Regular flattening

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Agenda

Representation and Fusion

Handling nested parallelism

Basic flattening rules

Incremental flattening

Multi-level parallelism

Final words as time permits

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Consider arrays of type [](i32, i8). Since an i32 is four bytes and a i8 is one byte, how is this stored in memory?

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0	1	2	3	4	5	6	7	8	9	10
	i32				i32				i8	

Problem?

Consider arrays of type [](i32, i8). Since an i32 is four bytes and a i8 is one byte, how is this stored in memory?

Problem? Unaligned accesses.

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Consider arrays of type [](i32, i8). Since an i32 is four bytes and a i8 is one byte, how is this stored in memory?

Problem? Unaligned accesses.

Problem? Waste of memory.

Tuples of arrays

Representation

```
An array [](t1, t2, t3...) is represented in memory as ([]t1, []t2, []t3...), i.e. as multiple arrays, each containing only primitive values.
```

0	1	2	3	4	5	6	7	8	9	10
i32								i 32		
i8	i8	i8	i8	i8	i8	i8	i8	i8	i8	

- Common (and crucial) optimisation.
- Called "struct of arrays" in legacy languages.
- Automatically done by the Futhark compiler.
- Only affects internal language.
- ...and also assumed for the rest of today's presentation.

"Unzipped" SOACs

Instead of

- In the compiler IR, All SOACs accept multiple array inputs and produce unzipped results.
- Arrays of tuples (or records, or sums) do not exist in the core language.
- Isomorphic to source language, but this form is much easier to work with in a compiler.

Loop fusion

```
let increment [n][m] (as: [n][m]i32) : [n]i32 =
  map (\r -> map (+2) r) a
let sum [n] (a: [n]i32) : i32 =
  reduce (+) 0 a
let sumrows [n][m] (as: [n][m]i32) : [n]i32 =
  map sum as
```

Let's say we wish to first call increment, then sumrows:

```
sumrows (increment a)
```

Naively Run increment, then call sumrows.

Problem Manifests intermediate matrix in memory.

Solution *Loop fusion*, which combines loops to avoid intermediate results.

An example of a fusion rule

The expression

is always equivalent to

map
$$(f \circ g)$$
 a

- This is an extremely powerful property that is only true in the absence of side effects.
- Fusion is the core optimisation that permits the efficient decomposition of a data-parallel program.
- A full fusion engine has much more awkward-looking rules (zip/unzip causes lots of bookkeeping), but safety is guaranteed.

A fusion example

```
sumrows (increment a) = (Initial expression)

map sum (increment a) = (Inline sumrows)

map sum (map (\lambda r \rightarrow \text{map}(+2) r) a) = (Inline increment)

map (sum \circ (\lambda r \rightarrow \text{map}(+2) r) a) = (Apply map-map fusion)

map (\lambda r \rightarrow \text{sum}(\text{map}(+2) r) a) = (Apply composition)
```

- We have avoided the temporary matrix, but the composition of sum and the map also holds an opportunity for fusion – specifically, reduce-map fusion.
- Will not cover in detail, but a reduce can efficiently apply a function to each input element before engaging in the actual reduction operation.
- Important to remember: a **map** going into a **reduce** is an efficient pattern.

A shorthand notation for sequences

$$\overline{z}^{(n)}=z_0,\cdots,z_{(n-1)}$$

- The *n* may be omitted.
- A separator may be implied by context.

$$f \, \overline{v}^{(n)} \equiv f \, v_1 \, \cdots \, v_n$$

or a tuple

$$(\overline{v}^{(n)}) \equiv (v_1, \ldots, v_n)$$

or a function type

$$\overline{\tau}^{(n)} \to \tau_{n+1} \equiv \tau_1 \to \cdots \to \tau_n \to \tau_{n+1}.$$

When not all terms under the bar are variant, subscript variant terms with i.

$$(\overline{[d]v_i}^{(n)}) = ([d]v_1, \dots, [d]v_n)$$

and

$$(\overline{[d_i]v_i}^{(n)})=([d_1]v_1,\ldots,[d_n]v_n)$$

Fused constructs

Convenient shorthands

- Emphasises that reduce/scan-map compositions can be considered as a single construct.
- We will see several examples where this is useful.

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Note:

```
\begin{array}{lll} \mathbf{reduce} \, \odot \, \overline{d} \, \overline{xs} \equiv & \mathbf{reduce} \, \odot \, \overline{d} \, (\mathsf{map} \, \mathsf{id} \, \overline{xs}) \equiv & \mathbf{redomap} \, \odot \, f \, \overline{d} \, \overline{xs} \\ \mathbf{scan} \, \odot \, \overline{d} \, \overline{xs} \equiv & \mathbf{scan} \, \odot \, \overline{d} \, (\mathsf{map} \, \mathsf{id} \, \overline{xs}) \equiv & \mathbf{scanomap} \, \odot \, f \, \overline{d} \, \overline{xs} \end{array}
```

Representation and Fusion

Handling nested parallelism

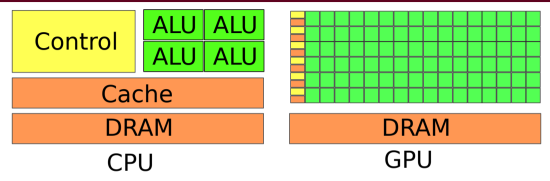
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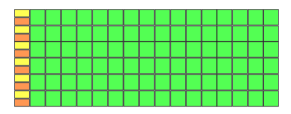
Final words as time permits

GPUs vs CPUs



- GPUs have thousands of simple cores and taking full advantage of their compute power requires tens of thousands of threads.
- GPU threads are very restricted in what they can do: no stack, no allocation, limited control flow, etc.
- Potential very high performance and lower power usage compared to CPUs, but programming them is hard.

