEL
 9
                   !$OMP
                            WORKSHARE
10
                              AA = BB
11
                   !$OMP
                              ATOMIC UPDATE
12
                                R = R + SUM(AA)
13
                              CC = DD
14
                            END WORKSHARE
                   ! SOMP
15
                   !$OMP END PARALLEL
16
17
                         END SUBROUTINE WSHARE3
                   Fortran WHERE and FORALL statements are compound statements, made up of a control
18
19
                   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
20
21
                   example shows the use of a WHERE statement in a workshare construct.
22
                   Each task gets worked on in order by the threads:
23
                   AA = BB then
24
                   CC = DD then
25
                   EE .ne. 0 then
26
                   FF = 1 / EE then
27
                   GG = HH
                   Example A.17.4f
28
29
                         SUBROUTINE WSHARE4 (AA, BB, CC, DD, EE, FF, GG, HH, N)
30
                         INTEGER N
31
                         REAL AA(N,N), BB(N,N), CC(N,N)
32
                         REAL DD(N,N), EE(N,N), FF(N,N)
33
                         REAL GG(N,N), HH(N,N)
34
35
                   ! SOMP
                          PARALLEL
36
                   !$OMP
                             WORKSHARE
37
                              AA = BB
38
                              CC = DD
39
                              WHERE (EE .ne. 0) FF = 1 / EE
40
                              GG = HH
41
                   ! SOMP
                            END WORKSHARE
42
                   !$OMP
                           END PARALLEL
43
44
                         END SUBROUTINE WSHARE4
```

,	Fortran	(cont.)	)
---	---------	---------	---

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 A.17.5f

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 A.17.6f

```
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 WORKSHARE
!$OMP END WORKSHARE
```

SUBROUTINE WSHARE6 WRONG(AA, BB, CC, DD, N)

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 A.17.7f

Fortran -

### A.18 The master Construct

The following example demonstrates the master construct (Section 2.12.1 on page 96). 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 A.18.1c
```

```
23
                    #include <stdio.h>
24
25
                    extern float average(float, float, float);
26
27
                    void master example (float* x, float* xold, int n, float tol)
28
29
                      int c, i, toobig;
30
                      float error, y;
31
                      c = 0;
32
                      #pragma omp parallel
33
34
                        do{
35
                          #pragma omp for private(i)
36
                          for( i = 1; i < n-1; ++i)
```

xold[i] = x[i];

```
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
```

```
}
#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++
```

Fortran

#### Example A.18.1f

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```
2
                           SUBROUTINE MASTER EXAMPLE ( X, XOLD, N, TOL )
 3
                           REAL X(*), XOLD(*), TOL
 4
                           INTEGER N
 5
                           INTEGER C, I, TOOBIG
 6
                           REAL ERROR, Y, AVERAGE
 7
                           EXTERNAL AVERAGE
 8
                           C = 0
 9
                           TOOBIG = 1
10
                     !SOMP PARALLEL
11
                             DO WHILE ( TOOBIG > 0 )
12
                     !$OMP
                               DO PRIVATE(I)
13
                                 DO I = 2, N-1
14
                                   XOLD(I) = X(I)
15
                                 ENDDO
16
                     ! SOMP
                               SINGLE
17
                                 TOOBIG = 0
18
                     !$OMP
                               END SINGLE
19
                     !$OMP
                               DO PRIVATE(I,Y,ERROR), REDUCTION(+:TOOBIG)
20
                                 DO I = 2, N-1
21
                                   Y = X(I)
22
                                   X(I) = AVERAGE(XOLD(I-1), X(I), XOLD(I+1))
23
                                   ERROR = Y-X(I)
24
                                   IF( ERROR > TOL .OR. ERROR < -TOL ) TOOBIG = TOOBIG+1</pre>
25
                                 ENDDO
26
                               MASTER
                     !$OMP
27
                                 C = C + 1
28
                                 PRINT *, 'Iteration ', C, 'TOOBIG=', TOOBIG
29
                               END MASTER
                     !$OMP
30
                             ENDDO
31
                     !$OMP END PARALLEL
32
                           END SUBROUTINE MASTER EXAMPLE
```

Fortran

### A.19 The critical Construct

The following example includes several **critical** constructs (Section 2.12.2 on page 97). 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 A.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 A.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 •
```

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# A.20 worksharing Constructs Inside a critical Construct

The following example demonstrates using a worksharing construct inside a **critical** construct (see Section 2.12.2 on page 97). This example is conforming because the worksharing **single** region is not closely nested inside the **critical** region (see Section 2.16 on page 158). 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++

C/C++

# Example A.20.1c

Appendix A Examples

#### **Fortran**

#### Example A.20.1f

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

Fortran

## A.21 Binding of barrier Regions

END SUBROUTINE CRITICAL WORK

The binding rules call for a **barrier** region to bind to the closest enclosing **parallel** region (see Section 2.12.3 on page 99).

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.

C/C++

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```

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```

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```
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 A.21.1c

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```

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```

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```

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```
Example A.21.1f
```

```
SUBROUTINE WORK (N)
        INTEGER N
      END SUBROUTINE WORK
      SUBROUTINE SUB3 (N)
      INTEGER N
        CALL WORK (N)
!$OMP
        BARRIER
        CALL WORK (N)
      END SUBROUTINE SUB3
      SUBROUTINE SUB2 (K)
      INTEGER K
        PARALLEL SHARED(K)
!$OMP
          CALL SUB3 (K)
!$OMP
        END PARALLEL
      END SUBROUTINE SUB2
      SUBROUTINE SUB1(N)
      INTEGER N
        INTEGER I
!$OMP
        PARALLEL PRIVATE(I) SHARED(N)
!$OMP
          DO I = 1, N
            CALL SUB2(I)
          END DO
!$OMP
        END PARALLEL
      END SUBROUTINE SUB1
      PROGRAM EXAMPLE
        CALL SUB1(2)
        CALL SUB2(2)
        CALL SUB3(2)
      END PROGRAM EXAMPLE
```

Fortran •

#### **A.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 (Section 2.12.6 on page 103).

5

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45

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 (see Section 2.12.2 on page 97) 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 A.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++
```

```
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 9
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23
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```

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#### Example A.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-1)
            Y(I) = Y(I) + WORK2(I-1)
          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-1, 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 -

1 The following example illustrates the read and write clauses for the atomic 2 directive. These clauses ensure that the given variable is read or written, respectively, as 3 a whole. Otherwise, some other thread might read or write part of the variable while the 4 current thread was reading or writing another part of the variable. Note that most 5 hardware provides atomic reads and writes for some set of properly aligned variables of 6 specific sizes, but not necessarily for all the variable types supported by the OpenMP 7 API. C/C++ Example A.22.2c 8 9 int atomic read(const int \*p) 10 11 int value; 12 /\* Guarantee that the entire value of \*p is read atomically. No part of 13 \* \*p can change during the read operation. 14 \*/ 15 #pragma omp atomic read 16 value = \*p; 17 return value; 18 } 19 20 void atomic\_write(int \*p, int value) 21 22 /\* Guarantee that value is stored atomically into \*p. No part of \*p can change 23 \* until after the entire write operation is completed. 24 \*/ 25 #pragma omp atomic write 26 \*p = value; 27 } C/C++ Fortran Example A.22.2f 28 29 function atomic read(p) 30 integer :: atomic read 31 integer, intent(in) :: p 32 ! Guarantee that the entire value of p is read atomically. No part of 33 ! p can change during the read operation. 34 35 !\$omp atomic read 36 atomic read = p 37 return 38 end function atomic read 39 40 subroutine atomic write(p, value) 41 integer, intent(out) :: p

```
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30
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32
33
34
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36
37
38
39
40
41
42
```

```
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 A.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();
```

```
1
                    #pragma omp flush
2
                        // Release the lock
3
                        fetch and add(&lock->turn);
4
                    }
                                                       C/C++ -
                                                      Fortran -
                   Example A.22.3f
5
6
                    function fetch and add(p)
7
                           integer:: fetch and add
8
                           integer, intent(inout) :: p
9
10
                    ! Atomically read the value of p and then increment it. The previous value is
11
                    ! returned. This can be used to implement a simple lock as shown below.
12
13
                    !$omp atomic capture
14
                           fetch and add = p
15
                           p = p + 1
16
                    !$omp end atomic
17
                           end function fetch and add
18
19
                    ! Use fetch and add to implement a lock
20
                           module m
21
                           interface
22
                             function fetch and add(p)
23
                               integer :: fetch and add
24
                               integer, intent(inout) :: p
25
                             end function
26
                             function atomic read(p)
27
                               integer :: atomic read
28
                               integer, intent(in) :: p
29
                             end function
30
                           end interface
31
                           type locktype
32
                              integer ticketnumber
33
                              integer turn
34
                           end type
35
                           contains
36
                           subroutine do locked work(lock)
37
                           type(locktype), intent(inout) :: lock
38
                           integer myturn
39
                           integer junk
40
                    ! obtain the lock
41
                            myturn = fetch and add(lock%ticketnumber)
42
                            do while (atomic read(lock%turn) .ne. myturn)
43
                              continue
44
                            enddo
45
46
                    ! Do some work. The flush is needed to ensure visibility of variables
47
                    ! not involved in atomic directives
```

```
1
                    !$omp flush
 2
                           call work
3
                    !$omp flush
4
                    ! Release the lock
6
                           junk = fetch and add(lock%turn)
7
                           end subroutine
8
                           end module
                                                       Fortran
9
10
```

### A.23 Restrictions on the atomic Construct

The following non-conforming examples illustrate the restrictions on the **atomic** construct given in Section 2.12.6 on page 103.

```
C/C++
Example A.23.1c
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 */
  }
}
                                - Fortran ·
Example A.23.1f
      SUBROUTINE ATOMIC WRONG()
        INTEGER:: I
```

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```
1
                            REAL:: R
2
                            EQUIVALENCE (I,R)
 3
4
                    !$OMP
                            PARALLEL
5
                    !$OMP
                              ATOMIC UPDATE
6
                                I = I + 1
7
                    ! SOMP
                              ATOMIC UPDATE
8
                                R = R + 1.0
9
                    ! incorrect because I and R reference the same location
10
                    ! but have different types
11
                            END PARALLEL
12
                          END SUBROUTINE ATOMIC WRONG

    Fortran -

                                                       C/C++ -
                    Example A.23.2c
13
14
                    void atomic_wrong2 ()
15
16
                     int x;
17
                     int *i;
18
                     float *r;
19
20
                     i = &x;
21
                     r = (float *)&x;
22
23
                    #pragma omp parallel
24
25
                    #pragma omp atomic update
26
                        *i += 1;
27
28
                    #pragma omp atomic update
29
                        *r += 1.0;
30
31
                    /* Incorrect because the atomic constructs reference the same location
32
                       through incompatible types */
33
34
35
                                                       C/C++ -
```

-	_			
-	- ^	rt	ra	٠
	٠.,		1 1	

1 2	The following example is non-conforming because ${\tt I}$ and ${\tt R}$ reference the same location but have different types.
3	Example A.23.2f
4 5 6	SUBROUTINE SUB() COMMON /BLK/ R REAL R
7 8 9	! \$OMP ATOMIC UPDATE  R = R + 1.0

```
!$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

1 Although the following example might work on some implementations, this is also nonconforming:

#### Example A.23.3f

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```
SUBROUTINE ATOMIC WRONG3
        INTEGER:: I
        REAL:: R
        EQUIVALENCE (I,R)
! SOMP
        PARALLEL
!$OMP
          ATOMIC UPDATE
            I = I + 1
! incorrect because I and R reference the same location
! but have different types
!$OMP
       END PARALLEL
! SOMP
       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

### A.24 The flush Construct without a List

The following example (for Section 2.12.7 on page 110) distinguishes the shared variables affected by a **flush** construct with no list from the shared objects that are not affected:

C/C++

```
28 Example A.24.1c
```

```
29
                    int x, *p = &x;
30
31
                    void f1(int *q)
32
33
                      *q = 1;
34
                      #pragma omp flush
35
                      /* x, p, and *g are flushed */
36
                      /* because they are shared and accessible */
37
                      /* q is not flushed because it is not shared. */
```

```
1
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33
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35
36
37
38
39
40
41
42
```

```
}
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++

```
Example A.24.1f
 1
 2
 3
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 6
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46
```

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49

```
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
!$OMP
       BARRIER
          ! a barrier implies a flush
          ! X, P and Q are flushed
          ! because they are shared and accessible
     END SUBROUTINE F2
     INTEGER FUNCTION G(N)
        COMMON /DATA/ X, P
        INTEGER, TARGET :: X
        INTEGER, POINTER :: P
        INTEGER N
        INTEGER I, J, SUM
        I = 1
        SUM = 0
!$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 -

# A.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. See Section 2.12.3 on page 99, Section 2.12.7 on page 110, Section 2.12.4 on page 101, and Section 2.11.2 on page 92.

· C/C++

#### Example A.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)
1
2
                      #pragma omp taskwait
3
                    /* incorrect as taskwait cannot be immediate substatement
4
                       of if statement */
5
6
                    }
                                                       C/C++ -
7
                    The following example is non-conforming, because the flush, barrier, taskwait,
8
9
                    and taskyield directives are stand-alone directives and cannot be the action
                    statement of an if statement or a labeled branch target.
10

    Fortran —

                    Example A.25.1f
11
12
                    SUBROUTINE STANDALONE WRONG()
13
                      INTEGER A
14
15
                      ! the FLUSH directive must not be the action statement
16
                      ! in an IF statement
17
                      IF (A .NE. 0) !$OMP FLUSH(A)
18
19
                      ! the BARRIER directive must not be the action statement
20
                      ! in an IF statement
21
                      IF (A .NE. 0) !$OMP BARRIER
22
23
                      ! the TASKWAIT directive must not be the action statement
24
                      ! in an IF statement
25
                      IF (A .NE. 0) !$OMP TASKWAIT
26
27
                      ! the TASKYIELD directive must not be the action statement
28
                      ! in an IF statement
29
                      IF (A .NE. 0) !$OMP TASKYIELD
30
31
                      GOTO 100
32
33
                      ! the FLUSH directive must not be a labeled branch target
34
                      ! statement
35
                      100 !$OMP FLUSH(A)
36
                      GOTO 200
37
38
                      ! the BARRIER directive must not be a labeled branch target
39
                      ! statement
40
                      200 !$OMP BARRIER
41
                      GOTO 300
42
43
                      ! the TASKWAIT directive must not be a labeled branch target
44
                      ! statement
```

