

Regular, Shape-polymorphic, Parallel Arrays in Haskell

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Abstract

We present a novel approach to regular, multi-dimensional arrays in Haskell. The main highlights of our approach are that it (1) is purely functional, (2) supports reuse through shape polymorphism, (3) avoids unnecessary intermediate structures rather than relying on subsequent loop fusion, and (4) supports transparent parallelisation.

We show how to embed two forms of shape polymorphism into Haskell’s type system using type classes and type families. In particular, we discuss the generalisation of regular array transformations to arrays of higher rank, and introduce a type-safe specification of array slices.

We discuss the runtime performance of our approach for three standard array algorithms. We achieve absolute performance comparable to handwritten C code. At the same time, our implementation scales well up to 8 processor cores.

Languages, Performance

1. Introduction

In purely functional form, array algorithms are often more elegant and easier to comprehend than their imperative, explicitly loop-based counterparts. The question is, can they also be efficient?

Experience with Clean, OCaml, and Haskell has shown that we can write efficient code if we sacrifice purity and use an *imperative array interface* based on reading and writing *individual array elements*, possibly wrapped in uniqueness types or monads Groningen [1997], Leroy et al. [2008], Launchbury and Peyton Jones [1994]. However, using impure features not only obscures clarity, but also forfeits the transparent exploitation of the data parallelism that is abundant in array algorithms.

In contrast, using a *purely-functional array interface* based on *collective operations*—such as maps, folds, and permutations—emphasises an algorithm’s high-level structure and often has an obvious parallel implementation. This observation was the basis for previous work on algorithmic skeletons and the use of the *Bird-Meertens Formalism (BMF)* for parallel algorithm design Rabhi and Gorlatch [2003]. Our own work on *Data Parallel Haskell (DPH)* is based on the same premise, but aims at irregular data parallelism which comes with its own set of challenges Peyton

Jones et al. [2008]. Other work on *byte arrays* Coutts et al. [2007b] also aims at high-performance, while abstracting over loop-based low-level code using a purely-functional combinator library.

We aim higher by supporting multi-dimensional arrays, more functionality, and transparent parallelism. We present a Haskell library of regular parallel arrays, which we call *Repa*¹ (Regular Parallel Arrays). While Repa makes use of the Glasgow Haskell Compiler’s many existing extensions, it is a pure library: it does not require any language extensions that are specific to its implementation. The resulting code is not only as fast as when using an imperative array interface, it approaches the performance of handwritten C code, and exhibits good parallel scalability on the configurations that we benchmarked.

In addition to good performance, we achieve a high degree of reuse by supporting *shape polymorphism*. For example, `map` works over arrays of arbitrary rank, while `sum` decreases the rank of an arbitrary array by one – we give more details in Section 4. The value of shape polymorphism has been demonstrated by the language Single Assignment C, or SAC Scholz [2003]. Like us, SAC aims at purely functional high-performance arrays, but SAC is a specialised array language based on a purpose-built compiler. We show how to embed shape polymorphism into Haskell’s type system.

The main contributions of the paper are the following:

- An API for purely-functional, collective operations over dense, rectangular, multi-dimensional arrays supporting shape polymorphism (Section 5).
- Support for various forms of constrained shape polymorphism in a Hindley-Milner type discipline with type classes and type families (Section 4).
- An aggressive loop fusion scheme based on a functional representation of delayed arrays (Section 6).
- A scheme to transparently parallelise array algorithms based on our API (Section 7)
- An evaluation of the sequential and parallel performance of our approach on the basis of widely used array algorithms (Section 8).

Before diving into the technical details of our contributions, the next section illustrates our approach to array programming by way of an example.

2. Our approach to array programming

A simple operation on two-dimensional matrices is transposition. With our library we express transposition in terms of a permutation operation that swaps the row and column indices of a matrix:

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¹Repa means “turnip” in Russian.

```
transpose2D :: Elt e => Array DIM2 e -> Array DIM2 e
transpose2D arr
  = backpermute new_extent swap arr
  where swap (Z :.i :.j) = Z :.j :.i
        new_extent      = swap (extent arr)
```

Like Haskell 98 arrays, our array type is parameterised by the array’s *index type*, here DIM2, and by its element type *e*. The index type gives the *rank* of the array, which we also call the array’s *dimensionality*, or *shape*.

Consider the type of `backpermute`, given in Figure 1. The first argument is the bounds (or *extent*) of the result array, which we obtain by swapping the row and column extents of the input array. For example transposing a 3×12 matrix gives a 12×3 matrix.² The `backpermute` function constructs a new array in terms of an existing array solely through an *index transformation*, supplied as its second argument, `swap`: given an index into the result matrix, `swap` produces the corresponding index into the argument matrix.

A more interesting example is matrix-matrix multiplication:

```
mmMult :: (Num e, Elt e)
       => Array DIM2 e -> Array DIM2 e
       -> Array DIM2 e
mmMult arr brr
  = sum (zipWith (*) arrRepl brrRepl)
  where
    trr      = transpose2D brr
    arrRepl  = replicate (Z :.All :.colsB :.All) arr
    brrRepl  = replicate (Z :.rowsA :.All :.All) trr
    (Z :.colsA :.rowsA) = extent arr
    (Z :.colsB :.rowsB) = extent brr
```

The idea is to expand both rank-two argument arrays into rank-three arrays by replicating them across a new dimension, or axis, as illustrated in Figure 2. The front face of the cuboid represents the array `arr`, which we replicate as often as `brr` has columns (`colsB`), producing `arrRepl`. The top face represents `trr` (the transposed `brr`), which we replicate as often as `arr` has rows (`rowsA`), producing `brrRepl`. As indicated by the figure, the two replicated arrays have the same extent, which corresponds to the index space of matrix multiplication:

$$(AB)_{i,j} = \sum_{k=1}^n A_{i,k} B_{k,j}$$

where *i* and *j* correspond to `rowsA` and `colsB` in our code. The summation index *k* corresponds to the innermost axis of the replicated arrays and to the left-to-right axis in Figure 2. Along this axis we perform the summation after an elementwise multiplication of the replicated elements of `arr` and `brr` by `zipWith (*)`.

