Arrows for Parallel Computations

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Abstract

Arrows are a general interface for computation and therefore form an alternative to Monads for API design. We express parallelism using this concept in a novel way: We define an arrows-based language for parallelism and implement it using multiple parallel Haskells. In this manner we are able to bridge across various parallel Haskells.

Additionally, our way of writing parallel programs has the benefit of being portable across multiple parallel Haskell implementations. Furthermore, as each parallel computation is an arrow, which means that they can be readily composed and transformed as such. We introduce some syntactic sugar to provide parallelism-aware arrow combinators.

To show that our arrow-based language has similar expressiveness to existing parallel languages, we also define several parallel skeletons with our framework. Benchmarks show that our framework does not induce too much overhead performance-wise. OL: Summarize conclusions

MB: Jedes Kapitel soll einmal ins Abstract. Conclusions sollen mit ins Abstract

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1 Introduction

One particular reason for this is the ease of introducing new sophisticated language concepts. Parallel functional languages have a long history of being used for experimenting with novel parallel programming paradigms including the expression of parallelism. Haskell, which we focus on in this paper, has several mature implementations. We regard here in-depth the GpH (a SMP implementation of GpH language), the Par Monad, and Eden, a distributed memory parallel Haskell. These languages represent orthogonal approaches. Some use a monad, even if only for the internal representation. Some introduce additional language constructs. A key novelty here is to use Arrows to represent parallel computations. They seem a natural fit as they are a generalization of the function \rightarrow and serve as general interface to computations.

We provide an Arrows-based typeclass and implementations for three parallel Haskells. Instead of introducing a new low-level parallel backend in order to implement our Arrows-based interface, we define a shallow-embedded DSL for Arrows. This DSL is defined as a common interface and varying implementations in existing parallel Haskells: GpH, *Par* Monad, and Eden. Thus, we not only define a parallel programming interface in a novel manner - we tame the zoo of parallel Haskells. We provide a common **Phil: quantify, e.g. less than 10** %, very low-penalty programming interface that allows to switch the parallel Haskell backends at will. Further backends based on HdpH or a Frege implementation (on the Java Virtual Machine) are viable, too.

Contributions We propose a new Arrow-based encoding for parallelism based on a new Arrow combinator $parEvalN :: [arr\ a\ b] \to arr\ [a]\ [b]$, which converts a list of Arrows into a new parallel Arrow. Because of this, we do not lose any benefits of using Arrows as the parallelism is encapsulated as another combinator instead of a different type. The resulting Arrow can be used in the same way as a potential serial version.

This is a big advantage as we do not introduce any new types as Monad solutions like the *Par* Monad do. We can just 'plug' in parallel parts into sequential Arrow-based programs

and libraries without having to change integral parts of the code. MB: mention Functions here? We have everything Eden has, but more.

MB: Überleitung auf Related Work hier machen

We do not reimplement all the parallel internals, as we host this functionality in the *ArrowParallel* typeclass, which abstracts all parallel implementation logic. This way, we can have multiple backends - currently a GpH, a *Par* Monad and an Eden version are supported - backends can easily be swapped, so we are not bound to any specific one. So as an example, during development, we can run the program in a simple GHC-compiled variant using a GpH backend and afterwards deploy it on a cluster by converting it into an Eden program, by just replacing the *ArrowParallel* instance and compiling with Eden's GHC variant.

We show that it is possible to define algorithmic skeletons with Arrows (Sections 6, 7) and finally demonstrate that Arrow Parallelism is a viable alternative to existing approaches and prove that only low overhead is introduced, i.e. less than x% MB: fix this after benchmarks are finally done as shown in Section 8.

MB: Mention Future work here?

2 Related Work

Parallel Haskells. Of course, the three parallel Haskell flavours we use as backends: the GpH (Trinder *et al.*, 1996, 1998) parallel Haskell dialect and its multicore version (Marlow *et al.*, 2009), the *Par* Monad (Marlow *et al.*, 2011; Foltzer *et al.*, 2012), and Eden (Loogen *et al.*, 2005; Loogen, 2012) are related to this work. We use these languages as backends: our DSL can switch from one to other at user's command.

HdpH (Maier *et al.*, 2014; Stewart *et al.*, 2016) is an extension of *Par* Monad to heterogeneous clusters. LVish (Kuper *et al.*, 2014) is a communication-centred extension of *Par* Monad. Further parallel Haskell approaches include pH (Nikhil & Arvind, 2001), research work done on distributed variants of GpH (Trinder *et al.*, 1996; Aljabri *et al.*, 2014, 2015), and low-level Eden implementation (Berthold, 2008; Berthold *et al.*, 2016). Skeleton composition (Dieterle *et al.*, 2016), communication (Dieterle *et al.*, 2010a), and generation of process networks (Horstmeyer & Loogen, 2013) are recent in-focus research topics in Eden. This also includes the definitions of new skeletons (Hammond *et al.*, 2003; Berthold & Loogen, 2006; Berthold *et al.*, 2009b,c; Dieterle *et al.*, 2010b; de la Encina *et al.*, 2011; Dieterle *et al.*, 2013; Janjic *et al.*, 2013).

More different approaches include data parallelism (Chakravarty *et al.*, 2007; Keller *et al.*, 2010), GPU-based approaches (Mainland & Morrisett, 2010; Svensson, 2011), software transactional memory (Harris *et al.*, 2005; Perfumo *et al.*, 2008). The Haskell–GPU bridge Accelerate (Chakravarty *et al.*, 2011; Clifton-Everest *et al.*, 2014; McDonell *et al.*, 2015) deserves a special mention. Accelerate is completely orthogonal to our approach. Marlow authored a recent book in 2013 on parallel Haskells.

Algorithmic skeletons. Algorithmic skeletons were introduced by Cole (1989). Early publications on this topic include (Darlington *et al.*, 1993; Botorog & Kuchen, 1996; Danelutto *et al.*, 1997; Gorlatch, 1998; Lengauer *et al.*, 1997). Rabhi & Gorlatch (2003) consolidated early reports on high-level programming approaches. The effort is ongoing,

including topological skeletons (Berthold & Loogen, 2006), special-purpose skeletons for computer algebra (Berthold et al., 2009c; Lobachev, 2011, 2012; Janjic et al., 2013), iteration skeletons (Dieterle et al., 2013). The idea of Linton et al. (2010) is to use a parallel Haskell to orchestrate further software systems to run in parallel. Dieterle et al. (2016) compare the composition of skeletons to stable process networks.

Arrows. Arrows were introduced by Hughes (2000), basically they are a generalised function arrow \rightarrow . Hughes (2005a) presents a tutorial on Arrows. Some theoretical details on Arrows (Jacobs et al., 2009; Lindley et al., 2011; Atkey, 2011) are viable. Paterson (2001) introduced a new notation for Arrows. Arrows have applications in information flow research (Li & Zdancewic, 2006, 2010; Russo et al., 2008), invertible programming (Alimarine et al., 2005), and quantum computer simulation (Vizzotto et al., 2006). But probably most prominent application of Arrows is Arrow-based functional reactive programming, AFRP (Nilsson et al., 2002; Hudak et al., 2003; Czaplicki & Chong, 2013). Liu et al. (2009) formally define a more special kind of Arrows that capsule the computation more than regular arrows do and thus enable optimizations. Their approach would allow parallel composition, as their special Arrows would not interfere with each other in concurrent execution. In contrast, we capture a whole parallel computation as a single entity: our main instantiation function parEvalN makes a single (parallel) Arrow out of list of Arrows. Huang et al. (2007) utilise Arrows for parallelism, but strikingly different from our approach. They basically use Arrows to orchestrate several tasks in robotics. We, however, propose a general interface for parallel programming, while remaining completely in Haskell.

Other languages. Although this work is centred on Haskell implementation of arrows, it is applicable to any functional programming language where parallel evaluation and arrows can be defined. Experiments with our approach in Frege language¹ (which is basically Haskell on the JVM) were quite successful. However, it is beyond the scope of this work.

Achten et al. (2004, 2007) use an arrow implementation in Clean for better handling of typical GUI tasks. Dagand et al. (2009) used arrows in OCaml in the implementation of a distributed system.

3 Background

3.1 Arrows

Arrows were introduced by Hughes (2000) as a general interface for computation. An Arrow arr a b represents a computation that converts an input a to an output b. This is defined in the Arrow type class shown in Fig. 1. Its arr operation is used to lift an ordinary function to the specified arrow type, analogous to the monadic return. The >>> operator is analogous to the monadic composition \gg and combines two arrows arr a b and arr b c by "wiring" the outputs of the first to the inputs to the second to get a new arrow arr a c. Lastly, the first operator takes the input arrow from b to c and converts it into an arrow on pairs with the

¹ GitHub project page at https://github.com/Frege/frege

```
class Arrow\ arr\ where arr::(a \rightarrow b) \rightarrow arr\ a\ b (>>>)::arr\ a\ b \rightarrow arr\ b\ c \rightarrow arr\ a\ c first::arr\ a\ b \rightarrow arr\ (a,c)\ (b,c) instance Arrow\ (\rightarrow) where arr\ f=f f>>>> g=g\circ f first\ f=\lambda(a,c) \rightarrow (f\ a,c) data Kleisli\ m\ a\ b=Kleisli\ \{run::a\rightarrow m\ b\} instance Monad\ m\Rightarrow Arrow\ (Kleisli\ m) where arr\ f=Kleisli\ (return\circ f) f>>>> g=Kleisli\ (\lambda a\rightarrow f\ a>=g) first\ f=Kleisli\ (\lambda(a,c)\rightarrow f\ a>=\lambda b\rightarrow return\ (b,c))
```

Figure 1: The definition of *Arrow* type class and its two most typical instances.

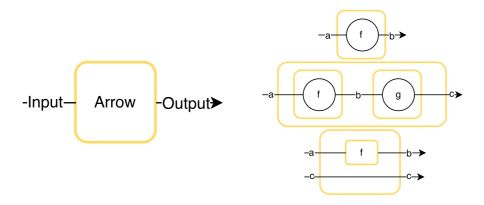


Figure 2: Schematic depiction of Arrow (left) and its basic combinators *arr*, >>> and *first* (right).

second argument untouched. It allows us to to save input across arrows. Figure 2 shows a graphical representation of the basic Arrow combinators. The most prominent instances of this interface are regular functions (\rightarrow) and the Kleisli type (Fig. 1), which wraps functions that return monadic values, e.g. $a \rightarrow mb$ with m being a Monad.