300 !\$OMP TASKWAIT

```
1
                      GOTO 400
 2
3
                      ! the TASKYIELD directive must not be a labeled branch target
4
                      ! statement
                      400 !$OMP TASKYIELD
6
                    END SUBROUTINE
                                                       Fortran
                    The following version of the above example is conforming because the flush,
9
                    barrier, taskwait, and taskyield directives are enclosed in a compound
10
                    statement.
                                                        C/C++
                    Example A.25.2c
11
12
                    void standalone ok()
13
                      int a = 1;
15
                      #pragma omp parallel
16
17
18
                         if (a != 0) {
19
                      #pragma omp flush(a)
20
                         }
21
                         if (a != 0) {
22
                      #pragma omp barrier
23
24
                         if (a != 0) {
25
                      #pragma omp taskwait
26
                         }
27
                      if (a != 0) {
28
                      #pragma omp taskyield
29
                          }
30
                      }
                    }
31
                                                        C/C++
                    The following example is conforming because the flush, barrier, taskwait, and
32
                    taskyield directives are enclosed in an if construct or follow the labeled branch
33
34
                    target.
                                                        Fortran
                    Example A.25.2f
35
36
                    SUBROUTINE STANDALONE OK()
37
                      INTEGER A
38
                      A = 1
```

```
1
                      IF (A .NE. 0) THEN
 2
                         !$OMP FLUSH(A)
 3
                      ENDIF
 4
                      IF (A .NE. 0) THEN
 5
                         !$OMP BARRIER
 6
                      ENDIF
 7
                      IF (A .NE. 0) THEN
8
                         !$OMP TASKWAIT
9
                      ENDIF
10
                      IF (A .NE. 0) THEN
                         !$OMP TASKYIELD
11
12
                      ENDIF
13
                      GOTO 100
14
                      100 CONTINUE
15
                      !$OMP FLUSH(A)
16
                      GOTO 200
17
                      200 CONTINUE
18
                      !SOMP BARRIER
19
                      GOTO 300
20
                      300 CONTINUE
21
                      !$OMP TASKWAIT
22
                      GOTO 400
23
                      400 CONTINUE
24
                       !$OMP TASKYIELD
25
                    END SUBROUTINE
```

Fortran -

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# 27 A.26 The ordered Clause and the ordered Construct

Ordered constructs (Section 2.12.8 on page 114) 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 A.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;
}</pre>
```

	Fortran —
1	Example A.26.1f
2	SUBROUTINE WORK(K)
3	INTEGER k
4	
5	!\$OMP ORDERED
6	WRITE(*,*) K
7	!\$OMP END ORDERED
8	
9	END SUBROUTINE WORK
10	
11	SUBROUTINE SUB(LB, UB, STRIDE)
12	INTEGER LB, UB, STRIDE
13	INTEGER I
14	
15	!\$OMP PARALLEL DO ORDERED SCHEDULE(DYNAMIC)
16	DO I=LB,UB,STRIDE
17	CALL WORK(I)
18	END DO
19	!\$OMP END PARALLEL DO
20	
21	END SUBROUTINE SUB
22	
23	PROGRAM ORDERED_EXAMPLE
24	CALL SUB(1,100,5)
25	END PROGRAM ORDERED_EXAMPLE

27 28

29

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:

```
C/C++
Example A.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);
  }
}
                                  C/C++
                                  Fortran
Example A.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
                                 Fortran •
```

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37

2 iteration will execute only one ordered region: C/C++ Example A.26.3c 3 4 void work(int i) {} 5 void ordered good(int n) 6 7 int i: 8 9 #pragma omp for ordered 10 for (i=0; i<n; i++) { 11 if (i <= 10) { 12 #pragma omp ordered 13 work(i); 14 15 16 if (i > 10) { 17 #pragma omp ordered 18 work(i+1); 19 20 } 21 C/C++ Fortran Example A.26.3f 22 23 SUBROUTINE ORDERED GOOD (N) 24 INTEGER N 25 26 ! SOMP DO ORDERED 27 DO I = 1,N28 IF (I <= 10) THEN 29 !\$OMP ORDERED 30 CALL WORK(I) 31 !\$OMP END ORDERED 32 ENDIF 33 34 IF (I > 10) THEN 35 ORDERED !\$OMP 36 CALL WORK (I+1) 37 !\$OMP END ORDERED 38 ENDIF 39 ENDDO 40 END SUBROUTINE ORDERED GOOD Fortran -

The following is a conforming example with more than one **ordered** construct. Each

## A.27 The threadprivate Directive

The following examples demonstrate how to use the **threadprivate** directive (Section 2.14.2 on page 126) to give each thread a separate counter.

```
C/C++
Example A.27.1c
int counter = 0;
#pragma omp threadprivate(counter)
int increment counter()
  counter++;
  return(counter);
                                 C/C++ -
                               - Fortran -
Example A.27.1f
      INTEGER FUNCTION INCREMENT COUNTER()
        COMMON/INC COMMON/COUNTER
! SOMP
       THREADPRIVATE(/INC COMMON/)
        COUNTER = COUNTER +1
        INCREMENT COUNTER = COUNTER
      END FUNCTION INCREMENT COUNTER
                                 Fortran -
                                 C/C++ ----
The following example uses threadprivate on a static variable:
Example A.27.2c
int increment counter 2()
  static int counter = 0;
  #pragma omp threadprivate(counter)
  counter++;
  return(counter);
```

3

6

7

9 10

11

12

13

15

16

17 18

19

20 21

22

23

24

25

27

28

29

30

}

7

8

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18 19 20

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30

31

32

33 34

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45

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 A.27.3c

```
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++
```

Appendix A

Examples

#### Fortran

The following examples show non-conforming uses and correct uses of the 1 2 threadprivate directive. For more information, see Section 2.14.2 on page 126 and Section 2.14.4.1 on page 148. 3 The following example is non-conforming because the common block is not declared local to the subroutine that refers to it: Example A.27.2f MODULE INC MODULE COMMON /T/ A 9 END MODULE INC MODULE 10 11 SUBROUTINE INC\_MODULE\_WRONG() 12 USE INC MODULE 13 !\$OMP THREADPRIVATE(/T/) !non-conforming because /T/ not declared in INC MODULE WRONG 14 15 END SUBROUTINE INC MODULE WRONG 16 17 The following example is also non-conforming because the common block is not 18 declared local to the subroutine that refers to it: 19 Example A.27.3f 20 SUBROUTINE INC WRONG() 21 COMMON /T/ A 22 ! SOMP THREADPRIVATE (/T/) 23 24 CONTAINS 25 SUBROUTINE INC\_WRONG\_SUB() 26 ! SOMP PARALLEL COPYIN(/T/)

!non-conforming because /T/ not declared in INC WRONG SUB

27

28

29

30

31

END PARALLEL

END SUBROUTINE INC WRONG

END SUBROUTINE INC WRONG SUB

!\$OMP

```
1
                  The following example is a correct rewrite of the previous example:
                  Example A.27.4f
 2
 3
                         SUBROUTINE INC GOOD ()
 4
                          COMMON /T/ A
 5
                  ! SOMP
                         THREADPRIVATE(/T/)
 6
 7
                          CONTAINS
8
                           SUBROUTINE INC GOOD SUB()
9
                             COMMON /T/ A
10
                  ! SOMP
                             THREADPRIVATE(/T/)
11
12
                  !$OMP
                            PARALLEL COPYIN(/T/)
13
                  !$OMP
                             END PARALLEL
14
                           END SUBROUTINE INC GOOD SUB
15
                         END SUBROUTINE INC GOOD
16
17
                  The following is an example of the use of threadprivate for local variables:
                  Example A.27.5f
18
19
                        PROGRAM INC GOOD2
20
                          INTEGER, ALLOCATABLE, SAVE :: A(:)
21
                          INTEGER, POINTER, SAVE :: PTR
22
                          INTEGER, SAVE :: I
23
                          INTEGER, TARGET :: TARG
24
                          LOGICAL :: FIRSTIN = .TRUE.
25
                  !$OMP
                          THREADPRIVATE(A, I, PTR)
26
27
                          ALLOCATE (A(3))
28
                          A = (/1,2,3/)
29
                          PTR => TARG
30
                          I = 5
31
32
                  ! SOMP
                         PARALLEL COPYIN(I, PTR)
33
                  !$OMP
                           CRITICAL
34
                              IF (FIRSTIN) THEN
35
                                TARG = 4
                                                 ! Update target of ptr
36
                               I = I + 10
37
                               IF (ALLOCATED(A)) A = A + 10
38
                               FIRSTIN = .FALSE.
39
                              END IF
40
41
                              IF (ALLOCATED(A)) THEN
42
                               PRINT *, 'a = ', A
43
                              ELSE
```

```
▼------ Fortran (cont.) ------
1
                                 PRINT *, 'A is not allocated'
2
                               END IF
3
4
                               PRINT *, 'ptr = ', PTR
5
                               PRINT *, 'i = ', I
6
                               PRINT *
7
8
                   ! SOMP
                           END CRITICAL
9
                   !$OMP END PARALLEL
10
                         END PROGRAM INC_GOOD2
11
12
                   The above program, if executed by two threads, will print one of the following two sets
13
                   of output:
14
15
                   a = 11 12 13
16
                   ptr = 4
17
                   i = 15
18
19
                   A is not allocated
20
                   ptr = 4
21
                  i = 5
22
                   or
23
24
                   A is not allocated
25
                   ptr = 4
26
                   i = 15
27
28
                   a = 1 2 3
29
                   ptr = 4
30
                   i = 5
31
32
                   The following is an example of the use of threadprivate for module variables:
                   Example A.27.6f
33
34
                         MODULE INC MODULE GOOD3
35
                           REAL, POINTER :: WORK(:)
36
                           SAVE WORK
37
                   ! $OMP THREADPRIVATE (WORK)
38
                         END MODULE INC MODULE GOOD3
39
40
                         SUBROUTINE SUB1(N)
41
                        USE INC_MODULE_GOOD3
42
                   !$OMP PARALLEL PRIVATE (THE SUM)
43
                           ALLOCATE (WORK (N))
```

```
1
                             CALL SUB2 (THE SUM)
 2
                            WRITE(*,*)THE SUM
 3
                     !$OMP
                             END PARALLEL
 4
                           END SUBROUTINE SUB1
 5
 6
                           SUBROUTINE SUB2 (THE SUM)
 7
                             USE INC MODULE GOOD3
 8
                             WORK(:) = 10
 9
                             THE SUM=SUM (WORK)
10
                           END SUBROUTINE SUB2
11
12
                           PROGRAM INC GOOD3
13
                             N = 10
14
                             CALL SUB1(N)
15
                           END PROGRAM INC GOOD3
                                                         Fortran
                                                         C/C++
                    The following example illustrates initialization of threadprivate variables for
16
                    class-type T. t1 is default constructed, t2 is constructed taking a constructor accepting
17
                    one argument of integer type, t3 is copy constructed with argument f():
18
                    Example A.27.4c
19
20
                    static T t1;
21
                    #pragma omp threadprivate(t1)
22
                     static T t2( 23 );
23
                     #pragma omp threadprivate(t2)
24
                     static T t3 = f();
25
                     #pragma omp threadprivate(t3)
26
                    The following example illustrates the use of threadprivate for static class
27
                    members. The threadprivate directive for a static class member must be placed
28
                    inside the class definition.
29
                    Example A.27.5c
30
31
                    class T {
32
                     public:
33
                       static int i;
34
                    #pragma omp threadprivate(i)
35
                     };
36
```

C/C++

C/C++

# A.28 Parallel Random Access Iterator Loop

The following example shows a parallel random access iterator loop.

```
Example A.28.1c
#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 //
    }
}</pre>
```

2	<b>A.29</b>	Fortran Restrictions on shared and	
3		private Clauses with Common Blocks	
4 5 6 7		When a named common block is specified in a <b>private</b> , <b>firstprivate</b> , or <b>lastprivate</b> 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. For more information, see Section 2.14.3 on page 130.	
8		The following example is conforming:	
9		Example A.29.1f	
10 11 12 13 14 15 16 17 18 19 20 21 22 23		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:	
24		Example A.29.2f	
25 26 27 28 29 30		SUBROUTINE COMMON_GOOD2()  COMMON /C/ X,Y  REAL X, Y  INTEGER I	
31 32 33 34 35 36 37		!\$OMP PARALLEL !\$OMP DO PRIVATE(/C/) DO I=1,1000 ! do work here ENDDO !\$OMP END DO !	

```
▼------Fortran (cont.) --------
 1
                   !$OMP
                           DO PRIVATE(X)
2
                            DO I=1,1000
3
                             ! do work here
                            ENDDO
5
                   !$OMP END DO
6
                   !$OMP END PARALLEL
7
                        END SUBROUTINE COMMON GOOD2
 9
                   The following example is conforming:
                   Example A.29.3f
10
11
                         SUBROUTINE COMMON GOOD3()
12
                          COMMON /C/ X,Y
13
                   !$OMP PARALLEL PRIVATE (/C/)
15
                           ! do work here
16
                   !$OMP END PARALLEL
17
18
                   !$OMP PARALLEL SHARED (/C/)
                           ! do work here
20
                   !$OMP END PARALLEL
21
                        END SUBROUTINE COMMON GOOD3
22
23
                   The following example is non-conforming because \mathbf{x} is a constituent element of \mathbf{c}:
                   Example A.29.4f
24
25
                         SUBROUTINE COMMON WRONG()
26
                          COMMON /C/ X,Y
27
                   ! Incorrect because X is a constituent element of C
28
                   !SOMP PARALLEL PRIVATE(/C/), SHARED(X)
29
                           ! do work here
30
                   !$OMP END PARALLEL
31
                       END SUBROUTINE COMMON WRONG
32
33
                   The following example is non-conforming because a common block may not be
34
                   declared both shared and private:
                   Example A.29.5f
35
36
                         SUBROUTINE COMMON WRONG2()
37
                          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
6
TEND SUBROUTINE COMMON_WRONG2
```

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Fortran -

## A.30 The default (none) Clause

The following example distinguishes the variables that are affected by the **default (none)** clause from those that are not. For more information on the **default** clause, see Section 2.14.3.1 on page 131.

C/C++

```
Example A.30.1c
12
13
                   #include <omp.h>
14
                   int x, y, z[1000];
15
                   #pragma omp threadprivate(x)
16
17
                   void default none(int a) {
18
                     const int c = 1;
19
                     int i = 0;
20
21
                     #pragma omp parallel default(none) private(a) shared(z)
22
23
                        int j = omp get num threads();
24
                             /* O.K. - j is declared within parallel region */
25
                                     /* O.K. - a is listed in private clause */
                                    /*
26
                                             - z is listed in shared clause */
27
                                     /* O.K. - x is threadprivate */
28
                                             - c has const-qualified type */
29
                        z[i] = y;
                                    /* Error - cannot reference i or y here */
30
31
                     #pragma omp for firstprivate(y)
32
                            /* Error - Cannot reference y in the firstprivate clause */
33
                        for (i=0; i<10; i++) {
34
                            z[i] = i; /* O.K. - i is the loop iteration variable */
35
36
37
                        z[i] = y;
                                   /* Error - cannot reference i or y here */
38
```

}

C/C++

#### Example A.30.1f

```
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
                   ! 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
```

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# 2 A.31 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 (see Section 2.14.3.2 on page 133). 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 A.31.1f

```
SUBROUTINE SHARED RACE
  INCLUDE "omp lib.h"
                            ! or USE OMP LIB
  REAL A(20)
  INTEGER MYTHREAD
!$OMP PARALLEL SHARED(A) PRIVATE(MYTHREAD)
  MYTHREAD = OMP GET THREAD NUM()
  IF (MYTHREAD .EQ. 0) THEN
     CALL SUB(A(1:10)) ! compiler may introduce writes to A(6:10)
     A(6:10) = 12
  ENDIF
!$OMP END PARALLEL
END SUBROUTINE SHARED RACE
SUBROUTINE SUB(X)
  REAL X(*)
  X(1:5) = 4
END SUBROUTINE SUB
```

Fortran -

## A.32 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. For more information on the **private** clause, see Section 2.14.3.3 on page 134.

