

# Lecture 12 — Autoparallelization

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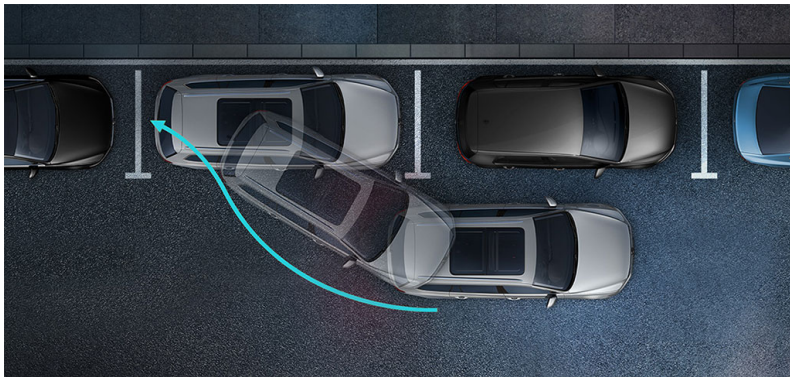


Image Credit: Volkswagen

Wait... autonomous vehicles is a different course...

- An intermediate code used by compilers for analysis and optimization.
- Statements represent one fundamental operation—we can consider each operation **atomic**.
- Statements have the form:  
$$result := operand_1 \operatorname{operator} operand_2$$
- Useful for reasoning about data races,  
and easier to read than assembly.  
(separates out memory reads/writes).

- GIMPLE is the three address code used by gcc.
- To see the GIMPLE representation of your code use the `-fdump-tree-gimple` flag.
- To see all of the three address code generated by the compiler use `-fdump-tree-all`. You'll probably just be interested in the optimized version.
- Use GIMPLE to reason about your code at a low level without having to read assembly.

A feature available as of C99: “The restrict type qualifier allows programs to be written so that translators can produce significantly faster executables.”

- gcc 5+ support C11 or better by default.

`restrict` means: you are promising the compiler that the pointer will never **alias** (another pointer will not point to the same data) for the lifetime of the pointer.

I, [insert your name], a PROFESSIONAL or AMATEUR [circle one] programmer recognize that there are limits to what a compiler can do. I certify that, to the best of my knowledge, there are no magic elves or monkeys in the compiler which through the forces of fairy dust can always make code faster. I understand that there are some problems for which there is not enough information to solve. I hereby declare that given the opportunity to provide the compiler with sufficient information, perhaps through some key word, I will gladly use said keyword and not bitch and moan about how "the compiler should be doing this for me."

In this case, I promise that the pointer declared along with the restrict qualifier is not aliased. I certify that writes through this pointer will not effect the values read through any other pointer available in the same context which is also declared as restricted.

\* Your agreement to this contract is implied by use of the restrict keyword ;)

Pointers declared with `restrict` must never point to the same data.

From Wikipedia:

---

```
void updatePtrs(int* ptrA, int* ptrB, int* val) {  
    *ptrA += *val;  
    *ptrB += *val;  
}
```

---

Would declaring all these pointers as `restrict` generate better code?

Let's look at the GIMPLE:

---

```
void updatePtrs(int* ptrA, int* ptrB, int* val) {  
    D.1609 = *ptrA;  
    D.1610 = *val;  
    D.1611 = D.1609 + D.1610;  
    *ptrA = D.1611;  
    D.1612 = *ptrB;  
    D.1610 = *val;  
    D.1613 = D.1612 + D.1610;  
    *ptrB = D.1613;  
}
```

---

- Could any operation be left out if all the pointers didn't overlap?



---

```
void updatePtrs(int* ptrA, int* ptrB, int* val) {  
    D.1609 = *ptrA;  
    D.1610 = *val;  
    D.1611 = D.1609 + D.1610;  
    *ptrA = D.1611;  
    D.1612 = *ptrB;  
    D.1610 = *val;  
    D.1613 = D.1612 + D.1610;  
    *ptrB = D.1613;  
}
```

---

- If `ptrA` and `val` are not equal, you don't have to reload the data on **line 7**.
- Otherwise, you would: there might be a call  
    `updatePtrs(&x, &y, &x);`

Hence, this markup allows optimization:

---

```
void updatePtrs(int* restrict ptrA,  
                int* restrict ptrB,  
                int* restrict val)
```

---

Note: you can get the optimization by just declaring `ptrA` and `val` as `restrict`; `ptrB` isn't needed for this optimization

- Use `restrict` whenever you know the pointer will not alias another pointer (also declared `restrict`)

It's hard for the compiler to infer pointer aliasing information; it's easier for you to specify it.