Hughes also defined some syntactic sugar (Fig. 3): The combinators *second*, *** and &&&. The combinator *second* is the mirrored version of *first* (Appendix A). The *** function combines *first* and *second* to handle two inputs in one arrow, is defined as

```
(***)::Arrow arr \Rightarrow arr\ a\ b \rightarrow arr\ c\ d \rightarrow arr\ (a,c)\ (b,d)
f ***g = first\ f >>> second\ g
```

while the &&& combinator, that constructs an arrow which outputs two different values like ***, but takes only one input, is:

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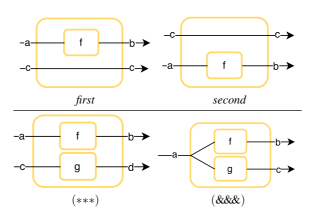


Figure 3: Visual depiction of syntactic sugar for Arrows.

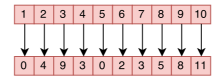


Figure 4: Schematic illustration of *parEvalN*. A list of inputs is transformed by different functions in parallel.

```
(&&&)::Arrow arr \Rightarrow arr\ a\ b \rightarrow arr\ a\ c \rightarrow arr\ a\ (b,c)
f &&& g = arr\ (\lambda a \rightarrow (a,a)) >>> (f *** g)
```

A first short example given by Hughes on how to use arrows is addition with arrows:

```
add:: Arrow arr \Rightarrow arr \ a \ Int \rightarrow arr \ a \ Int \rightarrow arr \ a \ Int
add \ f \ g = (f \&\&\& \ g) >>> arr \ (\lambda(u,v) \rightarrow u + v)
```

The more restrictive interface of Arrows allows for more elaborate composition and transformation combinators—a Monad can be *anything*, an Arrow is a process of doing something, a *computation*. This is exactly one of the key challenges in parallel computing.

3.2 Short introduction to parallel Haskells

In its purest form, parallel computation (on functions) can be looked at as the execution of some functions $a \to b$ in parallel or $parEvalN :: [a \to b] \to [a] \to [b]$, as also Figure 4 symbolically shows.

In this section, we will implement this non-Arrow version which will later be adapted for usage in our Arrow based parallel Haskell.

There exist several parallel Haskells already. Among the most important are probably GpH (based on *par & pseq* "hints") (Trinder *et al.*, 1996, 1998), the *Par* Monad (a monad for deterministic parallelism) (Marlow *et al.*, 2011; Foltzer *et al.*, 2012), Eden (Loogen *et al.*, 2005; Loogen, 2012) (a parallel Haskell for distributed memory), HdpH (a Template Haskell based parallel Haskell for distributed memory) (Maier *et al.*, 2014; Stewart *et al.*, 2016) and LVish (a *Par* extension with focus on communication) (Kuper *et al.*, 2014).

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main

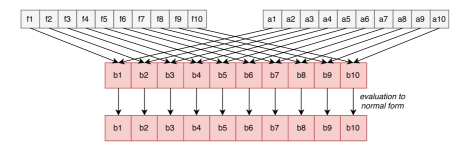


Figure 5: Dataflow of the GpH parEvalN version.

As the goal of this paper is not to reimplement yet another parallel runtime, but to represent parallelism with Arrows, we base our efforts on existing work which we wrap as backends behind a common interface. For this paper we chose GpH - for its simplicity, the *Par* Monad - to represent a monadic DSL, and Eden - as a distributed parallel Haskell.

LVish and HdpH were not chosen as the former does not differ from the original *Par* Monad with regard to how we would have used it in this paper, while the latter (at least in its current form) does not comply with our representation of parallelism due to its heavy reliance on TemplateHaskell.

We will now go into some detail on GpH, the *Par* Monad and Eden, and also give their respective implementations of the non-Arrow version of *parEvalN*.

3.2.1 Glasgow parallel Haskell - GpH

GpH (Marlow *et al.*, 2009; Trinder *et al.*, 1998) is one of the simplest ways to do parallel processing found in standard GHC.² Besides some basic primitives (*par* & *pseq* hints), it ships with parallel evaluation strategies for several types which can be applied with $using :: a \rightarrow Strategy \ a \rightarrow a$, which is exactly what is required for an implementation of parEvalN.

```
parEvalN :: (NFData\ b) \Rightarrow [a \rightarrow b] \rightarrow [a] \rightarrow [b]

parEvalN\ fs\ as = \mathbf{let}\ bs = zipWith\ (\$)\ fs\ as

\mathbf{in}\ bs\ `using\ `parList\ rdeepseq
```

In the above definition of parEvalN we just apply the list of functions $[a \rightarrow b]$ to the list of inputs [a] by zipping them with the application operator \$. We then evaluate this lazy list [b] according to a Strategy[b] with the $using: a \rightarrow Strategy[a \rightarrow a]$ operator. We construct this strategy with $parList: Strategy[a] \rightarrow Strategy[a]$ and $rdeepseq: NFData[a] \Rightarrow Strategy[a]$ where the latter is a strategy which evalutes to normal form. Fig. 5 shows a visual representation of this code.

² The Multicore implementation of GpH is available on Hackage under https://hackage.haskell.org/package/parallel-3.2.1.0, compiler support is integrated in the stock GHC.

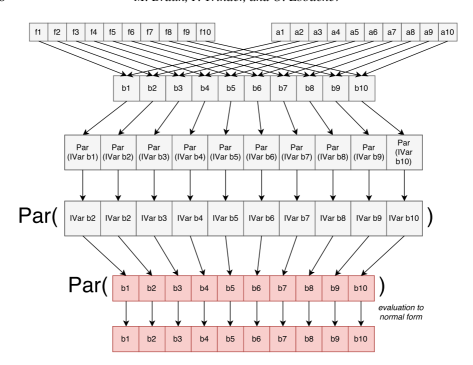


Figure 6: Dataflow of the Par Monad parEvalN version.

3.2.2 Par Monad

The *Par* Monad³ introduced by Marlow *et al.* (2011), is a Monad designed for composition of parallel programs. Let:

```
parEvalN :: (NFData\ b) \Rightarrow [a \rightarrow b] \rightarrow [a] \rightarrow [b]

parEvalN\ fs\ as = runPar\ \$

(sequenceA\ \$map\ (spawnP)\ \$zipWith\ (\$)\ fs\ as) > mapM\ get
```

The Par Monad version of our parallel evaluation function parEvalN is defined by zipping the list of $[a \rightarrow b]$ with the list of inputs [a] with the application operator \$ just like with GpH. Then, we map over this not yet evaluated lazy list of results [b] with $spawnP :: NFData \ a \Rightarrow a \rightarrow Par\ (IVar\ a)$ to transform them to a list of not yet evaluated forked away computations $[Par\ (IVar\ b)]$, which we convert to $Par\ [IVar\ b]$ with sequenceA. We wait for the computations to finish by mapping over the $IVar\ b$ values inside the Par Monad with get. This results in $Par\ [b]$. We execute this process with runPar to finally get [b]. Fig. 6 shows a graphical representation.

The Par monad can be found in the monad-par package on Hackage under https://hackage.haskell.org/package/monad-par-0.3.4.8/.

3.2.3 Eden

Eden (Loogen *et al.*, 2005; Loogen, 2012) is a parallel Haskell for distributed memory and comes with a MPI and a PVM backends.⁴ It is targeted towards clusters, but also functions well in a shared-memory setting with a further simple backend. However, in contrast to many other parallel Haskells, in Eden each process has its own heap. This seems to be a waste of memory, but with distributed programming paradigm and individual GC per process, Eden yields good performance results also on multicores (Berthold *et al.*, 2009a; Aswad *et al.*, 2009).

While Eden also comes with a Monad PA for parallel evaluation, it also ships with a completely functional interface that includes a spawnF:: $(Trans\ a, Trans\ b) \Rightarrow [a \rightarrow b] \rightarrow [a] \rightarrow [b]$ function that allows us to define parEvalN directly:

```
parEvalN :: (Trans\ a, Trans\ b) \Rightarrow [a \rightarrow b] \rightarrow [a] \rightarrow [b]

parEvalN = spawnF
```

Eden TraceViewer. To comprehend the efficiency and the lack thereof in a parallel program, an inspection of its execution is extremely helpful. While some large-scale solutions exist (Geimer *et al.*, 2010), the parallel Haskell community mainly utilises the tools Threadscope (Wheeler & Thain, 2009) and Eden TraceViewer⁵ (Berthold & Loogen, 2007). In the next sections we will present some *trace visualizations*, the post-mortem process diagrams of Eden processes and their activity.

The trace visualizations are color-coded. In such a visualization (Fig. 13), the *x* axis shows the time, the *y* axis enumerates the machines and processes. The visualization shows a running process in green, a blocked process is red. If the process is 'runnable', i.e. it may run, but does not, it is yellow. The typical reason for this is GC. An inactive machine, where no processes are started yet, or all are already terminated, shows as a blue bar. A comminication from one process to another is represented with a black arrow. A stream of communications, e.g. a transmitted list is shows as a dark shading between sender and receiver processes.

4 Parallel Arrows

Arrows are a general interface to computation. Here we introduce special Arrows as general interface to *parallel computations*. First, we present the interface and explain the reasonings behind it. Then, we discuss some implementations using exisiting parallel Haskells. Finally, we explain why using Arrows for expressing parallelism is beneficial.

⁴ The projects homepage can be found at http://www.mathematik.uni-marburg.de/~eden/.
The Hackage page is at https://hackage.haskell.org/package/edenmodules-1.2.0.0/.

⁵ See http://hackage.haskell.org/package/edentv on Hackage for the last available version of Eden TraceViewer.

```
data Conf a = Conf (Strategy a)

instance (NFData b, ArrowApply arr, ArrowChoice arr) \Rightarrow ArrowParallel arr a b () where

parEvalN\_fs =

listApp\ fs >>>

arr\ (withStrategy\ (parList\ rdeepseq))
```

Figure 7: Fully evaluating ArrowParallel instance for the GpH Haskell backend.

