

Programming Techniques for Supercomputers: Shared-memory parallel processing with OpenMP

Getting Started

Data Scoping

Worksharing

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Data Scoping

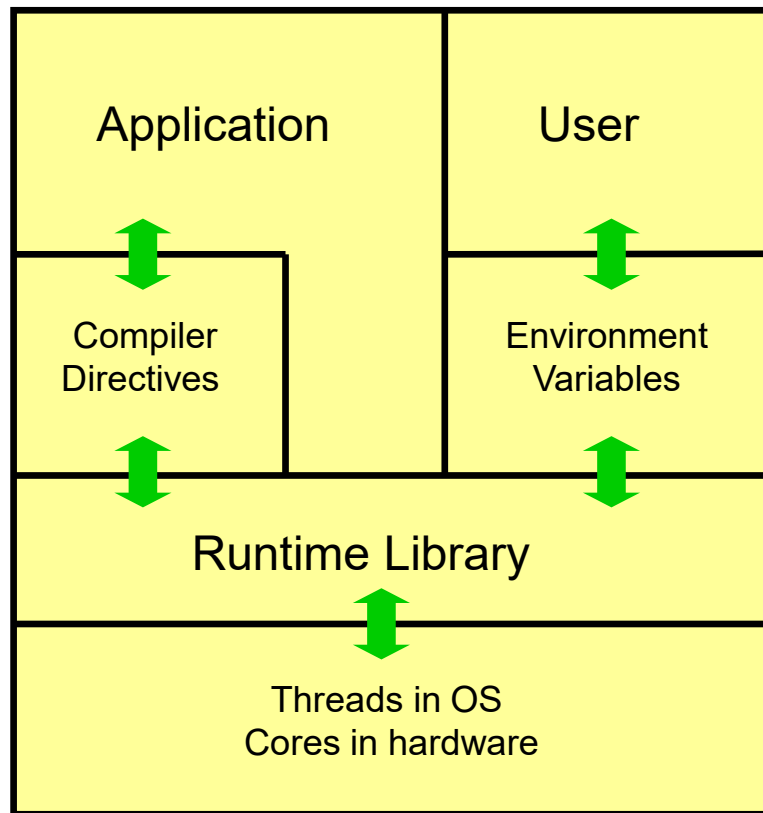
Worksharing



Introduction to OpenMP: Basics

- “Easy,” incremental and portable parallel programming of shared-memory computers: **OpenMP**
- **Original** design goal: Data-level shared memory parallelism – many **extensions**: Task parallelism, Accelerator offloading, SIMD support,...
- Standardized set of **compiler directives & library functions**:
<http://www.openmp.org/>
 - FORTRAN, C and C++ interfaces are defined
 - Supported by all current compilers
 - Free tools are available
- B. Chapman, G. Jost, R. v. d. Pas: Using OpenMP. MIT Press, 2007, ISBN 978-0262533027
- R. v. d. Pas, E. Stotzer, C. Terboven: Using OpenMP – The Next Step. MIT Press, 2017, ISBN 978-0-262-53478-9

