

Design and GPGPU Performance of Futhark’s Redomap Construct

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Abstract

This paper presents and evaluates a novel second-order operator, named `redomap`, that stems from `map-reduce` compositions in the context of the purely-functional array language Futhark, which is aimed at efficient GPGPU execution. Main contributions are: First, we demonstrate an aggressive fusion technique that is centered on the `redomap` operator. Second, we present a compilation technique for `redomap` that efficiently sequentializes the excess parallelism and ensures coalesced access to global memory, even for non-commutative `reduce` operators. Third, a detailed performance evaluation shows that Futhark’s automatically generated code matches or exceeds performance of hand-tuned Thrust code.

Our evaluation infrastructure is publicly available and we encourage replication and verification of our results.

Categories and Subject Descriptors D.1.3 [Concurrent Programming]: Parallel Programming; D.3.4 [Processors]: Compiler

Keywords GPGPU, map-reduce, autparallelization, functional language

1. Introduction

Commodity many-core hardware is now mainstream, driven in particular by the evolution of general purpose graphics programming units (GPGPUs) that support thousands of cores, but commodity programming is still falling short of efficiently utilizing this hardware: Principal reasons are that low-level programming APIs, such as CUDA and OpenCL, are difficult to use and the translation of a sequential program is quite tedious and requires specialized users.

For example, the implementation of parallel `reduce` and `scan` operators is not only algorithmically challenging, but also requires efficient use of the fast memory and barrier synchronizations. Libraries that provide generic implementations of common bulk-parallel operators, such as Thrust [18], enhance programmability, but at cost in performance, for example because:

- 1 A “naive” translation might not take advantage of the way in which, for example, `map` and `reduce` operators may be efficiently composed, and
- 2 Even when the library supports a wealth of such combined operators and the expert user take full advantage of those, per-

formance still remains sub-optimal due to generality-related constraints, e.g., the `reduce` operator requires an array-of-tuples representation, which results in less-coalesced accesses to global memory than a tuple-of-arrays representation would.

This context opens the door for data-parallel array languages [3, 15, 20] to emerge as the mainstream environment for programming massively-parallel hardware. In essence:

- 1 the non-specialist user is encouraged to “naively” write the application in its clearest form, in terms of a small and easy to understand set of simple operators, such as `map`, `reduce`, which
- 2 are optimized automatically by compiler analysis that scales to program-level to reach efficiencies that outperform highly-tuned code that relies on generic parallel libraries.

This paper analyses in detail the simple but common, hence important, case of `map-reduce` composition in the context of Futhark [17]: a purely-functional array language that supports nested parallelism on regular multidimensional arrays. We remark that Futhark adopts a non-restricted semantics for the `map` and `reduce` operators, for example (i) the mapped function may receive an array argument and return an array, and (ii) the `reduce` operator is arbitrary (user defined), may be only associative or also commutative, and its supported input type may be a tuple of scalars or even arrays in some cases. The contributions of this paper are:

First, we demonstrate in Section 3 how `map` and `reduce` operators are composed aggressively (at all nest-levels in the program) by a combination of producer-consumer and horizontal fusion. The result of fusion is an operator, named `redomap`, that is not exposed to the user language because (i) its semantics is non-trivial/non-intuitive, and (ii) because in all `map-reduce` examples we encountered so far the compiler reliably fuses such compositions.

Second, we present in Section 4 how the high-level operator is translated to a lower-level representation in a manner that optimizes both (i) efficient sequentialization of the parallelism in excess, and (ii) coalesced access to global memory, which in particular may have appeared as an artifact of efficient sequentialization. Intuitively, the latter is achieved by reshaping the array to have an inner dimension of the sequential-chunk size, and by transposing this dimension to the outermost level.

Third, we present in Section 5 a detailed evaluation of performance results on a mini-benchmark that demonstrate that the automatically optimized Futhark code outperforms tuned Thrust programs by a geometric-mean factor of $1.75\times$ (and as high as $8\times$).

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| | | |
|-----------------|---|---|
| $\bar{q}^{(n)}$ | $::= q_1, \dots, q_n$ | (notation) |
| s, x | $::= \text{id}$ | (variable names, s for scalar) |
| k | $::= \text{Ct} \mid [k_1, \dots, k_n]$ | (scalar or array value) |
| z | $::= \text{id} \mid \text{Ct}$ | (variable name or constant) |
| g, h | $::= \text{id}$ | (function names) |
| p | $::= \tau_1 \ x_1$ | (typed variable) |
| t | $::= \text{int} \mid \text{bool} \mid \text{f32} \mid \text{f64}$ | (basic types) |
| τ | $::= t \mid [\tau, s]$ | (size-dep array types) |
| ρ | $::= (\tau_1, \dots, \tau_n)$ | (tuple (of arrays) types) |
| ϕ | $::= \rho_1 \rightarrow \rho_2$ | (fun/lambda type) |
| l | $::= \text{fn } \bar{t}^{(n)} \ (\bar{p}^{(m)}) \Rightarrow e$ | (anonymous fun) |
| e | $::= z$ | (Variable or Value) |
| | $(\bar{y}^{(n)})$ | (n-tuple exp) |
| | $x[y_1, \dots, y_n]$ | (array indexing) |
| | $y_1 \odot y_2$ | (binop-call) |
| | $g(\bar{y})$ | (function-call) |
| | $\text{if } s \text{ then } e_1 \text{ else } e_2$ | (if binding) |
| | $\text{let } (\bar{p}) = e_1 \text{ in } e_2$ | (let-binding) |
| | $\text{iota}(s)$ | $[0, \dots, s-1]$ |
| | $\text{map}(l, \bar{x}^{(n)})$ | (n-ary map) |
| | $\text{reduce}(l, \bar{z}^{(n)}, \bar{x}^{(n)})$ | (reduce with n-ary op) |
| | $\text{scan}(l, \bar{z}^{(n)}, \bar{x}^{(n)})$ | (scan with n-ary op) |
| | $x \text{ with } [s_1, \dots, s_n] \leftarrow y$ | (in-place update) |
| | $\text{loop } (\bar{p}^{(n)} = \bar{y}^{(n)}) =$ | (sequential do-loop) |
| | $\text{for } s < S \text{ do } e$ | (next iter $\bar{p}^{(n)} \leftarrow e$) |
| P | $::= \text{fun } \rho \ g(p) = e; P \mid \epsilon$ | (named function def) |

Figure 1: Syntax of a Subset of Futhark’s Core Language.