4.1 The ArrowParallel typeclass

A parallel computation (on functions) in its purest form can be seen as execution of some functions $a \to b$ in parallel, as our parEvalN prototype shows (Sec. 3.2). Translating this into arrow terms gives us a new operator parEvalN that lifts a list of arrows $[arr\ a\ b]$ to a parallel arrow $arr\ [a]\ [b]$. This combinator is similar to our utility function listApp from Appendix A, but does parallel instead of serial evaluation.

```
parEvalN :: (Arrow \ arr) \Rightarrow [arr \ a \ b] \rightarrow arr \ [a] \ [b]
```

With this definition of *parEvalN*, parallel execution is yet another arrow combinator. But as the implementation may differ depending on the actual type of the arrow *arr* and we want this to be an interface for different backends, we introduce a new typeclass *ArrowParallel arr a b*:

```
class Arrow \ arr \Rightarrow Arrow Parallel \ arr \ a \ b where parEvalN :: [arr \ a \ b] \rightarrow arr \ [a] \ [b]
```

Sometimes parallel Haskells require or allow for additional configuration parameters, e.g. an information about the execution environment or the level of evaluation (weak head normal form vs. normal form). For this reason we also introduce an additional *conf* parameter to the function. We also do not want *conf* to be a fixed type, as the configuration parameters can differ for different instances of *ArrowParallel*.

```
class Arrow arr \Rightarrow ArrowParallel \ arr \ a \ b \ conf where parEvalN :: conf \rightarrow [arr \ a \ b] \rightarrow arr \ [a] \ [b]
```

We do not require the *conf* parameter in every implementation. If it is not needed, we usually just default the *conf* type parameter to () and even blank it out in the parameter list of the implemented *parEvalN*.

4.2 ArrowParallel instances

4.2.1 Glasgow parallel Haskell

The GpH implementation of *ArrowParallel* is implemented in a straightforward manner by using *listApp* (Appendix A) combined with the *withStrategy*:: *Strategy* $a \rightarrow a \rightarrow a$ combinators from GpH, where *withStrategy* is the same as *using*:: $a \rightarrow Strategy$ $a \rightarrow a$, but with flipped parameters. For most cases a fully evaluating version like in Fig. 7 would probably suffice, but as the GpH interface allows the user to specify the level of evaluation to be done via the *Strategy* interface, our DSL should allow for this. We therefore introduce

```
instance (NFData b,ArrowApply arr,ArrowChoice arr) ⇒
ArrowParallel arr a b (Conf b) where
parEvalN (Conf strat) fs =
  listApp fs >>>
  arr (withStrategy (parList strat)) && arr id >>>
  arr (uncurry pseq)
```

Figure 8: Configurable ArrowParallel instance for the GpH Haskell backend.

the *Conf a* data-type that simply wraps a *Strategy a*. With this definition in place, we can provide a delegating version of the non-configurable instance. We show this in Fig. B 1.

```
4.2.2 Par Monad
```

OL: introduce a newcommand for par-monad, "arrows", "parrows" and replace all mentions to them to ensure uniform typesetting done!, we write Arrows. also "Monad"? done! The Par Monad implementation (Fig. 9) makes use of Haskells laziness and Par Monad's $spawnP :: NFData \ a \Rightarrow a \rightarrow Par \ (IVar \ a)$ function. The latter forks away the computation of a value and returns an IVar containing the result in the Par Monad.

We therefore apply each function to its corresponding input value with and then fork the computation away with $arr \, spawnP$ inside a zipWithArr (Fig. A 3) call. This yields a list $[Par \, (IVar \, b)]$, which we then convert into $Par \, [IVar \, b]$ with $arr \, sequenceA$. In order to wait for the computation to finish, we map over the IVars inside the Par Monad with $arr \, (>= mapM \, get)$. The result of this operation is a $Par \, [b]$ from which we can finally remove the Monad again by running $arr \, runPar$ to get our output of [b].

```
instance (NFData b,ArrowApply arr,ArrowChoice arr) \Rightarrow ArrowParallel arr a b conf where parEvalN _fs = (arr $\lambda as \rightarrow (fs,as)) >>> zipWithArr (app >>> arr spawnP) >>> arr sequenceA >>> arr (>>=mapM get) >>> arr runPar
```

Figure 9: ArrowParallel instance for the Par Monad backend.

4.2.3 Eden

For both the GpH Haskell and *Par* Monad implementations we could use general instances of *ArrowParallel* that just require the *ArrowApply* and *ArrowChoice* typeclasses. With Eden this is not the case as we can only spawn a list of functions and we cannot extract simple functions out of arrows. While we could still manage to have only one class in the module by introducing a typeclass:



Figure 10: Schematic depiction of parEvalNLazy.

```
class (Arrow arr) \Rightarrow ArrowUnwrap arr where arr \ a \ b \rightarrow (a \rightarrow b)
```

However, we avoid doing so for aesthetic resons. For now, we just implement *ArrowParallel* for normal functions:

```
instance (Trans\ a, Trans\ b) \Rightarrow ArrowParallel\ (\rightarrow)\ a\ b\ conf\ where parEvalN\ \_fs\ as = spawnF\ fs\ as and the Kleisli type: 

instance (Monad\ m, Trans\ a, Trans\ b, Trans\ (m\ b)) \Rightarrow ArrowParallel\ (Kleisli\ m)\ a\ b\ conf\ where parEvalN\ conf\ fs = (arr\ parEvalN\ conf\ (map\ (\lambda(Kleisli\ f) \to f)\ fs)) >>> (Kleisli\ \$\ sequence)
```

4.3 Extending the Interface

With the *ArrowParallel* typeclass in place and implemented, we can now implement some further basic parallel interface functions. These are algorithmic skeletons that, however, mostly serve as a foundation to further, more specific algorithmic skeletons.

4.3.1 Lazy parEvalN

The function parEvalN is 100% strict, which means that it fully evaluates all passed arrows. Sometimes this might not be feasible, as it will not work on infinite lists of functions like e.g. $map\ (arr\circ(+))\ [1..]$ or just because we need the arrows evaluated in chunks. parEvalNLazy (Figs. 10, 11) fixes this. It works by first chunking the input from [a] to [[a]] with the given ChunkSize in $arr\ (chunksOf\ chunkSize)$. These chunks are then fed into a list $[arr\ [a]\ [b]]$ of parallel arrows created by feeding chunks of the passed ChunkSize into the regular parEvalN by using listApp. The resulting [[b]] is lastly converted into [b] with $arr\ concat$.

4.3.2 Heterogenous tasks

We have only talked about the paralellization arrows of the same type until now. But sometimes we want to paralellize heterogenous types as well. However, we can implement such a *parEval2* combinator (Figs. 12, B 12) which combines two arrows *arr* a b and *arr* c d

```
parEvalNLazy:: (ArrowParallel\ arr\ a\ b\ conf\ ,ArrowChoice\ arr\ ,ArrowApply\ arr\ ) \Rightarrow conf \to ChunkSize \to [arr\ a\ b] \to (arr\ [a]\ [b])
parEvalNLazy\ conf\ chunkSize\ fs = arr\ (chunksOf\ chunkSize\ ) >>> listApp\ fchunks >>> arr\ concat
where\ fchunks = map\ (parEvalN\ conf\ )\ chunksOf\ chunkSize\ fs
```

Figure 11: Definition of parEvalNLazy.



Figure 12: Schematic depiction of parEval2.

into a new parallel arrow arr(a,c)(b,d) quite easily with the help of the ArrowChoice typeclass. The idea is to use the +++ combinator which combines two arrows $arr\ a\ b$ and $arr\ c\ d$ and transforms them into $arr\ (Either\ a\ c)\ (Either\ b\ d)$ to get a common arrow type that we can then feed into parEvalN.

5 Futures

Consider a mock-up parallel arrow combinator:

```
someCombinator :: (Arrow arr) \Rightarrow [arr \ a \ b] \rightarrow [arr \ b \ c] \rightarrow arr \ [a] \ [c]
someCombinator \ fs1 \ fs2 = parEvalN \ () \ fs1 >>> rightRotate >>>> parEvalN \ () \ fs2
```

In a distributed environment, a resulting arrow of this combinator first evaluates all $[arr\ a\ b]$ in parallel, sends the results back to the master node, rotates the input once and then evaluates the $[arr\ b\ c]$ in parallel to then gather the input once again on the master node. Such situations arise, e.g. in scientific computations when the data distributed across the nodes needs to be transposed. A concrete example is 2D FFT computation (Gorlatch & Bischof, 1998; Berthold $et\ al.$, 2009c).

While the above example could be rewritten into only one *parEvalN* call by directly wiring the arrows properly together, it illustrates an important problem. When using a *ArrowParallel* backend that resides on multiple computers, all communication between the nodes is done via the master node, as shown in the Eden trace in Figure 13. This can become a serious bottleneck for larger amount of data and number of processes (as e.g. Berthold *et al.*, 2009c, showcases).

We should allow the nodes to communicate directly with each other. Eden already ships with "remote data" that enable this (Alt & Gorlatch, 2003; Dieterle *et al.*, 2010a). But as we want code with our DSL to be implementation agnostic, we have to wrap this context. We do this with the *Future* typeclass (Fig. 14). Since *RD* is only a type synonym for a

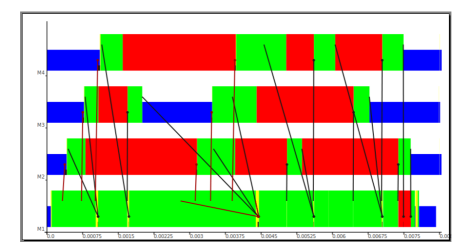


Figure 13: Communication between 4 Eden processes without Futures. All communication goes through the master node. Each bar represents one process. Black lines between processes represent communication. Colors: blue $\hat{=}$ idle, green $\hat{=}$ running, red $\hat{=}$ blocked, yellow $\hat{=}$ suspended.

OL: more practical and heavy-weight example! fft (I have the code)?

MB: Depends... Are the communications easy to read in such an example?

MB: Keep the description for the different colours, or link to the EdenTV description in 3.2.3

OL: ok as is OL: use the fft example (when it works)?

```
class Future fut a \mid a \rightarrow fut where

put :: (Arrow \ arr) \Rightarrow arr \ a \ (fut \ a)

get :: (Arrow \ arr) \Rightarrow arr \ (fut \ a) \ a
```

Figure 14: Definition of the *Future* typeclass.

communication type that Eden uses internally, we have to use some wrapper classes to fit that definition, though, as Fig. B 6 shows. Technical details are in Appendix, in Section B.