Introduction to OpenMP: Software Architecture

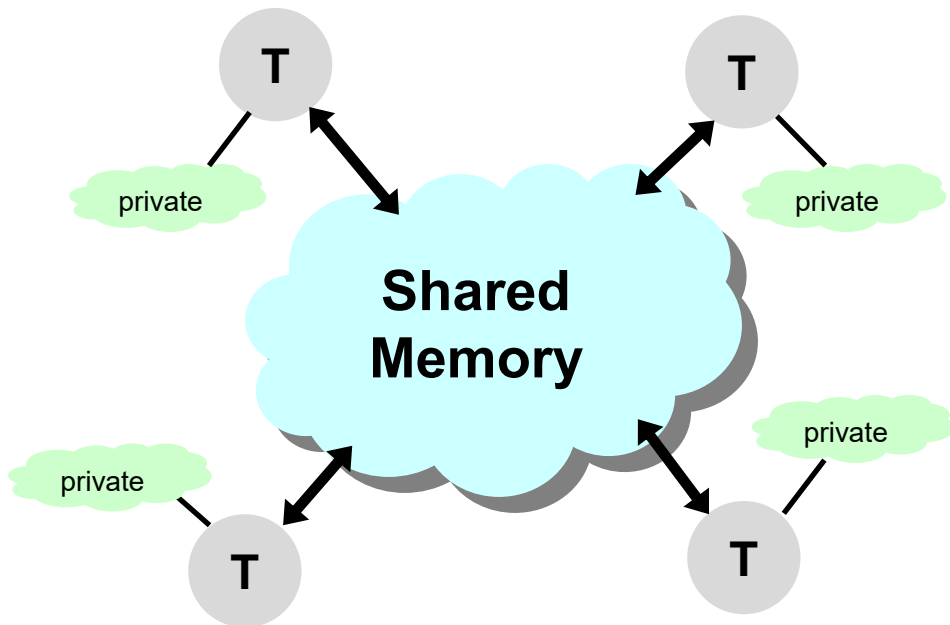


- Programmer's view:
 - **Directives/pragmas** in application code
 - (A few) **library routines**
- User's view (code execution):
 - **Environment variables** determine:
 - resource allocation
 - scheduling strategies and other (implementation-dependent) behavior
- Operating system view:
 - Parallel work done by OS **threads**

Introduction to OpenMP: shared-memory model

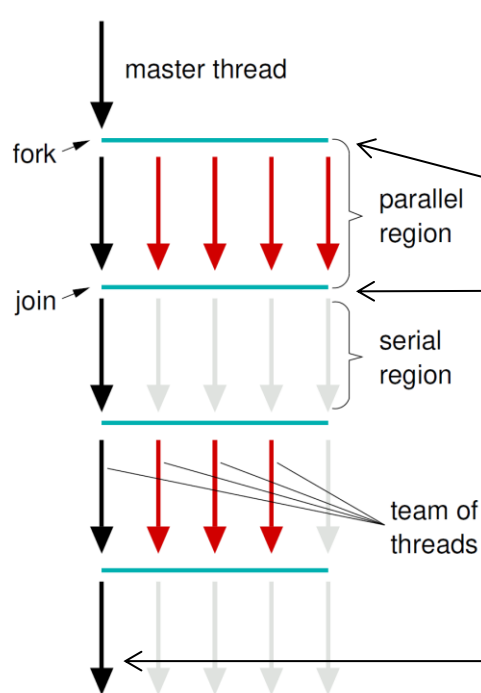
Central concept of OpenMP programming:

Threads



- **Threads:**
 - Spawned by a process
 - Local register set, instruction pointer, stack
 - **Shared global address space**
- **Data scope: shared or private**
 - shared data available to all threads
 - private data only available to thread that owns it
- **Data transfer between threads:**
 - transparent to programmer

Introduction to OpenMP: fork-join execution model



Program start:
one process (master thread) running

Parallel region: **team of threads is generated** (“fork”)
Synchronize when leaving parallel region (“join”)

Serial region:
only master executes

Thread # 0 1 2 3 4

Introduction to OpenMP: General syntax in C/C++

- Compiler directive:

```
#pragma omp [directive [clause ...]]  
    structured block
```

- If OpenMP is not enabled by compiler → treated like comment

- Include file for API calls: `#include <omp.h>`
- Conditional compilation: Compiler's OpenMP switch sets preprocessor macro (acts like `-D_OPENMP`)

```
#ifdef _OPENMP  
    t = omp_get_thread_num();  
#endif
```

Introduction to OpenMP: General syntax in Fortran

- Each directive starts with sentinel in column 1:
 - fixed source: `!$OMP` or `C$OMP` or `*$OMP`
 - free source: `!$OMP`followed by a **directive** and, optionally, **clauses**.
- API calls:
 - F77: include file `omp_lib.h`, F90+: module `omp_lib`
 - Conditional compilation of lines starting with `!$` or `C$` or `*$` to ensure compatibility with sequential execution

- Example:

```
myid = 0
!$ myid = omp_get_thread_num()
numthreads = 1
!$ numthreads = omp_get_num_threads()
```

Introduction to OpenMP: parallel region

- `#pragma omp parallel`

 - structured block**

 - Makes structured block a **parallel region**: All code executed between start and end of this region is executed **by all threads**
 - This includes subroutine calls within the region

```
#pragma omp parallel
  printf("Hello from %d of %d\n",
        omp_get_thread_num(), omp_get_num_threads());
```

API functions



ID of calling
thread 0...n-1

of threads
in region

- **END PARALLEL** required in Fortran

Introduction to OpenMP: compile and run

- Activate OpenMP directives
 - Intel: `-qopenmp`, GCC: `-fopenmp`
- Number of threads: Shell variable `OMP_NUM_THREADS`

```
$ icc -qopenmp hello.c
$ OMP_NUM_THREADS=4 ./a.out
Hello from 0 of 4
Hello from 3 of 4
Hello from 1 of 4
Hello from 2 of 4
```

- Ordering of output is not defined
- Avoid extensive output to stdout in parallel regions!