2. A Brief and Informal Introduction to Futhark

Futhark¹ is a monomorphic, statically typed, eagerly evaluated, pure functional language, in which nested parallelism is expressed via a set of second-order array combinators (SOACs), e.g., `map`, `reduce`, `scan`, `filter`, and which aims at efficient execution on GPGPUS. Figure 1 presents the abstract syntax of a subset of the Futhark *core language*, which is a restricted language used internally by the compiler. Whenever q is an object of some kind, we write $\bar{q}^{(n)}$ (or simply \bar{q}) to range over sequences of n objects of that kind, that is, q_1, \dots, q_n . Important details are:

- Similarly to the A-normal form representation [26], operands of compound expressions must be variables or values.
- The keyword `in` is optional before `let` (this is solely for aesthetic reasons).
- The source Futhark language uses the traditional array-of-tuples representation, supporting `zip/unzip` operators and in which, for example, `map` receives exactly one array argument.
- However, as shown in Figure 1, the core language does *not* support tuples because the program is (automatically) normalized to the tuple-of-arrays form. As a result, SOACs such as `map` take as input *several* arrays, and results in several arrays; our `map` can be seen as implicitly `zip`ping its input and `unzip`ping its output. This transformation is automatically performed early on in the compilation process, using a traditional array-of-tuples to tuple-of-array transformation [4].
- For any array variable defined in a let-binding pattern, its type is parameterised with the exact shape information, i.e., at array-

¹ The language is named after the first six letters of the runic alphabet, (i.e., “Futhark”). The `filter` operator is not covered in this paper.

```

loop ((x1, ..., xn) =      -- Equivalent function
      (a1, ..., an)) =   fun t f( int i
                          , int n
    for i < n do          , t1 x1
                          , ..., tn xn) =
      g(i, x1, ..., xn) =>
-- Equivalent to:        if i >= n then x
f(0, n, a1, ..., an)    else f(i+1, n, g(i, x1, ..., xn))

```

Figure 2: Loop to recursive function

creation point. For example, $[[\text{int}, m], n]$ denotes a two-dimensional $n \times m$ array of integers, where n and m must be constants or integer variables in scope.

- All arrays must be *regular*, that is, all rows of an array must have the same shape. For example array $[[4], [1.0]]$ is illegal; this is mostly verified through run-time checks.
- Futhark supports sequential do-loops, whose semantics is equivalent to a simple form of tail-recursive functions, as illustrated in Figure 2. The `do`-loop construct enables several important optimizations such as aggressive hoisting and `map`-loop interchange (outside the scope of this paper). The latter may significantly enhance the amount of parallelism that can be statically extracted.

The compiler performs producer-consumer and horizontal fusion, which is demonstrated in Section 3 in the simple(r) case of `map-reduce` compositions. Fusion is followed by extraction of flat-parallel kernels suitable for translation to GPU code. This step resembles the loop distribution performed by imperative approaches. Ultimately, several key lower-level optimizations are performed, such as changing the in-memory representation of arrays via transposition to promote *coalesced accesses*² to global GPGPU memory.

3. Redomap: Type, Semantics and Fusion Rules

The Futhark source language does not support the `redomap` second-order array combinator (SOAC), because this is synthesized automatically by fusion from `map-reduce` compositions.

In previous work [17] we have proposed an aggressive technique for fusing producer-consumer SOACs, without duplicating computation, even when the produced array is used in several places. Semantically, this is achieved by a bottom-up traversal of program’s dependency graph, in which SOACs are fused whenever a T2 reduction is possible (i.e., a SOAC can be fused if it is the source of only one dependency edge, and the target of the dependence is another (compatible) SOAC). The technique was further refined to solve some hindrances to fusion, for example, to move intervening transpositions before/after the fused kernel and to interchange an outer `reduce` with an inner `map` in order to create more fusion opportunities.

Using a Haskell notation, the `redomap` construct was inspired by the following rewriting of a `map-reduce` composition:

```

red ⊙ e . map f ≡ red ⊙ e . map (red ⊙ e . map f) . splitp
≡ red ⊙ e . map (foldl g e) . splitp ≡ redomap ⊙ g e

```

- the input array is `split` into a number of chunks that roughly equals the number of available processors
- each processor efficiently sequentializes the computation on its chunk by rewriting the inner $(\text{red} \odot e) . (\text{map } f)$ as `foldl g e`,
- and finally, the partial results are reduced across processors.

² Coalesced access requires that neighboring threads access neighboring memory words in a SIMD instruction.

As such, in previous work, `redomap` had type:

`redomap`:: $(\alpha \rightarrow \alpha \rightarrow \alpha) \rightarrow (\alpha \rightarrow \beta \rightarrow \alpha) \rightarrow \alpha \rightarrow [\beta] \rightarrow \alpha$
in which the first argument is a binary associative operator (\odot), the second is the folded function (g), the third is the neutral element (e) of the group induced by \odot , and the fourth is the input array.