The SIMT Programming Model



- GPUs are programmed using the SIMT model (*Single Instruction Multiple Thread*).
- Similar to SIMD (Single Instruction Multiple Data), but while SIMD has explicit vectors, we provide sequential scalar per-thread code in SIMT.

Each thread has its own registers, but they all execute the same instructions at the same time (i.e. they share their instruction pointer).

SIMT example

For example, to increment every element in an array a, we might use this code:

```
increment(a) {
  tid = get_thread_id();
  x = a[tid];
  a[tid] = x + 1;
}
```

- If a has n elements, we launch n threads, with get_thread_id() returning i for thread i.
- This is *data-parallel programming*: applying the same operation to different data.
- When we launch a GPU program (*kernel*), we say how many threads should be launched, *all running the same code*.

Branching

If all threads share an instruction pointer, what about branches?

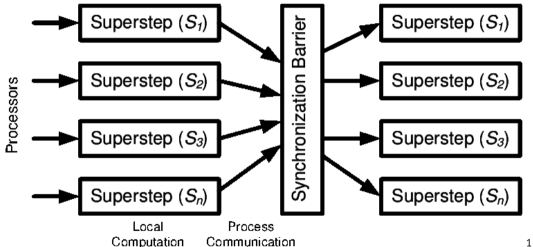
```
mapabs(a) {
   tid = get_thread_id();
   x = a[tid];
   if (x < 0) {
      a[tid] = -x;
   }
}</pre>
```

Masked Execution

Both branches are executed in all threads, but in those threads where the condition is false, a mask bit is set to treat the instructions inside the branch as no-ops.

Do GPUs exist in theory as well?

GPU programming is a close fit to *bulk synchronous parallelism*:



Supersteps are threads, which cannot talk to each other.

A SOAC-kernel correspondence

The compiler *knows*² that certain nests of perfect **map**s correspond to certain GPU basic blocks.

- maps containing scalar code is a kernel with one thread per iteration of the maps.
- maps containing a single **reduce** is a *segmented reduction*.
- maps containing a single scan is a segmented scan.
- maps containing a single scatter is a segmented scatter.
- ...see the pattern?

Crucial: the **map**s must be *perfectly nested*.

map (
$$x - x + y$$
) xs) xss ys

Suppose xss is of shape [n][m], then this can compile to a kernel with $n \times m$ threads, each doing a single x + y operation.

²Because it has taken PMPH.

Handling nested parallelism

Problem

Futhark permits nested (regular) parallelism, but GPUs need flat parallel kernels.

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Have the compiler rewrite program to perfectly nested **map**s containing sequential code, or known parallel patterns such as segmented reduction.

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Have the compiler rewrite program to perfectly nested **map**s containing sequential code, or known parallel patterns such as segmented reduction.

Flattening via loop fission

The classic map fusion rule:

$$\mathsf{map}\, f \circ \mathsf{map}\; g \Rightarrow \mathsf{map}\; (f \circ g)$$

³ Futhark: Purely Functional GPU-Programming with Nested Parallelism and In-Place Array Updates, PLDI 2017

Flattening via loop fission

The classic map fusion rule:

$$\mathsf{map}\, f \circ \mathsf{map}\; g \Rightarrow \mathsf{map}\; (f \circ g)$$

We can also apply it backwards to obtain fission:

$$\mathsf{map}\; (f\circ g)\Rightarrow \mathsf{map}\; f\circ \mathsf{map}\; g$$

This, along with other higher-order rules (see paper³, or just wait until later in the lecture), are applied by the compiler to extract perfect map nests.

³Futhark: Purely Functional GPU-Programming with Nested Parallelism and In-Place Array Updates, PLDI 2017

Example: (a) Initial program, we inspect the map-nest

```
let (asss. bss) =
  map (\() (ps: [m]i32) ->
         let ass = map (\((p: i32): [m]i32 ->
                           let cs = scan (+) 0 (iota p)
                           let r = reduce (+) 0 cs
                           in map (+r) ps) ps
         let bs = loop ws=ps for i < n do
                       map (\ as w: i32 \rightarrow
                              let d = reduce (+) 0 as
                              let e = d + w
                             in 2 * e) ass ws
         in (ass, bs)) pss
We assume the type of pss : \lceil m \rceil \lceil m \rceil i 32.
```

(b) Distribution

```
let asss: [m][m][m]i32 =
  map (\() (ps: [m]i32) ->
        let ass = map (\((p: i32): [m]i32 ->
                          let cs = scan (+) 0 (iota p)
                          let r = reduce (+) 0 cs
                          in map (+r) ps) ps
        in ass) pss
let bss: [m][m]i32 =
 map (\setminus ps ass ->
        let bs = loop ws=ps for i < n do
                     map (\setminus as w ->
                            let d = reduce (+) 0 as
                            let e = d + w
                            in 2 * e) ass ws
        in bs) pss asss
```

(c) Interchanging outermost map inwards

```
let asss: [m][m][m]i32 =
 map (\() (ps: [m]i32) ->
        let ass = map (\((p: i32): [m]i32 ->
                          let cs = scan (+) 0 (iota p)
                          let r = reduce (+) 0 cs
                          in map (+r) ps) ps
        in ass) pss
let bss: [m][m]i32 =
 map (\protect\ps ass ->
        let bs = loop ws=ps for i < n do
                     map (\setminus as w ->
                            let d = reduce (+) 0 as
                            let e = d + w
                            in 2 * e) ass ws
        in bs) pss asss
```

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                          let r = reduce (+) 0 cs
                          in map (+r) ps) ps
        in ass) pss
let bss: [m][m]i32 =
  loop wss=pss for i < n do
    map (\setminus ass ws ->
          let ws' = map (\setminus as w ->
                            let d = reduce (+) 0 as
                            let e = d + w
                            in 2 * e) ass ws
           in ws') asss wss
```

(d) Skipping scalar computation