Shared-memory parallel processing with OpenMP

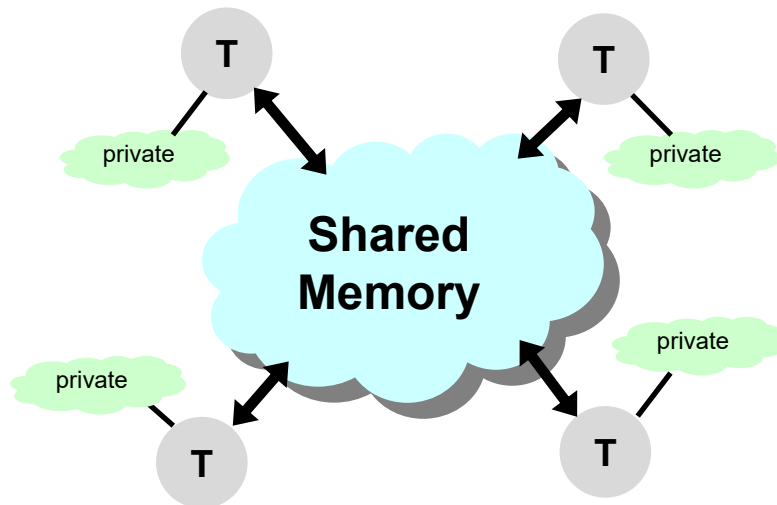
Getting Started
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Data scoping: Shared vs. private data

Data in a **parallel region** can be:

- **private** to each executing thread
→ each thread has its own **local copy** of data
- **shared** between threads
→ there is **only one instance** of data available to all threads
→ this does **not** mean that the instance is always **visible** to **all** threads!

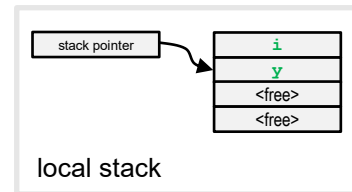
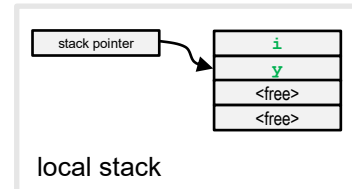
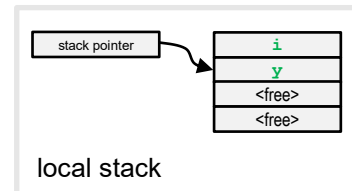
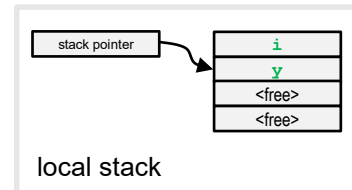
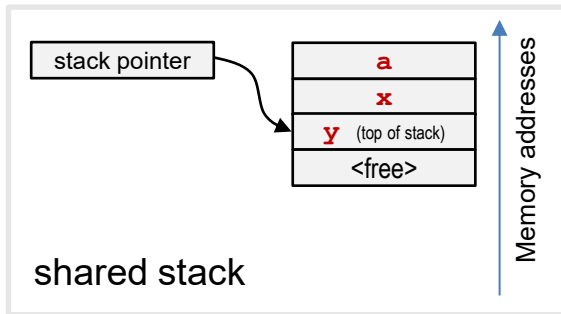


OpenMP **clause** specifies scope of variables:

```
#pragma omp parallel private(var1, tmp) shared(eps)
```

How is private data different from shared data?

```
void f() {  
    int a;  
    float x,y;  
    ...  
    #pragma omp parallel  
    {  
        int i;  
        float y; // masking shared y  
        ...  
    }  
}
```



- Local variables are kept on a stack (last-in first-out memory)
- Every thread has a private stack area
 - i.e., there is one global stack, plus one local stack for each thread
 - Private data goes to private stacks
 - Stack size is limited!