The result type of `redomap` is α , that is, the result (and input) type of the binary associative operator \odot . This is a *significant restriction* because, for example, it does not allow fusing the following code:

```
let x = map(f, a)
let r = reduce(+, 0.0, x)
in (r, x)
```

If the result of the map is used outside of the `reduce` then they cannot be fused because `redomap`'s type disallows the return of x .

Another *significant restriction* of the previous work was that horizontal fusion was not supported. The remaining of this section discusses our solution to solving the (two) observed limitations.

3.1 Extended Redomap: Type and Semantics

Denoting with $\bar{z}^{(n)}$ the sequence z_1, \dots, z_n , the type of the `redomap` (in the Futhark core language) was extended to:

$((\bar{\alpha}^{(p)}, \bar{\alpha}^{(p)}) \rightarrow \bar{\alpha}^{(p)}), ((\bar{\alpha}^{(p)}, \bar{\beta}^{(q)}) \rightarrow (\bar{\alpha}^{(p)}, \bar{\gamma}^{(r)})),$
 $\bar{\alpha}^{(p)}, [\beta_1, n], \dots, [\beta_q, n] \rightarrow (\bar{\alpha}^{(p)}, [\gamma_1, n], \dots, [\gamma_r, n])$

1^{st} argument is a binary associative operator, of type $(\bar{\alpha}^{(p)}, \bar{\alpha}^{(p)}) \rightarrow \bar{\alpha}^{(p)}$, where $\bar{\alpha}^{(p)}$ intuitively denotes a tuple of p elements,

2^{nd} argument is the folded function g of type: $((\bar{\alpha}^{(p)}, \bar{\beta}^{(q)}) \rightarrow (\bar{\alpha}^{(p)}, \bar{\gamma}^{(r)}))$. That is, it receives the accumulator of type $\bar{\alpha}^{(p)}$ and an arbitrary number q of (array) elements $\bar{\beta}^{(q)}$ and produces a new accumulator $\bar{\alpha}^{(p)}$ and an arbitrary number r of (array) elements $\bar{\gamma}^{(r)}$

3^{rd} argument is the neutral element of type $\bar{\alpha}^{(p)}$,

- The remaining arguments are q arrays of equal-size outermost dimension n (i.e., $[\beta_1, n], \dots, [\beta_q, n]$).
- The result has two components: (i) the reduced part of type $\bar{\alpha}^{(p)}$, and (ii) the mapped part $[\gamma_1, n], \dots, [\gamma_r, n]$ which corresponds to r arrays of outermost size equal to n whose elements were produced by each invocation of g .

The semantics of `redomap` is:

`redomap`($\oplus, g, \bar{e}^{(p)}, \bar{b}^{(q)}$) \equiv `let` ($\bar{a}^{(p)}, \bar{c}^{(r)}$) = `map`($g, \bar{b}^{(q)}$)
`in` (`reduce`($\oplus, \bar{e}^{(p)}, \bar{a}^{(p)}$), $\bar{c}^{(r)}$)

except that in practice it is executed very similar to the OPENMP-style parallel loop with reduction pragmas. That is, each processor computes a partial accumulator from a chunk of the mapped arrays, and the partial accumulators are then reduced across processors.

3.2 Horizontal Fusion

The extension of the `redomap` operator opened the door to eliminating two important limitations of the previous fusion engine:

First, fusion is allowed between two SOACs belonging to the same block of `let` statements even if the array produced by the first SOAC is used after the second³, i.e., as long as there is no unfused use of first SOAC's result array in between the two SOACs.

For example, even though in the code below x is used in the result:

```
let x = map(f, a) in
let r = reduce(+, 0.0, x)
in (r, x)
```

the `reduce` and the `map` are safely fused into:

³ This can be further relaxed by using a dependency-graph representation of block of `let` statements.

```
let (x, r) = redomap( +
, fn (f32, f32) (f32 e, f32 a) =>
let x = f(a) in (e+x, x)
, 0.0, a)
in (r, x)
```

Second, we allow horizontal fusion (i.e., when the two SOACs are not in a producer-consumer relation), whenever the two SOACs belong to the same block of `let` statements, their outermost sizes are equal, and as before, there is no (unfused) use of the result array in between the two SOACs. For example:

```
let x = reduce(+, 0.0, a)
let y = reduce(*, 1.0, a)
```

is fused horizontally as:

```
let (x, y) = reduce( fn (f32, f32) (f32 x1, f32 y1,
f32 x2, f32 y2) =>
let r1 = x1 + x2
let r2 = y1 * y2
in (r1, r2)
, 0.0, a, a)
```

3.3 Demonstrating Redomap Fusion

Figure 3 demonstrates the fusion engine on a contrived example whose structure resembles code from FinPar's Interest-Rate Calibration benchmark [1]. The original code is presented at the top, and `reduceComm` denotes a `reduce` in which the binary operator is declared to be also commutative (besides being associative). The previous fusion implementation would have succeeded in fusing the first three maps, denoted (7 – 9), but not the remaining SOACs (1 – 6). The rest of the figure demonstrates the enhanced fusion:

Step I: Since fusion proceeds bottom up, the last two maps, denoted (1) and (2) are fused *horizontally*, resulting in the map denoted (10), which receives two array arguments x and y and produces arrays v and w .

Step II: The obtained map (10) is fused *horizontally* with the `reduce` (3) resulting in the `redomap` (11), whose binary associative and commutative operator is `max`.

Step III: The obtained `redomap` (11) is fused *horizontally* with the other two `reduces`, (4 – 5), by extending the associative and commutative operator of the `redomap` to work over three-float tuples (and modifying accordingly the folded function).