```
let asss: [m][m][m]i32 =
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                            let e = d + w
                            in 2 * e) ass ws
           in ws') asss wss
```

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```

(e) Distributing reduction

```
let asss: [m][m][m]i32 =
  map (\() (ps: [m]i32) ->
        let ass = map (\((p: i32): [m]i32 ->
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                          let r = reduce (+) 0 cs
                          in map (+r) ps) ps
        in ass) pss
let bss: [m][m]i32 =
  loop wss=pss for i < n do
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          let ws' = map (\setminus as w ->
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                            let e = d + w
                            in 2 * e) ass ws
           in ws') asss wss
```

(e) Distributing reduction

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                            let cs = scan (+) 0 (iota p)
                            let r = reduce (+) 0 cs
                            in map (+r) ps) ps
         in ass) pss
let bss: [m][m]i32 =
  loop wss=pss for i < n do
    let dss: [m][m]i32 =
      map (\setminus ass ->
              map (\setminus as ->
                     reduce (+) 0 as) ass)
           asss
    in map (\ ws ds \rightarrow
               let ws' =
                  map (\backslash w d \rightarrow let e = d + w
                                 in 2 * e) ws ds
               in ws') asss dss
```

(f) Distributing inner map

(f) Distributing inner map

```
let rss: [m][m]i32 =
 map (\() (ps: [m]i32) ->
        let rss = map (\((p: i32): i32 ->
                          let cs = scan (+) 0 (iota p)
                          let r = reduce (+) 0 cs
                          in r) ps
        in rss) pss
let asss: [m][m][m]i32 =
 map (\((ps: [m]i32) (rs: [m]i32) \rightarrow
        map (\((r: i32): [m]i32 ->
              map (+r) ps) rs
      ) pss rss
let bss: [m][m]i32 = ...
```

(g) Cannot distribute as it would create irregular array

Array cs has type [p]i32, and p is variant to the innermost map nest.

(h) These statements are sequentialised

Array cs has type [p]i32, and p is variant to the innermost map nest.

Result

- From a single kernel with parallelism m to four kernels of parallelism m^2 , m^3 , m^3 , and m^2 .
- The last two kernels are executed *n* times each.

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Notation for flat parallelism

Instead of writing

```
map (\ps rs ->
  map (\r ->
  map (\p -> e)
     ps)
  rs)
  pss rss
```

We write

```
\begin{array}{l} \textbf{segmap} \; (\langle \texttt{ps}, \texttt{rs} \in \texttt{pss}, \texttt{rss} \rangle, \; \langle \texttt{r} \in \texttt{rs} \rangle, \; \langle \texttt{p} \in \texttt{ps} \rangle) \\ e \end{array}
```

Segmented flat parallel constructs

$$\Sigma = \Sigma', \langle \overline{x} \in \overline{y}
angle$$
 segmap Σ $e \equiv egin{array}{c} ext{map} \ (\lambda \overline{x_p}
ightarrow \ ext{map} \ (\lambda \overline{x_1}
ightarrow e) \overline{y_1}) \ \hline y_{\overline{p}-1} \end{pmatrix}$

- Conceptually a stack of maps with some parallel construct (here another map) inside.
- *These* are what triggers GPU code generation.
- Any SOACs left in e will be executed sequentially.

Similarly for reductions and scans

$$\begin{array}{lll} \operatorname{{\bf segred}} \Sigma \odot \overline{d} \ e \equiv & \operatorname{{\bf map}} \left(\lambda \overline{x_p} \to \\ & \operatorname{{\bf map}} \left(\lambda \overline{x_{p-1}} \to \ldots \right. \\ & \operatorname{{\bf redomap}} \odot \left(\lambda \overline{x_1} \to e \right) \overline{d} \ \overline{y_1} \right) \\ & \overline{y_p} \\ \\ \operatorname{{\bf segscan}} \Sigma \odot \overline{d} \ e \equiv & \operatorname{{\bf map}} \left(\lambda \overline{x_p} \to \\ & \operatorname{{\bf map}} \left(\lambda \overline{x_{p-1}} \to \ldots \right. \\ & \operatorname{{\bf scanomap}} \odot \left(\lambda \overline{x_1} \to e \right) \overline{d} \ \overline{y_1} \right) \\ & \overline{y_{p-1}} \right) \\ & \overline{y_p} \end{array}$$

Let us look at how one can rewrite SOAC nests to these segmented operations.

Example of rewrite rules

Rules describe how valid judgments can be formed.

Example with partial evaluation

$$\boxed{\mathcal{V} dash e_1 \Rightarrow e_2}$$
 where \mathcal{V} is a mapping from variable names v to values.

Example of rewrite rules

Rules describe how valid judgments can be formed.

Example with partial evaluation

 $\overline{\mathcal{V} dash e_1 \Rightarrow e_2}$ where $\mathcal V$ is a mapping from variable names v to values.

- Rewrite rules can be ambiguous (several may apply).
- Need a decision procedure in order to have an algorithm.