A naive implementation of the operations used in `mmMult` would result in very bad space and time performance. In particular, it would be very inefficient to compute explicit representations of the replicated matrices `arrRepl` and `brrRepl`. Indeed, a key principle of our design is to avoid generating explicit representations of the intermediate arrays that can be represented as the original arrays combined with a *transformation of the index space* (Section 3.2). A rigorous application of this principle results in aggressive loop fusion (Section 6) producing code that is similar to imperative code. As a consequence, this Haskell code has about the same performance as handwritten C code for the same computation. Even more importantly, we measured very good absolute speedup, $\times 7.2$ for 8 cores, on multicore hardware. For the C code, this can only be achieved through considerable additional effort or by employing special-purpose language extensions such as

²For now, just read the notation $(Z :.i :.j)$ as if it was the familiar pair (i, j) . The details are in Section 4 where we discuss shape polymorphism.

```
extent :: Array sh e -> sh
sum    :: (Shape sh, Elt e, Num e)
       => Array (sh :: Int) e -> Array sh e
zipWith :: (Shape sh, Elt e1, Elt e2, Elt e3)
        => (e1 -> e2 -> e3)
        -> Array sh e1 -> Array sh e2
        -> Array sh e3
backpermute :: (Shape sh, Shape sh', Elt e)
            => sh' -> (sh' -> sh)
            -> Array sh e -> Array sh' e
```

Figure 1. Types of library functions

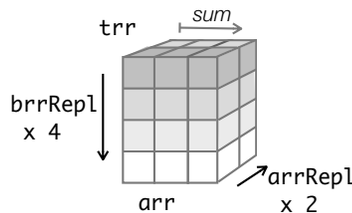


Figure 2. Matrix-matrix multiplication illustrated

OpenMP The OpenMP Architecture Review Board [2008] which often have difficulties with more complex programs.

3. Representing arrays

The representation of arrays is a central issue in any array library. Our library uses two devices to achieve good performance:

1. We represent array data as contiguously-allocated ranges of *unboxed* values.
2. We *delay* the construction of intermediate arrays to support constant-time index transformations and slices, and to combine these operations with traversals over successive arrays.

We describe these two techniques in the following sections.

3.1 Unboxed arrays

In Haskell 98 arrays are lazy: each element of an array is evaluated only when the array is indexed at that position. Although convenient, laziness is Very Bad Indeed for array-intensive programs:

- A lazy array of (say) `Float` is represented as an array of pointers to either heap-allocated thunks, or boxed `Float` objects, depending on whether they have been forced. This representation requires at least three times as much storage as a conventional, contiguous array of unboxed floats. Moreover, when iterating through the array, the lazy representation imposes higher memory traffic. This is due to the increased size of the individual elements, as well as their lower spacial locality.
- In a lazy array, evaluating one element does not mean that the other elements will be demanded. However, the overwhelmingly common case is that the programmer intends to demand the entire array, and wants it evaluated in parallel.

We can solve both of these problems simultaneously using a Haskell-folklore trick. We define a new data type of arrays, which we will call `UArr`, short for “unboxed array”. These arrays are

one-dimensional, indexed by `Int`, and are slightly stricter than Haskell 98 arrays: a `UArr` *as a whole* is evaluated lazily, but an attempt to evaluate any *element* of the array (e.g. by indexing) will cause evaluation of all the others, in parallel.

For the sake of definiteness we give a bare sketch of how `UArr` is implemented. However, this representation is not new; it is well established in the Haskell folklore, and we use it in Data Parallel Haskell (DPH) Peyton Jones et al. [2008], Chakravarty et al. [2007], so we do not elaborate the details.

```
class Elt e where
  data UArr e
  (!) :: Array e -> Int -> e
  ...more methods...

instance Elt Float where
  data UArr Float = UAF Int ByteArray#
  (UAF max ba) ! i
    | i < max    = F# (indexByteArray ba i)
    | otherwise  = error "Index error"
  ...more methods...

instance (Elt a, Elt b) => Elt (a :* b) where
  data UArr (a :* b) = UAP (UArr a) (UArr b)
  (UAP a b) ! i = (a!i :* b!i)
  ...more methods...
```

Here we make use of Haskell’s recently added *associated data types* Chakravarty et al. [2005b] to represent an array of `Float` as a contiguous array of unboxed floats (the `ByteArray#`), and an array of pairs as a pair of arrays. Because the representation of the array depends on the element type, indexing must vary with the element type too, which explains why the indexing operation (`!`) is in a type class `Elt`.

In addition to an efficient underlying array representation, we also need the infrastructure to operate on these arrays in parallel, using multiple processor cores. To that end we reuse part of our own parallel array library of Data Parallel Haskell. This provides us with an optimised implementation of `UArr` and the `Elt` class, and with parallel collective operations over `UArr`. It also requires us to represent pairs using the strict pair constructor (`*:`), instead of Haskell’s conventional `(,)`.

3.2 Delayed arrays

When using Repa, index transformations such as `transpose2D` (discussed in Section 2) are ubiquitous. As we expect index transformations to be cheap, it would be wrong to (say) copy a 100Mbyte array just to transpose it. It is much better to push the index transformation into the consumer, which can then consume the original, unmodified array.

We could do this transformation statically, at compile time, but doing so would rely on the consumer being able to “see” the index transformation. This could make it hard for the programmer to predict whether or not the optimisation would take place. In Repa we instead perform this optimisation dynamically, and offer a guarantee that index transformations perform no data movement. The idea is simple and well known: just represent an array by its indexing function, together with the array bounds (this is not our final array representation):

```
data DArray sh e = DArray sh (sh -> e)
```

With this representation, functions like `backpermute` (whose type signature appeared in Figure 1) are quite easy to implement:

```
backpermute sh' fn (Array sh ix1)
  = Array sh' (ix1 . fn)
```

We can also wrap a `UArr` as a `DArray`:

```
wrap :: (Shape sh, Elt e)
      => sh -> UArr e -> DArray sh e
wrap sh uarr = Array sh idx
  where idx i = uarr ! toIndex sh i
```

When wrapping a `DArray` over a `UArr`, we also take the opportunity to generalise from one-dimensional to multi-dimensional arrays. The index of these multi-dimensional arrays is of type `sh`, where the `Shape` class (to be described in Section 4) includes the method `toIndex :: Shape sh => sh -> sh -> Int`. This method maps the bounds and index of an `Array` to the corresponding linear `Int` index in the underlying `UArr`.

3.3 Combining the two

Unfortunately, there are at least two reasons why it is not always beneficial to delay an array operation. One is *sharing*, which we discuss later in Section 6. Another is *data layout*. In our `mmMult` example from Section 2, we want to delay the two applications of `replicate`, but not the application of `transpose2D`. Why? We store multi-dimensional arrays in row-major order (the same layout Haskell 98 uses for standard arrays). Hence, iterating over the second index of an array of rank 2 is more cache friendly than iterating over its first index.