```
C/C++
Example A.32.1c
#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 A.32.1f
      PROGRAM PRIV EXAMPLE
        INTEGER I, J
        I = 1
```

J = 2

3

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9 10

11 12

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20 21

22 23

24

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29 30

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35 36

```
1
 2
 3
 4
 5
 6
 7
 8
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15
16
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21
22
23
24
25
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27
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```

#### - 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 A.32.2c

```
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" */
    }
}</pre>
```

C/C++

#### Fortran

#### Example A.32.2f

```
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
          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.

2

3

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8

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```
C/C++
                   Example A.32.3c
 1
 2
                    #include <assert.h>
 3
                   void priv_example3()
4
5
                      int i, a;
6
7
                      #pragma omp parallel private(a)
8
9
10
                        #pragma omp parallel for private(a)
11
                          for (i=0; i<10; i++)
12
13
                           a = 2;
14
15
                        assert(a == 1);
16
17
                   }
                                                       C/C++
                                                     Fortran
                   Example A.32.3f
18
19
                          SUBROUTINE PRIV EXAMPLE3()
20
                            INTEGER I, A
21
22
                    !$OMP
                            PARALLEL PRIVATE(A)
23
24
                              PARALLEL DO PRIVATE(A)
                    !$OMP
25
                              DO I = 1, 10
26
                                A = 2
                              END DO
27
28
                    !$OMP
                              END PARALLEL DO
29
                            PRINT *, A ! Outer A still has value 1
30
                    !$OMP
                            END PARALLEL
31
                          END SUBROUTINE PRIV_EXAMPLE3
```

Fortran -

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23

24 25

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34 35

36

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# A.33 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 (see Section 2.14.3.3 on page 134).

! X is undefined ! Y is undefined

#### Example A.33.1f

```
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 A.33.2f
      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

PRINT \*,Y

END SUBROUTINE SUB

END PROGRAM PRIV RESTRICT2

```
▼------Fortran (cont.) -------
                  Example A.33.3f
 1
 2
                         PROGRAM PRIV RESTRICT3
 3
                         EQUIVALENCE (X,Y)
 4
                         X = 1.0
5
6
                  ! $OMP PARALLEL PRIVATE(X)
7
                                                    ! Y is undefined
                           PRINT *,Y
8
                           Y = 10
9
                           PRINT *,X
                                                  ! X is undefined
10
                  !$OMP END PARALLEL
11
                       END PROGRAM PRIV RESTRICT3
12
                  Example A.33.4f
13
14
                       PROGRAM PRIV RESTRICT4
15
                         INTEGER I, J
16
                         INTEGER A(100), B(100)
17
                         EQUIVALENCE (A(51), B(1))
18
19
                  !$OMP PARALLEL DO DEFAULT(PRIVATE) PRIVATE(I,J) LASTPRIVATE(A)
20
                           DO I=1,100
21
                              DO J=1,100
22
                                B(J) = J - 1
23
                              ENDDO
24
25
                              DO J=1,100
26
                               A(J) = J! B becomes undefined at this point
27
                              ENDDO
28
29
                              DO J=1,50
30
                                B(J) = B(J) + 1 ! B is undefined
31
                                         ! A becomes undefined at this point
32
                              ENDDO
33
                           ENDDO
34
                  !$OMP END PARALLEL DO ! The LASTPRIVATE write for A has
35
                                            ! undefined results
36
37
                          PRINT *, B ! B is undefined since the LASTPRIVATE
38
                                        ! write of A was not defined
39
                        END PROGRAM PRIV RESTRICT4
40
41
```

#### Example A.33.5f

1

```
2
3
                          SUBROUTINE SUB1(X)
4
                            DIMENSION X(10)
5
                            ! This use of X does not conform to the
7
                            ! specification. It would be legal Fortran 90,
8
                            ! but the OpenMP private directive allows the
9
                            ! compiler to break the sequence association that
10
                            ! A had with the rest of the common block.
11
12
                            FORALL (I = 1:10) X(I) = I
13
                          END SUBROUTINE SUB1
14
15
                          PROGRAM PRIV RESTRICT5
16
                            COMMON /BLOCK5/ A
17
18
                            DIMENSION B(10)
19
                            EQUIVALENCE (A,B(1))
20
21
                            ! the common block has to be at least 10 words
22
23
24
                    !$OMP PARALLEL PRIVATE(/BLOCK5/)
25
26
                              ! Without the private clause,
27
                              ! we would be passing a member of a sequence
28
                              ! that is at least ten elements long.
29
                              ! With the private clause, A may no longer be
30
                              ! sequence-associated.
31
32
                              CALL SUB1(A)
33
                    !$OMP
                              MASTER
34
                                PRINT *, A
35
                    !$OMP
                              END MASTER
36
37
                    !SOMP END PARALLEL
38
                          END PROGRAM PRIV RESTRICT5
                                                     - Fortran -
```

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#### C/C++

# A.34 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 (Section 2.14.3.4 on page 137). 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 A.34.1c
```

```
2
                    #include <assert.h>
3
4
                    int A[2][2] = \{1, 2, 3, 4\};
5
6
                    void f(int n, int B[n][n], int C[])
7
8
                      int D[2][2] = \{1, 2, 3, 4\};
9
                      int E[n][n];
10
11
                      assert(n >= 2);
12
                      E[1][1] = 4;
13
14
                      #pragma omp parallel firstprivate(B, C, D, E)
15
16
                        assert(sizeof(B) == sizeof(int (*)[n]));
17
                        assert(sizeof(C) == sizeof(int*));
18
                        assert(sizeof(D) == 4 * sizeof(int));
19
                        assert(sizeof(E) == n * n * sizeof(int));
20
21
                        /* Private B and C have values of original B and C. */
22
                        assert(&B[1][1] == &A[1][1]);
23
                        assert(&C[3] == &A[1][1]);
24
                        assert(D[1][1] == 4);
25
                        assert(E[1][1] == 4);
26
27
                    }
28
29
                    int main() {
30
                      f(2, A, A[0]);
31
                      return 0;
32
                                                        C/C++
```

## A.35 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 (Section 2.14.3.5 on page 139) so that the values of the variables are the same as when the loop is executed sequentially.

34

35

```
C/C++
                   Example A.35.1c
 1
 2
                   void lastpriv (int n, float *a, float *b)
 3
4
                     int i;
5
6
                     #pragma omp parallel
7
8
                       #pragma omp for lastprivate(i)
9
                       for (i=0; i<n-1; i++)
10
                         a[i] = b[i] + b[i+1];
11
12
13
                     a[i]=b[i];
                                  /* i == n-1 here */
14
                                                  C/C++ -
                                                   Fortran
                   Example A.35.1f
15
16
                         SUBROUTINE LASTPRIV(N, A, B)
17
18
                           INTEGER N
19
                           REAL A(*), B(*)
20
                           INTEGER I
21
22
                   !$OMP PARALLEL
23
                   !$OMP DO LASTPRIVATE(I)
24
25
                           DO I=1,N-1
26
                             A(I) = B(I) + B(I+1)
27
                           ENDDO
28
29
                   !$OMP END PARALLEL
30
31
                           A(I) = B(I)
                                       ! I has the value of N here
32
33
                         END SUBROUTINE LASTPRIV
                                                   - Fortran -
```

#### A.36 The reduction Clause

The following example demonstrates the **reduction** clause (Section 2.14.3.6 on page 142); 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 A.36.1c
#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 A.36.1f
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))
```

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```
1
                            C = MIN(C, Y(I))
 2
                            IF (D < X(I)) D = X(I)
 3
                          END DO
 4
 5
                    END SUBROUTINE REDUCTION1
                                                       Fortran
                    A common implementation of the preceding example is to treat it as if it had been
 6
                    written as follows:
 7
                                                       C/C++
                    Example A.36.2c
 8
 9
                    #include <limits.h>
10
                    #include <math.h>
11
                    void reduction2(float *x, int *y, int n)
12
13
                      int i, b, b_p, c, c_p;
14
                      float a, a_p, d, d_p;
15
                      a = 0.0f;
16
                      b = 0;
17
                      c = y[0];
18
                      d = x[0];
19
                      #pragma omp parallel shared(a, b, c, d, x, y, n) \
20
                                              private(a_p, b_p, c_p, d_p)
21
                      {
22
                        ap = 0.0f;
23
                        bp = 0;
24
                        c_p = INT_MAX;
25
                        d p = -HUGE VALF;
26
                        #pragma omp for private(i)
27
                        for (i=0; i<n; i++) {
28
                          ap += x[i];
29
                          b p ^= y[i];
30
                          if (c_p > y[i]) c_p = y[i];
31
                          dp = fmaxf(dp,x[i]);
32
33
                        #pragma omp critical
34
35
                          a += a_p;
                          b ^= b p;
36
37
                          if(c>cp)c=cp;
38
                          d = fmaxf(d,d p);
39
40
                      }
41
                    }
                                                       C/C++
```

#### 1 Example A.36.2f

```
2
                      SUBROUTINE REDUCTION2 (A, B, C, D, X, Y, N)
3
                        REAL :: X(*), A, D
                        INTEGER :: Y(*), N, B, C
5
                        REAL :: A_P, D P
6
                        INTEGER :: I, B P, C P
7
                        A = 0
8
                        B = 0
9
                        C = Y(1)
10
                        D = X(1)
11
                        !$OMP PARALLEL SHARED(X, Y, A, B, C, D, N) &
12
                        !$OMP&
                                        PRIVATE(A P, B P, C P, D P)
13
                          AP = 0.0
14
                          BP=0
15
                          C P = HUGE(C P)
16
                          D P = -HUGE(D P)
17
                          !$OMP DO PRIVATE(I)
18
                          DO I=1,N
19
                            A_P = A_P + X(I)
20
                            BP = IEOR(BP, Y(I))
21
                            C_P = MIN(C_P, Y(I))
22
                            IF (D P < X(I)) D P = X(I)
23
                          END DO
24
                          !$OMP CRITICAL
25
                            A = A + A P
26
                            B = IEOR(B, B P)
27
                            C = MIN(C, C_P)
28
                            D = MAX(D, D P)
29
                          !$OMP END CRITICAL
30
                        !$OMP END PARALLEL
31
                      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.

#### Example A.36.3f

```
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
```

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```
1
                     END PROGRAM REDUCTION WRONG
 2
3
                     SUBROUTINE SUB (M, I)
4
                        M = MAX(M, I)
5
                     END SUBROUTINE SUB
6
                    The following conforming program performs the reduction using the intrinsic procedure
7
                    name MAX even though the intrinsic MAX has been renamed to REN.
8
                    Example A.36.4f
9
10
                    MODULE M
11
                       INTRINSIC MAX
12
                    END MODULE M
13
14
                    PROGRAM REDUCTION3
15
                      USE M, REN => MAX
16
                       N = 0
17
                    !$OMP PARALLEL DO REDUCTION(REN: N) ! still does MAX
18
                       DO I = 1, 100
19
                          N = MAX(N,I)
20
                       END DO
21
                    END PROGRAM REDUCTION3
22
23
                    The following conforming program performs the reduction using intrinsic procedure
                    name MAX even though the intrinsic MAX has been renamed to MIN.
24
                    Example A.36.5f
25
26
                    MODULE MOD
27
                       INTRINSIC MAX, MIN
28
                    END MODULE MOD
29
30
                    PROGRAM REDUCTION4
31
                       USE MOD, MIN=>MAX, MAX=>MIN
32
                       REAL :: R
33
                       R = -HUGE(0.0)
34
35
                    !$OMP PARALLEL DO REDUCTION(MIN: R) ! still does MAX
                       DO I = 1, 1000
36
37
                          R = MIN(R, SIN(REAL(I)))
38
                       END DO
39
                       PRINT *, R
40
                    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.

### C/C++ -Example A.36.3c #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++ -

```
Fortran
                    Example A.36.6f
 1
2
                          INTEGER A, I
3
4
                    !$OMP PARALLEL SHARED(A) PRIVATE(I)
5
6
                    !$OMP MASTER
7
                          A = 0
8
                    !$OMP END MASTER
9
10
                          ! To avoid race conditions, add a barrier here.
11
12
                    !$OMP DO REDUCTION(+:A)
13
                          DO I= 0, 9
14
                             A = A + I
15
                          END DO
16
17
                    !$OMP SINGLE
18
                          PRINT *, "Sum is ", A
19
                    !$OMP END SINGLE
20
21
                    !$OMP END PARALLEL
22
                          END
                                                       Fortran
```

# A.37 The copyin Clause

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The **copyin** clause (see Section 2.14.4.1 on page 148) 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.

#### Example A.37.1c

```
#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();
   }
}</pre>
```

#### **Fortran**

#### Example A.37.1f

1

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36

```
2
                           MODULE M
 3
                             REAL, POINTER, SAVE :: WORK(:)
 4
                             INTEGER :: SIZE
 5
                             REAL :: TOL
 6
                     !$OMP
                             THREADPRIVATE (WORK, SIZE, TOL)
 7
                           END MODULE M
 8
 9
                           SUBROUTINE COPYIN EXAMPLE ( T, N )
10
                             USE M
11
                             REAL :: T
12
                             INTEGER :: N
13
                             TOL = T
14
15
                     !$OMP
                             PARALLEL COPYIN(TOL, SIZE)
16
                             CALL BUILD
17
                     !$OMP
                             END PARALLEL
18
                           END SUBROUTINE COPYIN_EXAMPLE
19
20
                           SUBROUTINE BUILD
21
                             USE M
22
                             ALLOCATE (WORK (SIZE))
23
                             WORK = TOL
24
                           END SUBROUTINE BUILD
```

Fortran

### A.38 The copyprivate Clause

The **copyprivate** clause (see Section 2.14.4.2 on page 149) 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 A.38.1c
 1
2
                    #include <stdio.h>
3
4
5
6
7
8
                    float x, y;
                    #pragma omp threadprivate(x, y)
                    void init(float a, float b ) {
                        #pragma omp single copyprivate(a,b,x,y)
9
                            scanf("%f %f %f %f", &a, &b, &x, &y);
10
11
                    }
                                                       C/C++ -
                                                     - Fortran -
                   Example A.38.1f
12
13
                          SUBROUTINE INIT(A,B)
14
                          REAL A, B
                            COMMON /XY/ X,Y
16
                    !$OMP
                            THREADPRIVATE (/XY/)
17
18
                    !$OMP
                            SINGLE
19
                              READ (11) A,B,X,Y
20
                    !$OMP
                           END SINGLE COPYPRIVATE (A,B,/XY/)
21
22
                          END SUBROUTINE INIT
                                                      Fortran
```

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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 A.38.2c
```

```
#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 A.38.2f

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```
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
! SOMP
        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.