Step IV: The resulting `redomap`, (12) consumes the array t produced by the map (16) and as such are fused together, and similarly for the maps producing arrays y , x and is , which are (7 – 9). At the end, the original program has been fused in one `redomap` construct, which is shown at the bottom of Figure 3.

The resulting code requires only the two result arrays v and w to be maintained in global memory. Because the input array `iota(n)` is optimized away by reference to the thread index space, the rest of the computation involves only scalars. The next section describes in detail how the `redomap` second-order operator is mapped to efficient GPGPU code.

4. Optimizing Sequentialization and Coalescing

The efficient implementation of reductions on GPUs is well studied in the literature [16, 21]. This section gives a short introduction to the subject to enable discussion of the implementation and optimization of the `redomap` construct.

To execute a reduction in parallel on an idealised perfectly parallel machine, we might use *tree reduction*: to reduce an n -element array $[x_1, \dots, x_n]$ using operator \oplus , we launch $n/2$ threads, with thread i computing $x_{2i} \oplus x_{2i+1}$. The initial n elements are thus reduced to $n/2$ elements. The process is repeated until just a

```

-- Original Program:
fun (f32, f32, f32, [f32], [f32])
main(f32 a, f32 b, int n) =
  let is = map(f32, iota(n)) -- (9)
  let x = map(*a, is) -- (8)
  let y = map(*b, x) -- (7)
  let t = map(+, zip(x,y)) -- (6)
  in
    let t0 = reduceComm(+, 0.0f32, t) -- (5)
    let t1 = reduceComm(min, inf(), x) -- (4)
    let t2 = reduceComm(max, 0.0f32, y) -- (3)
    in
      let v = map(*a, x) -- (2)
      let w = map(*b, y) -- (1)
      in (t0, t1, t2, v, w)

-- I. horizontal fusion (1) and (2) results in (10):
let ([f32,n] v, [f32,n] w) =
  map(fn (f32, f32) (f32 xi, f32 yi) => -- (10)
    let (f32 res_1) = xi * a
    let (f32 res_2) = yi * b
    in (res_1, res_2),
    , x, y)

-- II. horizontal fusion (10) and (3) results in (11):
let (f32 t2, [f32,n] v, [f32,n] w) =
  redomapComm( max -- (11)
    , fn (f32, f32, f32)
      (f32 acc, f32 yi, f32 xi) =>
        let (f32 res) = max(acc, yi)
        let (f32 vi) = xi * a
        let (f32 wi) = yi * b
        in (res, vi, wi)
    , (0.0f32), y, x)

-- III. horizontal fusion (11) and (4,5) results in (12):
let (f32 t0, f32 t1, f32 t2, [f32,n] v, [f32,n] w) =
  redomapComm( fn ( f32, f32, f32 ) -- (12)
    ( f32 x1, f32 x2, f32 x3
    , f32 y1, f32 y2, f32 y3 ) =>
    let (f32 z1) = x1 + y1
    let (f32 z2) = min(x2, y2)
    let (f32 z3) = max(x3, y3)
    in (z1, z2, z3),
    fn (f32, f32, f32, f32, f32)
      ( f32 acc1, f32 acc2, f32 acc3
      , f32 ti, f32 xi, f32 yi ) =>
    let (f32 res1) = acc1 + ti
    let (f32 res2) = min(acc2, xi)
    let (f32 res3) = max(acc3, yi)
    let (f32 vi) = xi * a
    let (f32 wi) = yi * b
    in (res1, res2, res3, vi, wi),
    (0.0f32, Infinityf32, 0.0f32), t, x, y)

-- IV. Producer-consumer fusion between (12) and (6),
-- then (7) then (8) then (9) results in (13):
let (f32 t0, f32 t1, f32 t2, [f32,n] v, [f32,n] w) =
  redomapComm( fn ( f32, f32, f32 ) -- (13)
    ( f32 x1, f32 x2, f32 x3
    , f32 y1, f32 y2, f32 y3 ) =>
    let (f32 z1) = x1 + y1
    let (f32 z2) = min(x2, y2)
    let (f32 z3) = max(x3, y3)
    in (z1, z2, z3),
    fn (f32, f32, f32, f32, f32)
      (f32 acc1, f32 acc2, f32 acc3, int i) =>
    let (f32 iflt) = f32(i)
    let (f32 xi) = iflt * a
    let (f32 yi) = xi * b
    let (f32 ti) = xi + yi
    let (f32 res1) = acc1 + ti
    let (f32 res2) = min(acc2, xi)
    let (f32 res3) = max(acc3, yi)
    let (f32 vi) = xi * a
    let (f32 wi) = yi * b
    in (res1, res2, res3, vi, wi),
    (0.0f32, Infinityf32, 0.0f32), iota(n) )

```

Figure 3: Fusion demonstrated on a simplified example, resembling the structure of FinPar’s Interest-Rate Calibration benchmark.

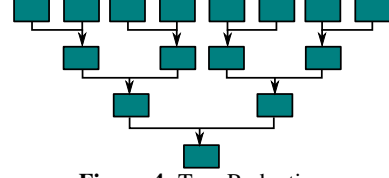


Figure 4: Tree Reduction

single value is left—the final result of the reduction. We perform $O(\log(n))$ partial reductions, each of which is perfectly parallel, resulting in a work depth of $O(\log(n))$. Figure 4 shows a simple example.

Tree reduction is optimal on an idealised perfectly parallel machine, but on real hardware, such as GPUs, it is inefficient. The inefficiency is caused by exposing more parallelism than needed to fully exploit the hardware. This *excess parallelism* means we pay an unnecessary overhead due to communication cost between threads. Efficient parallel execution relies on exposing as much parallelism as is needed to saturate the machine, but no more.