It is well known that the order of the loop nest in an imperative implementation of matrix-matrix multiplication has a dramatic effect on performance due to these cache effects. By forcing `transpose2D` to produce its result as an unboxed array in memory—we call this a *manifest* array—instead of leaving it as a delayed array, the code will traverse both matrices by iterating over the second index in the inner loop. Overall, we have the following implementation:

```
mmMult arr brr
  = sum (zipWith (*) arrRepl brrRepl)
  where
    trr      = force (transpose2D brr) -- New! force!
    arrRepl  = replicate (Z ::All  ::colsB ::All) arr
    brrRepl  = replicate (Z ::rowsA ::All  ::All) trr

    (Z ::colsA ::rowsA) = extent arr
    (Z ::colsB ::rowsB) = extent brr
```

We could implement `force` by having it produce a value of type `UArr` and then apply `wrap` to turn it into a `DArray` again, providing the appropriate memory layout for a cache-friendly traversal. This would work, but we can do better. The function `wrap` uses array indexing to access the underlying `UArr`. In cases where this indexing is performed in a tight loop, GHC can optimise the code more thoroughly when it is able to inline the indexing operator, instead of calling an anonymous function encapsulated in the data type `DArray`. For recursive functions, this also relies on the *constructor specialisation* optimisation Peyton Jones [1997]. However, as explained in Coutts et al. [2007a, Section 7.2], to allow this we must make the special case of a wrapped `UArr` explicit in the datatype, so the optimiser can see whether or not it is dealing directly with a manifest array.

Hence, we define regular arrays as follows:

```
data Array sh e = Manifest sh (UArr e)
                | Delayed   sh (sh -> e)
```

We can unpack an arbitrary `Array` into delayed form thus:

```
delay :: (Shape sh, Elt e)
       => Array sh e -> (sh, sh -> e)
delay (Delayed sh f)   = (sh, f)
delay (Manifest sh uarr) = delay (UArr uarr)
```

```

infixl 3 ..
data Z      = Z
data tail :: head = tail :: head

type DIM0 = Z
type DIM1 = DIM0 .. Int
type DIM2 = DIM1 .. Int
type DIM3 = DIM2 .. Int

class Shape sh where
  rank    :: sh -> Int
  size    :: sh -> Int      -- Number of elements
  toIndex :: sh -> sh -> Int -- Index into row-major
  fromIndex      -- representation
  :: sh -> Int -> sh -- Inverse of 'toIndex'
  <..and so on..>

instance          Shape Z          where ...
instance Shape sh => Shape (sh..Int) where ...

```

Figure 3. Definition of shapes

```

= (sh, \i -> uarr ! toIndex sh i)

```

This is the basis for a general `traverse` function that produces a delayed array after applying a transformation.

The transformation produced with `traverse` may include index space transformations or other computations:

```

traverse :: (Shape sh, Shape sh', Elt e)
  => Array sh e
  -> (sh -> sh')
  -> ((sh -> e) -> sh' -> e')
  -> Array sh' e'

traverse arr sh_fn elem_fn
  = Delayed (sh_fn sh) (elem_fn f)
  where (sh, f) = delay arr

```

We use `traverse` to implement many of the other operations of our library — for example, `backpermute` is implemented as:

```

backpermute :: (Shape sh, Shape sh', Elt e)
  => sh' -> (sh' -> sh) -> Array sh e
  -> Array sh' e

backpermute sh perm arr
  = traverse arr (const sh) (. perm)

```

We discuss the use of `traverse` in more detail in Sections 5 & 7.

4. Shapes and shape polymorphism

In Figure 1 we gave this type for `sum`:

```

sum :: (Shape sh, Num e, Elt e)
  => Array (sh..Int) e -> Array sh e

```

As the type suggests, `sum` is a *shape-polymorphic* function: it can sum the rightmost axis of an array of arbitrary rank. In this section we describe how shape polymorphism works in Repa. We will see that combination of parametric polymorphism, type classes, and type families enables us to track the rank of each array in its type, guaranteeing the absence of rank-related runtime errors. We can do this even in the presence of operations such as slicing and replication which change the rank of an array. However, bounds checks on indices are still performed at runtime — tracking them requires more sophisticated type system support Xi [2007], Swierstra and Altenkirch [2008].

4.1 Shapes and indices

Haskell’s tuple notation does not allow us the flexibility we need, so we introduce our own notation for indices and shapes. As defined in Figure 3, we use an inductive notation of tuples as heterogeneous *snoc* lists. On both the type-level and the value-level, we use the infix operator `(:.)` to represent *snoc*. The constructor `Z` corresponds to a rank zero shape, and we use it to mark the end of the list. Thus, a three-dimensional index with components `x`, `y` and `z` is written `(Z:.x:.y:.z)` and has type `(Z..Int..Int..Int)`. This type is the *shape* of the array. Figure 3 gives type synonyms for common shapes: a singleton array of shape `DIM0` represents a scalar value; an array of shape `DIM1` is a vector, and so on.

The motivation for using *snoc* lists, rather than the more conventional *cons* lists, is this: we store manifest arrays in row-major order, where the *rightmost* index is the most *rapidly-varying* when traversing linearly over the array in memory. For example, the value at index `(Z:.3:.8)` is stored adjacent to that at `(Z:.3:.9)`. This is the same convention adopted by Haskell 98 standard arrays.

We draw array indices from `Int` values only, so the shape of a rank-*n* array is:

$$Z \underbrace{:. \text{Int} :. \dots :. \text{Int}}_{n \text{ times}}$$

In principle, we could be more general and allow non-`Int` indices, like Haskell’s index type class `Ix`. However, this would complicate the library and the presentation, and is orthogonal to the contributions of this paper; so we do not consider it here. Nevertheless, shape types, such as `DIM2` etc, explicitly mention the `Int` type. This is for two reasons: firstly, it simplifies the transition to using the `Ix` class if that is desired; and secondly, in Section 4.4 we discuss more elaborate shape constraints that require an explicit index type.

The *extent* of an array is a value of the shape type:

```

extent :: Array sh e -> sh

```

The corresponding Haskell 98 function, `bounds`, returns an upper and lower bound, whereas `extent` returns only the upper bound. Repa uses zero-indexed arrays only, so the lower bound is always zero. For example, the extent `(Z:.4:.5)` characterises a 4×5 array of rank two containing 20 elements. The extent along each axis must be at least one.

The shape type of an array also types its indices, which range between zero and one less than the extent along the same axis. In other words, given an array with shape `(Z:.n1:...:.nm)`, its index range is from `(Z:.0:...:.0)` to `(Z:.n1-1:...:.nm-1)`. As indicated in Figure 3, the methods of the `Shape` type class determine properties of shapes and indices, very like Haskell’s `Ix` class. These methods are used to allocate arrays, index into their row-major in-memory representations, to traverse index spaces, and are entirely as expected, so we omit the details.