For our *Par* Monad and GpH Haskell backends, we can simply use *BasicFutures* (Fig. 15) which are just simple wrappers around the actual data, as in a shared memory setting we do not require Eden's sophisticated communication channels.

```
data BasicFuture\ a = BF\ a

instance (NFData\ a) \Rightarrow NFData\ (BasicFuture\ a) where

rnf\ (BF\ a) = rnf\ a

instance (NFData\ a) \Rightarrow Future\ BasicFuture\ a where

put = arr\ BF

get = arr\ (\lambda(BF\ a) \rightarrow a)
```

Figure 15: The *BasicFuture* type and its *Future* instance for the *Par* Monad and GpH Haskell backends.

In our communication example we can use this *Future* concept for direct communications between the nodes as shown in Fig. 16. In a distributed environment, this gives us a

```
\begin{split} some Combinator :: (Arrow \ arr) \Rightarrow [arr \ a \ b] \rightarrow [arr \ b \ c] \rightarrow arr \ [a] \ [c] \\ some Combinator \ fs1 \ fs2 = \\ parEvalN \ () \ (map \ (>>>put) \ fs1) >>> \\ rightRotate >>> \\ parEvalN \ () \ (map \ (get>>>>) \ fs2) \end{split}
```

Figure 16: The mock-up combinator in parallel.

communication scheme with messages going through the master node only if it is needed—similar to what is shown in the trace visualization in Fig. 17.OL: Fig. is not really clear. Do Figs with a lot of load? — fft?

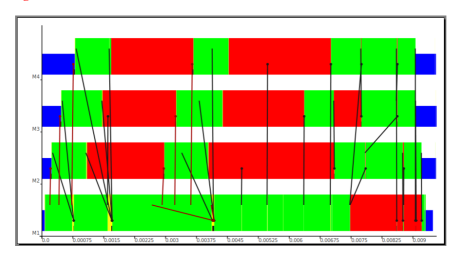


Figure 17: Communication between 4 Eden processes with Futures. Other than in Fig. 13, processes communicate directly (black lines between the bars) instead of always going through the master node (bottom bar).

6 Map-based Skeletons

Now we have developed Parallel Arrows far enough to define some useful algorithmic skeletons that abstract typical parallel computations.

Parallel *map* **and laziness.** The *parMap* skeleton (Figs. B 2, B 3) is probably the most common skeleton for parallel programs. We can implement it with *ArrowParallel* by repeating an arrow *arr a b* and then passing it into *parEvalN* to obtain an arrow *arr* [*a*] [*b*]. Just like *parEvalN*, *parMap* is 100% strict. As *parMap* is 100% strict it has the same restrictions as *parEvalN* compared to *parEvalNLazy*. So it makes sense to also have a *parMapStream* (Figs. B 4, B 5) which behaves like *parMap*, but uses *parEvalNLazy* instead of *parEvalN*. The code is quite straightforward, we show it in Appendix.

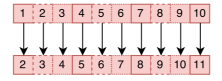


Figure 18: Schematic depiction of a farm, a statically load-balanced parMap.

```
\begin{array}{l} \textit{farm} :: (ArrowParallel\ arr\ a\ b\ conf\,, \\ ArrowParallel\ arr\ [a]\ [b]\ conf\,, ArrowChoice\ arr) \Rightarrow \\ conf \rightarrow \textit{NumCores} \rightarrow \textit{arr}\ a\ b \rightarrow \textit{arr}\ [a]\ [b] \\ \textit{farm}\ conf\ numCores\ f = \\ \textit{unshuffle}\ numCores >>> \\ \textit{parEvalN}\ conf\ (\textit{repeat}\ (\textit{mapArr}\ f)) >>> \\ \textit{shuffle} \end{array}
```

Figure 19: The definition of farm.

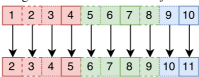


Figure 20: Schematic depiction of farmChunk.

Statically load-balancing parallel *map.* Our *parMap* spawns every single computation in a new thread (at least for the instances of *ArrowParallel* we gave in this paper). This can be quite wasteful and a statically load-balancing *farm* (Figs. 18, 19) that equally distributes the workload over *numCores* workers seems useful. The definitions of the helper functions *unshuffle*, *takeEach*, *shuffle* (Fig. B 7) originate from an Eden skeleton⁶.

Since a *farm* is basically just *parMap* with a different work distribution, it is, again, 100% strict. So we can define *farmChunk* (Figs. 20, B 10) which uses *parEvalNLazy* instead of *parEvalN*. It is basically the same definition as for *farm*, with *parEvalN* replaced with *parEvalNLazy*.

7 Topological Skeletons

Even though many algorithms can be expressed by parallel maps, some problems require more sophisticated skeletons. The Eden library leverages this problem and already comes with more predefined skeletons⁷, among them a *pipe*, a *ring*, and a *torus* implementations (Loogen *et al.*, 2003). These seem like reasonable candidates to be ported to our Arrowbased parallel Haskell. We aim to showcase that we can express more sophisticated skeletons with Parallel Arrows as well.

Available on Hackage under https://hackage.haskell.org/package/edenskel-2.1.0.0/docs/src/Control-Parallel-Eden-Map.html.

Available on Hackage: https://hackage.haskell.org/package/edenskel-2.1.0.0/docs/Control-Parallel-Eden-Topology.html.

If we used the original definition of *parEvalN*, however, these skeletons would produce an infinite loop with the GpH and Par Monad backends which during runtime would result in the program crashing. This materializes with the usage of *loop* of the *ArrowLoop* typeclass and is probably due to the way their respective parallelism engines work internally. **MB: okay so?** As these skeletons probably do not make any practical sense besides for testing with these backends anyways (because of the shared memory between the threads), we create an extra abstraction layer for the original *parEvalN* in these skeletons called *evalN* in the *FutureEval* typeclass. This allows us for selective enabling and disabling of parallelism.

```
class ArrowParallel arr a b conf \Rightarrow FutureEval arr a b conf where evalN :: (ArrowParallel \ arr \ a \ b \ conf) \Rightarrow conf \rightarrow [arr \ a \ b] \rightarrow arr \ [a] \ [b]
```

As Eden has no problems with the looping skeletons, we declare a delegating instance:

```
instance ArrowParallel arr a b conf \Rightarrow FutureEval arr a b conf where evalN = parEvalN
```

The Par Monad and GpH backends have parallelism disabled in their instance of *FutureEval*. This way the skeletons can still run without errors on shared-memory machines and still be used to test programs locally.

```
instance (Arrow arr, Arrow Choice arr, Arrow Apply arr,
Arrow Parallel arr a b conf) \Rightarrow Future Eval arr a b conf where
eval N_- = list App
```

7.1 Parallel pipe

The parallel *pipe* skeleton is semantically equivalent to folding over a list $[arr\ a\ a]$ of arrows with >>>, but does this in parallel, meaning that the arrows do not have to reside on the same thread/machine. We implement this skeleton using the ArrowLoop typeclass which gives us the $loop::arr\ (a,b)\ (c,b) \to arr\ a\ c$ combinator which allows us to express recursive fix-point computations in which output values are fed back as input. For example

```
loop (arr (\lambda(a,b) \rightarrow (b,a:b)))
```

which is the same as

```
loop (arr snd &&& arr (uncurry (:)))
```

defines an arrow that takes its input a and converts it into an infinite stream [a] of it. Using this to our advantage gives us a first draft of a pipe implementation (Fig. 21) by plugging in the parallel evaluation call *evalN* conf fs inside the second argument of && and then only picking the first element of the resulting list with arr last.

However, using this definition directly will make the master node a potential bottleneck in distributed environments as described in Section 5. Therefore, we introduce a more sophisticated version that internally uses Futures and obtain the final definition of *pipe* in Fig. 22.

Sometimes, this *pipe* definition can be a bit inconvenient, especially if we want to pipe arrows of mixed types together, i.e. *arr* a b and *arr* b c. By wrapping these two arrows inside a common type we obtain *pipe2* (Fig. 23).

```
\begin{array}{l} \textit{pipeSimple} :: (ArrowLoop\ arr, FutureEval\ arr\ a\ a\ conf) \Rightarrow \\ \textit{conf} \rightarrow [\textit{arr}\ a\ a] \rightarrow \textit{arr}\ a\ a \\ \textit{pipeSimple}\ conf\ fs = \\ \textit{loop}\ (\textit{arr}\ snd\ \&\&\& \\ \textit{(arr}\ (\textit{uncurry}\ (:) >>> \textit{lazy}) >>> \textit{evalN}\ conf\ fs)) >>> \\ \textit{arr}\ \textit{last} \end{array}
```

Figure 21: A first implementation of the *pipe* skeleton expressed with Parallel Arrows. Note that the use of lazy (Fig. B 8) is essential as without it programs using this definition would never halt. We need to enforce that the evaluation of the input [a] terminates before passing it into evalN.

```
pipe :: (ArrowLoop\ arr, FutureEval\ arr\ (fut\ a)\ (fut\ a)\ conf, Future\ fut\ a) \Rightarrow conf \to [arr\ a\ a] \to arr\ a\ a
pipe\ conf\ fs = unliftFut\ (pipeSimple\ conf\ (map\ liftFut\ fs))
```

Figure 22: Final definition of the *pipe* skeleton with Futures.

```
\begin{aligned} &\textit{pipe2} :: (ArrowLoop\ arr, ArrowChoice\ arr, Future\ fut\ (([a],[b]),[c]),\\ &\textit{FutureEval}\ arr\ (fut\ (([a],[b]),[c]))\ (fut\ (([a],[b]),[c]))\ conf) \Rightarrow\\ &\textit{conf} \rightarrow arr\ a\ b \rightarrow arr\ b\ c \rightarrow arr\ a\ c\\ &\textit{pipe2}\ conf\ f\ g =\\ &(arr\ return\ \&\&\&\ arr\ (const\ []))\ \&\&\&\ arr\ (const\ [])\ >>>\\ &\textit{pipe}\ conf\ (replicate\ 2\ (unifyf\ g))\ >>>\\ &\textit{arr\ snd}\ >>>\ arr\ head\ \textbf{where}\\ &\textit{unify} :: (ArrowChoice\ arr) \Rightarrow arr\ a\ b \rightarrow arr\ b\ c \rightarrow arr\ (([a],[b]),[c])\ (([a],[b]),[c])\\ &\textit{unify}\ g =\\ &(mapArr\ f\ ***\ mapArr\ g)\ ***\ arr\ (\ \_\rightarrow\ [])\ >>>\\ &arr\ (\lambda((a,b),c)\rightarrow ((c,a),b))\\ (|>>>|) :: (ArrowLoop\ arr, ArrowChoice\ arr, Future\ fut\ (([a],[b]),[c]),\\ &\textit{FutureEval}\ arr\ (fut\ (([a],[b]),[c]))\ (fut\ (([a],[b]),[c]))\ ()) \Rightarrow\\ &arr\ a\ b \rightarrow arr\ b\ c \rightarrow arr\ a\ c\\ (|>>>|) = pipe2\ ()\end{aligned}
```

Figure 23: Definition of *pipe2* and a parallel >>>.