Flattening rules

 $\sum \vdash e \Rightarrow e'$ In a map-nest context \sum , the source expression e can be translated into the target expression e'.

$$\begin{array}{c} e \text{ has inner SOACs} \\ \Sigma, \langle \overline{x} \in \overline{\mathsf{xs}} \rangle \vdash e \Rightarrow e_{\mathrm{flat}} \\ \hline \Sigma \vdash \mathsf{map} \ (\lambda \overline{x} \to e) \ \overline{\mathsf{xs}} \Rightarrow e_{\mathrm{flat}} \\ \hline \\ \text{no other rule applies} \\ \hline \bullet \vdash e \Rightarrow e \\ \hline \\ \Sigma \vdash e \Rightarrow \mathsf{segmap} \ \Sigma \ e \end{array}$$

$$\Sigma \vdash \mathbf{redomap} \odot (\lambda \overline{x} \rightarrow e) \overline{d} \overline{xs} \Rightarrow \mathbf{segred} (\Sigma, \overline{x} \in \overline{xs}) \odot \overline{d} e$$

Rule for map distribution

size of each array in
$$\overline{a_0}$$
 invariant to Σ

$$\Sigma = \langle \overline{x_p} \in \overline{y_p} \rangle, \dots, \langle \overline{x_1} \in \overline{y_1} \rangle \qquad \Sigma \vdash e_1 \Rightarrow e_1'$$

$$\overline{a_p}, \dots, \overline{a_1} \text{ fresh names} \qquad \Sigma' \vdash e_2 \Rightarrow e_2'$$

$$\Sigma' = \langle \overline{x_p} \ \overline{a_{p-1}} \in \overline{y_p} \ \overline{a_p} \rangle, \dots, \langle \overline{x_1} \ \overline{a_0} \in \overline{y_1} \ \overline{a_1} \rangle$$

$$\Sigma \vdash \mathbf{let} \ \overline{a_0} = e_1 \ \mathbf{in} \ e_2 \Rightarrow \mathbf{let} \ \overline{a_p} = e_1' \ \mathbf{in} \ e_2'$$

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$$\Sigma \vdash \mathbf{let} \overline{a_0} = e_1 \mathbf{in} e_2 \Rightarrow \mathbf{let} \overline{a_p} = e_1' \mathbf{in} e_2'$$

Example for

map (
$$\xs ->$$
 let $y =$ redomap (+) ($\xs ->$ x) 0 xs in map ($\xs ->$ x + y) xs)

Suppose already inside the outer map

$$\begin{split} \Sigma &= \langle xs \in xss \rangle \quad \Sigma' = \langle xs, y \in xss, ys \rangle \\ \Sigma &\vdash \mathbf{redomap} \ \ (+) \ \ (\lambda x \ \to \ x) \ \ \emptyset \ \ xs \Rightarrow \mathbf{segred} \ \ (\Sigma, \langle x \in xs \rangle) \ \ (+) \ \ 0 \ x \\ \Sigma' &\vdash \mathsf{map} \ \ (\lambda x \ \to \ x \ + \ y) \ \ xs \Rightarrow \mathbf{segmap} \ \ (\Sigma', \langle x \in xs \rangle) \ x \ + \ y \\ \Sigma &\vdash ... \Rightarrow \ \ \ \mathbf{let} \ \ ys = \mathbf{segred} \ \ \ (\langle xs \in xss \rangle, \langle x \in xs \rangle) \ \ \ (+) \ \ 0 \ x \\ &\quad \ \ \mathbf{in} \ \ \mathbf{segmap} \ \ \langle xs, y \in xss, ys \rangle \ x \ + \ y \end{split}$$

Handling transposition

rearrange (d_1, \dots, d_n) x is a generalization of **transpose** in that it rearranges the dimensions of d-dimensional array based on a permutation defined by the integer sequence d_1, \dots, d_n . E.g:

 $extbf{transpose} \equiv extbf{rearrange} (1,0)$

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$$extbf{transpose} \equiv extbf{rearrange} (1,0)$$

Flattening rule

$$egin{aligned} \Sigma \vdash \mathbf{rearrange} \ (0, 1 + k_1, \dots, 1 + k_n) \ y \Rightarrow e \ \hline \Sigma, \langle x \in y \rangle \vdash \mathbf{rearrange} \ (k_1, \dots, k_n) \ x \Rightarrow e \end{aligned}$$

Handling transposition

rearrange (d_1, \dots, d_n) x is a generalization of **transpose** in that it rearranges the dimensions of d-dimensional array based on a permutation defined by the integer sequence d_1, \dots, d_n . E.g:

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Flattening rule

$$\Sigma \vdash$$
 rearrange $(0, 1 + k_1, \dots, 1 + k_n) \ y \Rightarrow e$
 $\Sigma, \langle x \in y \rangle \vdash$ rearrange $(k_1, \dots, k_n) \ x \Rightarrow e$

Example

 $\bullet \vdash \text{map} \; (\lambda x \to \text{rearrange} \; (1,0) \; x) \; xs \Rightarrow \text{rearrange} \; (0,2,1) \; xs$

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Consider Matrix Multiplication

```
for i < n:
    for j < m:
        acc = 0
        for l < p:
        acc += xss[i,l] * yss[l,j]
        res[i,j] = acc</pre>
```

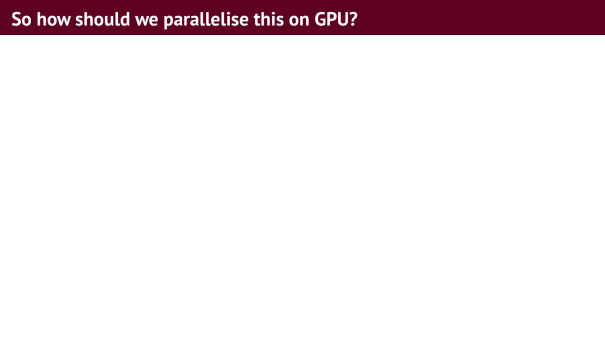
Turning it Functional

```
map (\xs ->
    map (\ys ->
        let zs = map (*) xs ys
        in reduce (+) 0 zs)
        (transpose yss))
xss
```

Using redomap notation

```
\mathbf{redomap} \, \odot \, f \, 0_{\odot} \, x \ \equiv \ \mathbf{reduce} \, \odot \, 0_{\odot} \, (\mathbf{map} \, f \, x)
```

Emphasises that a **map-reduce** composition can be turned into a fused tight sequential loop, or into a parallel reduction.



So how should we parallelise this on GPU?

