# OpenMP Loop Parallelism (2A)

- Loop
- •

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The simplest way to create parallelism in OpenMP is to use the **parallel** pragma.

A block preceded by the omp **parallel** pragma is called a **parallel region** ;

it is executed by a <u>newly created team</u> of threads.

This is an instance of the **SPMD** model: <u>all</u> **threads** execute the <u>same</u> segment of code.

```
#pragma omp parallel
{
    // this is executed by a team of threads
}
```

It would be pointless to have the block be executed identically by all threads.

One way to get a meaningful parallel code is to use the function **omp\_get\_thread\_num** , to find out which thread you are, and execute work that is individual to that thread.

There is also a function **omp\_get\_num\_threads** to find out the total number of threads.

Both these functions give a number relative to the current team; recall from figure 15.3 that new teams can be created recursively.

Immediately preceding the **parallel block**, <u>one</u> **thread** will be executing the code.

In the **main** program this is the **initial thread**.

At the <u>start</u> of the block, <u>a new team</u> of **threads** is <u>created</u>, and the **thread** that was <u>active before</u> the block becomes the **master thread** of that team.

After the block, only the master thread is active.

Inside the block there is team of threads:

each **thread** in the **team** <u>executes</u> the body of the block, and it will have <u>access</u> to all variables of the surrounding environment.

How many threads there are can be determined in a number of ways;

the **threads** that are <u>forked</u> are all **copies** of the **master thread** : they have <u>access</u> to all that was computed so far; this is their **shared data** 

Of course, if the **threads** were completely <u>identical</u> the parallelism would be pointless, so they also have **private data**, and they can <u>identify</u> themselves: they know their **thread number**.

this allows you to do meaningful parallel computations with threads.

### work sharing constructs In a team of threads,

<u>initially</u> there will be <u>replicated</u> <u>execution</u>; a **work sharing construct** <u>divides</u> available parallelism over the threads.

OpenMP uses **teams** of **threads**, and inside a **parallel region** the **work** is <u>distributed</u> over the **threads** with a **work sharing construct**. **threads** can <u>access</u> **shared data**, and they have some **private data**.

An important difference between **OpenMP** and **MPI** is that parallelism in **OpenMP** is **dynamically activated** by a **thread spawning** a **team** of **threads**.

Furthermore, the **number** of **threads** used can <u>differ</u> between **parallel regions**, and **threads** can <u>create</u> **threads** <u>recursively</u>.

This is known as as dynamic mode .

By contrast, in an **MPI** program the **number** of running processes is (mostly) <u>constant</u> throughout the run, and determined by factors <u>external</u> to the program.

OpenMP parallel loops are an example of OpenMP `worksharing' constructs

take an amount of work and distribute it over the available threads in a parallel region.

The parallel execution of a loop can be handled a number of different ways.

For instance, you can create a parallel region around the loop, and adjust the **loop bounds**:

#### #pragma omp **parallel**

ł

}

```
int threadnum = omp_get_thread_num(),
    numthreads = omp_get_num_threads();
```

int low = N\*threadnum/numthreads, high = N\*(threadnum+1)/numthreads;

for (i=**low**; i<**high**; i++) // do something with i

use the **parallel for** pragma:

```
#pragma omp parallel
#pragma omp for
for (i=0; i<N; i++) {
    // do something with i
}</pre>
```

you don't have to calculate the loop bounds for the threads yourself,

but you can also tell OpenMP to assign the loop iterations according to different schedules

```
#pragma omp parallel
{
    code1();
    #pragma omp for
    for (i=1; i<=4*N; i++) {
        code2();
    }
    code3();
}</pre>
```

The code before and after the loop is executed identically in each thread; t

he loop iterations are spread over the four threads.