Note that extensive use of *pipe2* over *pipe* with a hand-written combination data type will probably result in worse performance because of more communication overhead from the many calls to *parEvalN* inside of *evalN*. Nonetheless, we can define a version of parallel piping operator |>>>|, which is semantically equivalent to >>> similarly to other parallel syntactic sugar from Appendix C.

```
Another version of >>> is:
```

```
f > >> | g = (f \circ put) >>> (get \circ g)
```

It does not launch both arrows f and g in parallel, but allows for more smooth data communication between them. Basically, it is a *Future*-lifted *sequential* >>>>, a way to compose parallel Arrows efficiently.

Arrows for Parallel Computations

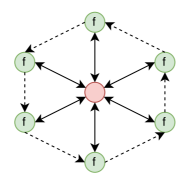


Figure 24: Schematic depiction of the ring skeleton.

```
\begin{array}{l} \textit{ring} :: (\textit{ArrowLoop arr}, \textit{Future fut } r, \textit{FutureEval arr } (i, \textit{fut } r) \ (o, \textit{fut } r) \ conf) \Rightarrow \\ \textit{conf} \rightarrow \textit{arr } (i, r) \ (o, r) \rightarrow \textit{arr } [i] \ [o] \\ \textit{ring conf } f = \\ \textit{loop (second (rightRotate >>> lazy) >>> arr (uncurry zip) >>> \\ \textit{evalN conf (repeat (second get >>> f >>> second put)) >>> arr unzip)} \end{array}
```

Figure 25: Final definition of the *ring* skeleton.

7.2 Ring skeleton

Eden comes with a ring skeleton⁸ (Fig. 24) implementation that allows the computation of a function $[i] \to [o]$ with a ring of nodes that communicate in a ring topology with each other. Its input is a node function $i \to r \to (o,r)$ in which r serves as the intermediary output that gets send to the neighbour of each node. This data is sent over direct communication channels, the so called 'remote data'. We depict it in Appendix, Fig. B 11.

We can rewrite this functionality easily with the use of *loop* as the definition of the node function, arr(i,r)(o,r), after being transformed into an arrow, already fits quite neatly into the *loop*'s $arr(a,b)(c,b) \rightarrow arr\ a\ c$. In each iteration we start by rotating the intermediary input from the nodes $[fut\ r]$ with $second\ (rightRotate >>> lazy)$ (Fig. B 8). Similarly to the pipe from Section 7.1 (Fig. 21), we have to feed the intermediary input into our lazy (Fig. B 8) arrow here, or the evaluation would fail to terminate. The reasoning is explained by Loogen (2012) as a demand problem.

Next, we zip the resulting $([i],[fut\,r])$ to $[(i,fut\,r)]$ with $arr\ (uncurry\,zip)$ so we can feed that into a our input arrow $arr\ (i,r)\ (o,r)$, which we transform into $arr\ (i,fut\,r)\ (o,fut\,r)$ before lifting it to $arr\ [(i,fut\,r)]\ [(o,fut\,r)]$ to get a list $[(o,fut\,r)]$. Finally we unzip this list into $([o],[fut\,r])$. Plugging this arrow $arr\ ([i],[fut\,r])\ ([o],fut\,r)$ into the definition of loop from earlier gives us $arr\ [i]\ [o]$, our ring arrow (Fig. 25). This combinator can, for example, be used to calculate the shortest paths in a graph using Warshall's algorithm.

⁸ Available on Hackage: https://hackage.haskell.org/package/edenskel-2.1.0.0/docs/Control-Parallel-Eden-Topology.html

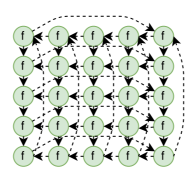


Figure 26: Schematic depiction of the *torus* skeleton.

7.3 Torus skeleton

If we take the concept of a *ring* from Section 7.2 one dimension further, we obtain a *torus* skeleton (Fig. 26, 27). Every node sends ands receives data from horizontal and vertical neighbours in each communication round. With our Parallel Arrows we re-implement the *torus* combinator⁹ from Eden—yet again with the help of the *ArrowLoop* typeclass.

Similar to the *ring*, we once again start by rotating the input (Fig. B 8), but this time not only in one direction, but in two. This means that the intermediary input from the neighbour nodes has to be stored in a tuple ([$[fut\ a]]$,[$[fut\ b]]$) in the second argument (loop only allows for two arguments) of our looped arrow $arr([[c]],([[fut\ a]],[[fut\ b]]))$ ([$[d]],([[fut\ a]],[[fut\ b]]))$ and our rotation arrow becomes

```
second ((mapArr rightRotate >>> lazy) *** (arr rightRotate >>> lazy))
```

instead of the singular rotation in the ring as we rotate $[[fut \ a]]$ horizontally and $[[fut \ b]]$ vertically. Then, we once again zip the inputs for the input arrow with

arr (uncurry3 zipWith3 lazyzip3)

from ([[c]], ([[fut a]], [[fut b]])) to [[(c, fut a, fut b)]], which we then feed into our parallel execution.

This action is, however, more complicated than in the ring case as we have one more dimension of inputs to be transformed. We first have to *shuffle* all the inputs to then pass it into *evalN conf* (repeat (ptorusf)) which yields [$(d,fut\ a,fut\ b)$]. We can then unpack this shuffled list back to its original ordering by feeding it into the specific unshuffle arrow we created one step earlier with $arr\ length >>> arr\ unshuffle$ with the use of $app::arr\ (arr\ a\ b,a)\ c$ from the arrowApply typeclass. Finally, we unpack this matrix [[[$(d,fut\ a,fut\ b)$]] with $arr\ (map\ unzip3) >>> arr\ unzip3 >>>> threetotwo\ to\ get\ ([[<math>[d]$], ([$[fut\ a]$], [$[fut\ b]$])).

As an example of using this skeleton (Loogen *et al.*, 2003) showed the matrix multiplication using the Gentleman algorithm (1978). An adapted version can be found in Fig. 28. If we compare the trace from a call using our arrow definition of the torus (Fig. 29) with the Eden version (Fig. 30) we can see that the behaviour of the arrow version and execution

⁹ Available on Hackage: https://hackage.haskell.org/package/edenskel-2.1.0.0/docs/Control-Parallel-Eden-Topology.html.

```
torus:: (ArrowLoop arr, ArrowChoice arr, ArrowApply arr, Future fut a, Future fut b,
   FutureEval arr (c, fut \ a, fut \ b) \ (d, fut \ a, fut \ b) \ conf) \Rightarrow
   conf \rightarrow arr(c,a,b)(d,a,b) \rightarrow arr[[c]][[d]]
torus conf f =
   loop (second ((mapArr rightRotate >>> lazy) *** (arr rightRotate >>> lazy)) >>>
   arr (uncurry3 (zipWith3 lazyzip3)) >>>
   (arr length >>> arr unshuffle) &&& (shuffle >>> evalN conf (repeat (ptorus f)) >>> app >>>
   arr (map unzip3) >>> arr unzip3 >>> threetotwo)
ptorus :: (Arrow arr, Future fut a, Future fut b) \Rightarrow
   arr(c,a,b)(d,a,b) \rightarrow arr(c,fut\ a,fut\ b)(d,fut\ a,fut\ b)
ptorus f = arr (\lambda \sim (c, a, b) \rightarrow (c, get \ a, get \ b)) >>> f >>> arr (\lambda \sim (d, a, b) \rightarrow (d, put \ a, put \ b))
```

Figure 27: Definition of the torus skeleton. The definitions of lazyzip3, uncurry3 and threetotwo have been omitted and can be found in Fig. B 9

```
type Matrix = [[Int]]
prMM\_torus :: Int \rightarrow Int \rightarrow Matrix \rightarrow Matrix \rightarrow Matrix
prMM_torus numCores problemSizeVal m1 m2 =
   combine $ torus () (mult torusSize) $ zipWith (zipWith (,)) (split m1) (split m2)
   \textbf{where} \ \textit{torusSize} = (\textit{floor} \circ \textit{sqrt}) \ \$ \textit{fromIntegral} \ \textit{numCores}
           combine = concat \circ (map (foldr (zipWith (++)) (repeat [])))
           split = splitMatrix (problemSizeVal 'div' torusSize)
   -- Function performed by each worker
mult :: Int \rightarrow ((Matrix, Matrix), [Matrix], [Matrix]) \rightarrow (Matrix, [Matrix], [Matrix])
mult\ size\ ((sm1,sm2),sm1s,sm2s) = (result,toRight,toBottom)
   where toRight = take (size - 1) (sm1 : sm1s)
     toBottom = take (size - 1) (sm2' : sm2s)
     sm2' = transpose \ sm2
     sms = zipWith \ prMMTr \ (sm1 : sm1s) \ (sm2' : sm2s)
     result = foldl1' matAdd sms
```

Figure 28: Adapted matrix multiplication in Eden using a the torus skeleton. prMM_torus is the parallel matrix multiplication. *mult* is the function performed by each worker. *prMMTr* calculates AB^T and is used for the (sequential) calculation in the chunks. splitMatrix splits the Matrix into chunks. matAdd calculates A + B. Omitted definitions can be found in B 13.

times are comparable. We discuss further examples on larger clusters and in a more detail in the next section.