On a GPU, a simple improvement is to have every thread in a *workgroup*⁴ read a single value from the initial array, after which the threads cooperate in reducing their individual values to a partial result per workgroup. Thus, if we use size- k workgroups, each parallel level will result in a factor- k shrinkage, instead of the factor-2 performed by the naive algorithm.

We can still do better: as mentioned earlier, a concrete GPU can only use up to a certain amount of parallelism. The amount depends on the hardware and exact form of the reduction, but suppose that w groups of size k each are sufficient. Then, instead of spawning a number of threads dependent on the input size n , we always spawn $w \times k$ threads, organised into w workgroups. Each thread sequentially reduces a chunk of the input consisting of $\frac{n}{w \times k}$ elements, producing a per-thread intermediate result, which is then reduced inside each workgroup (using tree reduction) to one result per workgroup. If w is less than the maximum workgroup size, we can then launch a new reduction with one workgroup of size w that reduces the w per-workgroup results into the final result of the reduction. If n is not divisible by $w \times k$, not all threads will have the exact same chunk size, but it is not hard to partition the array among the threads. One interesting consequence is that in the sequential stage, the function need not be a proper reduction operator, but can be a fold function with no associativity requirements – this is what we exploit in `redomap`.

This is an efficient algorithm: it uses only as much parallelism as is necessary, and it requires only two kernel launches in total. The only caveat is that we have to be careful about how the threads traverse their assigned chunk of the input. On a GPU, we must ensure that memory accesses are *coalesced* in order to fully utilise the memory bus. For reductions, which are bandwidth-bound for most operators, this is an important concern. If each thread sequentially accesses neighboring elements in the input array, as on figure 5a, the resulting memory accesses will be non-coalesced, severely impacting performance. The solution is to access with a stride, as on figure 5b: if the per-thread chunk size is wk , then thread i should access the input at indices $i, i + 2wk, i + 3wk$, and so forth. This means that in the same cycle, neighboring threads i and $i + 1$ within the same warp will access elements at neighboring indices $i + jwk$ and $i + 1 + jwk$ (where j is the sequential iteration), thus obtaining coalesced memory access.

The strided access pattern just mentioned will result in the correct result if the operator is commutative, although we are in

⁴ This paper follows the OpenCL terminology. In NVIDIA’s CUDA, a workgroup is called a *thread block*.

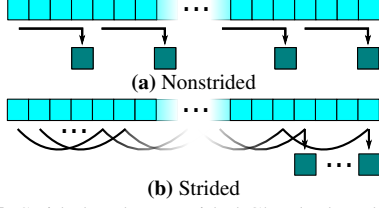


Figure 5: Strided and Nonstrided Chunked Reduction

effect accessing the array in a *transposed* form. Even if the input array is one-dimensional, we can pretend that it is a $wk \times \frac{n}{wk}$ array which is then transposed – if the input size is not divisible by wk we will have to pad the input, but we will ignore this issue in the present paper.

Many interesting operators are only *associative*, and swapping the order of application will produce a different (wrong) result. This is for example the case in the **maximum segment sum problem**, which is discussed in section 5. In such cases, we must first transpose the array in memory prior to the reduction – the strided access in the reduction will then “undo” the first transposition and restore the original evaluation order. Whilst transposition is not a free operation, it can be implemented on the GPU to run at essentially the speed of the memory bus.

This procedure is sketched on figure 6, where we assume that the reduction operator f is not commutative. The `redomap` on figure 6a is being turned into two kernels, the first of which is shown on figure 6b. The second, single-workgroup kernel is not particularly interesting, as the per-thread chunk size is always 1. In our intermediate language, we represent a reduction kernel via a `reduceKernel` construct. This construct is much like a `redomap`, except that instead of a fold function operating on an accumulator and single element, we have a function that processes an entire *chunk* of the input array with an explicit sequential loop.

In Futhark, the `transpose` operation is by default *delayed* (i.e., not made manifest in memory) – instead, the code generator will replace accesses to a transposed array with accesses to the original array, with the indices swapped. This is exploited in figure 6b – the sequential loop accesses to `chunk[i]`, really access `xs'[t,i]` (where t is the thread index), corresponding to `xs_tr_manifested[i,t]`. This array is itself a manifested version of `xs_tr_delayed`, which is a (delayed) transposed version of the original `xs`. Thus, `xs'[t,i] == xs[t,i]`, but with `xs'` represented in transposed form in memory.⁵

The mapped part of the `redomap` is always produced in transposed form by the `reduceKernel`, again to ensure coalescing. We explicitly transpose the mapped part to restore the intended order. If the reduction operator is commutative, and there is no mapped part in the `redomap`, the transposition is not actually performed in memory – this is for example the case when compiling particularly simple reductions such as `reduce(+, 0, as)`.

During the parallel part of the reduction, intermediate results are kept in arrays stored in fast local memory on the GPU. We use one such array for every input array. Due to the array-of-tuples transformation mentioned in section 2, a reduction of an array of pairs turns into a reduction of two arrays, meaning that we use two local buffers during the reduction.