### Without #pragma omp for

code1	code1	code1	code1
code2	code2	code2	code2
(1 :N)	(1 :N)	(1 :N)	(1 :N)
code2	code2	code2	code2
(N+1	(N+1	(N+1	(N+1
:2N)	:2N)	:2N)	:2N)
code2	code2	code2	code2
(2N+1	(2N+1	(2N+1	(2N+1
:3N)	:3N)	:3N)	:3N)
code2	code2	code2	code2
(3N+1	(3N+1	(3N+1	(3N+1
:4N)	:4N)	:4N)	:4N)
code3	code3	code3	code3

#### With #pragma omp for



Note that the **parallel do** and **parallel for** pragmas do not create a team of threads: they take the team of threads that is active, and divide the loop iterations over them.

This means that the **omp for** or **omp do** directive needs to be <u>inside</u> a **parallel region**. It is also possible to have a <u>combined</u> **omp parallel for** or **omp parallel do** directive.

If your parallel region only contains a loop, you can combine the pragmas for the parallel region and distribution of the loop iterations:

#pragma omp parallel for
for (i=0; .....



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#pragma omp parallel for
for (i=0; .....

# Loop Schedules (1)

more iterations in a loop than **threads** several ways to <u>assign</u> loop iterations to the **threads** OpenMP lets you specify this with the **schedule clause**.

#pragma omp for schedule(....)

#### **Static schedules**

the iterations are assigned purely based on the <u>number of iterations</u> and the <u>number of threads</u> (and the **chunk parameter**; see later).

#### **Dynamic schedules**

iterations are assigned to threads that are <u>unoccupied</u>.

when iterations take an <u>unpredictable</u> amount of time, so **load balancing** is needed.

### Loop Schedules (2)



W4	W5
w3	W6
W2	W7
W1	W8

https://pages.tacc.utexas.edu/~eijkhout/pcse/html/omp-loop.html

### **OpenMP Loop (2A)**

15

Young Won Lim 2/13/21

# Loop Schedules (3)

Static													
	thr0 thr1 thr2 thr3												
Static,	Static, n												
thr0	thr1	thr2	thr3	thr0	thr1	thr2	thr3	thrO	thr1	thr2			
Dynam	nic												
thr0	thr1	thr2	thr3	thr1	thr0	thr2	thr1	thr3	thr2	thr1			
Guideo	ł												
thrO	thi	1	thr2	thr3	thrO	thr1	thr2	thr3	thrO	thr1			

# Loop Schedules (4)

assume that each core gets assigned two (blocks of) iterations and these blocks take gradually less and less time.

thread 1 gets two fairly long blocks, whereas thread 4 gets two short blocks,

Thread 1 finishes much earlier. Imbalance : unequal amounts of work

#### load balancing

thread 4 gets block 5, since it finishes the first set of blocks early. The effect is a perfect





## Loop Schedule – Static (1)

The **default static schedule** is to assign one consecutive **block** of iterations to each **thread**.

#pragma omp for schedule(static)
#pragma omp for schedule(static, chunk)

With **static scheduling**, the compiler will <u>split up</u> the loop iterations <u>at compile time</u>,

When the iterations take <u>roughly</u> the <u>same</u> amount of time, this is the most <u>efficient</u> at runtime.

#### Static

### Loop Schedule – Static (2)

#pragma omp for schedule(static,chunk)

If you want <u>different</u> <u>sized blocks</u> you can defined a **chunk size chunk** 

The choice of a **chunk size** is often a balance between the **low overhead** of having only <u>a few chunks</u>, (big chunks) versus the **load balancing effect** of having <u>smaller chunks</u>. (many chunks)



https://pages.tacc.utexas.edu/~eijkhout/pcse/html/omp-loop.html

Static, n

### Loop Schedule – Static (3)

#pragma omp for schedule(static,chunk)

OpenMP <u>divides</u> the iterations into chunks of size <u>chunk-size</u>

distributes the chunks to **threads** in a circular order.

When <u>no</u> chunk-size is specified, OpenMP divides iterations into chunks that are approximately <u>equal in size</u> and distributes <u>at most</u> one **chunk** to each **thread**.