8 Performance results

8.1 Hardware

We have tested our parallel DSL and algorithmic skeletons implemented in it. Benchmarks were conducted both in a shared and in a distributed memory setting. All benchmarks were done on the "Glasgow grid", consisting of 16 machines with 2 Intel® Xeon® E5-2640 v2 and 64 GB of DDR3 RAM each. Each processor has 8 cores and 16 (hyperthreaded) threads

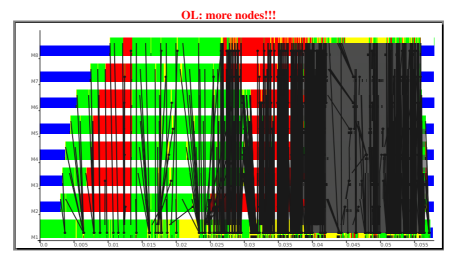


Figure 29: Matrix Multiplication with torus (PArrows).

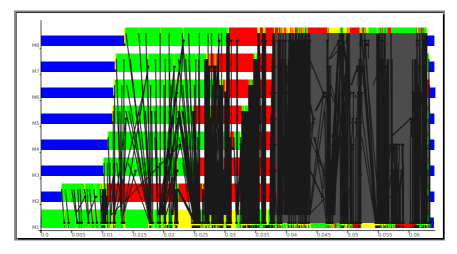


Figure 30: Matrix Multiplication with torus (Eden).

with a base frequency of 2 GHz and a turbo frequency of 2.50 GHz. This results in a total of 256 cores and 512 threads for the whole cluster. The operating system was Ubuntu 14.04 LTS with Kernel 3.19.0-33. Non-surprisingly, we found that hyperthreaded 32 cores do not behave in the same manner as real 16 cores (numbers here for a single machine). We disregarded the hyperthreading ability in most of the cases.

We used a single node with 16 real cores as a shared memory testbed and the whole grid with 256 real cores as a device to test our distributed memory software.

8.2 Test programs

We used multiple tests that originated from different sources. Most of them are parallel mathematical computations, initially implemented in Eden. Table 1 summarizes.

Arrows for Parallel Computations

Table 1: The benchmarks we use in this paper.

Name	Area	Туре	Origin	Citation
Rabin-Miller test	Mathematics	parMap + reduce	Eden	Lobachev (2012)
Jacobi sum test	Mathematics	workpool + reduce	Eden	Lobachev (2012)
Gentleman	Mathematics	torus	Eden	Loogen et al. (2003)
Sudoku	Puzzle	parMap	Par Monad	Marlow et al. (2011) 10

Rabin–Miller test is a probabilistic primality test that iterates multiple (32–256 here) "subtests". Should a subtest fail, the input is definitely not a prime. If all n subtest pass, the input is composite with the probability of $1/4^n$.

Jacobi sum test or APRCL is also a primality test, that however, guarantees the correctness of the result. It is probabilistic in the sense that its run time is not certain. Unlike Rabin–Miller test, the subtests of Jacobi sum test have very different durations. Lobachev (2011) **OL: discuss some optimisations of parallel APRCL.** Generic parallel implementation of Rabin–Miller test and APRCL were presented in Lobachev (2012).

"Gentleman" is a standard Eden test program, developed for their *torus* skeleton. It implements a parallel matrix multiplication (Gentleman, 1978). We ported an Eden based version (Loogen *et al.*, 2003) to PArrows.

A parallel Sudoku solver was used by Marlow *et al.* (2011) to compare *Par* Monad to GpH Haskell. We ported it to PArrows.

8.3 What parallel Haskells run where

The *Par* monad and GpH Haskell - in its multicore version ((Marlow *et al.*, 2009)) - can be executed on a shared memory machines only. Although GpH is available on distributed memory clusters, and newer distributed memory Haskells such as HdpH exist, current support of distributed memory in PArrows is limited to Eden. We used the MPI backend of Eden in a distributed memory setting. However, for shared memory Eden features a "CP" backend that merely copies the memory blocks between distributed heaps. In this mode Eden still operates in the "nothing shared" setting, but is adapted better to multicore machines. We label this version of Eden in the plots as "Eden CP".

8.4 Effect of hyperthreading

The PArrows version of Rabin–Miller test on a single node of the Glasgow grid showed almost linear speedup (Fig. 31). The speedup of 64-task PArrows/Eden at 16 real cores version was 13.65, the efficiency was 85.3%. However, if we increase the number of requested cores to be 32—i.e. if we use hyperthreading on 16 real cores—the speedup does not increase that well. It is merely 15.99 for 32 tasks with PArrows/Eden. It is worse for other backends. As for 64 tasks, we obtain the speedup of 16.12 with PArrows/Eden at 32

¹⁰ actual code from: http://community.haskell.org/ simonmar/par-tutorial.pdf and https://github.com/simonmar/parconc-examples

hyperthreaded cores and only 13.55 with PArrows/GpH. Efficiency is 50.4% and 42.3%, respectively. The Eden version used here was Eden CP, the "share nothing" SMP build.

In the distributed memory setting the same effect ensues. We obtain plummeting speedup of 124.31 at 512 hyperthreaded cores, whereas it was 213.172 for 256 real cores. Apparently, hyperthreading in the Glasgow grid fails to execute two parallel Haskell processes with full-fledged parallelism. For this reason, we did not regard hyperthreaded cores in our speedup plots in Figs. 31-35.

8.5 Benchmark results

The difference between, say, PArrows with *Par* Monad backend and a genuine *Par* Monad benchmark is very small. To give an example, it is 0.4s in favour of PArrows for 16 cores (10.8s vs. 11.2s) and -0.8s in favour of the *Par* monad for 8 cores (16.1s vs. 16.9s) for the Sudoku benchmark in the shared memory setting. It is almost invisible in speedup and (non shown) run time plots. We thus show only the results for the PArrows-enabled versions.

To show that PArrows induce very small overhead in a distributed context as well, we compare the original Eden versions of the benchmark to its PArrows-enabled counterpart in the Rabin–Miller test, Gentleman and Jacobi sum test benchmarks. We plot execution time differences between measurements for PArrows and the corresponding backend in a separate plot (Figs. 32, 34, 33). As an example, the differences range in about 0.5 seconds for the execution time of 46 seconds on 256 cores for distributed Rabin–Miller test with PArrows and Eden. For these comparisons, the plots show absolute time differences that are not relative w.r.t. the total execution time. Furthermore, the error bars ends were computed from pointwise maximum of both standard deviations from both measurements for PArrows and non-PArrows versions. These are the values provided by the *bench* package that we used for benchmarking. We call a difference between two versions significant when the border of the error bar of absolute time difference is above or below zero. In other words: the time difference is significant if it is above measurement error.

8.5.1 Rabin–Miller test

OL: THE ACTUAL TEXT IS MISSING. What do we see in the plots? Why is it good?

The multicore version of our parallel Rabin–Miller test benchmark is depicted in Figure 31. We executed the test with 32 and 64 tasks. The plot shows the PArrows-enabled versions with corresponding backends. The performance of PArrows/Eden CP in shared memory is slightly better than for SMP variants such as PArrows/GpH and PArrows/Par Monad but most of the time the performance is still comparable with the GpH backend performing slightly worse than the other two in terms of speedup. In particular, the speedups for 16 cores fall behind quite noticeably with 11.69 for 32 tasks as Eden and the Par Monad both score 13.05 for 32 tasks. While the GpH backend scores a bit better in the 64 tasks benchmark, its speedup of 12.70 is still behind the 13.65 and 13.33 of Eden and Par Monad, respectively.

Comparing the PArrows version of the Rabin–Miller test with the original from Eden with the MPI backend in a distributed memory setting, we see an almost linear speedup of Rabin–Miller test with 256 tasks and input $2^{444497} - 1$ in both versions. The sequential

Arrows for Parallel Computations

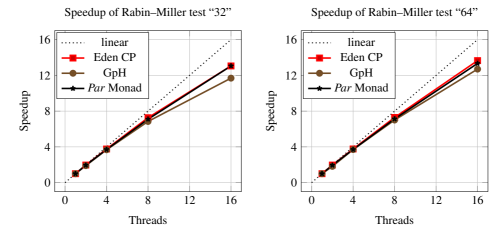


Figure 31: Relative speedup of Rabin–Miller test on a multicore machine. We used the same PArrows-based implementation with different backends on the same hardware. Measurements were performed on a single node of the Glasgow grid; it has 16 real cores. Input was $2^{11213} - 1$, we used 32 (left) or 64 (right) tasks. The closer to linear speedup the better.

run time was computed as the mean of three consecutive executions on a single core—the single run took two hours 43 minutes. The zero difference between PArrows/Eden and Eden almost always lies on the error bar of the measurement. The only exception, where the PArrows and Eden versions differ and it was it was significant, was for 64 cores, where it performed 0.49s better. This corresponds to 0.30% relative time difference. Otherwise, PArrows was 0.23s (or 0.0007%) slower for 32 cores, 0.26s (or 0.0030%) slower for 128 cores, and 0.09s faster (or 0.0019%) faster for 256 cores.

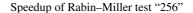
8.5.2 Jacobi sum test

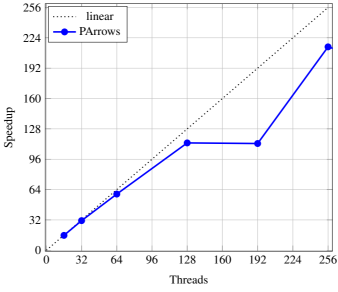
Continuing, the results of the Jacobi sum test in 33 are as follows:

MB: FIXME: redo this test with 2⁴⁴⁴⁹⁷

8.5.3 Gentleman

Next is the Gentleman benchmark. The results of the comparison of vanilla Eden to our PArrows-based version can be found in 34. While we do not see too much speedup for more than 16 cores, we can prove that the difference between the Eden and PArrows version are again marginal. The difference between PArrows and Eden is only significant for 16 and 64 cores where it ran 1.7% and 2.7% slower which corresponds to a real-time difference of 0.12s and 0.13s. For 256 cores PArrows performed 0.2% slower which corresponds to 0.01s overhead.