4.2 Shape polymorphism

We call functions that operate on a variety of shapes *shape polymorphic*. Some such functions work on arrays of any shape at all. For example, here is the type of `map`:

```

map :: (Shape sh, Elt a, Elt b)
  => (a -> b)
  -> Array sh a -> Array sh b

```

The function `map` applies its functional argument to all elements of an array without any concern for the shape of the array. The type class constraint `Shape sh` merely asserts that the type variable `sh` ought to be a shape. It does not constrain the shape of that shape in any way.

4.3 At-least constraints and rank generalisation

With indices as *snoc* lists, we can impose a lower bound on the rank of an array by fixing a specific number of lower dimensions, but keeping the tail of the resulting *snoc* list variable. For example, here is the type of `sum`:

```
sum :: (Shape sh, Num e, Elt e)
    => Array (sh:.Int) e -> Array sh e
```

This says that `sum` takes an array of any rank $n \geq 1$ and returns an array of rank $n - 1$. For a rank-1 array (a vector), `sum` adds up the vector to return a scalar. But what about a rank-2 array? In this case, `sum` adds up all the rows of the matrix *in parallel*, returning a vector of the sums. Similarly, given a three-dimensional array `sum` adds up each row of the array in parallel, returning a two-dimensional array of sums.

Functions like `sum` impose a lower bound on the rank of an array. We call such constraints shape polymorphic *at-least constraints*. Every shape-polymorphic function with an at-least constraint is implicitly also a data-parallel map over the unspecified dimensions. This is a major source of parallelism in Repa. We call the process of generalising the code defined for the minimum rank to higher ranks *rank generalisation*.

The function `sum` only applies to the rightmost index of an array. What if we want to reduce the array across a different dimension? In that case we simply perform an index permutation, which is guaranteed cheap, to bring the desired dimension to the rightmost position:

```
sum2 :: (Shape sh, Elt e, Num e)
    => Array (sh:.Int:.Int) e -> Array (sh:.Int) e
sum2 a = sum (backpermute new_extent swap2 a)
  where
    new_extent      = swap2 (extent a)
    swap2 (is :.i2 :.i1) = is :.i1 :.i2
```

In our examples so far, we have sometimes returned arrays of a different rank than the input, but their extent in any one dimension has always been unchanged. However, shape-polymorphic functions can also change the extent:

```
selEven :: (Shape sh, Elt e)
    => Array (sh:.Int) e -> Array (sh:.Int) e
selEven arr = backpermute new_extent expand arr
  where
    (ns :.n)      = extent arr
    new_extent    = ns :.(n 'div' 2)
    expand (is :.i) = is :.(i * 2)
```

As we can see from the calculation of `new_extent`, the array returned by `selEven` is half as big as the input array, in the rightmost dimension. The index calculation goes in the opposite direction, selecting every alternate element from the input array.

Note carefully that the extent of the new array is calculated from the extent of the old array, *but not from the data in the array*. This guarantees we can do rank generalisation and still have a rectangular array. To see the difference, consider:

```
filter :: Elt e => (e -> Bool)
    -> Array DIM1 e -> Array DIM1 e
```

The `filter` function is not, and cannot be, shape-polymorphic. If we filter each row of a matrix based on the element values, then each new row may have a different length. This gives no guarantee that the resulting matrix is rectangular. In contrast, we have carefully chosen our shape-polymorphic primitives to guarantee the rectangularity of the output.

```
data All      = All
data Any sh   = Any

type family FullShape ss
type instance FullShape Z      = Z
... FullShape (Any sh)        = sh
... FullShape (s1 :. Int)     = FullShape s1 :. Int
... FullShape (s1 :. All)     = FullShape s1 :. Int

type family SliceShape ss
type instance SliceShape Z     = Z
... SliceShape (Any sh)       = sh
... SliceShape (s1 :. Int)    = SliceShape s1
... SliceShape (s1 :. All)    = SliceShape s1 :. Int

class Slice ss where
  sliceOfFull :: ss -> FullShape ss -> SliceShape ss
  fullOfSlice :: ss -> SliceShape ss -> FullShape ss

instance Slice Z      where ...
instance Slice (Any sh) where ...

instance Slice s1 => Slice (s1 :. Int) where
  sliceOfFull (fsl :. _) (ssl :. _)
    = sliceOfFull fsl ssl

  fullOfSlice (fsl :. n) ssl
    = fullOfSlice fsl ssl :. n

instance Slice s1 => Slice (s1 :. All) where
  sliceOfFull (fsl :. All) (ssl :. s)
    = sliceOfFull fsl ssl :. s

  fullOfSlice (fsl :. All) (ssl :. s)
    = fullOfSlice fsl ssl :. s

replicate :: ( Slice s1, Elt e
             , Shape (FullShape s1)
             , Shape (SliceShape s1)
             => s1 -> Array (SliceShape s1) e
             -> Array (FullShape s1) e
replicate s1 arr
  = backpermute (fullOfSlice s1 (extent arr))
    (sliceOfFull s1) arr

slice :: ( Slice s1, Elt e
        , Shape (FullShape s1)
        , Shape (SliceShape s1)
        => Array (FullShape s1) e
        -> s1 -> Array (SliceShape s1) e
slice arr s1
  = backpermute (sliceOfFull s1 (extent arr))
    (fullOfSlice s1) arr
```

Figure 4. Definition of slices

4.4 Slices and slice constraints

Shape types characterise a *single* shape. However, some collective array operations require a relationship between *pairs* of shapes. One such operation is `replicate`, which we used in `mmMult`. The function `replicate` takes an array of arbitrary rank and replicates it along one or more additional dimensions. Note that we cannot uniquely determine the behaviour of `replicate` from the shape of the original and resulting arrays alone. For example, suppose

that we want to expand a rank-2 array into a rank-3 array. There are three ways of doing this, depending on which dimension of the resulting array is to be duplicated. Indeed, the two calls to `replicate` in `mmMult` performed replication along two different dimensions, corresponding to different sides of the cuboid in Figure 2.

It should be clear that `replicate` needs an additional argument, a *slice specifier*, that expresses exactly how the shape of the result array depends on the shape of the argument array. A slice specifier has the same format as an array index, but some index positions may use the value `All` instead of a numeric index.

```
data All = All
```

In `mmMult`, we use `replicate (Z:.All:.colsB:.All) arr` to indicate that we replicate `arr` across the second innermost axis, `colsB` times. We use `replicate (Z:.rowsA:.All:.All) trr` to specify that we replicate `trr` across the outermost axis, `rowsA` times.