- All parallelism exploited
- Some communication overhead
- Best if outer maps don't saturate GPU

So how should we parallelise this on GPU?

Full flattening

- All parallelism exploited
- Some communication overhead
- Best if outer maps don't saturate GPU
- There is no one size fits all.
- Both situations may be encountered at program runtime.

Moderate flattening

- Only outer parallelism
- The redomap can be block tiled
- Best if outer maps saturate GPU

The essence of incremental flattening

From a single source program, for each parallel construct generate multiple semantically equivalent parallelisations, and generate a single program that at runtime picks the least parallel that still saturates the hardware.

- Implemented in the Futhark compiler.
- ...but technique is applicable to any (regular) nested parallelism expressed with the common Bird-Meertens-style array constructs (map, reduce, scan, etc).

Simple Incremental Flattening

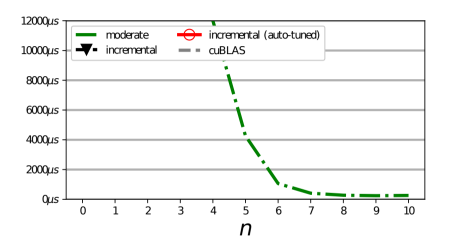
At every level of map-nesting we have two options:

- 1. Continue flattening inside the map, exploiting the parallelism there.
- 2. Sequentialise the map body; exploiting only the parallelism on top.
- Full flattening in the Blelloch style will do the former, maximising utilised parallelism.
- Incremental flattening generates both versions and uses a predicate to pick at runtime.

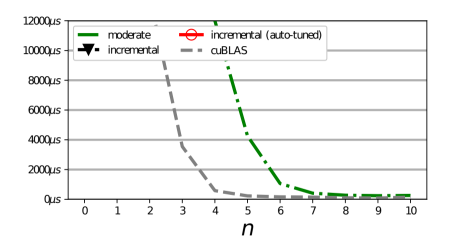
Multi-versioned matrix multiplication

```
xss : [n][p]i32
yss : [p][m]i32.
if n * m > t_0 then
  map (\setminus xs \rightarrow
           map (\setminus vs ->
                     redomap (+) (*) 0 xs ys)
                 (transpose vss))
        XSS
else
  map (\setminus xs \rightarrow
           map (\setminus vs ->
                     redomap (+) (*) 0 xs ys)
                 (transpose yss))
        XSS
```

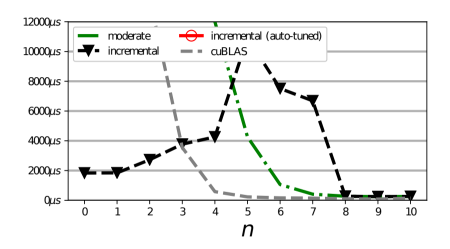
The t_0 threshold parameter is used to select between the two versions—and should be auto-tuned on the concrete hardware.



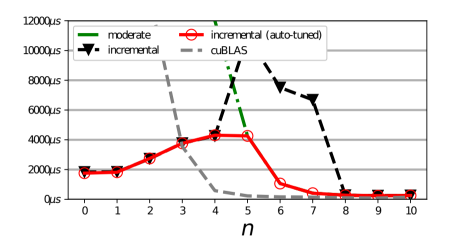
Multiplying matrices of size $2^n \times 2^m$ and $2^m \times 2^n$, where m = 25 - 2n, meaning that work is constant as we vary n.



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Incremental flattening rule

Example for

```
map (\ xs -> redomap (+) (\ x -> x) 0 xs) xss
```

Incremental flattening rule

$$egin{aligned} \Sigma' &= \Sigma, \langle \overline{x} \in \overline{ imes s}
angle & \Sigma' \vdash e \Rightarrow e_{ ext{flat}} \ \hline \Sigma \vdash \mathsf{map} \left(\lambda \overline{x}
ightarrow e
ight) \overline{ imes s} \Rightarrow \ & ext{if } \operatorname{Par}(\Sigma') \geq t_{ ext{top}} \ & ext{then segmap } \Sigma' \ e \ & ext{else } e_{ ext{flat}} \end{aligned}$$

Example for

```
\begin{split} \Sigma = \bullet \quad & \Sigma' = \langle \mathsf{xs} \in \mathsf{xss} \rangle \\ & \Sigma' \vdash e \Rightarrow \mathbf{segred} \left( \langle \mathsf{xs} \in \mathsf{xss} \rangle, \langle \mathsf{x} \in \mathsf{xs} \rangle \right) (\texttt{+}) \ 0 \ \mathsf{x} \\ & \qquad \qquad \quad & \mathbf{if} \ \mathsf{length} (\mathsf{xss}) \geq t_\mathsf{top} \\ & \Sigma \vdash ... \Rightarrow \qquad & \mathbf{then} \ \mathsf{segmap} \ \langle \mathsf{xs} \in \mathsf{xss} \rangle \ (\mathsf{redomap} \ (\texttt{+}) \ (\lambda \mathsf{x} \ \to \ \mathsf{x}) \ 0 \ \mathsf{xs}) \\ & \qquad \qquad & \mathbf{else} \ \mathsf{segred} \ (\langle \mathsf{xs} \in \mathsf{xss} \rangle, \langle \mathsf{x} \in \mathsf{xs} \rangle) \ (\texttt{+}) \ 0 \ \mathsf{x} \end{split}
```

map ($\xs \rightarrow \text{redomap}$ (+) ($\xs \rightarrow x$) 0 xs) xss

Autotuning

- An incrementally flattened program may have dozens of threshold parameters, t_i , used to select versions at runtime.
- As we have seen, the default value (2¹⁶) is often not optimal.

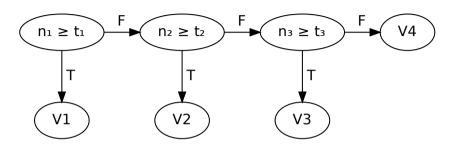
A configuration P maps each t_i to an integer $P(t_i)$.