### Loop Schedule – Static (4)

Static scheduling is used when you know that each thread will do the approximately same amount of work at the **compile time**.

the following code can be parallelized using OMP. (assume only 4 threads)

```
float A[100][100];
```

```
for(int i = 0; i < 100; i++)
{
    for(int j = 0; j < 100; j++)
    {
        A[i][j] = 1.0f;
    }
}</pre>
```

100 \* 100 iterations 10000 / 4 iterations / thread



## Loop Schedule – Static (5)

### float A[100][100];

```
#pragma omp for schedule(static)
```

```
for(int i = 0; i < 100; i++)
{
    for(int j = 0; j < 100; j++)
    {
        A[i][j] = 1.0f;
    }
}</pre>
```

i	i	i	i
(0:24)	(25:49)	(50:74)	(75 :99)
j	j	j	j
(0:99)	(0:99)	(0:99)	(0:99)

to use the default static scheduling, place pragma on the <u>outer for loop</u>,

then each thread will do 25% of the outer loop (i) work and eqaul amount of inner loop (j) work

Hence, the total amount of work done by each thread is <u>same</u>.

Hence, we could simply stick with the default static scheduling to give optimal load balancing.

https://stackoverflow.com/questions/15508128/using-omp-schedule-with-pragma-omp-for-parallel-scheduleruntime

#### **OpenMP Loop (2A)**

### Loop Schedule – Static (6)

a for loop with 64 iterations 4 threads each row represents a thread.

**schedule(static)** has 16 iterations in the first row. 64 iterations and 4 threads  $\rightarrow 64 / 4 = 16$ 

the first tread executes iterations 1, 2, 3, ..., 15 and 16. the second thread executes iterations 17, 18, 19, ..., 31, 32. Similar applies to the threads three and four.

For **schedule(static)**, OpenMP divides iterations into four chunks of size 16 and it distributes them to four threads.

For schedule(static, 4) and schedule(static, 8)

OpenMP divides iterations into chunks of size 4 and 8, respectively.

The static scheduling type is appropriate when all iterations have the same computational cost.

### Loop Schedule – Static (6)

#### schedule(static, 4)

0	1	2	-3	16	17	18	19	32	33	34	-35	48	49	50	51
4	5	6	-7	20	21	22	23	36	37	38	-39	52	53	54	55
8	9	10	-11	24	25	26	27	40	41	42	-43	56	57	58	59
12	13	14	15	28	29	30	31	44	45	46	-47	60	61	62	63

#### schedule(static, 8)

0	1	2	3	-4	5	6	-7	32	33	34	35	36	37	38	-39
8	9	10	-11	12	13	14	-15	40	41	42	43	44	45	46	47
16	17	18	19	20	21	22	23	48	49	50	51	52	53	54	55
24	25	26	27	28	29	30	-31	56	57	58	59	60	61	62	63

#### schedule(static)

Ο	1	2	2	1	5	6	7	8	g	10	11	12	13	14	_15
U	-	2	J	-	5	U		U	5	TO		12	10	<b>T</b> -	10
16	17	10	10	20	21	ງງ	22	24	25	26	27	20	20	20	21
10	<u>т</u>	то	19	20	<b>Z</b> 1	22	23	24	25	20	21	20	29	30	- <b>3</b> 1
22	22	24	25	26	27	20	20	10	11	12	12	11	15	16	17
32	33	34	35	30	37	30	29	40	41	42	43	44	45	40	47
10	40	50	<b>E</b> 1	<b>E</b> 2	E2	<b>E</b> 1	55	FG	57	БО	50	60	61	60	62
40	49	50	ЭТ	52	55	54	55	50	57	50	29	00	01	02	03

http://jakascorner.com/blog/2016/06/omp-for-scheduling.html

### **OpenMP Loop (2A)**

# Loop Schedule – Dynamic (1)

In **dynamic scheduling** OpenMP will put **blocks** of iterations in a **task queue**, (the **default chunk size** is **1**) and the **threads** take **one** of these tasks whenever they are finished with the previous.