The type of the slice specifier `(Z :.All :.colsB :.All)` is `(Z :.All :.Int :.All)`. This type is sufficiently expressive to determine the shape of *both* the original array, before it gets replicated, *and* of the replicated array. More precisely, both of these types are a function of the slice specifier type. In fact, we derive these shapes using *associated type families*, a recent extension to the Haskell type system Chakravarty et al. [2005a], Schrijvers et al. [2008], using the definition for the `Slice` type class shown in Figure 4.

A function closely related to `replicate` is `slice`, which extracts a slice along multiple axes of an array. The full types of `replicate` and `slice` appear in Figure 4. We chose their argument order to match that used for lists: `replicate` is a generalisation of `Data.List.replicate`, while `slice` is a generalisation of `Data.List.(!!)`.

Finally, to enable rank generalisation for `replicate` and `slice`, we add a last slice specifier, namely `Any`, which is also defined in Figure 4. It is used in the tail position of a slice, just like `Z`, but gives a shape variable for rank generalisation. With its aid we can write `repN` which replicates an arbitrary array `n` times, with the replication being on the rightmost dimension of the result array:

```
repN :: Int -> Array sh e -> Array (sh:.Int) e
repN n a = replicate (Any:.n) a
```

5. Rectangular arrays, purely functional

As mentioned in Section 3, the type class `Elt` determines the set of types that can be used as array elements. We adopt `Elt` from the library of unboxed one-dimensional arrays in `Data.Parallel.Haskell`. With this library, array elements can be of the basic numeric types, `Bool`, and pairs formed from the strict pair constructor:

```
data a :: b = !a :: !b
```

We have also extended this to support index types, formed from `Z` and `(:.)`, as array elements. Although it would be straightforward to allow other product and enumeration types as well, support for general sum types appears impractical in a framework based on regular arrays. Adding this would require irregular arrays and nested data parallelism Peyton Jones et al. [2008].

Table 1 summarises the central functions of our library `Repa`. They are grouped according to the structure of the implemented array operations. We discuss the groups and their members in the following sections.

5.1 Structure-preserving operations

The simplest group of array operations are those that apply a transformation on individual elements without changing the shape, array

size, or order of the elements. We have the plain `map` function, `zip` for element-wise pairing, and a family of `zipWith` functions that apply workers of different arity over multiple arrays in lockstep. In the case of `zip` and `zipWith`, we determine the shape value of the result by intersecting the shapes of the arguments — that is, we take the minimum extent along every axis. This behaviour is the same as Haskell’s `zip` functions when applied to lists.

The function `map` is implemented as follows:

```
map :: (a -> b) -> Array sh a -> Array sh b
map f arr = traverse arr id (f .)
```

The various `zip` functions are implemented in a similar manner, although they also use a method of the `Shape` type class to compute the intersection shape of the arguments.

5.2 Reductions

Our library, `Repa`, provides two kinds of reductions: (1) generic reductions, such as `foldl`, and (2) specialised reductions, such as `sum`. In a purely sequential implementation, the latter would be implemented in terms of the former. However, in the parallel case we must be careful.

Reductions of an n element array can be computed with parallel tree reduction, providing $\log n$ asymptotic step complexity in the ideal case, but only if the reduction operator is associative. Unfortunately, Haskell’s type system does not provide a way to express this side condition on the first argument of `foldl`. Hence, the generic reduction functions must retain their sequential semantics to remain deterministic. In contrast, for specialised reductions such as `sum`, when we know that the operators they use meet the associativity requirement, we can use parallel tree reduction.

As outlined in Section 4.3, all reduction functions are defined with a shape polymorphic at-least constraint and admit rank generalisation. Therefore, even generic reductions, with their sequential semantics, are highly parallel if used with rank generalisation.

Rank generalisation also affects specialised reductions, as they can be implemented in one of the following two ways. Firstly, if we want to maximise parallelism, then we can use a segmented tree reduction that conceptually performs multiple parallel tree reductions concurrently. Alternatively, we can simply use the same scheme as for general reductions, and perform all rank one reductions in parallel. We follow the latter approach and sacrifice some parallelism, as tree reductions come with some sequential overhead.

In summary, when applied to an array of rank one, generic reductions (`foldl` etc.) execute purely sequentially with an asymptotic step complexity of n , whereas specialised reductions (`sum` etc.) execute in parallel using a tree reduction with an asymptotic step complexity of $\log n$. In contrast, when applied to an array of rank strictly greater than one, both generic and specialised reductions use rank generalisation to execute many sequential reductions on one-dimensional subarrays concurrently.

5.3 Index space transformations

The structure-preserving operations and the reductions transform array elements, whereas index space transformations only alter the index at which an element is placed — that is, they rearrange and possibly drop elements. A prime example of this group of operations is `reshape`, which imposes a new shape on the elements of an array. A precondition of `reshape` is that the size of the extent of the old and new array is the same, meaning that the number of elements stays the same:

```
reshape :: Shape sh
=> sh -> Array sh' e -> Array sh e
reshape sh' (Manifest sh ua)
= assert (size sh == size sh') $
  Manifest sh' ua
```

Structure-preserving operations

<code>map</code>	<code>:: (Shape sh, Elt a, Elt b)</code> <code>=> (a -> b) -> Array sh a -> Array sh b</code>	Apply function to every array element.
<code>zip</code>	<code>:: (Shape sh, Elt a, Elt b)</code> <code>=> Array sh a -> Array sh b -> Array sh (a *: b)</code>	Elementwise pairing.
<code>zipWith</code>	<code>:: (Shape sh, Elt a, Elt b, Elt c)</code> <code>=> (a -> b -> c) -> Array sh a -> Array sh b -> Array sh c</code> (Other map-like operations: <code>zipWith3</code> , <code>zipWith4</code> , and so on)	Apply a function elementwise to two arrays. (the resulting shape is the intersection)

Reductions

<code>foldl</code>	<code>:: (Shape sh, Elt a, Elt b)</code> <code>=> (a -> b -> a) -> a -> Array (sh:.Int) b -> Array sh a</code> (Other reduction schemes: <code>foldr</code> , <code>foldl1</code> , <code>foldr1</code> , <code>scanl</code> , <code>scanr</code> , <code>scanl1</code> & <code>scanr1</code>)	Left fold.
<code>sum</code>	<code>:: (Shape sh, Elt e, Num e) => Array (sh:.Int) e -> Array sh a</code> (Other specific reductions: <code>product</code> , <code>maximum</code> , <code>minimum</code> , and <code>& or</code>)	Sum an array along its innermost axis.