⁵Futhark by default represents arrays in row-major form, so we could also say that the transposed array is column-major.

```
let (int p, [int,n] vs) =
  redomap(f,
    fn (int, int) (int acc, int x) =>
      let (int res) = f(acc, x)
      let (int v) = x * a
      (res, v),
      (0), xs)
```

(a) The original redomap

```
let xs_tr_delayed = transpose(xs)
let xs_tr_manifested = manifest(xs_tr_delayed)
let xs' = transpose(xs_tr_manifested)
let ([int,w] group_res, [int,n] vs_trans) =
  reduceKernel(f, -- reduction operator
    fn (int,[int,c]) (int c, [int,c] chunk) =>
      let init_ws = replicate(c, 0) in
      loop ((acc, ws') = (0, init_ws)) =
        for i < c do
          let x = chunk[i]
          let (int res) = f(acc, x)
          let (int v) = x * a
          let ([int,c] ws'') =
            ws' with [i] <- v
          in (res, ws''),
      (0), xs')
```

(b) The first of the two resulting kernels

Figure 6: Compilation of `redomap` to GPU kernels

5. Performance Analysis

5.1 Experimental Methodology and Hardware

Reported running times are averaged across one hundred runs within a loop. An initial iteration is executed without timing in order to avoid warmup and delayed initialization affecting the measured runtime. Running times represent core computation time. That is, it does not account for initial device-host memory transfers, GPU context creation, build time, reading/writing data from/to files, etc. Futhark programs are compiled and run with no special compiler flags (the default, i.e., no tuning). Thrust programs are compiled with `nvcc -O3` and `-arch=sm_35` where this did not hurt performance⁶, and we use the Git version of Thrust⁷ at commit ID 00315ad, as this was substantially faster than the pre-installed version. Running times are measured on a GeForce GTX 780 Ti with 3GiB of global memory and 2880 cores running at 1.08GHz. We use GCC 4.8.4 and CUDA 6.0 to compile C++/C and OpenCL/CUDA code (although as mentioned above, we use a newer version of Thrust than the one bundled with CUDA 6.0).

Our experimental setup is publicly available at the URL:

<https://github.com/HIPERFIT/futhark-array16>

We strongly believe in the value of reproducible results, and have attempted to automate the reproduction of our experiments. We have also documented common technical problems with running the experiments.

5.2 Mini-Benchmark Description

Figure 7 shows the Futhark code of the nine simple programs we have used for performance evaluation:

- 1 ReducePlus sums up the elements of an array of integers.

⁶Most of the benchmarks required the `-arch=sm_35` option in order to compile successfully, but `BlackScholes` and `MSSP` suffered significant slowdown when this option was used. We have no explanation for this. The reported runtimes for these two benchmarks are without the option `-arch=sm_35`.

⁷<https://github.com/thrust/thrust>

```

-- 1. ReducePlus --
-----
fun int main([int] a) = reduce(+, 0, a)

-- 2. ReduceMax --
-----
fun int main([int,n] as) = reduceComm(max, 0, as)
fun int max(int x, int y) = if x < y then y else x

-- 3. IndexOfMax --
-----
fun int main([int,n] as) =
  let (_,i)=reduceComm(maxWithIndex,(0,0),zip(as,iota(n)))
  in i
fun (int,int) maxWithIndex((int,int) x, (int,int) y) =
  let (xv, xi) = x in let (yv, yi) = y
  in if xv < yv then y
    else if yv < xv then x
    else if xi < yi then x else y

-- 4. IndexOfMaxPack --
-----
fun int main([int,n] as) =
  let i = reduceComm(maxWithIndex, 0i64,
    map(1, zip( map(<<32i64,map(i64,as))
      , map(i64,iota(n)) ) ) )
  in int(i)
fun i64 maxWithIndex(i64 x, i64 y) =
  let (xv, xi) = (int(x >> 32i64) & 0xFFFFFFFF, int(x))
  let (yv, yi) = (int(y >> 32i64) & 0xFFFFFFFF, int(y))
  in if xv < yv then y
    else if yv < xv then x
    else if xi < yi then x else y

-- 5. Reduce2x2MM --
-----
fun int main([int] a) =
  loop(s = 1) = for i < 42 do
    let a' = map(+s, a)
    in reduce(twoByTwoMult, 0xFF00FF00, a')
  in s

fun int twoByTwoMult(int x, int y) =
  let (x11, x12, x21, x22) = unpackInt(x)
  let (y11, y12, y21, y22) = unpackInt(y)
  let z11 = x11 * y11 + x12 * y21
  let z12 = x11 * y12 + x12 * y22
  let z21 = x21 * y11 + x22 * y21
  let z22 = x21 * y12 + x22 * y22
  in packInt(z11, z12, z21, z22)

fun (i8, i8, i8, i8) unpackInt(int x) =
  (i8(x >>> 24), i8(x >>> 16), i8(x >>> 8), i8(x >>> 0))
fun int packInt(i8 a, i8 b, i8 c, i8 d) =
  (((i32(a)&0xFF) << 24) | ((i32(b)&0xFF) << 16) |
  ((i32(c)&0xFF) << 8) | ((i32(d)&0xFF) << 0))

-- 6. MSSP (maximum segment sum problem) --
-----
fun int main([int,n] xs) =
  let (x, _, _, _) =
    reduce(redOp, (0,0,0,0), map(mapOp, xs)) in
  x
fun (int,int,int,int) redOp((int,int,int,int) x,
  (int,int,int,int) y) =
  let (mssx, misx, mcx, tsx) = x in
  let (mssy, misy, mcsy, tsy) = y in
  ( max(mssx, max(mssy, mcx + misy))
  , max(misx, tsx+misy), max(mcsy, mcx+tsy), tsx + tsy)

fun (int,int,int,int) mapOp (int x) =
  (max(x,0), max(x,0), max(x,0), x)

-- 7. ScanPlus --
-----
fun [int] main([int] a) = scan(+, 0, a)

-- 8. RedomapNT: resembles fusion's running example --
-- 9. Black Scholes: not shown --

```

Figure 7: Mini-Benchmark used for evaluating the Redomap construct. RedomapNT has a structure similar to the one in Figure 3.