The search problem

Find the P that minimises the cost function F(P), where the the cost function runs the program on a set of user-provided representative datasets and sums the observed runtimes.

- Other cost functions are also possible, e.g. average runtime over datasets.
- Note: recompilation is not necessary.

Briefly on our search procedure⁴



- Suppose we are given training data sets D_j , j < k, each of which provide a value $v_{i,j}$ for each threshold parameter n_i .
- Starting from the deepest comparison (t_3) , for each D_j find an (x_j, y_j) that minimises runtime, take the intersection of the intervals, and use that to determine threshold value.
- Tuning time is linear in the number of comparisons.

⁴https://futhark-lang.org/student-projects/svend-msc-thesis.pdf

Using incremental flattening

Compile with a GPU backend (opencl or cuda):

\$ futhark opencl matmul.fut

To autotune:

\$ futhark autotune -v --backend=opencl matmul.fut

Produces matmul .fut .tuning, which is automatically picked up by futhark bench (use --no-tuning to stop this).

Use futhark dev -s --extract-kernels matmul.fut to see IR.

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Final words as time permits

Confession

I lied when I claimed that GPU threads were completely isolated.

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- Most hardware has useful (fixed) levels of parallelism.
- An ideal flattening algorithm maps levels of application parallelism (any number) to hardware parallelism (fixed number) in a way that exploits locality well.

High-example: a system consists of multiple *datacenters*, that each contain multiple *computers*, that each contain multiple *GPUs*, that each contain multiple *SMs* (next slide), that each run some number of threads.

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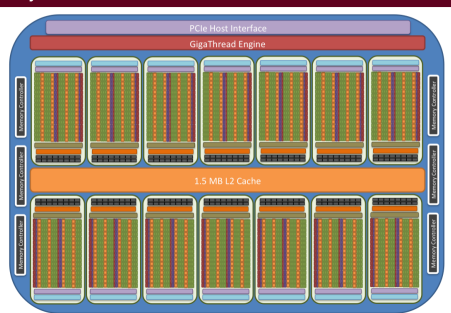
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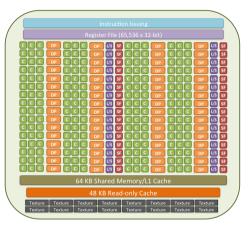
General principle

"Tasks" at the same hardware level cannot communicate, but can "launch" tasks at a lower level.

K20 GPU layout



Streaming Multiprocessor (SM) layout



single precision/integer CUDA core smemory load/store unit double precision FP unit special function unit

Level-aware segmented operations

 $l \in \text{thread}, \text{group}$

- *Group* is the same as a CUDA *thread block*
- Each segmented operation then tagged with the level at which its *body* executes.

$$\begin{array}{l} \operatorname{\mathtt{segmap}}^l \Sigma \ e \\ \operatorname{\mathtt{segscan}}^l \Sigma \ \odot \ \overline{d} \ e \\ \operatorname{\mathtt{segred}}^l \Sigma \ \odot \ \overline{d} \ e \end{array}$$

Restrictions

Both thread and group can occur at top level, but a group construct can contain only thread constructs, and thread cannot any segmented constructs.

Examples

Each thread transposes part of an array

$$segmap^{thread} \langle x \in xs \rangle$$
 (transpose x)

Each workgroup transposes part of an array

$$segmap^{group} \langle x \in xs \rangle$$
 (transpose x)

These are both equivalent to map transpose xs.

Each workgroup sums the row of an array

$$extsf{segmap}^{group} \langle xs \in xss \rangle \text{ (segred}^{thread} \langle x \in xs \rangle \text{ (+) 0 x)}$$

Equivalent to map (reduce (+) 0) xss.

Tags carry no semantic meaning; used solely for code generation.

Example: LocVolCalib

The following is the essential core of the LocVolCalib benchmark from the FinPar suite.

```
map (\xss ->
    map (\xs ->
        let bs = scan \oplus d_{\oplus} xs
    let cs = scan \otimes d_{\otimes} bs
    in scan \odot d_{\odot} cs)
    xss)
```

How can we map the application parallelism to hardware parallelism?

Option I: sequentialise the inner scans

```
\begin{array}{l} \textbf{segmap}^{\text{thread}} \; (\langle \texttt{xss} \in \texttt{xsss} \rangle, \langle \texttt{xs} \in \texttt{xss} \rangle) \\ \textbf{let} \; \texttt{bs} = \textbf{scan} \oplus \; d_{\oplus} \; \texttt{xs} \\ \textbf{let} \; \texttt{cs} = \textbf{scan} \otimes \; d_{\otimes} \; \texttt{bs} \\ \textbf{in} \; \textbf{scan} \; \odot \; d_{\odot} \; \textit{cs} \end{array}
```

scan is relatively expensive in parallel, so this is a good option if the outer dimensions provide enough parallelism.

Option II: flatten and parallelise inner scans

Flattening uses *loop distribution* (or *fission*) to create **map** nests:

```
map (\xss ->
    map (\xs ->
        let bs = scan \oplus d_{\oplus} xs
    let cs = scan \otimes d_{\otimes} bs
    in scan \odot d_{\odot} cs)
    xss)
```

Option II: flatten and parallelise inner scans

```
\label{eq:letbsss} \begin{split} &\textbf{segscan}^{\text{thread}} \; \big( \langle \texttt{xss} \in \texttt{xsss} \rangle, \langle \texttt{xs} \in \texttt{xss} \rangle,, \langle \texttt{x} \in \texttt{xs} \rangle \big) \; \oplus \; d_{\oplus} \; \texttt{x} \\ &\textbf{let} \; \texttt{csss} = \\ &\textbf{segscan}^{\text{thread}} \; \big( \langle \texttt{bss} \in \texttt{bsss} \rangle, \langle \texttt{bs} \in \texttt{bss} \rangle,, \langle \texttt{b} \in \texttt{bs} \rangle \big) \; \oplus \; d_{\oplus} \; \texttt{b} \\ &\textbf{in} \\ &\textbf{segscan}^{\text{thread}} \; \big( \langle \texttt{css} \in \texttt{csss} \rangle, \langle \texttt{cs} \in \texttt{css} \rangle,, \langle \texttt{c} \in \texttt{cs} \rangle \big) \; \oplus \; d_{\oplus} \; \texttt{c} \end{split}
```

This is what full flattening will do.