Index space transformations

<code>reshape</code>	<code>:: (Shape sh, Shape sh', Elt e)</code> <code>=> sh -> Array sh' e -> Array sh e</code>	Impose a new shape on the same elements.
<code>replicate</code>	<code>:: (Slice sl, Shape (FullShape sl), Shape (SliceShape sl))</code> <code>=> sl -> Array (SliceShape sl) e -> Array (FullShape sl) e</code>	Extend an array along new dimensions.
<code>slice</code>	<code>:: (Slice sl, Shape (FullShape sl), Shape (SliceShape sl))</code> <code>=> Array (FullShape sl) e -> sl -> Array (SliceShape sl) e</code>	Extract a subarray according to a slice specification.
<code>(+:+)</code>	<code>:: Shape sh => Array sh e -> Array sh e -> Array sh e</code>	Append a second array to the first.
<code>backpermute</code>	<code>:: (Shape sh, Shape sh', Elt e)</code> <code>=> sh' -> (sh' -> sh) -> Array sh e -> Array sh' e</code>	Backwards permutation.
<code>backpermuteDft</code>	<code>:: (Shape sh, Shape sh', Elt e)</code> <code>=> Array sh' e -> (sh' -> Maybe sh) -> Array sh e</code> <code>-> Array sh' e</code>	Default backwards permutation.
<code>unit</code>	<code>:: Elt e => e -> Array Z e</code>	Wrap a scalar into a singleton array.
<code>(!::)</code>	<code>:: (Shape sh, Elt e) => Array sh e -> sh -> e</code>	Extract an element at a given index.

General traversal

<code>traverse</code>	<code>:: (Shape sh, Shape sh', Elt e)</code> <code>=> Array sh e -> (sh -> sh') -> ((sh -> e) -> sh' -> e')</code> <code>-> Array sh' e'</code>	Unstructured traversal.
<code>force</code>	<code>:: (Shape sh, Elt e) => Array sh e -> Array sh e</code>	Force a delayed array into manifest form.
<code>extent</code>	<code>:: Array sh e -> sh</code>	Obtain size in all dimensions of an array.

Table 1. Summary of array operations

```
reshape sh' (Delayed sh f)
  = assert (size sh == size sh') $
    Delayed sh' (f . fromIndex sh . toIndex sh')
```

The functions `index` and `fromIndex` are methods of the class `Shape` from Figure 3. The functions `replicate` and `slice` were already discussed in Section 4.4, and `unit` and `(!::)` are defined as follows:

```
unit :: e -> Array Z e
unit = Delayed Z . const
```

```
(!::) :: (Shape sh, Elt e) => Array sh e -> sh -> e
arr !: ix = snd (delay arr) ix
```

A simple operator to rearrange elements is the function `(+:+)`; it appends its second argument to the first and can be implemented with `traverse` by adjusting shapes and indexing.

In contrast, general shuffle operations, such as backwards permutation, require the detailed mapping of target to source indices.

We have seen this in the example `transpose2D` in Section 2. Another example is the following function that extracts the diagonal of a square matrix.

```
diagonal :: Elt e => Array DIM2 e -> Array DIM1 e
diagonal arr = assert (width == height) $
  backpermute width (\x -> (x, x)) arr
  where
    _ .. height .. width = extent arr
```

Code that uses `backpermute` appears more like element-based array processing. However, it is still a collective operation with a clear parallel interpretation.

Backwards permutation is defined in terms of the general `traverse` as follows:

```
backpermute :: sh' -> (sh' -> sh) -> Array sh e
             -> Array sh' e
backpermute sh perm arr
  = traverse arr (const sh) (. perm)
```

The variant `backpermuteDft`, known as default backwards permutation, operates in a similar manner, except that the target index is partial. When the target index maps to `Nothing`, the corresponding element from the default array is used. Overall, `backpermuteDft` can be interpreted as a means to bulk update the contents of an array. As we are operating on purely functional, immutable arrays, the original array is still available and the repeated use of `backpermuteDft` is only efficient if large part of the array are updated on each use.

5.4 General traversal

The most general form of array traversal is `traverse`, which supports an arbitrary change of shape and array contents. Nevertheless, it is still represented as a delayed computation as detailed in Section 3.3. Although for efficiency reasons it is better to use specific functions such as `map` or `backpermute`, it is always possible to fall back on `traverse` if a custom computational structure is required.

For example, `traverse` can be used to implement stencil-based relaxation methods, such as the following update function to solve the Laplace equation in a two dimensional grid Mathews and Fink [1999]:

$$u'(i, j) = (u(i-1, j) + u(i+1, j) + u(i, j-1) + u(i, j+1)) / 4$$

To implement this stencil, we use `traverse` as follows:

```
stencil :: Array DIM2 Double -> Array DIM2 Double
stencil arr
  = traverse arr id update
  where
    _ :: height :: width = extent arr

    update get d@(sh :: i :: j)
      = if isBoundary i j
        then get d
        else (get (sh :: (i-1) :: j)
              + get (sh :: i :: (j-1))
              + get (sh :: (i+1) :: j)
              + get (sh :: i :: (j+1))) / 4

    isBoundary i j
      = (i == 0) || (i >= width - 1)
        || (j == 0) || (j >= height - 1)
```

As the shape of the result array is the same as the input, the second argument to `traverse` is `id`. The third argument is the update function that implements the stencil, while taking the grid boundary into account. The function `get`, passed as the first argument to `update`, is the lookup function for the input array.

To solve the Laplace equation we would set boundary conditions along the edges of the grid and then iterate `stencil` until the inner elements converge to their final values. However, for benchmarking purposes we simply iterate it a fixed number of times:

```
laplace :: Int
         -> Array DIM2 Double -> Array DIM2 Double
laplace steps arr = go steps arr
  where
    go s arr
      | s == 0    = arr
      | otherwise = go (s-1) (force $ stencil arr)
```

The use of `force` after each recursion is important, as it ensures that all updates are applied and that we produce a manifest array. Without it, we would accumulate a long chain of delayed computations with a rather non-local memory access pattern. In Repa, the function `force` triggers all computation, and as we will discuss in

Section 7, the size of forced array determines the amount of parallelism in an algorithm.

6. Delayed arrays and loop fusion

We motivated the use of delayed arrays in Section 3.2 by the desire to avoid superfluous copying of array elements during index space transformation, such as in the definition of `backpermute`. However, another major benefit of delayed arrays is that it gives *by-default automatic loop fusion*. Recall the implementation of `map`:

```
map :: (a -> b) -> Array sh a -> Array sh b
map f arr = traverse arr id (f .)
```

Now, imagine evaluating `(map f (map g a))`. If you consult the definition of `traverse` (Section 3.3) it should be clear that the two `maps` simply build a delayed array whose indexing function first indexes `a`, then applies `g`, and then applies `f`. No intermediate arrays are allocated and, in effect, the two loops have been fused. Moreover, this fusion does not require a sophisticated compiler transformation, nor does it require the two calls of `map` to be statically juxtaposed; fusion is a property of the data representation.