- 2 ReduceMax computes the maximum of an integral array.
- 3 IndexOfMax computes the index of the maximal element of an integral array. The difference with the previous benchmarks is that the reduce operator is defined over index-value *tuples*, rather than basic-type scalars.
- 4 IndexOfMaxPack uses a 64-bit integer to pack together the index and the value into a scalar.
- 5 Reduce2x2MM consists of a loop of count 42, in which the loop-variant integral scalar *s* is added to each element of an input array, and then the array is reduced with the 2-by-2-matrix-multiplication operator, where a 32-bit integer is assumed to pack the four byte-size elements of the matrix. The result of the reduce becomes the value of *s* for the next iteration. Since the reduce operator is not associative, the Thrust version is implemented as a `transform_inclusive_scan`, which fuses together the map and the scan and selects the last element of the scanned array.
- 6 MSSP is the maximum-segment-sum problem: it looks for the segment of consecutive array elements whose element-wise sum is the largest across all possible segments, and returns the sum. The reduce operator is not commutative and operates on four-integer tuples, hence the Thrust implementation uses a `transform_inclusive_scan`.
- 7 ScanPlus is a prefix-sum computation on an array of integers.
- 8 RedomapNT is a code whose structure resembles the one in Figure 3, except that the mapped functions are more computationally expensive, e.g., involve division and exp. We note that Thrust does not support an operator that can express the fully-fused code, as in Futhark.
- 9 BlackScholes [20] implements a simple pricing engine for European options, and can be represented by one redomap in which the mapped arrays are consumed by the reduce (i.e., no arrays are returned).

5.3 Mini-Benchmark Evaluation

Figure 8 shows the running time (in milliseconds) of the Futhark and Thrust code of the eight mini programs, when the input arrays have sizes: 10^2 , $5 \cdot 10^4$, 10^5 , $5 \cdot 10^5$, 10^6 , $5 \cdot 10^6$, and 10^7 . Important observations are:

- ReducePlus and ReduceMax operates on 32-bit-integer scalars and exhibit very similar behavior: Futhark is about $3\times$ faster on sizes up to one million, and the gap narrows after that, e.g., only $1.55\times$ faster on the largest dataset. IndexOfMaxPack operates on 64-bit integers and shows a similar trend: Futhark is about $1.3\times$ faster on the largest array size.
- IndexOfMax operates on two-integer *tuples* which seems to significantly affect Thrust's performance: on arrays of size fifty thousand, Futhark is about $4\times$ faster, and the speedup reaches about $14\times$ for the largest size. This is likely due to genericity-related constraints, such as using an array of tuple representation (we use Thrust's zip-iterator). ThrustOpt denotes the version of the code that uses Thrust's built-in, hence optimized, `max_element` operator, but this is still $2\times$ slower than Futhark.
- MSSP performs a reduction with a *non-commutative* operator on a four-integer *tuple*. We have implemented this as a scan in Thrust. On smaller sizes, Thrust version is slightly faster, but Futhark is about $2.5\times$ faster on the largest dataset.
- Reduce2by2MM also performs a reduction with non-commutative operator, but on integer scalars. Also, the array being reduced is invariant to the 42-iteration outer loop, which permits Futhark to amortise the cost of the manifested transposition that is re-