Run time differences for Rabin–Miller test "256"

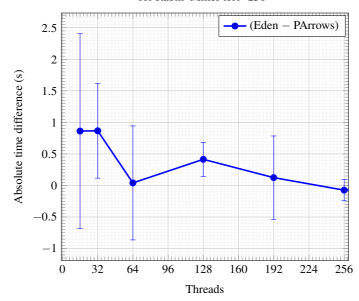
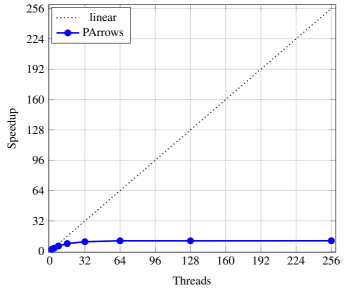


Figure 32: Parallel performance of Rabin–Miller test on the Glasgow grid consisting of 256 cores. Input was $2^{44497}-1$, we used 256 tasks. The top plot shows absolute speedup in a distributed memory setting. The closer to linear speedup the better. Time (and hence speedup) measurements for PArrows with Eden backend and Eden almost coincide. Hence, bottom plot shows absolute time differences for this benchmark. The higher the value, the better for PArrowsOL: CHECKME.

Arrows for Parallel Computations

Speedup of Jacobi sum test "3217"



Run time differences for Jacobi sum test "3217"

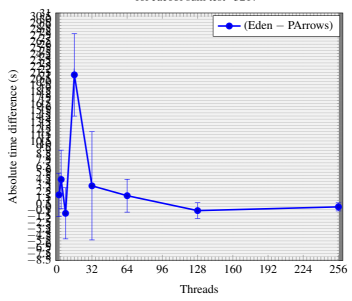


Figure 33: Parallel performance of Rabin–Miller test on the Glasgow grid consisting of 256 cores. Input was $2^{1279}-1$, we used 256 tasks. The top plot shows absolute speedup in a distributed memory setting. The closer to linear speedup the better. Time (and hence speedup) measurements for PArrows with Eden backend and Eden almost coincide. Hence, bottom plot shows absolute time differences for this benchmark. The higher the value, the better for PArrowsOL: CHECKME.

benchmark missing, in the works benchmark missing, in the works

Figure 34: Parallel performance of Gentleman on the Glasgow grid consisting of 256 cores. Input was a matrix size of 1024. The top plot shows absolute speedup in a distributed memory setting. The closer to linear speedup the better. Time (and hence speedup) measurements for PArrows with Eden backend and Eden almost coincide. Hence, bottom plot shows absolute time differences for this benchmark. The higher the value, the better for PArrowsOL: CHECKME.

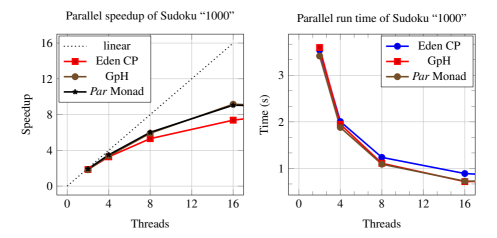


Figure 35: Relative speedup of Sudoku on a multicore machine. We used the same PArrows-based implementation with different backends on the same hardware and the *parMap* version from the *Par* Monad examples. Measurements were performed on a single node of the Glasgow grid; it has 16 real cores and 32 threads. Input was a file of 1000 Sudokus. The closer to linear speedup the better.

8.5.4 Sudoku

As the last Benchmark in this paper we present the Sudoku in 35 running in a shared memory setting. Here we see all three SM backends performing similarly again like in the Rabin–Miller test SM benchmarks in Figs. 35-36. However, we notice that the GpH backend seems to choke on a bigger input 36. This is due to the benchmark only using *parMap* instead of a chunking variant (however we did not change that for simplicity's sake) and is reflected by debug output which shows, that of 16000 sparks being created (one for each sudoku) only 8365 were converted (executed) with the rest (7635) overflowing the runtime spark pool. Another remarkable finding is that the Eden backend seems to lack behind for \leq 16 threads, but manages to pull ahead noticeably with all 32 threads of the system in use.

9 Conclusion

Arrows are a generic concept that allows for powerful composition combinators. To our knowledge we are first to represent *parallel* computation with Arrows. Arrows turn out to be a useful tool for composing in parallel programs. We do not have to introduce new

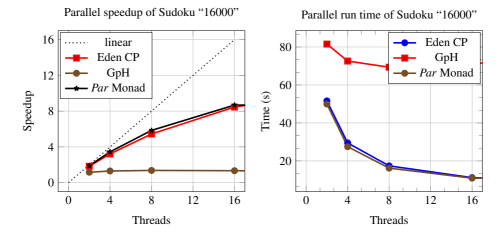


Figure 36: Relative speedup of Sudoku on a multicore machine. We used the same PArrows-based implementation with different backends on the same hardware and the *parMap* version from the *Par* Monad examples. Measurements were performed on a single node of the Glasgow grid; it has 16 real cores and 32 threads. Input was a file of 16000 Sudokus. The closer to linear speedup the better. The GpH version shows signs of choking with too many sparks being created.

monadic types that wrap the computation. Instead, we use Arrows in the same manner one uses sequential pure functions. This work features multiple parallel backends: the already available parallel Haskell flavours. Parallel Arrows, as presented here, feature an implementation of the *ArrowParallel* typeclass for GpH Haskell, *Par* Monad, and Eden. With our approach parallel programs can be ported across these flavours with little to no effort. It is quite straightforward to add further backends. Performance-wise, Parallel Arrows are on par with existing parallel Haskells, as they do not introduce any notable overhead. The benefit is, however, the greatly increased portability of parallel programs.

9.1 Future Work

Our PArrows DSL can be expanded to further parallel Haskells. More specifically we target HdpH (Maier *et al.*, 2014) for this future extension. HdpH is a modern distributed Haskell that would benefit from our Arrows notation. Further Future-aware versions of Arrow combinators can be defined. Existing combinators could also be improved. We would look into more transparency of our DSL, basically it should infuse as little overhead as possible.

More experiences with seamless porting of parallel PArrows-based programs across the backends are welcome. Of course, we are working ourselves on expanding both our skeleton library and the number of skeleton-based parallel programs that use our DSL to be portable across flavours of parallel Haskells. It would also be interesting to see a hybrid of PArrows and Accelerate (McDonell *et al.*, 2015). Ports of our approach to other languages like Frege or Java directly are in an early development stage.

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A Utility Arrows

Following are definitions of some utility Arrows used in this paper that have been lef out for brevity. We start with the *second* combinator from Hughes (2000), which is a mirrored version of *first*, which is for example used in the definition of ***:

```
second :: Arrow arr \Rightarrow arr ab \rightarrow arr(c,a)(c,b)
second f = arr swap >>> first <math>f >>> arr swap
where swap(x,y) = (y,x)
```

Next, we also define map, foldl and zipWith on Arrows. The mapArr combinator (Fig. A 1) lifts any arrow $arr\ a\ b$ to an arrow $arr\ [a]\ [b]$ (Hughes, 2005b). Similarly, we can also define foldlArr (Fig. A 2) that lifts any arrow $arr\ (b,a)\ b$ with a neutral element b to $arr\ [a]\ b$.

```
\begin{split} \mathit{mapArr} &:: Arrow Choice \ \mathit{arr} \Rightarrow \mathit{arr} \ \mathit{a} \ \mathit{b} \rightarrow \mathit{arr} \ [\mathit{a}] \ [\mathit{b}] \\ \mathit{mapArr} \ \mathit{f} &= \\ \mathit{arr} \ \mathit{list} \mathit{case} >>> \\ \mathit{arr} \ (\mathit{const} \ []) \parallel (\mathit{f} *** \mathit{mapArr} \ \mathit{f} >>> \mathit{arr} \ (\mathit{uncurry} \ (:))) \\ \mathit{list} \mathit{case} \ [] &= \mathit{Left} \ () \\ \mathit{list} \mathit{case} \ (x : xs) &= \mathit{Right} \ (x , xs) \end{split}
```

Figure A 1: The definition of map over Arrows and the listcase helper function.

Finally, with the help of mapArr (Fig. A 1), we can define zipWithArr (Fig. A 3) that lifts any arrow arr(a,b)c to an arrow arr([a],[b])[c].

These combinators make use of the ArrowChoice type class which provides the \parallel combinator. It takes two arrows $arr\ a\ c$ and $arr\ b\ c$ and combines them into a new arrow $arr\ (Either\ a\ b)\ c$ which pipes all $Left\ a$'s to the first arrow and all $Right\ b$'s to the second arrow:

```
 \begin{split} & \textit{foldlArr} :: (\textit{ArrowChoice arr}, \textit{ArrowApply arr}) \Rightarrow \textit{arr } (b, a) \ b \rightarrow b \rightarrow \textit{arr } [a] \ b \\ & \textit{foldlArr} \ f \ b = \\ & \textit{arr listcase} >>> \\ & \textit{arr } (\textit{const } b) \parallel \\ & \textit{(first } (\textit{arr } (\lambda a \rightarrow (b, a)) >>>> f >>> \textit{arr } (\textit{foldlArr} \ f)) >>>> \textit{app}) \end{split}
```

Figure A 2: The definition of *foldl* over Arrows.

```
zipWithArr::ArrowChoice\ arr \Rightarrow arr\ (a,b)\ c \rightarrow arr\ ([a],[b])\ [c]
zipWithArr\ f = (arr\ \lambda(as,bs) \rightarrow zipWith\ (,)\ as\ bs) >>> mapArr\ f
```

Figure A 3: zipWith over arrows.

```
(\parallel \parallel)::ArrowChoice arr a c \rightarrow arr b c \rightarrow arr (Either <math>a b) c
```

With the zipWithArr combinator we can also write a combinator listApp, that lifts a list of arrows $[arr\ a\ b]$ to an arrow $arr\ [a]\ [b]$.

```
listApp :: (ArrowChoice\ arr, ArrowApply\ arr) \Rightarrow [arr\ a\ b] \rightarrow arr\ [a]\ [b]
listApp\ fs = (arr\ \lambda as \rightarrow (fs, as)) >>> zipWithArr\ app
```

Note that this additionally makes use of the ArrowApply typeclass that allows us to evaluate arrows with $app::arr(arr\ a\ b,a)\ c$.