Guaranteed, automatic fusion sounds too good to be true — and so it is. The trouble is that we cannot *always* use the delayed representation for arrays. One reason not to delay arrays is data layout, as we discussed in Section 3.3. Another is parallelism: `force` triggers data-parallel execution (Section 7). But the most immediately pressing problem with the delayed representation is *sharing*. Consider the following:

```
let b = map f a
in mmMult b b
```

Every access to an element of `b` will apply the (arbitrarily-expensive) function `f` to the corresponding element of `a`. It follows that *these arbitrarily-expensive computations will be done at least twice*, once for each argument of `mmMult`, quite contrary to the programmer's intent. Indeed, if `mmMult` itself consumes elements of its arguments in a non-linear way, accessing them more than once, the computation of `f` will be performed each time. If instead we say:

```
let b = force (map f a)
in mmMult b b
```

then the now-manifest array `b` ensures that `f` is called only once for each element of `a`. In effect, a manifest array is simply a memo table for a delayed array. Here is how we see the situation:

- In most array libraries, every array is *manifest by default*, so that sharing is guaranteed. However, loop fusion is difficult, and must often be done manually, doing considerable violence to the structure of the program.
- In Repa every array is *delayed by default*, so that fusion is guaranteed. However, sharing may be lost; it can be restored manually by adding calls to `force`. These calls do not affect the structure of the program.

By using `force`, Repa allows the programmer tight control over some crucial aspects of the program: sharing, data layout, and parallelism. The cost is, of course, that the programmer must exercise that control to get good performance. Ignoring the issue altogether can be disastrous, because it can lead to arbitrary loss of sharing. In further work, beyond the scope of this paper, we are developing a compromise approach that offers guaranteed sharing with aggressive (but not guaranteed) fusion.

7. Parallelism

As described in Section 3.1, all elements of a Repa array are demanded simultaneously. This is the source of all parallelism

in the library. In particular, an application of the function `force` triggers the parallel evaluation of a delayed array, producing a manifest one. Assuming that the array has n elements and that we have P parallel *processing elements* (PEs) available to perform the work, each PE is responsible for computing n/P consecutive elements in the row-major layout of the manifest array. In other words, the structure of parallelism is always determined by the layout and partitioning of a forced array. The execution strategy is based on gang parallelism and is described in detail in Chakravarty et al. [2007].

Let us re-consider the function `mmMult` from Section 3.3 and Figure 2 in this light. We assume that `arr` is a manifest array, and know that `trr` is manifest because of the explicit use of `force`. The rank-2 array produced by the rank-generalised application of `sum` corresponds to the right face of the cuboid from Figure 2. Hence, if we `force` the result of `mmMult`, the degree of available parallelism is proportional to the number of elements of the resulting array — 8 in the figure. As long as the hardware provides a sufficient number of PEs, each of these elements may be computed in parallel. Each involves the element-wise multiplication of a row from `arr` with a row from `trr` and the summation of these products. If the hardware provides fewer PEs, which is usually the case, the evaluation is evenly distributed over the available PEs.

Let’s now turn to a more sophisticated parallel algorithm, the three-dimensional fast Fourier transform (FFT). Three-dimensional FFT works on one axis at a time: we apply the one dimensional FFT to all vectors along one axis, then the second and then the third. Instead of writing a separate transform for each dimension, we implement one-dimensional FFT as a shape polymorphic function that operates on the innermost axis. We combine it with a three-dimensional rotation, `rotate3D`, which allows us to cover all three axes one after another:

```
fft3D :: Array DIM3 Complex -- roots of unity
      -> Array DIM3 Complex -- data to transform
      -> Array DIM3 Complex
fft3D rofu = fftTrans . fftTrans . fftTrans
  where
    fftTrans = rotate3D . fft1D rofu
```

The first argument, `rofu`, is an array of complex roots of unity, which are constants that we wish to avoid recomputing for each call. The second is the three-dimensional array to transform, and we require both arrays to have the same shape. We also require each dimension to have a size which is a power of 2.

If the result of `fft3D` is `forced`, evaluation by P PEs is again on P consecutive segments of length n^3/P of the row-major layout of the transformed cube, where n is the side length of the cube. However, the work that needs to be performed for each of the elements is harder to characterise than for `mmMult`, as the computations of the individual elements of the result are not independent and as `fft1D` uses `force` internally.

Three-dimensional rotation is easily defined based on the function `backpermute` which we discussed previously:

```
rotate3D :: Array DIM3 Complex
         -> Array DIM3 Complex
rotate3D arr = backpermute (Z:.m:.k:.l) f
  where
    (Z:.k:.l:.m) = extent arr
    f (Z:.m:.k:.l) = (Z:.k:.l:.m)
```

The one-dimensional fast Fourier transform is more involved: it requires us to recursively split the input vector in half and apply the transform to the split vectors. To facilitate the splitting, we first define a function `halve` that drops half the elements of a vector,

where the elements to pick of the original are determined by a selector function `sel`.

```
halve :: (sh:.Int -> sh:.Int)
      -> Array (sh:.Int) Complex
      -> Array (sh:.Int) Complex
halve sel arr
  = backpermute (sh :. n 'div' 2) sel arr
  where
    sh:.n = extent arr
```

By virtue of rank generalisation, this shape polymorphic function will split all rows of a three-dimensional cube at once and in the same manner.

The following two convenience functions use `halve` to extract all elements in even and odd positions, respectively.

```
evenHalf, oddHalf :: Array (sh:.Int) Complex
                  -> Array (sh:.Int) Complex
evenHalf = halve (\(ix:.i) -> ix :. 2*i)
oddHalf  = halve (\(ix:.i) -> ix :. 2*i+1)
```

Now, the definition of the one-dimensional transform is a direct encoding of the Cooley-Tukey algorithm:

```
fft1D :: Array (sh:.Int) Complex
      -> Array (sh:.Int) Complex
      -> Array (sh:.Int) Complex
fft1D rofu v
  | n > 2 = (left +^ right) :+ (left -^ right)
  | n == 2 = traverse v id swivel
  where
    (_ :. n) = extent v

    swivel f (ix:.0) = f (ix:.0) + f (ix:.1)
    swivel f (ix:.1) = f (ix:.0) - f (ix:.1)

    rofu' = evenHalf rofu
    left  = force . fft1D rofu' . evenHalf $ v
    right = force . (*^ rofu) . fft1D rofu' . oddHalf $ v

    (+^) = zipWith (+)
    (-^) = zipWith (-)
    (*^) = zipWith (*)
```

All the index space transformations that are implemented in terms of `backpermute`, as well as the elementwise arithmetic operations based on `zipWith` produce delayed arrays. It is only the use of `force` in the definition of `left` and `right` that triggers the parallel evaluation of subcomputations. In particular, as we `force` the recursive calls in the definition of `left` and `right` separately, these calls are performed in sequence. The rank-generalised input vector `v` is halved with each recursive call, and hence, the amount of available parallelism decreases.