### #pragma omp for schedule(dynamic[,chunk])

While this schedule may give **good load balancing** if the iterations take very <u>differing</u> amounts of time to execute, it does carry **runtime overhead** for managing the **queue** of iteration tasks.

#### Dynamic

thr0	thr1	thr2	thr3	thr1	thr0	thr2	thr1	thr3	thr2	thr1
------	------	------	------	------	------	------	------	------	------	------

# Loop Schedule – Dynamic (2)

<u>large</u> chunks carry the <u>least</u> overhead, but <u>smaller</u> chunks are <u>better</u> for **load balancing**.

If you don't want to decide on a schedule in your code, you can specify the schedule will then **at runtime** be read from the OMP\_SCHEDULE environment variable.

You can even just leave it to the runtime library by specifying **omp\_set\_schedule**.

#### Dynamic

thr0	thr1	thr2	thr3	thr1	thr0	thr2	thr1	thr3	thr2	thr1
------	------	------	------	------	------	------	------	------	------	------

## Loop Schedule – Dynamic (3)

#### for schedule(dynamic, chunk-size)

the dynamic scheduling type

OpenMP divides the iterations into chunks of size chunk-size.

Each thread <u>executes</u> a chunk of iterations and then <u>requests</u> another chunk <u>until</u> there are <u>no</u> more chunks available. There is <u>no particular order</u> in which the chunks are distributed to the threads.

The order changes each time when we execute the for loop.

If we do <u>not</u> specify **chunk-size**, it defaults to <u>one</u>.

#### Dynamic

thr0	thr1	thr2	thr3	thr1	thr0	thr2	thr1	thr3	thr2	thr1
------	------	------	------	------	------	------	------	------	------	------

## Loop Schedule – Dynamic (4)

**Dynamic scheduling** is used when you know that each thread will <u>not</u> do <u>same</u> amount of work by using **static scheduling**.

float A[100][100];

```
for(int i = 0; i < 100; i++)
{
    for(int j = 0; j < i; j++)
    {
        A[i][j] = 1.0f;
    }
}</pre>
```







# Loop Schedule – Dynamic (5)

### float A[100][100];

```
for(int i = 0; i < 100; i++)
{
    for(int j = 0; j < i; j++)
    {
        A[i][j] = 1.0f;
    }
}</pre>
```



The inner loop variable **j** is <u>dependent</u> on the **i**.

### the default static scheduling,

- the outer loop (i) work might be divided equally between the 4 threads,
- but the inner loop (j) work will be large for some threads.
- not equal amount of work
- not optimal load balancing



# Loop Schedule – Dynamic (6)

the dynamic scheduling

this scheduling is done at the run time can make sure optimal **load balance**.

Note:

you can also specify the chunk\_size for scheduling. It depends on the **loop size**.

# Loop Schedule – Dynamic (7)

for **schedule(dynamic)** and **schedule(dynamic, 1)** OpenMP determines similar scheduling. the size of chunks is equal to **1** in both instances. the distribution of chunks between the threads is <u>arbitrary</u>.

For **schedule(dynamic, 4)** and **schedule(dynamic, 8)** OpenMP divides iterations into chunks of size **4** and **8**, respectively. the distribution of chunks to the threads has <u>no pattern</u> (abitrary)

The dynamic scheduling type is appropriate when the iterations require different computational costs. I.e, when the iterations are poorly balanced between each other.

the dynamic scheduling type has higher **overhead** than the static scheduling type because it dynamically distributes the iterations during the runtime.