Option III: Mapping innermost parallelism to the workgroup level

```
\begin{array}{c} \text{map} \ (\backslash \ \mathsf{xss} \ - \gt \\ \quad \mathsf{map} \ (\backslash \ \mathsf{xs} \ - \gt \\ \quad \mathsf{let} \ \ \mathsf{bs} \ = \ \mathsf{scan} \ \oplus \ d_{\oplus} \ \ \mathsf{xs} \\ \quad \mathsf{let} \ \ \mathsf{cs} \ = \ \mathsf{scan} \ \otimes \ d_{\otimes} \ \ \mathsf{bs} \\ \quad \mathsf{in} \ \ \ \mathsf{scan} \ \odot \ d_{\odot} \ \ \mathsf{cs} \,) \\ \quad \mathsf{xsss} \,) \\ \\ \mathsf{xsss} \end{array}
```

Option III: Mapping innermost parallelism to the workgroup level

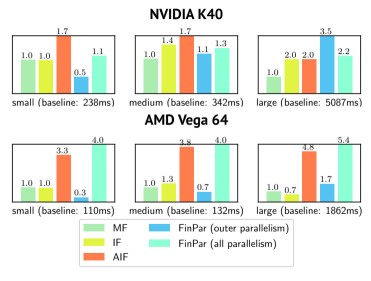
```
\begin{split} &\textbf{segmap}^{group} \; \big( \langle \texttt{xss} \in \texttt{xsss} \rangle, \langle \texttt{xs} \in \texttt{xss} \rangle \big) \\ &\textbf{let} \; \texttt{bs} = \textbf{segscan}^{thread} \; \langle \texttt{x} \in \texttt{xs} \rangle \; \oplus \; d_{\oplus} \; \texttt{x} \\ &\textbf{let} \; \texttt{cs} = \textbf{segscan}^{thread} \; \langle \texttt{b} \in \texttt{bs} \rangle \otimes \; d_{\otimes} \; \texttt{b} \\ &\textbf{in} \; \; \textbf{segscan}^{thread} \; \langle \texttt{c} \in \texttt{cs} \rangle \otimes \; d_{\otimes} \; \texttt{c} \end{split}
```

- Iterations of outer **segmap**s assigned to GPU workgroups⁵.
- Each segscan^{thread} is executed collaboratively by a workgroup and in local memory⁶.
- Only works if the innermost parallelism fits in a workgroup.

⁵Thread block in CUDA

⁶Shared memory in CUDA

LocVolCalib speedup (higher is better)



Sequential scans (MF) is the baseline.

Level-aware incremental flattening

 $\sum \vdash^l e \Rightarrow e'$ In a map-nest context Σ , the source expression e can be translated at machine level l into the target expression e'.

In the Futhark compiler, only two levels are handled (thread, group), but we believe the idea generalises well.

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Final words as time permits

Block tiling

Level-aware constructs can also be used for expressing other powerful optimisations.

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Level-aware constructs can also be used for expressing other powerful optimisations.



Threads accessing same memory can cooperatively cache it in on-chip memory.

Motivation for block tiling

map (
$$\x -> \text{redomap} (+) (\y -> y + x) 0 xs$$
) xs

After flattening we get this inner-sequential version:

$$\textbf{segmap}^{thread} \ \langle x \in xs \rangle \ (\textbf{redomap} \ (+) \ (\lambda y \rightarrow y + x) \ 0 \ xs)$$

Operation One thread for each element of xs, and each sequentially traverses xs. Problem ?

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$$\x -> \text{redomap} (+) (\y -> y + x) 0 xs$$
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Operation One thread for each element of xs, and each sequentially traverses xs. Problem Poor utilisation of memory bus.

- Many threads simultaneously read same address, which is redundant.
- **Better:** *cooperatively* copy *block* into on-chip memory and iterate from there.

Strip mining/chunking the outer segmap

$$\textbf{segmap}^{thread} \ \langle x \in xs \rangle \ (\textbf{redomap} \ (+) \ (\lambda y \rightarrow y + x) \ 0 \ xs)$$

Assuming we can split xs into m equally sized *tiles* each of size t, giving xss: [m][t]f32, then we can rewrite to

$$\begin{array}{l} \textbf{segmap}^{group} \; \langle \textbf{xs'} \in \textbf{xss} \rangle \\ \textbf{segmap}^{thread} \; \langle \textbf{x} \in \textbf{xs'} \rangle \\ \textbf{redomap} \; (+) \; (\lambda \textbf{y} \rightarrow \textbf{y} + \textbf{x}) \; \textbf{0} \; \textbf{xs} \end{array}$$

Question: does this compute the same value as the original?

Strip mining/chunking the outer segmap

$$\textbf{segmap}^{thread}~\langle x \in xs \rangle~(\textbf{redomap}~(+)~(\lambda y \rightarrow y + x)~0~xs)$$

Assuming we can split xs into m equally sized *tiles* each of size t, giving xss: [m][t]f32, then we can rewrite to

```
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```

Question: does this compute the same value as the original?

- No—the original expression had type [n]f32, while this has type [m][t]f32
- This can be flattened away.