Data scoping: Shared vs. private data

- **Default:** All data in a parallel region is **shared**
This includes global data (global/static variables, C++ class variables)
- **Exceptions:**
 1. **Loop variables** of parallel (“sliced”) loops are **private** (cf. workshare constructs)
 2. **Local (stack) variables** within parallel region
 3. **Local** data within enclosed **function calls** are **private unless** declared **static**
- **Stack size limits** → may be necessary to make large arrays static
 - If not possible → use heap [i.e., `malloc()`, `new[]`, `allocate()`]
 - **OMP_STACKSIZE** shell variable allows to set per-thread stack size

```
$ export OMP_STACKSIZE=100M
```

Data scoping: private data example

C:

```
include <omp.h>
...
int myid = 0, numthreads = 1;
#pragma omp parallel \
    private(myid, numthreads)
{
#ifdef _OPENMP
    myid      = omp_get_thread_num();
    numthreads = omp_get_num_threads();
#endif
    printf("I am %d of %d\n",
          myid, numthreads);
}
```

Fortran 90+:

```
use omp_lib
integer myid, numthreads
...
myid = 0
numthreads = 1
!$omp parallel private(myid,numthreads)
!$ myid = omp_get_thread_num()
!$ numthreads = omp_get_num_threads()
print *, "I am ", myid, &
        " of ", numthreads
!$omp end parallel
```

Data scoping: alternative in C

```
include <omp.h>
...
#pragma omp parallel
{
    int myid = 0, numthreads = 1;
#ifdef _OPENMP
    myid = omp_get_thread_num();
    numthreads = omp_get_num_threads();
#endif
    printf("I am %d of %d\n",
        myid, numthreads);
}
```

Local variables in structured block are automatically private! → less need for private clauses in C

Caveat: local variables are destroyed (go out of scope) at end of block!

Data scoping: important side effects

- What happens if a variable is unintentionally shared?
 - Nothing if it is just read
 - Possibly **hazardous** if **at least one thread writes** to it

```
float x = 0.0;
#pragma omp parallel
{
    x += some_work(...);
}
```

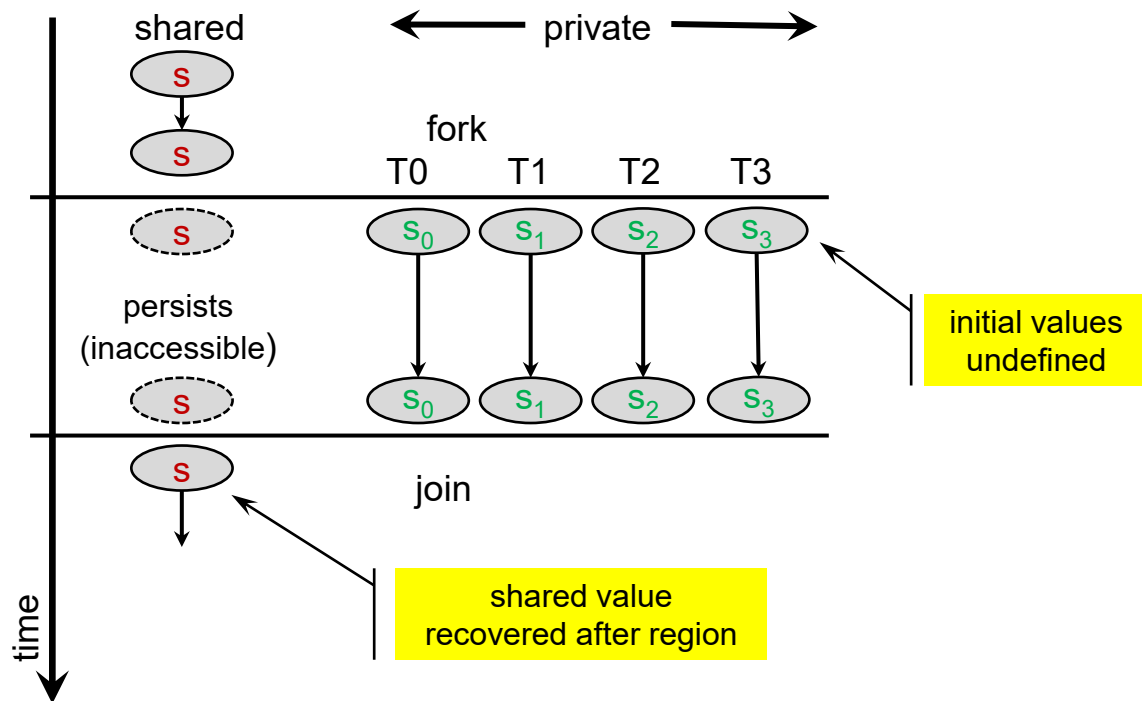
“Race condition”