B Omitted Function Definitions

We have omitted some function definitions in the main text for brevity, and redeem this here

To begin with, we give the actual implementation of the non-configurable instance of the *ArrowParallel* instance:

Next, we warp Eden's build-in Futures in PArrows as in Figure B 6, where *rd* is the accessor function for the *RD* wrapped inside *RemoteData*. Furthermore, in order for these *Future* types to fit with the *ArrowParallel* instances we gave earlier, we have to give the necessary *NFData* and *Trans* instances, the latter are only needed in Eden. The *Trans* instance does not have any functions declared as the default implementation suffices here. Furthermore, because *MVar* already ships with a *NFData* instance, we only have to supply a simple delegating *NFData* instance for our *RemoteData* type, where *rd* simply unwraps

```
 \begin{array}{l} \textbf{instance} \; (NFData \; b, ArrowApply \; arr, ArrowChoice \; arr) \Rightarrow ArrowParallel \; arr \; a \; b \; () \; \textbf{where} \\ parEvalN \; \_fs = parEvalN \; (hack \; fs) \; fs \\ \textbf{where} \\ hack :: (NFData \; b) \Rightarrow [arr \; a \; b] \rightarrow Conf \; b \\ hack \; \_ = Conf \; rdeepseq \\ \end{array}
```

Figure B 1: The actual default implementation of GpH's ArrowParallel.

RD. The *Trans* instance does not have any functions declared as the default implementation suffices:

```
instance NFData (RemoteData a) where rnf = rnf \circ rd instance Trans (RemoteData a)
```

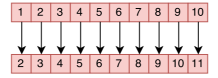


Figure B 2: Schematic depiction of parMap.

 $parMap :: (ArrowParallel \ arr \ a \ b \ conf) \Rightarrow conf \rightarrow (arr \ a \ b) \rightarrow (arr \ [a] \ [b])$ $parMap \ conf \ f = parEvalN \ conf \ (repeat \ f)$



Figure B 3: Definition of parMap.

Figure B 4: Schematic depiction of parMapStream.

7 8

9

10 11

5 6

```
parMapStream :: (ArrowParallel\ arr\ a\ b\ conf\ , ArrowChoice\ arr\ , ArrowApply\ arr) \Rightarrow conf\ \rightarrow ChunkSize\ \rightarrow arr\ a\ b\ \rightarrow arr\ [a]\ [b] parMapStream\ conf\ chunkSize\ f\ =\ parEvalNLazy\ conf\ chunkSize\ (repeat\ f)
```

Figure B 5: Definition of parMapStream.

Figures B 2–B 5 show the definitions and a visualizations of two parallel *map* variants, defined using *parEvalN* and its lazy counterpart.

Arrow versions of Eden's *shuffle*, *unshuffle* and the definition of *takeEach* are in Figure B 7. Similarly, Figure B 8 contains the definition of arrow versions of Eden's *lazy* and *rightRotate* utility functions. Fig. B 9 contains Eden's definition of *lazyzip3* together with the utility functions *uncurry3* and *threetotwo*. The full definition of *farmChunk* is in Figure B 10. Eden definition of *ring* skeleton is in Figure B 11. It follows Loogen (2012).

```
data RemoteData a = RD \{ rd :: RD a \}

instance (Trans \ a) \Rightarrow Future \ RemoteData \ a \ where

put = arr \ (\lambda a \rightarrow RD \{ rd = release \ a \})

get = arr \ rd >>> arr \ fetch
```

Figure B 6: RD-based RemoteData version of Future for the Eden backend.

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```
shuffle :: (Arrow arr) \Rightarrow arr [[a]] [a]
shuffle = arr (concat \circ transpose)
unshuffle :: (Arrow \ arr) \Rightarrow Int \rightarrow arr \ [a] \ [[a]]
unshuffle n = arr(\lambda xs \rightarrow [takeEach \ n \ (drop \ i \ xs) \mid i \leftarrow [0..n-1]])
takeEach :: Int \rightarrow [a] \rightarrow [a]
takeEach\ n\ [\ ]=[\ ]
takeEach\ n\ (x:xs) = x:takeEach\ n\ (drop\ (n-1)\ xs)
```

Figure B 7: Definitions of shuffle, unshuffle, takeEach.

```
lazy :: (Arrow arr) \Rightarrow arr [a] [a]
lazy = arr (\lambda \sim (x:xs) \rightarrow x: lazy xs)
rightRotate :: (Arrow arr) \Rightarrow arr [a] [a]
rightRotate = arr \$ \lambda list \rightarrow \mathbf{case} \ list \ \mathbf{of}
   [] \rightarrow []
   xs \rightarrow last \ xs : init \ xs
```

Figure B 8: Definitions of lazy and rightRotate.

The parEval2 skeleton is defined in Figure B 12. We start by transforming the (a,c) input into a two-element list [Either a c] by first tagging the two inputs with Left and Right and wrapping the right element in a singleton list with return so that we can combine them with arr (uncurry (:)). Next, we feed this list into a parallel arrow running on two instances of f + + + g as described above. After the calculation is finished, we convert the resulting [Either b d] into ([b], [d]) with arr partition Eithers. The two lists in this tuple contain only one element each by construction, so we can finally just convert the tuple to (b,d) in the last step. Furthermore, Fig. B 13 contains the ommitted definitions of prMMTr, - which calculates AB^T for two matrices A and B, splitMatrix - which splits the a matrix into chunks, and lastly matAdd, that calculates A + B for two matrices A and B.

C Syntactic Sugar

Finally, we also give the definitions for some syntactic sugar for PArrows, namely *** and &&&. For basic arrows, we have the *** combinator (Fig. 3) which allows us to combine two arrows $arr \ a \ b$ and $arr \ c \ d$ into an arrow $arr \ (a,c) \ (b,d)$ which does both computations at once. This can easily be translated into a parallel version *** with the use of parEval2, but for this we require a backend which has an implementation that does not require any configuration (hence the () as the *conf* parameter):

```
(|***|) :: (ArrowChoice\ arr, ArrowParallel\ arr\ (Either\ a\ c)\ (Either\ b\ d)\ ())) \Rightarrow
  arr\ a\ b \rightarrow arr\ c\ d \rightarrow arr\ (a,c)\ (b,d)
(|***|) = parEval2()
```

We define the parallel &&& in a similar manner to its sequential pendant &&& (Fig. 3):

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```
\begin{aligned} &lazyzip3::[a] \rightarrow [b] \rightarrow [c] \rightarrow [(a,b,c)] \\ &lazyzip3 \ as \ bs \ cs = zip3 \ as \ (lazy \ bs) \ (lazy \ cs) \\ &uncurry3::(a \rightarrow b \rightarrow c \rightarrow d) \rightarrow (a,(b,c)) \rightarrow d \\ &uncurry3f \ (a,(b,c)) = f \ a \ b \ c \\ &threetotwo::(Arrow \ arr) \Rightarrow arr \ (a,b,c) \ (a,(b,c)) \\ &threetotwo = arr \$ \lambda \sim (a,b,c) \rightarrow (a,(b,c)) \end{aligned}
```

Figure B 9: Definitions of *lazyzip3*, *uncurry3* and *threetotwo*.

```
\begin{split} & \textit{farmChunk} :: (\textit{ArrowParallel arr } a \ b \ \textit{conf} \ , \textit{ArrowParallel arr } [a] \ [b] \ \textit{conf}, \\ & \textit{ArrowChoice } \textit{arr}, \textit{ArrowApply } \textit{arr}) \Rightarrow \\ & \textit{conf} \rightarrow \textit{ChunkSize} \rightarrow \textit{NumCores} \rightarrow \textit{arr } a \ b \rightarrow \textit{arr } [a] \ [b] \\ & \textit{farmChunk conf } \textit{chunkSize } \textit{numCores} \ f = \\ & \textit{unshuffle } \textit{numCores} \ >>> \\ & \textit{parEvalNLazy } \textit{conf } \textit{chunkSize } (\textit{repeat } (\textit{mapArr} \ f)) \ >>> \\ & \textit{shuffle} \end{split}
```

Figure B 10: Definition of farmChunk.

```
(|\&\&\&|) :: (ArrowChoice\ arr, ArrowParallel\ arr\ (Either\ a\ a)\ (Either\ b\ c)\ ()) \Rightarrow arr\ a\ b \to arr\ a\ c \to arr\ a\ (b,c) 
 (|\&\&\&|)\ f\ g = (arr\ \lambda a \to (a,a)) >>> f\ |***|\ g
```

Arrows for Parallel Computations

```
\begin{array}{l} ringSimple :: (Trans\ i, Trans\ o, Trans\ r) \Rightarrow (i \rightarrow r \rightarrow (o,r)) \rightarrow [i] \rightarrow [o] \\ ringSimple\ f\ is = os \\ \textbf{where}\ (os, ringOuts) = unzip\ (parMap\ (toRD\ uncurry\ f)\ (zip\ is\ lazy\ ringIns)) \\ ringIns = rightRotate\ ringOuts \\ toRD :: (Trans\ i, Trans\ o, Trans\ r) \Rightarrow ((i,r) \rightarrow (o,r)) \rightarrow ((i,RD\ r) \rightarrow (o,RD\ r)) \\ toRDf\ (i,ringIn) = (o,release\ ringOut) \\ \textbf{where}\ (o,ringOut) = f\ (i,fetch\ ringIn) \\ rightRotate :: [a] \rightarrow [a] \\ rightRotate\ xs = last\ xs : init\ xs \\ lazy :: [a] \rightarrow [a] \\ lazy \sim (x:xs) = x: lazy\ xs \end{array}
```

Figure B 11: Eden's definition of the *ring* skeleton.

```
parEval2 :: (ArrowChoice\ arr, \\ ArrowParallel\ arr\ (Either\ a\ c)\ (Either\ b\ d)\ conf) \Rightarrow \\ conf \to arr\ a\ b \to arr\ c\ d \to arr\ (a,c)\ (b,d) \\ parEval2\ conf\ f\ g = \\ arr\ Left *** (arr\ Right >>> arr\ return) >>> \\ arr\ (uncurry\ (:)) >>> \\ parEvalN\ conf\ (replicate\ 2\ (f+++g)) >>> \\ arr\ partitionEithers >>> \\ arr\ head *** arr\ head
```

Figure B 12: Definition of *parEval2*.

```
\begin{split} prMMTr \ ml \ m2 &= [[sum \ (zipWith \ (*) \ row \ col) \ | \ col \leftarrow m2] \ | \ row \leftarrow m1] \\ splitMatrix :: Int \rightarrow Matrix \rightarrow [[Matrix]] \\ splitMatrix \ size \ matrix &= map \ (transpose \circ map \ (chunksOf \ size)) \ \ chunksOf \ size \ \ matrix \\ matAdd &= chunksOf \ (dimX \ x) \ \ sipWith \ (+) \ (concat \ x) \ (concat \ y) \end{split}
```

Figure B 13: Definition of prMMTr, splitMatrix and matAdd.

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