⇒ compiler can better optimize your code (more perf!)



Caveat: don't lie to the compiler, or you will get **undefined behaviour**.

Aside: `restrict` is not the same as `const`. `const` data can still be changed through an alias.

# Automatic Parallelization of Example Code

Let's try automatic parallelization.

Compiling with `solarisstudio` and automatic parallelization yields the following:

---

```
% solarisstudio -cc -O3 -xautopar -xloopinfo omp_vector.c  
"omp_vector.c", line 5: PARALLELIZED, and serial version generated  
"omp_vector.c", line 15: not parallelized, call may be unsafe
```

---

How will this code compare to our manual efforts?  
(If you weren't in class, you'll have to try it yourself.)

**Note:** `solarisstudio` generates two versions of the code, and decides, at runtime, if the parallel code would be faster.

# Example Code to Parallelize

---

```
#include <stdlib.h>

void setup(double *vector, int length) {
    int i;
    for (i = 0; i < length; i++)
    {
        vector[i] += 1.0;
    }
}

int main()
{
    double *vector;
    vector = (double*) malloc(sizeof(double)*1024*1024);
    for (int i = 0; i < 1000; i++)
    {
        setup (vector, 1024*1024);
    }
    // if you don't read vector, compiler NOPs everything
    printf("%f\n", vector[0]);
}
```

---

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# Autoparallelization implementation: OpenMP

Under the hood, most parallelization frameworks use OpenMP, which we'll see next lecture.

For now: you can control the number of threads with the `OMP_NUM_THREADS` environment variable.



gcc (since 4.3) can also auto-parallelize loops via Graphite.

Parallelization has been getting better,  
but not super well-maintained.

I think gcc can insert a runtime check of loop iteration count.

Magic incantation:

```
gcc -O2 -floop-parallelize-all  
    -ftree-parallelize-loops=4 -fopt-info
```

-floop-parallelize-all: parallelize all the things  
-ftree-parallelize-loops=N: use N threads

**Note:** gcc uses OpenMP but overrides OMP\_NUM\_THREADS with N above.

## Sample -fopt-info output

```
$ gcc L11-omp_vector.c -O2 \  
-floop-parallelize-all -ftree-parallelize-loops=4 -fopt-info  
L11-omp_vector.c:14:3: note: parallelizing inner loop 1  
L11-omp_vector.c:23:3: note: loop nest optimized  
L11-omp_vector.c:15:15: note: parallelizing inner loop 4  
  
line 14: for (i = 0; i < length; i++) {  
line 15: vector[i] += 1.0;  
line 23: for (int i = 0; i < 1000; i++) {
```

# Understanding Automatic Parallelization in gcc

Want to better understand?

Look at the assembly code to see the parallelizations  
(obviously, impractical for a large project).

```
$ gcc -std=c99 -O3 -ftree-parallelize-loops=4 \  
    omp_vector_gcc.c -S -o omp_vector_gcc_auto.s
```

[Or, you can use `objdump` on files compiled with `-g`.]

Older gcc generated a .s file with:

---

```
call    GOMP_parallel_start
leaq    80(%rsp), %rdi
call    setup._loopfn.0
call    GOMP_parallel_end
```

---

**Note:** gcc also parallelizes `main._loopfn.2` and `main._loopfn.3`, although it looks like it serves little purpose.

# Loops That gcc's Automatic Parallelization Can Handle

Single loop:

---

```
for (i = 0; i < 1000; i++)  
    x[i] = i + 3;
```

---

Nested loops with simple dependency:

---

```
for (i = 0; i < 100; i++)  
    for (j = 0; j < 100; j++)  
        x[i][j] = x[i][j] + y[i-1][j];
```

---

Single loop with not-very-simple dependency:

---

```
for (i = 0; i < 10; i++)  
    x[2*i+1] = x[2*i];
```

---

# More Loops That gcc's Automatic Parallelization Can Handle

Single loop with if statement:

---

```
for (j = 0; j <= 10; j++)  
    if (j > 5) X[i] = i + 3;
```

---

Triangle loop:

---

```
for (i = 0; i < 100; i++)  
    for (j = i; j < 100; j++)  
        X[i][j] = 5;
```

---

Examples from: <http://gcc.gnu.org/wiki/AutoparRelated>

# Summary of Conditions for Automatic Parallelization

From Chapter 10 of Oracle's *Fortran Programming Guide*<sup>1</sup> translated to C, a loop must:

- have a recognized loop style, e.g., for loops with bounds that don't vary per-iteration;
- have no dependencies between data accessed in loop bodies for each iteration;
- not conditionally change scalar variables read after the loop terminates, or change any scalar variable across iterations; and
- have enough work in the loop body to make parallelization profitable.

---

<sup>1</sup><http://download.oracle.com/docs/cd/E19205-01/819-5262/index.html>

# Example Code to Parallelize (once more)

---

```
#include <stdlib.h>

void setup(double *vector, int length) {
    int i;
    for (i = 0; i < length; i++)
    {
        vector[i] += 1.0;
    }
}

int main()
{
    double *vector;
    vector = (double*) malloc(sizeof(double)*1024*1024);
    for (int i = 0; i < 1000; i++)
    {
        setup (vector, 1024*1024);
    }
    // if you don't read vector, compiler NOPs everything
    printf("%f\n", vector[0]);
}
```

---



# Manually Parallelizing the Example Code

What can we do to parallelize this code?

**Option 1:**

**Option 2:**

**Option 3:**

# Manually Parallelizing the Example Code

What can we do to parallelize this code?

**Option 1:** horizontal 

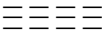
- Create 4 threads; each thread does 1000 iterations on its own sub-array.

**Option 2:**

**Option 3:**

# Manually Parallelizing the Example Code

What can we do to parallelize this code?

**Option 1:** horizontal      

- Create 4 threads; each thread does 1000 iterations on its own sub-array.

**Option 2:** bad horizontal      

- 1000 times, create 4 threads which each operate once on the sub-array.

**Option 3:**

# Manually Parallelizing the Example Code

What can we do to parallelize this code?

**Option 1:** horizontal      ≡≡≡≡

- Create 4 threads; each thread does 1000 iterations on its own sub-array.

**Option 2:** bad horizontal      ≡≡≡≡

- 1000 times, create 4 threads which each operate once on the sub-array.

**Option 3:** vertical      |||| |||| |||| ||||

- Create 4 threads; for each element, the owning thread does 1000 iterations on that element.

Methodology: compiling with `solarisstudio`,  
flags `-O3 -lpthread`.

Which manual option performs better?

# Comparing Parallelization Results

How does autoparallelization compare to manual parallelization?

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- Relative ordering: **Option 3** > Automatic > **Option 1**
- Automatic parallelization of **Option 1** was better than manual, why?

How does autparallelization compare to manual parallelization?

- Relative ordering: **Option 3** > Automatic > **Option 1**
- Automatic parallelization of **Option 1** was better than manual, why?
- Manual **Option 3** performed better, even though both used the same number of threads, why?



# Lingering Questions about Runtimes

What happened here?

≡≡≡≡

horizontal good:

create 4 threads to do 1000 iterations on sub-arrays.

≡≡≡≡

horizontal bad:

1000 times, create 4 threads to iterate on sub-array.

|||| |||| |||| ||||

vertical:

create 4 threads, handle 1 element at a time.

In 2015, `perf stat -r 5` gave following task-clocks (in seconds):

	H good	H bad	V	auto
gcc, no opt	2.794	2.953	2.799	
gcc, -O3	0.588	1.490	0.980	
solaris, no opt	3.175	3.291	2.966	
solaris, -xO4	0.494	1.453	2.739	0.688

## Observations:

- Good runs had 5 to 7 cpu-migrations; bad had 4000.
- # cycles varied from 2B to 9.7B (no opt).
- Branch misses varied from 8k to 208k.