- Clause for specifying default scope: **default(shared|private|none)**
- **Recommendation: Use**
`#pragma omp parallel default(none)`
 - to not overlook anything
 - compiler complains about every variable that has no explicit scoping attribute

Data scoping: private variables and masking

```
double s;  
  
s = ...;  
#pragma omp parallel private(s)  
{  
    s = ...;  
    ... = ... + s;  
}  
... = ... + s;
```

Masking privatized variables defined in scope outside the parallel region



But what happens if the initial value is required within the parallel region?

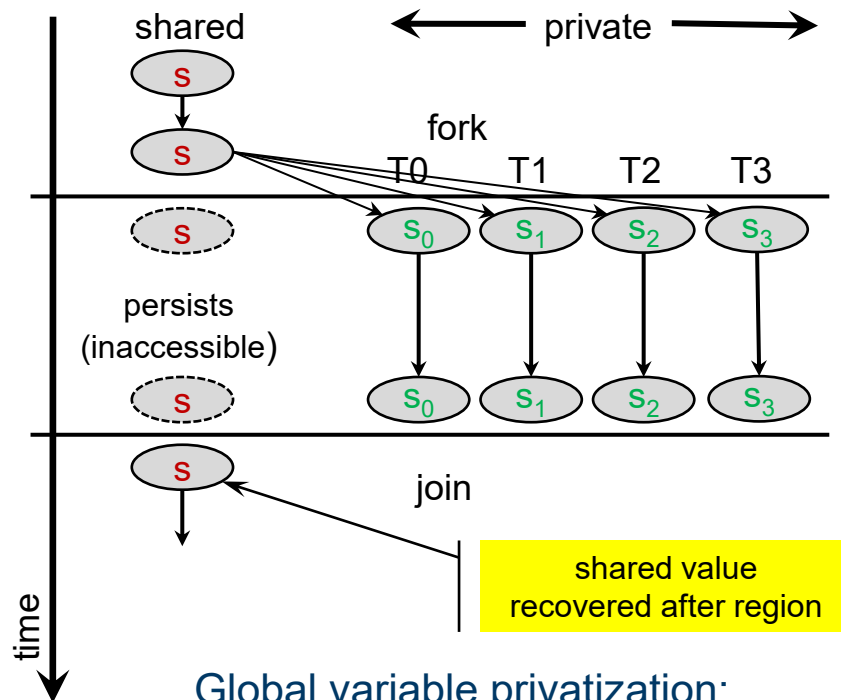
The firstprivate clause

```
double s;  
  
s = ...;  
#pragma omp parallel firstprivate(s)  
{  
    s += ...;  
    ... = ... + s;  
}  
... = ... + s;
```

Extension of private:

value of master copy is transferred to private variables

Restrictions: not a pointer, not assumed shape, not a subobject, master copy not itself private etc.



Global variable privatization:

`threadprivate`, `copyprivate` clauses

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Worksharing: manual loop scheduling

- Work distribution by thread ID
- Only works so easily for canonical loops
- Load balancing very hard
- Complex code

→ don't do it.

```
#include
    int tid, numth, bstart, bend, blen, N;
    double a[N], b[N], c[N], d[N];
    ...
#pragma omp parallel private(tid, numth, bstart, bend, blen)
{
    tid=0; numth=1;
#ifdef _OPENMP
    tid = omp_get_thread_num();
    numth = omp_get_num_threads();
#endif
    blen = N/numth;
    if(tid < N % numth) {
        ++blen; bstart = blen * tid;
    } else
        bstart = blen * tid + N % numth;
    bend=bstart+blen-1;
    for(int i=bstart; i<=bend; ++i)
        a[i] = b[i] + c[i] * d[i];
}
```

One consecutive
chunk of iterations
per thread

Actual work

Worksharing: parallel loop

- `#pragma omp for [clauses]` declares that the following loop iterations are to be distributed among threads
 - Active only if encountered within a parallel region

```
int i, N;
double a[N], b[N], c[N], d[N];
...
#pragma omp parallel          // parallel threads
{
  #pragma omp for            // parallelize loop
  for(i=0; i<N; ++i)
    a[i] = b[i] + c[i] * d[i];
}
```

barriers here!