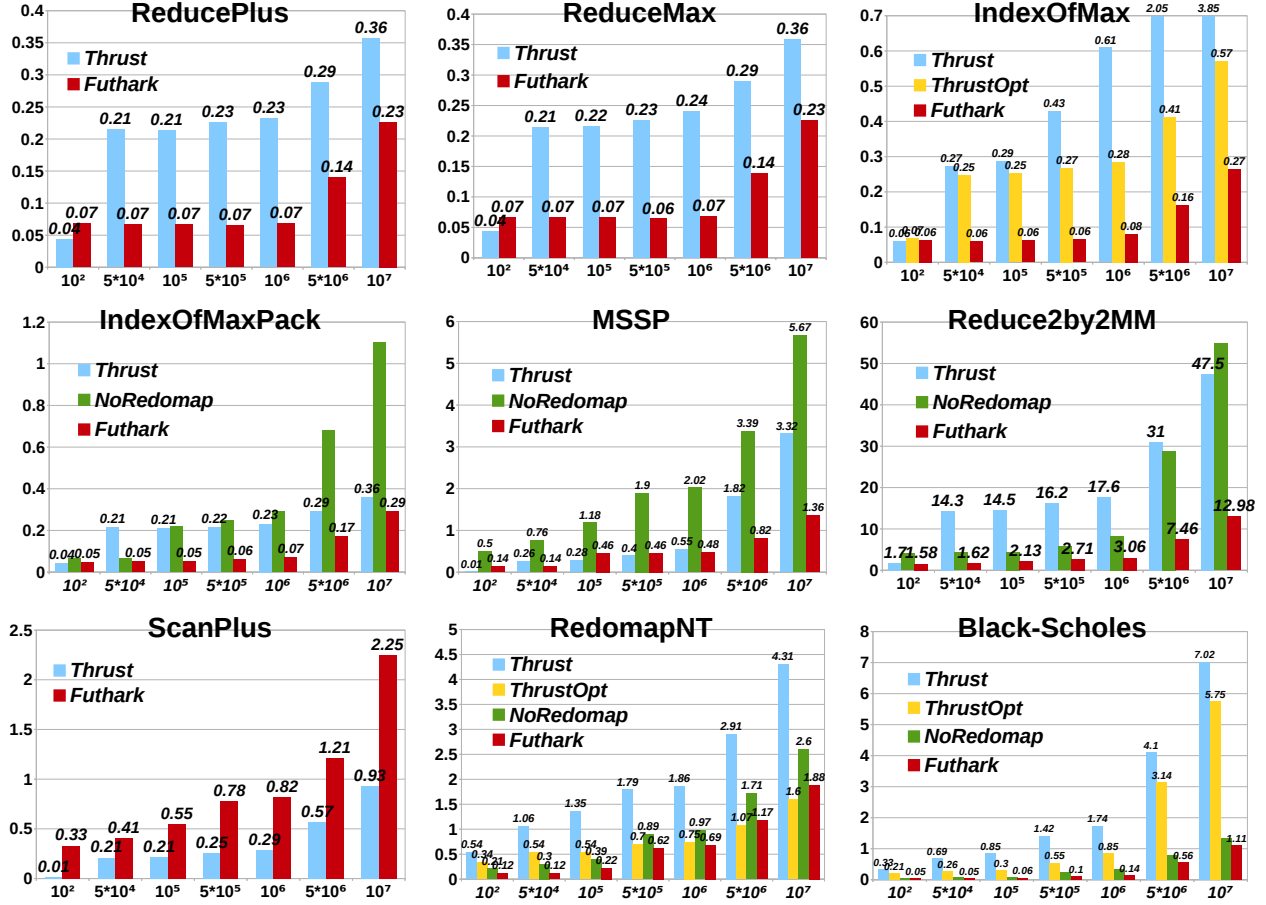


Figure 8: Running Times for Thrust and Futhark on GPGPU. X-Axis is the Array Size; Y-Axis is the runtime (in milliseconds). Thrust/ThrustOpt are “natural/tuned” versions of Thrust; NoRedomap/Futhark are Futhark-generated code where redomap fusion is off/on.

quired when executing non-commutative reductions. Futhark is about $8.8\times$ faster on an array of size fifty thousand and speedup decreases to only $3.6\times$ on the largest size. This suggests that Thrust’s inefficiencies are related to a larger extent with the use of tuples rather than to the use of `scan`.

- In fact, `ScanPlus` shows that Thrust’s `scan` operator is up to $2.4\times$ faster than Futhark’s. This is because Futhark’s `scan` is implemented in a manner similar to `redomap` (i.e., efficiently sequentializes the computation at the cost of transposing both the input and result arrays (to preserve coalesced accesses)). The overhead introduced the transposition is however too high, as it reaches two thirds of the total `scan` runtime.
- On `RedomapNT` the Futhark compiler has fused all operators into one `redomap`, without duplicating any computation. Since Thrust does not support an operator capable of expressing this, the Thrust optimized version corresponds to the code being fused in two kernels: semantically a `redomap` that “consumes” the input arrays and another `map` that duplicates some of the computation performed in `redomap`. Thrust unoptimized version corresponds to the unfused code. We observe that Futhark is about $2.3\times$ faster than the unoptimized Thrust and only $1.2\times$ slower than the optimized Thrust code. This is due to some redundant-copy inefficiencies in our translation, but even so, Futhark would still win if the map computations would be heavier (as it is typically the case in real-world applications).

- On `Black-Scholes`, the unoptimized and optimized Thrust versions correspond to unfused and fully fused code. The latter is possible since the `reduce` consumes the mapped arrays, and is implemented in Thrust by the `transform_reduce` and `transform_iterator` operators. Futhark is about $5\times$ faster on the largest dataset, which seems to indicate that the Thrust’s code manifests some of the intermediary arrays (i.e., it is memory bound rather than compute bound as in Futhark).

Finally, the code version denoted `NoRedomap` is produced by the Futhark compiler when the `redomap` fusion is disabled. `Redomap` fusion has significant impact ($\sim 4\times$) in the cases of `indexOfMaxPack`, `MSSP` and `Reduce2by2MM`, and smaller impact for `RedomapNT` and `Black-Scholes`.

6. Related Work

One strand of related work is embedded domain-specific languages targeting GPGPU execution, such as Nikola [19], Accelerate [20], Obsidian [6], and SPL [10]. Such embedded solutions often suffer significant limitations imposed by the host language, for example they do not support nested parallelism. Perhaps more related to Futhark is the work on SAC [11, 12], which is a standalone functional language that, like Futhark, also seeks a common ground with imperative approaches, but, unlike Futhark, does not support tuple types and operators such as `filter` and `scan` in the intermediate representation. The *with*-loops of SAC can fulfill the same

role as *redomap* [13], although the fusion algorithm in which they are used is very different: in SAC, producer-consumer and horizontal fusion is combined in a general framework of *with-loop fusion*.

Another strand of related work is concerned with automatic parallelization of imperative code. Solution include (i) static dependency analyses by polyhedral analysis [14, 25], and (ii) dynamic extraction of parallelism, for example by means of thread-level speculation [7, 24]. The latter, however, have not been found suitable for GPGPU execution.

Finally, a rich body of work has studied interoperability solutions between array languages, for example, (i) for inter-operating across computer-algebra systems [5, 23] or (ii) for efficient integration of option pricing in banks' IT infrastructure [22], or (iii) for integrating a contract-specification language [2] with dynamical graphical user interface [9]. Since Futhark is intended as an accelerator language, work is in progress to extend the compiler with various backends and friendly foreign function interfaces for various mainstream languages, such as Python, Java, C++, etc, as well as using Futhark in conjunction with TAIL [8] as an intermediate language for an APL compiler.

7. Conclusions

This paper has presented a new data-parallel core language construct, *redomap*, that can be used to perform both horizontal and producer-consumer fusion in a variety of complex cases, and which enables efficient mapping to parallel hardware.

Furthermore, we have validated our fully-automatic compiler on a range of microbenchmarks that perform both simple and nontrivial reductions, several with non-commutative operators. We show that we obtain performance comparable to or exceeding that of the Thrust library in most cases involving reductions, thus empirically demonstrating the potential for efficient compilation of the *redomap* construct.

We have also shown the value of supporting reductions with non-commutative operators, as well as demonstrated that such reductions can be compiled efficiently on modern GPU hardware.

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