## Case Study 2: Multiplying a Matrix by a Vector

Let's see how automatic parallelization does on a more complicated program (could we parallelize this?):

---

```
void matVec (double **mat, double *vec, double *out,
             int *row, int *col)
{
    int i, j;
    for (i = 0; i < *row; i++)
    {
        out[i] = 0;
        for (j = 0; j < *col; j++)
        {
            out[i] += mat[i][j] * vec[j];
        }
    }
}
```

---

$$\text{Reminder: } \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \end{bmatrix} \begin{bmatrix} 1 \\ 2 \\ 3 \end{bmatrix} = \begin{bmatrix} 14 \\ 32 \end{bmatrix}$$

# Case Study: Automatic Parallelization, Attempt 1

Well, based on our knowledge, we could parallelize the outer loop.

Let's see what `solarisstudio` will do for us...

---

```
% solarisstudio -cc -xautopar -xloopinfo -O3 -c fploop.c  
"fploop.c", line 5: not parallelized, not a recognized for loop  
"fploop.c", line 8: not parallelized, not a recognized for loop
```

---

...it refuses to do anything, guesses?

## Case Study: Automatic Parallelization, Attempt 2

- The loop bounds are not constant, since one of the variables may alias row or col, even though `int`  $\neq$  `double`.

So, let's add `restrict` to row and col and see what happens...

---

```
% solarisstudio -cc -O3 -xautopar -xloopinfo -c fploop.c  
"fploop.c", line 5: not parallelized, unsafe dependence  
"fploop.c", line 8: not parallelized, unsafe dependence
```

---

Now it recognizes the loop, but still won't parallelize it. Why?

# Case Study: Automatic Parallelization, Attempt 3

- out might alias mat or vec, which would make this unsafe

Let's add another restrict to out:

---

```
% solarisstudio -cc -O3 -xautopar -xloopinfo -c fploop.c  
"fploop.c", line 5: PARALLELIZED, and serial version  
generated  
"fploop.c", line 8: not parallelized, unsafe dependence
```

---

Now, we can get the outer loop to parallelize.

- Parallelizing the outer loop is almost always better than inner loops, and usually it's a waste to do both, so we're done.

**Note:** We can parallelize the inner loop as well (it's similar to the assignment). We'll see that solarisstudio can do it automatically.

- Reductions combine input data into a smaller (summary) set.
- We'll see a more complete definition when we touch on functional programming.
- Simplest instance: computing the sum of an array.

Consider the following code:

---

```
double sum (double *array, int length)
{
    double total = 0;

    for (int i = 0; i < length; i++)
        total += array[i];
    return total;
}
```

---

Can we parallelize this?

Barriers to parallelization:

- 1 value of total depends on previous iterations;
- 2 addition is actually non-associative for floating-point values (is this a problem?)

Recall that “associative” means:

$$a + (b + c) = (a + b) + c.$$

In this case, the program probably isn't sensitive to rounding, but you should always consider if an operation is associative.



# Automatic Parallelization via Reduction

If we compile the program with `solarisstudio` and add the flag `-xreduction`, it will parallelize the code:

---

```
% solarisstudio -cc -xautopar -xloopinfo -xreduction -O3 -c sum.c  
"sum.c", line 5: PARALLELIZED, reduction, and serial version  
generated
```

---

**Note:** If we try to do the reduction on `fploop.c` with `restricts` added, we'll get the following:

---

```
% solarisstudio -cc -O3 -xautopar -xloopinfo -xreduction -c fploop.c  
"fploop.c", line 5: PARALLELIZED, and serial version generated  
"fploop.c", line 8: not parallelized, not profitable
```

---

- A general function could have arbitrary side effects.
- Production compilers tend to avoid parallelizing any loops with function calls.

Some built-in functions, like `sin ( )`, are “pure”, have no side effects, and are safe to parallelize.

**Note:** this is why functional languages are nice for parallel programming: impurity is visible in type signatures.

# Dealing with Function Calls in solarisstudio

- For solarisstudio you can use the `-xbuiltin` flag to make the compiler use its whitelist of “pure” functions.
- The compiler can then parallelize a loop which uses `sin()` (you shouldn't replace built-in functions with your own if you use this option).

Other options which may work:

- 1 Crank up the optimization level (`-xO4`).
- 2 Explicitly tell the compiler to inline certain functions (`-xinline=`, or use the `inline` keyword).

# Summary of Automatic Parallelization

To help the compiler, we can:

- use `restrict` (make a restricted copy); and,
- make sure that loop bounds are constant (temporary variables).

Some compilers automatically create different versions for the alias-free case and the (parallelized) aliased case.

At runtime, the program runs the aliased case if correct.