# Example A.38.3c

```
#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;
}
```

Example A.38.3f 1 2 FUNCTION NEW LOCK() 3 USE OMP LIB ! or INCLUDE "omp lib.h" 4 INTEGER (OMP LOCK KIND), POINTER :: NEW LOCK 5 6 !\$OMP SINGLE 7 ALLOCATE (NEW LOCK) 8 CALL OMP INIT LOCK (NEW LOCK) 9 END SINGLE COPYPRIVATE (NEW LOCK) !\$OMP 10 END FUNCTION NEW LOCK 11 12 Note that the effect of the copyprivate clause on a variable with the allocatable 13 attribute is different than on a variable with the **pointer** attribute. The value of **A** is 14 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 15 16 parallel region. Example A.38.4f 17 18 SUBROUTINE S(N) 19 INTEGER N 20 21 REAL, DIMENSION(:), ALLOCATABLE :: A 22 REAL, DIMENSION(:), POINTER :: B 23 24 ALLOCATE (A(N)) 25 !\$OMP SINGLE 26 ALLOCATE (B(N)) 27 READ (11) A,B 28 END SINGLE COPYPRIVATE(A,B) !\$OMP 29 ! Variable A is private and is 30 ! assigned the same value in each thread 31 ! Variable B is shared 32 33 !\$OMP BARRIER 34 SINGLE !\$OMP 35 DEALLOCATE (B) 36 ! SOMP END SINGLE NOWAIT

END SUBROUTINE S

37

Fortran

## A.39 Nested Loop Constructs

The following example of loop construct nesting (see Section 2.16 on page 158) is conforming because the inner and outer loop regions bind to different **parallel** regions:

```
C/C++
Example A.39.1c
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++
```

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#### Fortran -

#### Example A.39.1f

1

```
2
                          SUBROUTINE WORK(I, J)
 3
                          INTEGER I, J
4
                          END SUBROUTINE WORK
5
                          SUBROUTINE GOOD NESTING(N)
7
                          INTEGER N
8
9
                             INTEGER I
10
                            PARALLEL DEFAULT (SHARED)
                    !$OMP
11
                    !$OMP
12
                               DO I = 1, N
13
                    !$OMP
                                 PARALLEL SHARED (I, N)
14
                    !$OMP
15
                                   DO J = 1, N
16
                                     CALL WORK(I,J)
17
                                   END DO
18
                    !$OMP
                                 END PARALLEL
19
                               END DO
20
                    !$OMP
                            END PARALLEL
21
                          END SUBROUTINE GOOD_NESTING
```

- Fortran -

```
1
```

# Example A.39.2c

```
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)</pre>
```

#pragma omp for

work1(i, n);

}

}

for (i=0; i<n; i++)

C/C++

```
Fortran
                    Example A.39.2f
 1
2
                          SUBROUTINE WORK(I, J)
3
                          INTEGER I, J
4
                          END SUBROUTINE WORK
5
6
                          SUBROUTINE WORK1(I, N)
7
                          INTEGER J
8
                    !$OMP PARALLEL DEFAULT(SHARED)
9
                    !$OMP DO
10
                            DO J = 1, N
11
                              CALL WORK(I,J)
12
                            END DO
13
                    !$OMP END PARALLEL
14
                          END SUBROUTINE WORK1
15
16
                          SUBROUTINE GOOD NESTING2 (N)
17
                          INTEGER N
18
                    !$OMP PARALLEL DEFAULT (SHARED)
19
                    !$OMP DO
20
                          DO I = 1, N
21
                             CALL WORK1(I, N)
22
23
```

#### A<sub>-</sub>40 **Restrictions on Nesting of Regions**

END SUBROUTINE GOOD NESTING2

!\$OMP END PARALLEL

24

26

27

The examples in this section illustrate the region nesting rules. For more information on region nesting, see Section 2.16 on page 158.

Fortran -

```
1
                    The following example is non-conforming because the inner and outer loop regions are
 2
                    closely nested:
                                                        C/C++
                    Example A.40.1c
 3
4
                    void work(int i, int j) {}
5
6
                    void wrong1(int n)
7
8
                      #pragma omp parallel default(shared)
9
10
                        int i, j;
11
                        #pragma omp for
12
                        for (i=0; i<n; i++) {
13
                            /* incorrect nesting of loop regions */
14
                           #pragma omp for
15
                             for (j=0; j<n; j++)
16
                               work(i, j);
17
18
                      }
19
                                                        C/C++ -
                                                        Fortran
                    Example A.40.1f
20
21
                          SUBROUTINE WORK(I, J)
22
                          INTEGER I, J
23
                          END SUBROUTINE WORK
24
25
                          SUBROUTINE WRONG1(N)
26
                          INTEGER N
27
28
                            INTEGER I,J
29
                            PARALLEL DEFAULT (SHARED)
                    !$OMP
30
                    !$OMP
31
                              DO I = 1, N
32
                    !$OMP
                                                 ! incorrect nesting of loop regions
33
                                 DO J = 1, N
34
                                   CALL WORK(I,J)
35
                                 END DO
36
                              END DO
37
                            END PARALLEL
                    ! SOMP
38
                          END SUBROUTINE WRONG1
                                                       Fortran -
```

The following orphaned version of the preceding example is also non-conforming: 1 C/C++ Example A.40.2c 2 3 void work(int i, int j) {} 4 void work1(int i, int n) 5 6 int j; 7 /\* incorrect nesting of loop regions \*/ 8 #pragma omp for 9 for (j=0; j< n; j++)10 work(i, j); 11 } 12 13 void wrong2(int n) 14 15 #pragma omp parallel default(shared) 16 17 int i; 18 #pragma omp for 19 for (i=0; i<n; i++) 20 work1(i, n); 21 } 22 } C/C++ Fortran Example A.40.2f 23 24 SUBROUTINE WORK1(I,N) 25 INTEGER I, N 26 INTEGER J 27 ! SOMP ! incorrect nesting of loop regions 28 DO J = 1, N 29 CALL WORK(I,J) 30 END DO 31 END SUBROUTINE WORK1 32 SUBROUTINE WRONG2 (N) 33 INTEGER N 34 INTEGER I 35 !\$OMP PARALLEL DEFAULT (SHARED) 36 !\$OMP 37 DO I = 1, N38 CALL WORK1(I,N) 39 END DO 40 !\$OMP END PARALLEL 41 END SUBROUTINE WRONG2 Fortran

```
1
                    The following example is non-conforming because the loop and single regions are
 2
                    closely nested:
                                                        C/C++
                    Example A.40.3c
 3
4
                    void work(int i, int j) {}
5
                    void wrong3(int n)
6
7
                      #pragma omp parallel default(shared)
8
9
                        int i;
10
                        #pragma omp for
11
                          for (i=0; i<n; i++) {
12
                    /* incorrect nesting of regions */
13
                            #pragma omp single
14
                              work(i, 0);
15
                          }
16
                      }
17
                    }
                                                        C/C++
                                                       Fortran
                    Example A.40.3f
18
19
20
                          SUBROUTINE WRONG3 (N)
21
                          INTEGER N
22
23
                            INTEGER I
24
                    !$OMP
                            PARALLEL DEFAULT (SHARED)
25
                    !$OMP
                              DO
26
                              DO I = 1, N
27
                    !$OMP
                                SINGLE
                                                    ! incorrect nesting of regions
28
                                   CALL WORK(I, 1)
29
                    !$OMP
                                END SINGLE
30
                              END DO
31
                    ! SOMP
                            END PARALLEL
32
                          END SUBROUTINE WRONG3
                                                       Fortran
```

```
The following example is non-conforming because a barrier region cannot be closely
2
                    nested inside a loop region:
                                                        C/C++
                    Example A.40.4c
3
4
                    void work(int i, int j) {}
5
                    void wrong4(int n)
6
7
8
                      #pragma omp parallel default(shared)
9
10
                        int i;
11
                        #pragma omp for
12
                          for (i=0; i<n; i++) {
13
                            work(i, 0);
14
                    /* incorrect nesting of barrier region in a loop region */
15
                            #pragma omp barrier
16
                            work(i, 1);
17
                          }
18
                      }
19
                                                       C/C++ =
                                                       Fortran
                    Example A.40.4f
20
21
22
                          SUBROUTINE WRONG4 (N)
23
                          INTEGER N
24
25
                            INTEGER I
26
                    ! SOMP
                            PARALLEL DEFAULT (SHARED)
27
                    !$OMP
                              DO
28
                              DO I = 1, N
29
                                CALL WORK(I, 1)
30
                    ! incorrect nesting of barrier region in a loop region
31
                    !$OMP
                                BARRIER
32
                                CALL WORK(I, 2)
33
                              END DO
34
                            END PARALLEL
                    ! SOMP
35
                          END SUBROUTINE WRONG4
                                                       Fortran -
```

```
1
 2
 3
 4
 5
 6
 7
8
9
10
11
12
13
14
15
16
17
                         }
18
                       }
19
20
21
                              INTEGER N
22
23
                       ! SOMP
24
                       !$OMP
25
26
27
                       !$OMP
28
29
                       ! SOMP
```

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:

```
C/C++
Example A.40.5c
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);
                                   C/C++
                                  Fortran
Example A.40.5f
      SUBROUTINE WRONG5 (N)
        PARALLEL DEFAULT (SHARED)
          CRITICAL
            CALL WORK (N, 1)
! incorrect nesting of barrier region in a critical region
            BARRIER
            CALL WORK (N, 2)
          END CRITICAL
!$OMP
        END PARALLEL
      END SUBROUTINE WRONG5
```

Fortran •

30

```
The following example is non-conforming because the barrier region cannot be
 1
 2
                    closely nested inside the single region. If this were permitted, it would result in
 3
                    deadlock due to the fact that only one thread executes the single region:
                                                         C/C++
                    Example A.40.6c
 4
 5
                    void work(int i, int j) {}
 6
                    void wrong6(int n)
 7
8
                      #pragma omp parallel
 9
10
                         #pragma omp single
11
12
                           work(n, 0);
13
                    /* incorrect nesting of barrier region in a single region */
14
                           #pragma omp barrier
15
                           work(n, 1);
16
                         }
17
                      }
18
                    }
                                                         C/C++
                                                        Fortran
                    Example A.40.6f
19
20
                          SUBROUTINE WRONG6 (N)
21
                           INTEGER N
22
23
                    ! SOMP
                             PARALLEL DEFAULT (SHARED)
24
                    !$OMP
                               SINGLE
25
                                 CALL WORK (N, 1)
26
                    ! incorrect nesting of barrier region in a single region
27
                    !$OMP
                                 BARRIER
28
                                 CALL WORK (N, 2)
29
                    ! SOMP
                               END SINGLE
30
                    !$OMP
                             END PARALLEL
31
                           END SUBROUTINE WRONG6
```

Fortran -

# A.41 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 (Section 3.2.7 on page 169), and omp set num threads (Section 3.2.1 on page 162).