# Loop Schedule – Dynamic (6)

schedule(dynamic, 4) -14 15 -39 

schedule(dynamic, 8)

	16	17	18	19	20	21	22	23
	24	25	26	27	28	29	30	-31
•	0	1	2	3	-4	5	6	-7
	8	9	10	11	12	13	14	15

	40	41	42	43	44	45	46	47
•	32	33	34	35	36	37	38	<del>3</del> 9
	56	57	58	59	60	61	62	63
	48	49	50	51	52	53	54	55

### schedule(dynamic)

16	A	50	35	-36	5	22	λ	56	41	10	27	60	61	14	15
48	33	34	51	52	37	<del>3</del> 8	55	40	57	42	59	44	29	46	<b>-</b> 47
0	49	2	19	20	21	6	23	8-	_9	58	11	28	13	30	63
32	17-	18	3	-4	53	54	39	24	25	26	43	12	45	62	31

# Loop Schedule – Guided (1)

The **guided** scheduling type is similar to the **dynamic** scheduling type.

OpenMP again divides the iterations into chunks.

Each thread executes a chunk of iterations and then requests another chunk <u>until</u> there are no more chunks available. The difference is in the size of chunks.

The size of a chunk is proportional to the number of unassigned iterations divided by the number of the threads.

Therefore **the size of the chunks** <u>decreases</u> as the execution goes on

#### Guided

thr0 thr1 thr2 thr3 thr0 thr1 thr2 thr3 thr0 thr1 thr2 thr3 thr0 th	thr0	1 thr2	thr3	thrO	thr1	thr2	thr3	thr0 t	thr1
---	------	--------	------	------	------	------	------	--------	------

# Loop Schedule – Guided (2)

### The minimum size of a chunk

is set by '**chunk-size'** in the scheduling clause:

### for schedule(guided, chunk-size).

the chunk which contains the last iterations may have <u>smaller</u> size than chunk-size.

If we do not specify **chunk-size**, it defaults to **one**.

#### Guided

thr0 thr1 thr2 thr3 thr0 thr1 thr2 thr3 t	r0 thr1
---	---------

## Loop Schedule – Guided (3)

the **size** of the **chunks** is <u>decreasing</u>. first chunk has always 16 iterations. 64 iterations and 4 threads  $\rightarrow 64 / 4 = 16$ 

the **minimum chunk size** is determined in the schedule clause. The only exception is the **last chunk**. Its size might be lower then the prescribed minimum size.

The guided scheduling type is appropriate when the iterations are poorly balanced between each other.

The initial chunks are larger, because they reduce overhead. The smaller chunks fills the schedule towards the end of the computation and improve load balancing. This scheduling type is especially appropriate when poor load balancing occurs toward the end of the computation.

# Loop Schedule – Guided (3)

schedule	e(guided)	schedule	(guided, 4)	schedule(guided, 8)					
64	64 / 4 = 16	64	64 / 4 = 16	64	64 / 4 = 16				
(64-16)	48 / 4 = 12	(64-16)	48 / 4 = 12	(64-16)	48 / 4 = 12				
(48-12)	36 / 4 = 9	(48-12)	36 / 4 = 9	(48-12)	36 / 4 = 9				
(36-9)	27 / 4 = 7	(36-9)	27 / 4 = 7	(36-9)	27 / 4 = 7 < 8				
(27-7)	20 / 4 = 5	(27-7)	20 / 4 = 5	(27-8)	19 8				
(20-5)	15 / 4 = 4	(20-5)	15 / 4 = 4	(19-8)	11 8				
(15-4)	11 / 4 = 3	(15-4)	11 / 4 = 3 < 4	(11-8)	3 3				
(11-3)	8 / 4 = 2	(11- <b>4</b> )	7 4	(3-3)	0				
(8-2) (6-2)	6 / 4 = 2 4 / 4 = 1	(7-4) (3-3)	3 3 0	(0 0)	0				
(4-1) (3-1) (2-1)	3 / 4 = 1 2 / 4 = 1 1 / 4 = 1								