- Loop counter of parallel loop is declared **private** implicitly
- **Implicit** thread synchronization (**barrier**) at end of `parallel` and at end of `for`
- Fortran: `!$omp do [clauses]`

Worksharing: combined construct

- `#pragma omp parallel for`
structured block

```
int i, N;  
double a[N], b[N], c[N], d[N];  
...  
#pragma omp parallel for  
for(i=0; i<N; ++i)  
    a[i] = b[i] + c[i] * d[i];
```

- Just easier to type...
- Fortran: `!$omp parallel do / $!omp end parallel do`

Worksharing constructs

`#pragma omp for`

- Only the loop **immediately following** the directive is workshared
- Restrictions on parallel loops
 - trip count must be **computable** (**no do ... while**)
 - loop body with **single entry and single exit** point (no breaking out of loop)
- C++ **random access iterator** loops are supported:

```
#pragma omp for
for(auto i=v.begin(); i!=v.end(); ++i) {
    (*i) *= 2.0;
}
```

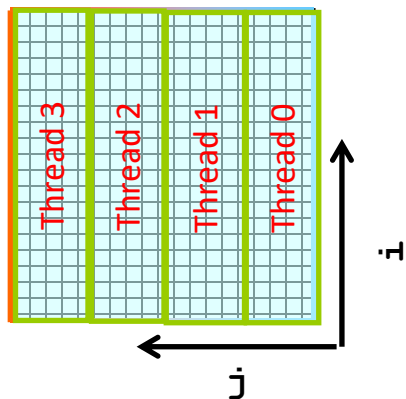
Worksharing constructs in general

- Distribute the execution of the enclosed code region among the members of the team
 - Must be enclosed dynamically **within a parallel region**
 - No implied barrier on entry
 - Implicit barrier at end of worksharing (unless `nowait` clause is specified)
- Directives
 - **for** directive (C/C++), **do** directive (Fortran)
 - **section(s)** directives (ignored here)
 - **workshare** directive (Fortran 90 only – ignored here)

 - **Tasking** (advanced)

Worksharing constructs example

Example: matrix processing with nested loop structure



Only these loops are parallel!

synchronization

```
double a[ndim][ndim], b[ndim][ndim];
...
#pragma omp parallel
{
  #pragma omp for
  for(int j=1; j<ndim-1; ++j) {
    for(int i=1; i<ndim-1; ++i)
      a[j][i] = ( b[j][i+1]+b[j][i-1]
                  +b[j+1][i]+b[j-1][i] ) * 0.25;
  }
  #pragma omp for
  for(int j=1; j<ndim-1; ++j) {
    for(int i=1; i<ndim-1; ++i)
      b[j][i] = ( a[j][i+1]+a[j][i-1]
                  +a[j+1][i]+a[j-1][i] ) * 0.25;
  }
}
```

Some workshare construct clauses

- Examples for workshare construct clauses:
 - `private`, `firstprivate`, `lastprivate`
 - `nowait`
 - `collapse (n)`
 - `schedule (type [, chunk])` [see next slide]
 - `reduction (operator : list)` [see later]
 - There are some more...
- **Implicit barrier** at the end of loop unless `nowait` is specified (barrier may be costly!)
- **collapse**: Fuse nested loops to a single (larger one) and parallelize it
- **schedule** clause specifies how iterations of the loop are distributed among the threads of the team.

Loop worksharing: the `schedule` clause

Within `schedule (type [, chunk])`, `type` can be one of the following:

- **static**: Iterations are divided into pieces of a size specified by `chunk`. The pieces are statically assigned to threads in the team in a round-robin fashion in the order of the thread number.
Default chunk size: one contiguous piece for each thread.
- **dynamic**: Iterations are broken into pieces of a size specified by `chunk`. As each thread finishes a piece of the iteration space, it dynamically obtains the next set of iterations. *Default chunk size: 1.*
- **guided**: The chunk size is reduced in an exponentially decreasing manner with each dispatched piece of the iteration space.
`chunk` specifies the smallest piece (except possibly the last).
Default chunk size: 1. Initial chunk size is implementation dependent.
- **runtime**: The decision regarding scheduling is deferred until run time. The schedule type and chunk size can be chosen at run time by setting the `OMP_SCHEDULE` environment variable.
- **auto**: Compiler/runtime decides
- Default **schedule**: implementation dependent

Loop worksharing: the `schedule` clause