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 (see Algorithm 2.1 on page 40). 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 A.41.1c

```
#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);
}
}
```

#### Fortran

#### Example A.41.1f

1

33

34

35 36

```
2
                          SUBROUTINE DO BY 16(X, IAM, IPOINTS)
3
                             REAL X(*)
4
                             INTEGER IAM, IPOINTS
5
                          END SUBROUTINE DO_BY_16
6
7
                           SUBROUTINE DYNTHREADS (X, NPOINTS)
8
9
                             INCLUDE "omp lib.h"
                                                     ! or USE OMP LIB
10
11
                             INTEGER NPOINTS
12
                             REAL X (NPOINTS)
13
14
                             INTEGER IAM, IPOINTS
15
16
                             CALL OMP SET DYNAMIC (.FALSE.)
17
                             CALL OMP SET NUM THREADS (16)
18
19
                    !$OMP
                            PARALLEL SHARED (X, NPOINTS) PRIVATE (IAM, IPOINTS)
20
                               IF (OMP GET_NUM_THREADS() .NE. 16) THEN
21
22
                                 STOP
23
                               ENDIF
24
25
                               IAM = OMP GET THREAD NUM()
26
                               IPOINTS = NPOINTS/16
27
                               CALL DO BY 16(X, IAM, IPOINTS)
28
29
                    !$OMP
                             END PARALLEL
30
31
                          END SUBROUTINE DYNTHREADS
```

Fortran -

# A.42 The omp\_get\_num\_threads Routine

In the following example, the <code>omp\_get\_num\_threads</code> call (see Section 3.2.2 on page 163) returns 1 in the sequential part of the code, so <code>np</code> will always be equal to 1. To determine the number of threads that will be deployed for the <code>parallel</code> region, the call should be inside the <code>parallel</code> region.

```
C/C++
Example A.42.1c
#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++
                               Fortran
Example A.42.1f
      SUBROUTINE WORK(I)
      INTEGER I
        I = I + 1
      END SUBROUTINE WORK
      SUBROUTINE INCORRECT()
                            ! or USE OMP LIB
        INCLUDE "omp lib.h"
        INTEGER I, NP
                                    !misplaced: will return 1
       NP = OMP GET NUM THREADS()
       PARALLEL DO SCHEDULE (STATIC)
!$OMP
         DO I = 0, NP-1
            CALL WORK(I)
         ENDDO
     END PARALLEL DO
      END SUBROUTINE INCORRECT
```

Fortran –

1

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3 4

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20 21

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27

28

29

30

```
The following example shows how to rewrite this program without including a query for
 1
 2
                    the number of threads:
                                                       C/C++
                    Example A.42.2c
 3
 4
                    #include <omp.h>
 5
                    void work(int i);
6
7
                    void correct()
8
9
                      int i;
10
11
                      #pragma omp parallel private(i)
12
13
                        i = omp_get_thread_num();
14
                        work(i);
15
                      }
16
                    }
                                                       C/C++ -
                                                     - Fortran -
                    Example A.42.2f
17
18
                          SUBROUTINE WORK(I)
19
                            INTEGER I
20
21
                            I = I + 1
22
23
                          END SUBROUTINE WORK
24
25
                          SUBROUTINE CORRECT()
26
                            INCLUDE "omp lib.h"
                                                 ! or USE OMP LIB
27
                            INTEGER I
28
29
                    !$OMP
                             PARALLEL PRIVATE(I)
30
                              I = OMP_GET_THREAD_NUM()
31
                              CALL WORK(I)
32
                            END PARALLEL
                    !$OMP
33
34
                          END SUBROUTINE CORRECT
                                                       Fortran -
```

# A.43 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** (Section 3.3.1 on page 198).

```
C/C++
Example A.43.1c
#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 A.43.1f
      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
! SOMP
       END PARALLEL DO
```

Fortran -

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23 24

25 26

27

28

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31 32

END FUNCTION NEW\_LOCKS

## A.44 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 <code>omp\_unset\_lock</code> 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 <code>omp\_unset\_lock</code> 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++

C/C++

#### Example A.44.1c

```
#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;
```

#### **Fortran**

#### Example A.44.1f

```
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
```

Fortran

22

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14

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16 17

18

19 20

21

# A.45 Simple Lock Routines

In the following example (for Section 3.3 on page 196), 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 <code>omp\_set\_lock</code> function blocks, but the <code>omp\_test\_lock</code> function does not, allowing the work in <code>skip</code> to be done.

13

14 15

16 17

18 19

20 21

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27

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32

33 34

35 36 37

38 39

40

Note that the argument to the lock routines should have type <code>omp\_lock\_t</code>, and that there is no need to flush it.

#### Example A.45.1c

```
#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++

33 34

35 36

37 38 Note that there is no need to flush the lock variable.

```
Example A.45.1f
```

```
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
```

- Fortran -

## A.46 Nestable Lock Routines

2

3

The following example (for Section 3.3 on page 196) 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 A.46.1c
4
5
                    #include <omp.h>
6
                    typedef struct {
7
                          int a,b;
8
                          omp_nest_lock_t lck; } pair;
9
10
                    int work1();
11
                    int work2();
12
                    int work3();
13
                   void incr a(pair *p, int a)
14
15
                      /* Called only from incr pair, no need to lock. */
16
                     p->a += a;
17
18
                   void incr b(pair *p, int b)
19
20
                      /* Called both from incr pair and elsewhere, */
21
                      /* so need a nestable lock. */
22
23
                      omp set nest lock(&p->lck);
24
                      p->b+=b;
25
                      omp unset nest lock(&p->lck);
26
27
                    void incr pair(pair *p, int a, int b)
28
29
                      omp set nest lock(&p->lck);
30
                      incr a(p, a);
31
                      incr b(p, b);
32
                      omp_unset_nest_lock(&p->lck);
33
34
                    void nestlock(pair *p)
35
36
                      #pragma omp parallel sections
37
38
                        #pragma omp section
39
                          incr pair(p, work1(), work2());
40
                        #pragma omp section
41
                          incr b(p, work3());
42
43
                    }
                                                       C/C++
```

#### 1 Example A.46.1f

```
2
                          MODULE DATA
3
                            USE OMP LIB, ONLY: OMP NEST LOCK KIND
                            TYPE LOCKED PAIR
5
                              INTEGER A
6
                              INTEGER B
7
                              INTEGER (OMP NEST LOCK KIND) LCK
8
                           END TYPE
9
                          END MODULE DATA
10
                          SUBROUTINE INCR A(P, A)
12
                            ! called only from INCR PAIR, no need to lock
13
                            USE DATA
14
                            TYPE (LOCKED PAIR) :: P
15
                            INTEGER A
                            P%A = P%A + A
17
                          END SUBROUTINE INCR A
18
19
                          SUBROUTINE INCR B(P, B)
20
                            ! called from both INCR PAIR and elsewhere,
21
                            ! so we need a nestable lock
22
                            USE OMP LIB ! or INCLUDE "omp lib.h"
23
                            USE DATA
24
                            TYPE (LOCKED PAIR) :: P
25
                            INTEGER B
26
                            CALL OMP_SET_NEST_LOCK (P%LCK)
27
                            P%B = P%B + B
28
                            CALL OMP UNSET NEST LOCK (P%LCK)
29
                          END SUBROUTINE INCR B
30
31
                          SUBROUTINE INCR PAIR(P, A, B)
32
                            USE OMP_LIB ! or INCLUDE "omp_lib.h"
33
                            USE DATA
34
                            TYPE (LOCKED PAIR) :: P
35
                            INTEGER A
36
                            INTEGER B
37
38
                            CALL OMP SET NEST LOCK (P%LCK)
39
                            CALL INCR A(P, A)
40
                            CALL INCR B(P, B)
41
                            CALL OMP UNSET NEST LOCK (P%LCK)
42
                          END SUBROUTINE INCR PAIR
44
                          SUBROUTINE NESTLOCK (P)
45
                            USE OMP LIB
                                         ! or INCLUDE "omp lib.h"
46
                            USE DATA
47
                            TYPE (LOCKED PAIR) :: P
48
                            INTEGER WORK1, WORK2, WORK3
49
                            EXTERNAL WORK1, WORK2, WORK3
50
```

1	!\$OMP	PARALLEL SECTIONS
2		
3	!\$OMP	SECTION
4		CALL INCR_PAIR(P, WORK1(), WORK2())
5	!\$OMP	SECTION
6		CALL INCR_B(P, WORK3())
7	!\$OMP	END PARALLEL SECTIONS
8		
9		END SUBROUTINE NESTLOCK
	<b>A</b>	Fortron

## 1 APPENDIX **B**

2

3

# Stubs for Runtime Library Routines

4 5 6 7 8	This section provides stubs for the runtime library routines defined in the OpenMP API. The stubs are provided to enable portability to platforms that do not support the OpenMP API. On these platforms, OpenMP programs must be linked with a library containing these stub routines. The stub routines assume that the directives in the OpenMP program are ignored. As such, they emulate serial semantics.
9 10 11	Note that the lock variable that appears in the lock routines must be accessed exclusively through these routines. It should not be initialized or otherwise modified in the user program.
12 13 14 15	In an actual implementation the lock variable might be used to hold the address of an allocated memory block, but here it is used to hold an integer value. Users should not make assumptions about mechanisms used by OpenMP implementations to implement locks based on the scheme used by the stub procedures.
	Fortran —
16 17 18 19	<b>Note</b> — In order to be able to compile the Fortran stubs file, the include file omp_lib.h was split into two files: omp_lib_kinds.h and omp_lib.h and the omp_lib_kinds.h file included where needed. There is no requirement for the implementation to provide separate files.
	Fortran —

## B.1 C/C++ Stub Routines

```
#include <stdio.h>
#include <stdlib.h>
#include "omp.h"
void omp_set_num_threads(int num_threads)
}
int omp get num threads(void)
   return 1;
int omp_get_max_threads(void)
    return 1;
}
int omp get thread num(void)
    return 0;
int omp_get_num_procs(void)
   return 1;
int omp_in_parallel(void)
{
    return 0;
void omp_set_dynamic(int dynamic_threads)
}
int omp_get_dynamic(void)
    return 0;
int omp_get_cancellation(void)
   return 0;
}
```

```
1
                    void omp set nested(int nested)
2
3
                    }
4
5
                    int omp_get_nested(void)
6
7
                        return 0;
8
                    }
9
10
                    void omp set schedule(omp sched t kind, int modifier)
                    {
11
12
                    }
13
14
                    void omp get schedule(omp sched t *kind, int *modifier)
15
16
                        *kind = omp sched static;
17
                        *modifier = 0;
18
                    }
19
20
                    int omp_get_thread_limit(void)
21
22
                        return 1;
23
24
25
                    void omp set max active levels(int max active levels)
26
27
                    }
28
29
                    int omp_get_max_active_levels(void)
30
31
                        return 0;
32
33
34
                    int omp_get_level(void)
35
36
                        return 0;
37
                    }
38
39
                    int omp get ancestor thread num(int level)
40
41
                        if (level == 0)
42
43
                            return 0;
44
                        }
45
                        else
46
47
                            return -1;
48
49
                    }
50
```

```
1
                    int omp_get_team_size(int level)
2
3
                        if (level == 0)
4
5
                            return 1;
6
7
                        else
8
9
                            return -1;
10
                    }
11
12
13
                    omp proc bind t omp get proc bind(void)
14
15
                        return omp_proc_bind_false;
16
17
18
                    int omp get active level(void)
19
20
                        return 0;
                    }
21
22
23
                    int omp_in_final(void)
24
25
                        return 1;
26
27
                    struct __omp_lock
28
29
30
                        int lock;
31
                    };
32
33
                    enum { UNLOCKED = -1, INIT, LOCKED };
34
35
                    void omp init lock(omp lock t *arg)
36
37
                        struct omp lock *lock = (struct omp lock *)arg;
38
                        lock->lock = UNLOCKED;
39
                    }
40
41
                    void omp destroy lock(omp lock t *arg)
42
43
                        struct omp lock *lock = (struct omp lock *)arg;
44
                        lock->lock = INIT;
45
                    }
46
```

```
void omp_set_lock(omp_lock_t *arg)
 1
 2
 3
                        struct omp lock *lock = (struct omp lock *)arg;
4
                        if (lock->lock == UNLOCKED)
5
6
                            lock->lock = LOCKED;
7
8
                        else if (lock->lock == LOCKED)
9
10
                            fprintf(stderr,
                               "error: deadlock in using lock variable\n");
11
12
                            exit(1);
13
                        }
14
                        else
15
                        {
16
                            fprintf(stderr, "error: lock not initialized\n");
17
                            exit(1);
18
19
                    }
20
21
22
                    void omp unset lock(omp lock t *arg)
23
24
                        struct omp lock *lock = (struct __omp_lock *)arg;
25
                        if (lock->lock == LOCKED)
26
27
                            lock->lock = UNLOCKED;
28
                        }
29
                        else if (lock->lock == UNLOCKED)
30
31
                            fprintf(stderr, "error: lock not set\n");
32
                            exit(1);
33
                        }
34
                        else
35
36
                            fprintf(stderr, "error: lock not initialized\n");
37
                            exit(1);
38
                    }
39
```

```
1
                    int omp_test_lock(omp_lock_t *arg)
2
3
                        struct omp lock *lock = (struct omp lock *)arg;
4
                        if (lock->lock == UNLOCKED)
5
6
                            lock->lock = LOCKED;
7
                            return 1;
8
9
                        else if (lock->lock == LOCKED)
10
11
                            return 0;
12
13
                        else
14
15
                            fprintf(stderr, "error: lock not initialized\n");
16
                            exit(1);
                        }
17
18
                    }
19
20
                    struct __omp_nest_lock
21
22
                        short owner;
23
                        short count;
24
                    };
25
26
                    enum { NOOWNER = -1, MASTER = 0 };
27
28
                   void omp_init_nest_lock(omp_nest_lock_t *arg)
29
30
                        struct omp nest lock *nlock=(struct omp nest lock *)arg;
31
                        nlock->owner = NOOWNER;
32
                       nlock->count = 0;
33
                    }
34
35
36
                   void omp destroy nest lock(omp nest lock t *arg)
37
                        struct __omp_nest_lock *nlock=(struct __omp_nest_lock *)arg;
38
39
                        nlock->owner = NOOWNER;
40
                       nlock->count = UNLOCKED;
41
42
```

```
1
                    void omp set nest lock(omp nest lock t *arg)
2
3
                        struct omp nest lock *nlock=(struct omp nest lock *)arg;
4
                        if (nlock->owner == MASTER && nlock->count >= 1)
5
6
                            nlock->count++;
7
8
                        else if (nlock->owner == NOOWNER && nlock->count == 0)
9
10
                            nlock->owner = MASTER;
11
                            nlock->count = 1;
12
                        }
13
                        else
14
                        {
15
                            fprintf(stderr,
16
                               "error: lock corrupted or not initialized\n");
17
                            exit(1);
18
                        }
19
                    }
20
21
                   void omp unset nest lock(omp nest lock t *arg)
22
23
                        struct __omp_nest_lock *nlock=(struct __omp_nest_lock *)arg;
24
                        if (nlock->owner == MASTER && nlock->count >= 1)
25
26
                            nlock->count--;
27
                            if (nlock->count == 0)
28
29
                                nlock->owner = NOOWNER;
30
                            }
31
                        }
32
                        else if (nlock->owner == NOOWNER && nlock->count == 0)
33
34
                            fprintf(stderr, "error: lock not set\n");
35
                            exit(1);
36
                        }
37
                        else
38
39
                            fprintf(stderr,
40
                               "error: lock corrupted or not initialized\n");
41
                            exit(1);
42
                        }
43
                    }
44
45
                    int omp test nest lock(omp nest lock t *arg)
46
47
                        struct __omp_nest_lock *nlock=(struct __omp_nest_lock *)arg;
48
                        omp set nest lock(arg);
49
                        return nlock->count;
50
                    }
51
```

```
1
                    double omp_get_wtime(void)
2
3
4
5
6
7
                    /* This function does not provide a working
                     * wallclock timer. Replace it with a version
                     * customized for the target machine.
                        return 0.0;
8
9
10
                    double omp_get_wtick(void)
11
12
                    /* This function does not provide a working
13
                     * clock tick function. Replace it with
14
                     * a version customized for the target machine.
15
                        return 365. * 86400.;
16
17
                    }
18
```