```
\begin{array}{c} \textbf{segmap}^{\text{group}} \ \langle \textbf{xs'} \in \textbf{xss} \rangle \\ \textbf{segmap}^{\text{thread}} \ \langle \textbf{x} \in \textbf{xs'} \rangle \\ \textbf{redomap} \ (+) \ (\lambda \textbf{y} \rightarrow \textbf{y} + \textbf{x}) \ \textbf{0} \ \textbf{xs} \end{array}
```

Chunking/strip-mining the **redomap**, we get

```
\begin{array}{l} \textbf{segmap}^{\text{group}} \ \langle \textbf{xs'} \in \textbf{xss} \rangle \\ \textbf{segmap}^{\text{thread}} \ \langle \textbf{x} \in \textbf{xs'} \rangle \\ \textbf{loop} \ \text{acc} \ = \ 0 \ \textbf{for} \ \textbf{ys} \ \textbf{in} \ \textbf{xss} \ \textbf{do} \\ \textbf{redomap} \ (+) \ (\lambda \textbf{y} \rightarrow \textbf{y} + \textbf{x}) \ \text{acc} \ \textbf{ys} \end{array}
```

```
\begin{array}{c} \textbf{segmap}^{\text{group}} \left< x \textbf{s} \text{'} \in x \textbf{s} \textbf{s} \right> \\ \textbf{segmap}^{\text{thread}} \left< x \in x \textbf{s} \text{'} \right> \\ \textbf{redomap} \left( + \right) \left( \lambda \textbf{y} \rightarrow \textbf{y} + \textbf{x} \right) \textbf{0} \ \textbf{xs} \end{array}
```

Chunking/strip-mining the **redomap**, we get

```
egin{align*} \mathsf{segmap}^{\mathsf{group}}\left\langle \mathsf{xs'} \in \mathsf{xss} \right
angle \\ \mathsf{segmap}^{\mathsf{thread}}\left\langle \mathsf{x} \in \mathsf{xs'} \right
angle \\ \mathsf{loop}\ \mathsf{acc}\ =\ 0\ \mathsf{for}\ \mathsf{ys}\ \mathsf{in}\ \mathsf{xss}\ \mathsf{do} \\ \mathsf{redomap}\left(+\right)\left(\lambda \mathsf{y} 
ightarrow \mathsf{y} + \mathsf{x} \right)\ \mathsf{acc}\ \mathsf{ys} \end{aligned}
```

 $egin{aligned} extsf{segmap}^{ ext{group}}\left\langle extsf{xs'} \in extsf{xss}
ight
angle \ extsf{loop} \ ext{accs} = ext{replicate t 0} \end{aligned}$

Distributing and interchanging segmap thread gives

 $\begin{array}{l} \textbf{for ys in xss do} \\ \textbf{segmap}^{thread} \; \langle \textbf{x}, \texttt{acc} \in \textbf{xs'}, \texttt{accs} \rangle \\ \textbf{redomap} \; (+) \; (\lambda \textbf{y} \rightarrow \textbf{y} + \textbf{x}) \; \texttt{acc ys} \end{array}$

```
\begin{array}{l} \textbf{segmap}^{\text{group}} \ \langle \textbf{xs'} \in \textbf{xss} \rangle \\ \textbf{loop} \ \text{accs} = \ \textbf{replicate} \ \textbf{t} \ \textbf{0} \\ \textbf{for} \ \textbf{ys} \ \textbf{in} \ \textbf{xss} \ \textbf{do} \\ \textbf{segmap}^{\text{thread}} \ \langle \textbf{x}, \textbf{acc} \in \textbf{xs'}, \textbf{accs} \rangle \\ \textbf{redomap} \ (+) \ (\lambda \textbf{y} \rightarrow \textbf{y} + \textbf{x}) \ \textbf{acc} \ \textbf{ys} \end{array}
```

Collectively copy ys to shared/local memory

```
segmap<sup>group</sup> \langle xs' \in xss \rangle
loop accs = replicate t 0
for ys in xss do
let ys' = copy ys in
segmap<sup>thread</sup> \langle x, acc \in xs', accs \rangle
redomap (+) (\lambda y \rightarrow y + x) acc ys'
```

- Now the many iterations of the **redomap** read from fast on-chip memory rather than slower global memory!
- copy done collectively by all threads in group

The fine print

```
map (\x -> \text{redomap} (+) (\y -> y + x) 0 xs) xs to
```

```
\begin{array}{l} \textbf{segmap}^{group} \; \langle \texttt{xs'} \in \texttt{xss} \rangle \\ \textbf{loop} \; \texttt{accs} \; = \; \texttt{replicate} \; \texttt{t} \; \texttt{0} \\ \textbf{for} \; \texttt{ys} \; \textbf{in} \; \texttt{xss} \; \textbf{do} \\ \textbf{let} \; \texttt{ys'} \; = \; \textbf{copy} \; \texttt{ys} \; \textbf{in} \\ \textbf{segmap}^{thread} \; \langle \texttt{x}, \texttt{acc} \in \texttt{xs'}, \texttt{accs} \rangle \\ \textbf{redomap} \; (+) \; (\lambda \texttt{y} \to \texttt{y} + \texttt{x}) \; \texttt{acc} \; \texttt{ys'} \end{array}
```

- Very simple case (e.g. xss traversed in both loops)
- 2D tiling much more complex
- The *tile size* t is a sensitive tuning parameter; in this case it should coincide with workgroup size
- Appreciate what a compiler can do for you

Summary

- There is no *one size fits all:* for optimal performance, we need different amounts of parallelisation for different workloads.
- Incremental flattening generates a single program that for varying datasets exploits only as much parallelism as profitable.
- Autotuning for specific hardware and program is needed to select the optimal version at runtime.
- A good IR is as crucial to a compiler as a good language is to a human.