### Loop Schedule – Static (6)

scł	nedu	lle <b>(s</b>	tatic	:)													schedu	le(guided)	
0	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		64	64 / 4 = 16	
32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47		(64-16)	48 / 4 = 12	
48	49	50	51	52	53	54	55	56	57	58	59	60	61	62	63		(48-12)	36 / 4 = 9	
																	(36-9)	27 / 4 = 7	
																	(27-7)	20 / 4 = 5	
	_	-	-	-	_		_										(20-5)	15/4 - 4	
0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	44	45_46	47 48 58 59	63
16	17	18	19	20	21	22	23	24	25	26	27	49	50	51	52	60	(11-3)	8 / 4 = 2	
28	29	30	31	32	33	34	35	36	53	54	55	61					(8-2)	6 / 4 = 2	
37	38	39	40	41	42	43	56	57	62								(6-2)	4 / 4 = 1	
										-							(4-1)	3 / 4 = 1	
																	(3-1)	2 / 4 = 1	
0	1	2	3	Δ	5	6	7	8	9	10	11	12	13	14	15	63	(2-1)	1 / 4 = 1	
16	17	 18	10	20	21	22	23	24	25	26	27	53	54	55	60	00			
28	20	30	21	20	23	21	25	24	2J	50	51	52	56	57	61				
20	29	30	JT .	52	33	54	35	30	49	50	JT	JZ	50	57	01				
37	38	39	40	41	42	43	44	45	46	47	48	58	59	62					

### Loop Schedule – Static (6)

sch	nedu	le <b>(s</b>	tatic	:)													schedule	e(guic	led, 4)
0	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		64	64 /	/ 4 = 16
32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47		(64-16)	48 /	/ 4 = 12
48	49	50	51	52	53	54	55	56	57	58	59	60	61	62	63		(48-12)	36 /	/ 4 = 9
																	(36-9)	27 /	/ 4 = 7
																	(27-7)	20 /	/ 4 = 5
																	(20-5)	15	14 - 1
0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	44	45 46	471	48 61 <mark>&lt; /</mark>
16	17	18	19	20	21	22	23	24	25	26	27	49	50	51	52	62	(11-4)	7	4
28	29	30	31	32	33	34	35	36	53	54	55	56	63				(7-4)	3	3
37	38	39	40	41	42	43	57	58	59	60							(3-3)	0	-

http://jakascorner.com/blog/2016/06/omp-for-scheduling.html

38

### Loop Schedule – Auto

The auto scheduling type <u>delegates</u> the decision of the scheduling to the **compiler** and/or **runtime** system.



# Loop Schedule – Runtime

The schedule(runtime) clause tells it to set the schedule using the environment variable.

The environment variable can be set to any other scheduling type.

It can be set by

setenv OMP\_SCHEDULE "dynamic,5"





### Loop Schedules – Runtime

The **runtime** scheduling type <u>defers</u> the decision about the scheduling until the **runtime**.

different ways of specifying the scheduling type in this case.

One option is with the environment variable **OMP\_SCHEDULE** 

and the other option is with the function omp\_set\_schedule.

If the scheduling-type (in the schedule clause of the loop construct) is equal to runtime then OpenMP determines the scheduling by the internal control variable run-sched-var. We can set this variable by setting the environment variable OMP\_SCHEDULE to the desired scheduling type. For example, in bash-like terminals, we can do

\$ export OMP\_SCHEDULE=sheduling-type

Another way to specify run-sched-var is to set it with omp\_set\_schedule function.

omp\_set\_schedule(sheduling-type);

http://jakascorner.com/blog/2016/06/omp-for-scheduling.html

. . .

# Nested Parallelism (1)

```
void fun1()
{
  for (int i=0; i<80; i++)
      . . .
}
main()
  #pragma omp parallel
    #pragma omp for
    for (int i=0; i<100; i++)
        - - -
    #pragma omp for
    for (int i=0; i<10; i++)
       fun1();
  }
```

the 2nd loop in **main** can only be distributed to **10** threads

80 loop iterations in fun1which will be called 10 times in main loop.

total 800 iterations in fun1 and the main loop

This gives much more parallelism potential if parallelism can be added in both levels.