## **B.2** Fortran Stub Routines

```
2
                    C23456
 3
                           subroutine omp_set_num_threads(num_threads)
4
                              integer num threads
5
                             return
6
                           end subroutine
7
8
                           integer function omp get num threads()
9
                             omp_get_num_threads = 1
10
                             return
11
                           end function
12
13
                           integer function omp get max threads()
14
                             omp_get_max_threads = 1
15
                             return
16
                           end function
17
18
                           integer function omp get thread num()
19
                             omp get thread num = 0
20
                             return
21
                           end function
22
23
                           integer function omp get num procs()
24
                             omp_get_num_procs = 1
25
                             return
26
                           end function
27
28
                           logical function omp in parallel()
29
                             omp in parallel = .false.
30
                             return
31
                           end function
32
33
                           subroutine omp set dynamic (dynamic threads)
34
                             logical dynamic threads
35
                             return
36
                           end subroutine
37
38
                           logical function omp get dynamic()
39
                             omp get dynamic = .false.
40
                             return
41
                           end function
42
43
                           logical function omp get cancellation()
44
                                omp get cancellation = .false.
45
                                return
46
                           end function
47
48
49
```

```
1
                           subroutine omp set nested(nested)
2
                             logical nested
3
                             return
4
                           end subroutinelogical function omp get nested()
5
                             omp get nested = .false.
6
                             return
7
                           end function
8
9
                           subroutine omp set schedule(kind, modifier)
10
                             include 'omp lib kinds.h'
11
                             integer (kind=omp sched kind) kind
12
                             integer modifier
13
                             return
14
                           end subroutine
15
16
                           subroutine omp get schedule(kind, modifier)
                             include 'omp lib kinds.h'
17
18
                             integer (kind=omp sched kind) kind
19
                             integer modifier
20
21
                             kind = omp sched static
22
                             modifier = 0
23
                             return
24
                           end subroutine
25
26
                           integer function omp_get_thread_limit()
27
                             omp get thread limit = 1
28
                             return
29
                           end function
30
31
                           subroutine omp set max active levels ( level )
32
                             integer level
33
                           end subroutine
34
35
                           integer function omp get max active levels()
36
                             omp get max active levels = 0
37
                             return
38
                           end function
39
40
                           integer function omp get level()
41
                             omp get level = 0
42
                             return
43
                           end function
44
                           integer function omp_get_ancestor_thread_num( level )
45
46
                             integer level
47
                             if (level .eq. 0) then
48
                                omp get ancestor thread num = 0
49
                             else
50
                                omp_get_ancestor_thread_num = -1
51
                             end if
52
                             return
53
                           end function
54
```

```
1
                           integer function omp get team size( level )
2
                             integer level
3
                             if (level .eq. 0) then
4
                                omp get team size = 1
5
6
                                omp get team size = -1
7
                             end if
8
                             return
9
                           end function
10
11
                           integer function omp get active level()
12
                             omp get active level = 0
13
                             return
14
                           end function
15
16
                           logical function omp in final()
17
                             omp in final = .true.
18
                             return
19
                           end function
20
21
                           function omp get proc bind()
22
                             include 'omp lib kinds.h'
23
                             integer (kind=omp_proc_bind_kind) omp_get_proc_bind
24
                             omp get proc bind = omp proc bind false
25
                           end function omp get proc bind
26
27
                           subroutine omp init lock(lock)
28
                             ! lock is 0 if the simple lock is not initialized
29
                                      -1 if the simple lock is initialized but not set
30
                                       1 if the simple lock is set
31
                             include 'omp lib kinds.h'
32
                             integer(kind=omp lock kind) lock
33
34
                             lock = -1
35
                             return
36
                           end subroutine
37
38
                           subroutine omp destroy lock(lock)
39
                             include 'omp lib kinds.h'
40
                             integer(kind=omp lock kind) lock
41
42
                             lock = 0
43
                             return
44
                           end subroutine
45
46
                           subroutine omp set lock(lock)
47
                             include 'omp lib kinds.h'
48
                             integer(kind=omp lock kind) lock
49
50
                             if (lock .eq. -1) then
51
                               lock = 1
52
                             elseif (lock .eq. 1) then
53
                               print *, 'error: deadlock in using lock variable'
54
                               stop
```

```
1
                             else
2
                               print *, 'error: lock not initialized'
3
4
                             endif
5
                             return
6
                           end subroutine
7
                           subroutine omp unset lock(lock)
8
                             include 'omp lib kinds.h'
9
                             integer(kind=omp lock kind) lock
10
11
                             if (lock .eq. 1) then
12
                               lock = -1
13
                             elseif (lock .eq. -1) then
14
                               print *, 'error: lock not set'
15
                               stop
16
                             else
17
                               print *, 'error: lock not initialized'
18
                               stop
19
                             endif
20
21
                             return
22
                           end subroutine
23
24
                           logical function omp test lock(lock)
25
                             include 'omp lib kinds.h'
26
                             integer(kind=omp_lock_kind) lock
27
28
                             if (lock .eq. -1) then
29
                               lock = 1
30
                               omp test lock = .true.
31
                             elseif (lock .eq. 1) then
32
                               omp test lock = .false.
33
34
                               print *, 'error: lock not initialized'
35
                               stop
36
                             endif
37
38
                             return
39
                           end function
40
41
                           subroutine omp init nest lock(nlock)
42
                             ! nlock is
43
                             ! 0 if the nestable lock is not initialized
44
                             ! -1 if the nestable lock is initialized but not set
45
                             ! 1 if the nestable lock is set
46
                             ! no use count is maintained
47
                             include 'omp lib kinds.h'
48
                             integer(kind=omp nest lock kind) nlock
49
50
                             nlock = -1
51
52
                             return
53
                           end subroutine
54
```

```
subroutine omp destroy nest lock(nlock)
 1
 2
                             include 'omp lib kinds.h'
 3
                             integer(kind=omp nest lock kind) nlock
 4
 5
                             nlock = 0
6
7
                             return
8
                           end subroutine
9
10
                           subroutine omp set nest lock(nlock)
11
                             include 'omp lib kinds.h'
12
                             integer(kind=omp nest lock kind) nlock
13
14
                             if (nlock .eq. -1) then
15
                               nlock = 1
16
                             elseif (nlock .eq. 0) then
17
                               print *, 'error: nested lock not initialized'
18
                               stop
19
                             else
20
                               print *, 'error: deadlock using nested lock variable'
21
                               stop
22
                             endif
23
24
                             return
25
                           end subroutine
26
27
                           subroutine omp unset nest lock(nlock)
28
                             include 'omp lib kinds.h'
29
                             integer(kind=omp_nest_lock_kind) nlock
30
31
                             if (nlock .eq. 1) then
32
                               nlock = -1
33
                             elseif (nlock .eq. 0) then
34
                               print *, 'error: nested lock not initialized'
35
                               stop
36
                             else
37
                               print *, 'error: nested lock not set'
38
                               stop
39
                             endif
40
41
                             return
42
                           end subroutine
43
```

```
1
                           integer function omp test nest lock(nlock)
2
                             include 'omp lib kinds.h'
3
                             integer(kind=omp nest lock kind) nlock
4
5
                             if (nlock .eq. -1) then
6
                               nlock = 1
7
                               omp test nest lock = 1
8
                             elseif (nlock .eq. 1) then
9
                               omp test nest lock = 0
10
11
                               print *, 'error: nested lock not initialized'
12
                               stop
13
                             endif
14
15
                             return
16
                           end function
17
18
19
                           double precision function omp get wtime()
20
                             ! this function does not provide a working
21
                             ! wall clock timer. replace it with a version
22
                             ! customized for the target machine.
23
24
                             omp get wtime = 0.0d0
25
26
                             return
27
                           end function
28
29
                           double precision function omp_get_wtick()
30
                             ! this function does not provide a working
31
                             ! clock tick function. replace it with
32
                             ! a version customized for the target machine.
33
                             double precision one year
34
                             parameter (one_year=365.d0*86400.d0)
35
36
                             omp get wtick = one year
37
38
                             return
39
                           end function
```

## 1 APPENDIX **C**

# **OpenMP C and C++ Grammar**

3

2

## C.1 Notation

The grammar rules consist of the name for a non-terminal, followed by a colon, 5 6 followed by replacement alternatives on separate lines. 7 The syntactic expression  $term_{opt}$  indicates that the term is optional within the 8 replacement. The syntactic expression  $term_{optseq}$  is equivalent to  $term-seq_{opt}$  with the following 9 additional rules: 10 11 term-seq: 12 term 13 term-seq term 14 term-seq, term

# C.2 Rules

2 The notation is described in Section 6.1 of the C standard. This grammar appendix shows the extensions to the base language grammar for the OpenMP C and C++ 3 directives. /\* in C++ (ISO/IEC 14882:1998) \*/ statement-seq: statement openmp-directive 10 statement-seq statement 11 statement-seq openmp-directive 12 13 /\* in C90 (ISO/IEC 9899:1990) \*/ 14 15 statement-list: 16 statement 17 openmp-directive 18 statement-list statement 19 statement-list openmp-directive 20 21 22 /\* in C99 (ISO/IEC 9899:1999) \*/ 23 block-item: 24 declaration 25 statement 26 openmp-directive

1	
2	statement:
3	/* standard statements */
4	openmp-construct
5	openmp-construct:
6	parallel-construct
7	for-construct
8	sections-construct
9	single-construct
10	parallel-for-construct
11	parallel-sections-construct
12	task-construct
13	master-construct
14	critical-construct
15	atomic-construct
16	ordered-construct
17	openmp-directive:
18	barrier-directive
19	taskwait-directive
20	taskyield-directive
21	flush-directive
22	structured-block:
23	statement
24	parallel-construct:
25	parallel-directive structured-block
26	parallel-directive:
27	# pragma omp parallel $parallel$ - $clause_{optseq}$ $new$ - $line$
28	

```
parallel-clause:
 1
                         unique-parallel-clause
                        data-default-clause
                        data-privatization-clause
                        data-privatization-in-clause
                        data-sharing-clause
                        data-reduction-clause
                     unique-parallel-clause:
                         if ( expression )
10
                        num_threads ( expression )
11
                        copyin ( variable-list )
12
                     for-construct:
13
                        for-directive iteration-statement
                     for-directive:
14
                         \# pragma omp for for\text{-}clause_{optseq} new\text{-}line
15
16
                     for-clause:
                         unique-for-clause
17
18
                        data-privatization-clause
19
                        data-privatization-in-clause
                        data-privatization-out-clause
20
21
                        data-reduction-clause
22
                        nowait
23
                     unique-for-clause:
24
                         ordered
25
                        schedule (schedule-kind)
26
                        schedule (schedule-kind, expression)
27
                        collapse ( expression )
28
```

1	schedule-kind:
2	static
3	dynamic
4	guided
5	auto
6	runtime
7	sections-construct:
8	sections-directive section-scope
9	sections-directive:
10	# pragma omp sections $sections$ -clause $_{optseq}$ $new$ -line
11	sections-clause:
12	data-privatization-clause
13	data-privatization-in-clause
14	data-privatization-out-clause
15	data-reduction-clause
16	nowait
17	section-scope:
18	{ section-sequence }
19	section-sequence:
20	section-directive <sub>opt</sub> structured-block
21	section-sequence section-directive structured-block
22	section-directive:
23	# pragma omp section new-line
24	single-construct:
25	single-directive structured-block
26	single-directive:
27	# pragma omp single $single$ - $clause_{optseq}$ $new$ - $line$
28	

```
1
                       single-clause:
                          unique-single-clause
                         data-privatization-clause
 3
                         data-privatization-in-clause
                         nowait
                       unique-single-clause:
                          copyprivate (variable-list )
                       task-construct:
                          task-directive structured-block
10
                      task-directive:
                          # pragma omp task task-clause<sub>optsea</sub> new-line
11
12
                       task-clause:
13
                         unique-task-clause
14
                         data-default-clause
15
                         data-privatization-clause
16
                         data-privatization-in-clause
                         data-sharing-clause
17
18
                       unique-task-clause:
19
                          if ( scalar-expression )
20
                          final(scalar-expression)
21
                         untied
22
                         mergeable
23
                      parallel-for-construct:
24
                          parallel-for-directive iteration-statement
25
                      parallel-for-directive:
26
                           \verb|# pragma omp parallel for \textit{parallel-for-clause}_{\textit{optseq}} \textit{new-line} \\
```

1	parallel-for-clause:
2	unique-parallel-clause
3	unique-for-clause
4	data-default-clause
5	data-privatization-clause
6	data-privatization-in-clause
7	data-privatization-out-clause
8	data-sharing-clause
9	data-reduction-clause
10	parallel-sections-construct:
11	parallel-sections-directive section-scope
12	parallel-sections-directive:
13	# pragma omp parallel sections $parallel$ -sections-clause $_{optseq}$ $new$ -line
14	parallel-sections-clause:
15	unique-parallel-clause
16	data-default-clause
17	data-privatization-clause
18	data-privatization-in-clause
19	data-privatization-out-clause
20	data-sharing-clause
21	data-reduction-clause
22	master-construct:
23	master-directive structured-block
24	master-directive:
25	# pragma omp master new-line
26	critical-construct:
27	critical-directive structured-block

1	critical-directive:
2	# pragma omp critical $region-phrase_{opt}$ $new-line$
3	region-phrase:
4	( identifier )
5	
6	barrier-directive:
7	# pragma omp barrier new-line
8	taskwait-directive:
9	<pre># pragma omp taskwait new-line</pre>
10	taskyield-directive:
11	# pragma omp taskyield new-line
12	atomic-construct:
13	atomic-directive expression-statement
14	atomic-directive structured block
15	atomic-directive:
16	# pragma omp atomic atomic-clause <sub>opt</sub> new-line
17	atomic-clause:
18	read
19	write
20	update
21	capture
22	flush-directive:
23	# pragma omp flush flush-varson new-line
24	flush-vars:
25	( variable-list )
26	ordered-construct:
27	ordered-directive structured-block
28	

```
ordered-directive:
1
 2
                       # pragma omp ordered new-line
                    declaration:
 3
                       /* standard declarations */
4
                       threadprivate-directive
5
                       declare-reduction-directive
6
7
                    threadprivate-directive:
8
                       # pragma omp threadprivate (variable-list) new-line
                    declare-reduction-directive:
9
10
                       # pragma omp declare reduction (reduction-identifier:
                       reduction-type-list: expression) [initializer-clause] new-line
11
12
                    reduction-identifier:
                       /* in C */
13
14
                       identifier
                       /* in C++ */
15
16
                       id-expression
                       /* in C/C++ */
17
                       one of: + * - & ^ | && || min max
18
19
                    reduction-type-list:
20
                       type-id
21
                       reduction-type-list, type-id
22
                    initializer-clause:
                       /* in C */
23
24
                       initializer ( identifier = initializer )
25
                       initializer (identifier (argument-expression-list) )
                       /* in C++ */
26
27
                       initializer ( identifier initializer )
28
                       initializer (id-expression (expression-list))
```

```
1
2
                    data-default-clause:
                       default ( shared )
 5
                       default ( none )
                    data-privatization-clause:
                       private ( variable-list )
8
                    data-privatization-in-clause:
9
                       firstprivate (variable-list)
                    data-privatization-out-clause:
10
11
                       lastprivate (variable-list)
12
                    data-sharing-clause:
13
                       shared (variable-list)
14
                    data-reduction-clause:
                       reduction ( reduction-identifier : variable-list )
15
                    /* in C */
16
17
                    variable-list:
18
                       identifier
```

variable-list , identifier

1	/* in C++ */
2	variable-list:
3	id-expression
4	variable-list , id-expression

## 1 APPENDIX **D**

# **Interface Declarations**

This appendix gives examples of the C/C++ header file, the Fortran include file and
Fortran module that shall be provided by implementations as specified in Chapter 3. It
also includes an example of a Fortran 90 generic interface for a library routine. This is a
non-normative section, implementation files may differ.

## D.1 Example of the omp.h Header File

```
#ifndef _OMP H DEF
#define _OMP_H_DEF
 * define the lock data types
typedef void *omp lock t;
typedef void *omp nest lock t;
 * define the schedule kinds
typedef enum omp sched t
    omp sched static = 1,
    omp sched dynamic = 2,
    omp sched guided = 3,
    omp sched auto = 4
/* , Add vendor specific schedule constants here */
} omp sched t;
 * define the proc bind values
typedef enum omp proc bind t
    omp proc bind false = 0,
    omp_proc_bind_true = 1,
    omp proc bind master = 2,
    omp proc bind close = 3,
    omp proc bind spread = 4
} omp proc bind t;
 * exported OpenMP functions
#ifdef __cplusplus
extern
                "C"
#endif
extern void
              omp set num threads(int num threads);
extern int
               omp get num threads (void);
extern int
               omp get max threads(void);
extern int
               omp get thread num(void);
extern int
               omp get num procs(void);
extern int
              omp in parallel(void);
extern void
              omp set dynamic(int dynamic threads);
```

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```
1
                   extern int
                                  omp get dynamic (void);
2
                  extern void omp set nested(int nested);
3
                   extern int
                                  omp get nested(void);
4
                   extern int
                                  omp get thread limit(void);
5
                   extern void
                                 omp set max active levels(int max active levels);
6
                                  omp get max active levels(void);
                   extern int
7
                   extern int
                                  omp get level(void);
8
                   extern int
                                  omp get ancestor thread num(int level);
9
                   extern int
                                  omp get team size(int level);
10
                   extern int
                                  omp get active level(void);
11
                   extern int
                                omp in final(void);
12
                  extern void
                                 omp set schedule(omp sched t kind, int modifier);
13
                   extern void
                                 omp get schedule(omp sched t *kind, int *modifier);
14
                   extern omp proc bind t omp get proc bind(void);
15
16
                                 omp init lock(omp lock t *lock);
                  extern void
17
                   extern void
                                 omp destroy lock(omp lock t *lock);
18
                  extern void omp set lock(omp lock t *lock);
19
                   extern void
                                 omp unset lock(omp lock t *lock);
20
                   extern int
                                  omp test lock(omp lock t *lock);
21
22
                                 omp init nest lock(omp_nest_lock_t *lock);
                  extern void
23
                   extern void
                                 omp_destroy_nest_lock(omp_nest_lock_t *lock);
24
                   extern void
                                 omp set nest lock(omp nest lock t *lock);
25
                   extern void
                                 omp unset nest lock(omp nest lock t *lock);
26
                  extern int
                                  omp test nest lock(omp nest lock t *lock);
27
28
                  extern double omp get wtime(void);
29
                  extern double omp_get_wtick(void);
30
                   #ifdef cplusplus
31
32
33
                   #endif
34
```

#endif

# D.2 Example of an Interface Declaration include File

```
omp lib kinds.h:
       integer
                   omp lock kind
                   omp nest lock kind
       integer
! this selects an integer that is large enough to hold a 64 bit integer
       parameter ( omp lock kind = selected int kind( 10 ) )
       parameter ( omp nest lock kind = selected int kind( 10 ) )
                   omp sched kind
! this selects an integer that is large enough to hold a 32 bit integer
       parameter ( omp sched kind = selected int kind( 8 ) )
       integer ( omp sched kind ) omp sched static
       parameter ( omp sched static = 1 )
       integer ( omp_sched_kind ) omp_sched_dynamic
       parameter ( omp sched dynamic = 2 )
       integer ( omp sched kind ) omp sched guided
       parameter ( omp_sched_guided = 3 )
       integer ( omp sched kind ) omp sched auto
       parameter ( omp sched auto = 4 )
       integer omp proc bind kind
       parameter ( omp proc bind kind = selected int kind( 8 ) )
       integer ( omp proc bind kind ) omp proc bind false
       parameter ( omp proc bind false = 0 )
       integer ( omp proc bind kind ) omp proc bind true
       parameter ( omp proc bind true = 1 )
       integer ( omp_proc_bind_kind ) omp_proc_bind_master
       parameter ( omp proc bind master = 2 )
       integer ( omp proc bind kind ) omp proc bind close
       parameter ( omp proc bind close = 3 )
       integer ( omp proc bind kind ) omp proc bind spread
       parameter ( omp proc bind spread = 4 )
omp_lib.h:
! default integer type assumed below
! default logical type assumed below
! OpenMP API v4.0
       include 'omp lib kinds.h'
       integer
                   openmp version
       parameter ( openmp version = 201211 )
       external omp set num threads
       external omp get num threads
       integer omp get num threads
       external omp_get_max_threads
       integer omp get max threads
       external omp get thread num
       integer omp get thread num
       external omp get num procs
```

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```
1
                          integer omp get num procs
2
                          external omp in parallel
3
                          logical omp in parallel
4
                          external omp set dynamic
5
                          external omp get dynamic
                          logical omp_get_dynamic
6
7
                          external omp set nested
8
                          external omp get nested
9
                          logical omp get nested
10
                          external omp set schedule
11
                          external omp get schedule
12
                          external omp get thread limit
13
                          integer omp get thread limit
14
                          external omp set max active levels
15
                          external omp get max active levels
16
                          integer omp get max active levels
17
                          external omp get level
18
                          integer omp get level
19
                          external omp get ancestor thread num
20
                          integer omp get ancestor thread num
21
                          external omp get team size
22
                          integer omp get team size
23
                          external omp_get_active_level
24
                          integer omp get active level
25
26
                          external omp in final
27
                          logical omp in final
28
29
                          integer ( omp proc bind kind ) omp get proc bind
30
                          external omp get proc bind
31
32
                          external omp init lock
33
                          external omp_destroy_lock
34
                          external omp set lock
35
                          external omp unset lock
36
                          external omp test lock
37
                          logical omp test lock
38
39
                          external omp init nest lock
40
                          external omp destroy nest lock
41
                          external omp set nest lock
42
                          external omp unset nest lock
43
                          external omp test nest lock
44
                          integer omp_test_nest_lock
45
46
                          external omp get wtick
47
                          double precision omp_get_wtick
48
                          external omp get wtime
49
                          double precision omp_get_wtime
```

# D.3 Example of a Fortran Interface Declaration

module

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32 33

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36 37

38 39

40

41

42 43

44

45

46 47

48

```
the "!" of this comment starts in column 1
123456
        module omp lib kinds
        integer, parameter :: omp lock kind = selected int kind( 10 )
       integer, parameter :: omp nest lock kind = selected int kind( 10 )
       integer, parameter :: omp sched kind = selected int kind( 8 )
       integer(kind=omp sched kind), parameter ::
          omp sched static = 1
        integer(kind=omp sched kind), parameter ::
          omp sched dynamic = 2
       integer(kind=omp sched kind), parameter ::
          omp sched guided = 3
        integer(kind=omp sched kind), parameter ::
          omp sched auto = 4
      integer, parameter :: omp proc bind kind = selected int kind( 8 )
      integer (kind=omp proc bind kind), parameter ::
          omp proc bind false = 0
       integer (kind=omp proc bind kind), parameter ::
          omp proc bind true = 1
      integer (kind=omp proc bind kind), parameter ::
          omp proc bind master = 2
      integer (kind=omp proc bind kind), parameter ::
          omp proc bind close = 3
      integer (kind=omp_proc_bind_kind), parameter ::
          omp proc bind spread = 4
        end module omp lib kinds
      module omp lib
         use omp lib kinds
                                      OpenMP API v4.0
         integer, parameter :: openmp version = 201211
        interface
         subroutine omp set num threads (number of threads expr)
          integer, intent(in) :: number of threads expr
         end subroutine omp set num threads
         function omp get num threads ()
          integer :: omp get num threads
         end function omp get num threads
         function omp get max threads ()
          integer :: omp get max threads
         end function omp get max threads
```

```
1
 2
                             function omp get thread num ()
 3
                              integer :: omp get thread num
 4
                             end function omp get thread num
 5
6
 7
                             function omp get num procs ()
8
                              integer :: omp get num procs
                             end function omp_get_num_procs
9
10
11
                             function omp in parallel ()
12
                              logical :: omp in parallel
13
                             end function omp in parallel
14
15
                             subroutine omp set dynamic (enable expr)
16
                              logical, intent(in) ::enable expr
17
                             end subroutine omp set dynamic
18
19
                             function omp get dynamic ()
20
                              logical :: omp get dynamic
21
                             end function omp get dynamic
22
23
                             subroutine omp_set_nested (enable_expr)
24
                              logical, intent(in) :: enable expr
25
                             end subroutine omp set nested
26
27
                             function omp get nested ()
28
                              logical :: omp get nested
29
                             end function omp_get_nested
30
31
                             subroutine omp set schedule (kind, modifier)
32
                              use omp lib kinds
33
                              integer(kind=omp sched kind), intent(in) :: kind
34
                              integer, intent(in) :: modifier
35
                             end subroutine omp set schedule
36
37
                             subroutine omp get schedule (kind, modifier)
38
                              use omp lib kinds
39
                              integer(kind=omp sched kind), intent(out) :: kind
40
                              integer, intent(out)::modifier
41
                             end subroutine omp get schedule
42
43
                             function omp get thread limit()
44
                              integer :: omp get thread limit
45
                             end function omp get thread limit
46
47
                             subroutine omp set max active levels(var)
48
                              integer, intent(in) :: var
49
                             end subroutine omp set max active levels
50
51
                             function omp get max active levels()
52
                              integer :: omp get max active levels
53
                             end function omp get max active levels
54
```

```
1
                             function omp get level()
2
                              integer :: omp get level
3
                             end function omp get level
4
5
                             function omp get ancestor thread num(level)
6
                              integer, intent(in) :: level
7
                              integer :: omp get ancestor thread num
8
                             end function omp get ancestor thread num
9
10
                             function omp get team size(level)
11
                              integer, intent(in) :: level
12
                              integer :: omp get team size
13
                             end function omp get team size
14
15
                             function omp get active level()
16
                              integer :: omp get active level
17
                             end function omp_get_active_level
18
19
                            function omp in final()
20
                              logical omp in final
                            end function omp in final
21
22
23
                             function omp_get_proc_bind( )
24
                               include 'omp lib kinds.h'
25
                               integer (kind=omp proc bind kind) omp get proc bind
26
                               omp_get_proc_bind = omp_proc_bind_false
27
                              end function omp get proc bind
28
29
                             subroutine omp_init_lock (var)
30
                              use omp lib kinds
31
                              integer (kind=omp lock kind), intent(out) :: var
32
                             end subroutine omp init lock
33
34
                             subroutine omp destroy lock (var)
35
                              use omp lib kinds
36
                              integer (kind=omp lock kind), intent(inout) :: var
37
                             end subroutine omp destroy lock
38
39
                             subroutine omp set lock (var)
40
                              use omp lib kinds
41
                              integer (kind=omp lock kind), intent(inout) :: var
42
                             end subroutine omp set lock
43
44
                             subroutine omp unset lock (var)
45
                              use omp lib kinds
46
                              integer (kind=omp lock kind), intent(inout) :: var
47
                             end subroutine omp unset lock
48
49
                             function omp test lock (var)
50
                              use omp lib kinds
51
                              logical :: omp test lock
52
                              integer (kind=omp lock kind), intent(inout) :: var
53
                             end function omp_test_lock
54
```

```
1
 2
 3
                             subroutine omp init nest lock (var)
4
                              use omp lib kinds
5
                              integer (kind=omp_nest_lock_kind), intent(out) :: var
6
                             end subroutine omp init nest lock
7
8
                             subroutine omp destroy nest lock (var)
9
                              use omp lib kinds
10
                              integer (kind=omp nest lock kind), intent(inout) :: var
                             end subroutine omp destroy nest lock
11
12
13
                             subroutine omp set nest lock (var)
14
                              use omp lib kinds
15
                              integer (kind=omp nest lock kind), intent(inout) :: var
16
                             end subroutine omp set nest lock
17
18
                             subroutine omp unset nest lock (var)
19
                              use omp lib kinds
20
                              integer (kind=omp nest lock kind), intent(inout) :: var
21
                             end subroutine omp unset nest lock
22
23
                             function omp_test_nest_lock (var)
24
                              use omp lib kinds
25
                              integer :: omp test nest lock
26
                              integer (kind=omp nest lock kind), intent(inout) :: var
27
                             end function omp test nest lock
28
29
                             function omp_get_wtick ()
30
                               double precision :: omp get wtick
31
                             end function omp get wtick
32
33
                             function omp get wtime ()
34
                               double precision :: omp get wtime
35
                             end function omp get wtime
36
37
                             end interface
38
39
                          end module omp lib
```

# D.4 Example of a Generic Interface for a Library Routine

Any of the OpenMP runtime library routines that take an argument may be extended with a generic interface so arguments of different KIND type can be accommodated.