# Nested Parallelism (2)



may either have <u>insufficient threads</u> for the 1st main loop as it has <u>larger loop count</u>, or

create exploded number of threads for the 2nd main loop when OMP\_NESTED=TRUE.

The simple solution is to <u>split</u> the parallel region in main and create separate ones for each loop with a distinct thread number specified.

# Nested Parallelism (3)



don't have to worry about the thread number changes in 1st and 2nd main loops.

Even though you still have a small amount of (10) threads allocated for 2nd main loop, the rest available threads will be able to be distributed through omp **taskloop** in fun1.



### Nested Parallelism (4)

**nested parallel regions** is a way to <u>distribute</u> **tasks** by <u>creating</u> / <u>forking</u> more **threads**.

parallel region is the only construct determines execution thread <u>number</u> and <u>controls</u> thread affinity

Using **nested parallel regions** means each **thread** in **parent region** will yield multiple **threads** in enclosed regions, which in turn create a <u>product</u> of **thread number**.



### Nested Parallelism (5)

**omp tasking** shows another way to explore parallelism by adding more **tasks**, instead of **threads**.

though the **thread number** is <u>unchanged</u> as specified <u>at the entry</u> of the **parallel region**, the <u>increased</u> **tasks** from the **nested tasking** constructs can be <u>distributed</u> and <u>executed</u> by any available/idle **threads** in the current team of the same parallel region.

This gives opportunities to fully use all threads' capability, and improve balance of workloads automatically.

# Implicit task (1)

In addition to **explicit tasks** specified using the **task** directive, the OpenMP specification version 3.0 introduces the notion of **implicit tasks**.

An **implicit task** is a task generated

- by the implicit parallel region,
- when a **parallel construct** is encountered during execution.

The **code** for each **implicit task** is the code <u>inside</u> the **parallel construct**.

Each implicit task is

- assigned to a <u>different</u> thread in the team and is tied;
- <u>always executed</u> from <u>beginning</u> to <u>end</u> by the **thread** to which it is <u>initially assigned</u>.

https://docs.oracle.com/cd/E19205-01/820-7883/6nj43o69j/index.html

# Implicit task (2)

All **implicit tasks** <u>generated</u> when a **parallel construct** is encountered are guaranteed to be <u>complete</u> when the **master thread** exits the **implicit barrier** at the <u>end</u> of the **parallel region**.

all **explicit tasks** generated <u>within</u> a **parallel region** are guaranteed to be <u>complete</u> on <u>exit</u> from the <u>next</u> **implicit** or **explicit barrier** <u>within</u> the **parallel region**.

https://docs.oracle.com/cd/E19205-01/820-7883/6nj43o69j/index.html

# Implicit task (3)

When an **if clause** is present on a **task construct** and the value of the scalar-expression evaluates to **false**, the thread that encounters the task <u>must immediately</u> execute the task.

The **if clause** can be used to <u>avoid</u> the **overhead** of generating many **finely grained tasks** and placing them in the **conceptual pool**.

https://docs.oracle.com/cd/E19205-01/820-7883/6nj43o69j/index.html

# Implicit barrier

Implicit BarriersSeveral OpenMP\* constructs have implicit barriers

- parallel
- for
- single

Unnecessary barriers hurt performance

• Waiting threads accomplish no work!

Waiting threads accomplish no work! Suppress implicit barriers, when safe, with the nowait

https://www.intel.com/content/dam/www/public/apac/xa/en/pdfs/ssg/Programming\_with\_OpenMP-Linux.pdf

### References

- [1] en.wikipedia.org
- [2] M Harris, http://beowulf.lcs.mit.edu/18.337-2008/lectslides/scan.pdf