The **OMP\_SET\_NUM\_THREADS** interface could be specified in the **omp\_lib** module as the following:

396

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## 1 APPENDIX **E**

# OpenMP Implementation-Defined Behaviors

This appendix summarizes the behaviors that are described as implementation defined in this API. Each behavior is cross-referenced back to its description in the main specification. An implementation is required to define and document its behavior in these cases.

- **Memory model**: the minimum size at which a memory update may also read and write back adjacent variables that are part of another variable (as array or structure elements) is implementation defined but is no larger than required by the base language (see Section 1.4.1 on page 15).
- Internal control variables: the initial values of *nthreads-var*, *dyn-var*, *run-sched-var*, *def-sched-var*, *bind-var*, *place-partition-var*, *stacksize-var*, *wait-policy-var*, *thread-limit-var*, and *max-active-levels-var* are implementation defined (see Section 2.3.2 on page 32).
- **Dynamic adjustment of threads**: providing the ability to dynamically adjust the number of threads is implementation defined. Implementations are allowed to deliver fewer threads (but at least one) than indicated in Algorithm 2-1 even if dynamic adjustment is disabled (see Section 2.5.1 on page 40).
- Loop directive: the integer type or kind used to compute the iteration count of a collapsed loop is implementation defined. The effect of the schedule(runtime) clause when the run-sched-var ICV is set to auto is implementation defined. See Section 2.7.1 on page 47.
- **sections construct**: the method of scheduling the structured blocks among threads in the team is implementation defined (see Section 2.7.2 on page 53).
- **single construct**: the method of choosing a thread to execute the structured block is implementation defined (see Section 2.7.3 on page 56).

1 2 3 4 5 6	
7 8 9	
10 11 12 13 14	
15 16 17 18	
19 20 21	
22 23 24 25	
26 27 28	
29 30	
31 32	
33 34 35	
36 37	
38 39 40 41	

- atomic construct: a compliant implementation may enforce exclusive access between atomic regions which update different storage locations. The circumstances under which this occurs are implementation defined (see Section 2.12.6 on page 103).
- omp\_set\_num\_threads routine: if the argument is not a positive integer the behavior is implementation defined (see Section 3.2.1 on page 162).
- omp\_set\_schedule routine: for implementation specific schedule types, the values and associated meanings of the second argument are implementation defined. (see Section 3.2.12 on page 176).
- omp\_set\_max\_active\_levels routine: when called from within any explicit parallel region the binding thread set (and binding region, if required) for the omp\_set\_max\_active\_levels region is implementation defined and the behavior is implementation defined. If the argument is not a non-negative integer then the behavior is implementation defined (see Section 3.2.15 on page 180).
- omp\_get\_max\_active\_levels routine: when called from within any explicit parallel region the binding thread set (and binding region, if required) for the omp\_get\_max\_active\_levels region is implementation defined (see Section 3.2.16 on page 182).
- OMP\_SCHEDULE environment variable: if the value of the variable does not conform to the specified format then the result is implementation defined (see Section 4.1 on page 210).
- OMP\_NUM\_THREADS environment variable: if any value of the list specified in the OMP\_NUM\_THREADS environment variable leads to a number of threads that is greater than the implementation can support, or if any value is not a positive integer, then the result is implementation defined (see Section 4.2 on page 211).
- OMP\_PROC\_BIND environment variable: if the value is not true, false, or a comma separated list of master, close, or spread, the behavior is implementation defined (see Section 4.4 on page 213).
- OMP\_DYNAMIC environment variable: if the value is neither true nor false the behavior is implementation defined (see Section 4.3 on page 212).
- OMP\_NESTED environment variable: if the value is neither true nor false the behavior is implementation defined (see Section on page 213).
- OMP\_STACKSIZE environment variable: if the value does not conform to the specified format or the implementation cannot provide a stack of the specified size then the behavior is implementation defined (see Section 4.7 on page 216).
- OMP\_WAIT\_POLICY environment variable: the details of the ACTIVE and PASSIVE behaviors are implementation defined (see Section 4.8 on page 217).
- OMP\_MAX\_ACTIVE\_LEVELS environment variable: if the value is not a nonnegative integer or is greater than the number of parallel levels an implementation can support then the behavior is implementation defined (see Section 4.9 on page 217).

1 2 3 4 5	
6 7 8	
9 10 11 12 13	
14 15	
16 17 18 19	
20 21 22 23 24	

- OMP\_THREAD\_LIMIT environment variable: if the requested value is greater than the number of threads an implementation can support, or if the value is not a positive integer, the behavior of the program is implementation defined (see Section 4.10 on page 218).
- OMP\_PLACES environment variable: if any numerical values cannot be mapped to a processor on the target platform the behavior is implementation defined. The behavior is also implementation defined when the OMP\_PLACES environment variable is defined using an abstract name.
- When creating a place list of *n* elements by appending the number *n* to an abstract name, the determination of which resources to include in the place list and their respective order is implementation defined. When requesting more resources than available, the length of the place list is also implementation defined.
- Thread affinity policy: if the request for threads in one or more places in the *place-partition-var* ICV for a parallel construct cannot be fulfilled, the behavior of the thread affinity policy is implementation defined for that parallel construct.

#### Fortran -

- threadprivate directive: if the conditions for values of data in the threadprivate objects of threads (other than the initial thread) to persist between two consecutive active parallel regions do not all hold, the allocation status of an allocatable array in the second region is implementation defined (see Section 2.14.2 on page 126).
- **shared clause**: passing a shared variable to a non-intrinsic procedure may result in the value of the shared variable being copied into temporary storage before the procedure reference, and back out of the temporary storage into the actual argument storage after the procedure reference. Situations where this occurs other than those specified are implementation defined (see Section 2.14.3.2 on page 133).
- Runtime library definitions: it is implementation defined whether the include file omp\_lib.h or the module omp\_lib (or both) is provided. It is implementation defined whether any of the OpenMP runtime library routines that take an argument are extended with a generic interface so arguments of different KIND type can be accommodated (see Section 3.1 on page 160).

Fortran

### 1 APPENDIX **F**

# **Features History**

This appendix summarizes the major changes between recent versions of the OpenMP API since version 2.5.

### F.1 Version 3.1 to 4.0 Differences

- The **taskgroup** construct (see Section 2.12.5 on page 102) was added to support more flexible deep task synchronization.
- The **reduction** clause (see Section 2.14.3.6 on page 142) was extended to support user defined reductions.
- The **declare reduction** construct (see Section 2.15 on page 153) was added to allow user defined reductions.
- The proc\_bind clause (see Section 2.5.2 on page 42) and the OMP\_PLACES environment variable (see Section 4.5 on page 213) were added to support thread affinity policies.
- Eliminated possibility of implementation defined task scheduling points for untied tasks (see Section 2.11.3 on page 94).
- Added the concept of SIMD parallelism and associated constructs (see Section 2.8 on page 60).

## F.2 Version 3.0 to 3.1 Differences

• The **final** and **mergeable** clauses (see Section 2.11.1 on page 88) were added to the **task** construct to support optimization of task data environments.

- 1 • The taskyield construct (see Section 2.11.2 on page 92) was added to allow user-2 defined task scheduling points. 3 • The atomic construct (see Section 2.12.6 on page 103) was extended to include read, write, and capture forms, and an update clause was added to apply 4 the already existing form of the atomic construct. 5 6 • Data environment restrictions were changed to allow intent(in) and const-7 qualified types for the **firstprivate** clause (see Section 2.14.3.4 on page 137). 8 Data environment restrictions were changed to allow Fortran pointers in 9 firstprivate (see Section 2.14.3.4 on page 137) and lastprivate (see 10 Section 2.14.3.5 on page 139). 11 • New reduction operators min and max were added for C and C++ 12 The nesting restrictions in Section 2.16 on page 158 were clarified to disallow 13 closely-nested OpenMP regions within an atomic region. This allows an atomic 14 region to be consistently defined with other OpenMP regions so that they include all the code in the atomic construct. 15 16 added to support specialization of final task regions. 17 18
  - The omp in final runtime library routine (see Section 3.2.21 on page 188) was
  - The nthreads-var ICV has been modified to be a list of the number of threads to use at each nested parallel region level. The value of this ICV is still set with the **OMP NUM THREADS** environment variable (see Section 4.2 on page 211), but the algorithm for determining the number of threads used in a parallel region has been modified to handle a list (see Section 2.5.1 on page 40).
  - The bind-var ICV has been added, which controls whether or not threads are bound to processors (see Section 2.3.1 on page 31). The value of this ICV can be set with the **OMP PROC BIND** environment variable (see Section 4.4 on page 213).
  - Descriptions of examples (see Appendix A on page 221) were expanded and clarified.
  - Replaced incorrect use of omp integer kind in Fortran interfaces (see Section D.3 on page 392 and Section D.4 on page 396) with selected int kind(8).

#### **F**.3 Version 2.5 to 3.0 Differences

The concept of tasks has been added to the OpenMP execution model (see Section 1.2.4 on page 8 and Section 1.3 on page 13).

- The task construct (see Section 2.11 on page 88) has been added, which provides a mechanism for creating tasks explicitly.
- The taskwait construct (see Section 2.12.4 on page 101) has been added, which causes a task to wait for all its child tasks to complete.

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1 2 3	<ul> <li>The OpenMP memory model now covers atomicity of memory accesses (see Section 1.4.1 on page 15). The description of the behavior of volatile in terms of flush was removed.</li> </ul>
4 5 6 7 8 9	• In Version 2.5, there was a single copy of the nest-var, dyn-var, nthreads-var and run-sched-var internal control variables (ICVs) for the whole program. In Version 3.0, there is one copy of these ICVs per task (see Section 2.3 on page 30). As a result, the omp_set_num_threads, omp_set_nested and omp_set_dynamic runtime library routines now have specified effects when called from inside a parallel region (see Section 3.2.1 on page 162, Section 3.2.7 on page 169 and Section 3.2.10 on page 173).
11 12 13	• The definition of active <b>parallel</b> region has been changed: in Version 3.0 a <b>parallel</b> region is active if it is executed by a team consisting of more than one thread (see Section 1.2.2 on page 2).
14 15	• The rules for determining the number of threads used in a parallel region have been modified (see Section 2.5.1 on page 40).
16 17	• In Version 3.0, the assignment of iterations to threads in a loop construct with a <b>static</b> schedule kind is deterministic (see Section 2.7.1 on page 47).
18 19 20	• In Version 3.0, a loop construct may be associated with more than one perfectly nested loop. The number of associated loops may be controlled by the <b>collapse</b> clause (see Section 2.7.1 on page 47).
21 22	<ul> <li>Random access iterators, and variables of unsigned integer type, may now be used as loop iterators in loops associated with a loop construct (see Section 2.7.1 on page 47).</li> </ul>
23 24 25	• The schedule kind <b>auto</b> has been added, which gives the implementation the freedom to choose any possible mapping of iterations in a loop construct to threads in the team (see Section 2.7.1 on page 47).
26	• Fortran assumed-size arrays now have predetermined data-sharing attributes (see

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- utes (see Section 2.14.1.1 on page 122).
- In Fortran, firstprivate is now permitted as an argument to the default clause (see Section 2.14.3.1 on page 131).
- For list items in the **private** clause, implementations are no longer permitted to use the storage of the original list item to hold the new list item on the master thread. If no attempt is made to reference the original list item inside the parallel region, its value is well defined on exit from the parallel region (see Section 2.14.3.3 on page 134).
- In Version 3.0, Fortran allocatable arrays may appear in private, firstprivate, lastprivate, reduction, copyin and copyprivate clauses. (see Section 2.14.2 on page 126, Section 2.14.3.3 on page 134, Section 2.14.3.4 on page 137, Section 2.14.3.5 on page 139, Section 2.14.3.6 on page 142, Section 2.14.4.1 on page 148 and Section 2.14.4.2 on page 149).
- In Version 3.0, static class members variables may appear in a threadprivate directive (see Section 2.14.2 on page 126).

- Version 3.0 makes clear where, and with which arguments, constructors and destructors of private and threadprivate class type variables are called (see Section 2.14.2 on page 126, Section 2.14.3.3 on page 134, Section 2.14.3.4 on page 137, Section 2.14.4.1 on page 148 and Section 2.14.4.2 on page 149)
- The runtime library routines omp\_set\_schedule and omp\_get\_schedule have been added; these routines respectively set and retrieve the value of the run-sched-var ICV (see Section 3.2.12 on page 176 and Section 3.2.13 on page 178).
- The *thread-limit-var* ICV has been added, which controls the maximum number of threads participating in the OpenMP program. The value of this ICV can be set with the OMP\_THREAD\_LIMIT environment variable and retrieved with the omp\_get\_thread\_limit runtime library routine (see Section 2.3.1 on page 31, Section 3.2.14 on page 179 and Section 4.10 on page 218).
- The max-active-levels-var ICV has been added, which controls the number of nested active parallel regions. The value of this ICV can be set with the OMP\_MAX\_ACTIVE\_LEVELS environment variable and the omp\_set\_max\_active\_levels runtime library routine, and it can be retrieved with the omp\_get\_max\_active\_levels runtime library routine (see Section 2.3.1 on page 31, Section 3.2.15 on page 180, Section 3.2.16 on page 182 and Section 4.9 on page 217).
- The *stacksize-var* ICV has been added, which controls the stack size for threads that the OpenMP implementation creates. The value of this ICV can be set with the **OMP\_STACKSIZE** environment variable (see Section 2.3.1 on page 31 and Section 4.7 on page 216).
- The wait-policy-var ICV has been added, which controls the desired behavior of waiting threads. The value of this ICV can be set with the OMP\_WAIT\_POLICY environment variable (see Section 2.3.1 on page 31 and Section 4.8 on page 217).
- The omp\_get\_level runtime library routine has been added, which returns the number of nested parallel regions enclosing the task that contains the call (see Section 3.2.17 on page 183).
- The omp\_get\_ancestor\_thread\_num runtime library routine has been added, which returns, for a given nested level of the current thread, the thread number of the ancestor (see Section 3.2.18 on page 184).
- The omp\_get\_team\_size runtime library routine has been added, which returns, for a given nested level of the current thread, the size of the thread team to which the ancestor belongs (see Section 3.2.19 on page 185).
- The omp\_get\_active\_level runtime library routine has been added, which returns the number of nested, active parallel regions enclosing the task that contains the call (see Section 3.2.20 on page 187).
- In Version 3.0, locks are owned by tasks, not by threads (see Section 3.3 on page 196).

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