However, keep in mind that —by virtue of rank generalisation— we perform the one-dimensional transform in parallel on all vectors of a cuboid. That is, if we apply `fft3D` to a $64 \times 64 \times 64$ cube, then `fft1D` still operates on $64 * 64 * 2 = 8192$ complex numbers in one parallel step at the base case, where $n = 2$.

8. Benchmarks

In this section, we discuss the performance of three programs presented in this paper: matrix-matrix multiplication from Section 3.3, the Laplace solver from Section 5.4 and the fast Fourier transform from Section 7. We ran the benchmarks on two different machines:

- a 2x Quad-core 3GHz Xeon server and
- a 1.4GHz UltraSPARC T2.

		GCC	Repa	
		4.2.1	1 thread	fastest parallel
Matrix mult	1024×1024	3.8s	4.6s	0.64s
Laplace	300×300	0.70s	1.7s	0.68s
FFT	128×128×128	0.24s	8.8s	2.0s

Figure 5. Performance on the Xeon

		GCC	Repa	
		4.1.2	1 thread	fastest parallel
Matrix mult	1024×1024	53s	92s	2.4s
Laplace	300×300	6.5s	32s	3.8s
FFT	128×128×128	2.4s	98s	7.7s

Figure 6. Performance on the SPARC

The first machine is a typical x86-based server with good single-core performance but frequent bandwidth problems in memory-intensive applications. The bus architecture directly affects the scalability of some of our benchmarks, namely, the Laplace solver, which cannot utilise multiple cores well due to bandwidth limitations.

The SPARC-based machine is more interesting. The T2 processor has 8 cores and supports up to 8 hardware threads per core. This allows it to effectively hide memory latency in massively multi-threaded programs. Thus, despite a significantly worse single-core performance than the Xeon it exhibits much better scalability which is clearly visible in our benchmarks.

8.1 Absolute performance

Before discussing the parallel behaviour of our benchmarks, let us investigate how Repa programs compare to hand-written C code when executed with only one thread. The C matrix-matrix multiplication and Laplace solver are straightforwardly written programs, while the FFT uses FFTW 3.2.2 Frigo and Johnson [2005] in “estimate” mode.

Figure 5 shows the single threaded results together with the fastest running times obtained through parallel execution. For matrix multiplication and Laplace on the Xeon, Repa is slower than C when executed sequentially, but not by much. FFTW uses a finely tuned in-place algorithm, which is significantly faster but more complicated than our own direct encoding of the Cooley-Tukey algorithm. We include the numbers with FFTW for comparative purposes, but note that parallelism is no substitute for a more efficient algorithm.

Compared with the Xeon, the results on the SPARC (Figure 6) are quite different. The SPARC T2 is a “throughput” machine, designed to execute workloads consisting of many concurrent threads. It has half the clock rate of the Xeon, and does not exploit instruction level parallelism. This shows in the fact that the single threaded C programs run about 10x slower than their Xeon counterparts. The SPARC T2 also does not perform instruction reordering or use speculative execution. GHC does not perform compile time scheduling to account for this, which results in a larger gap between the single threaded C and Repa programs than on the Xeon.

We have also compared the performance of the Laplace solver to an alternative, purely sequential Haskell implementation based on unboxed, mutable arrays running in the IO Monad (IOUArray). This version was about two times slower than the Repa program, probably due to the overhead introduced by bounds checking, which is currently not supported by our library. Note, however, that bounds checking is unnecessary for many collective opera-

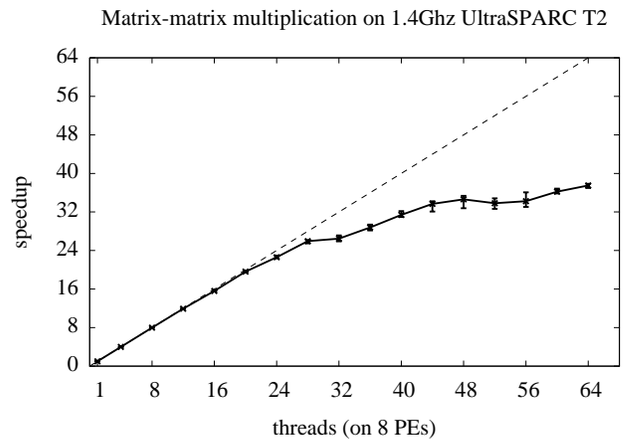
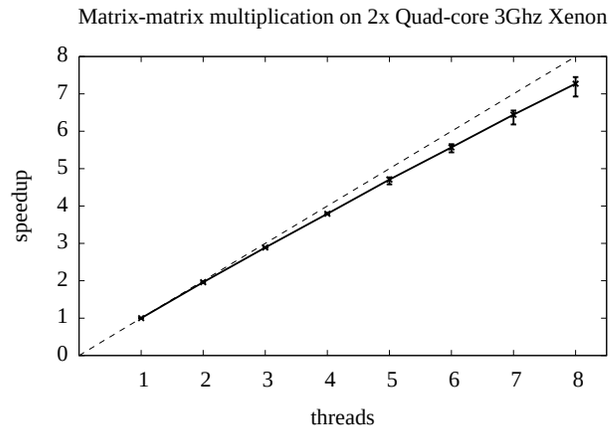


Figure 7. Matrix-matrix multiplication, size 1024x1024.

tions such as map and sum, so even after we introduce it in Repa we still expect to see better performance than a low-level, imperative implementation based on mutable arrays.

8.2 Parallel behaviour

The parallel performance of matrix multiplication is shown in Figure 7.³ Each point shows the lowest, average, and highest speedups for ten consecutive runs. Here, we get excellent scalability on both machines. On the Xeon, we achieve a speedup of 7.2 with 8 threads. On the SPARC, it scales up to 64 threads with a peak speedup of 38.

Figure 8 shows the relative speedups for the Laplace solver. This program achieves good scalability on the SPARC, reaching a speedup of 8.4 with 14 threads but performs much worse on the Xeon, stagnating at a speedup of 2.5. As Laplace is memory bound, we attribute this behaviour to insufficient bandwidth on the Xeon machine. There is also some variation in the speedups from run to run, which is more pronounced when using specific numbers of threads. We attribute this to scheduling effects, in the hardware, OS, and GHC runtime system.

Finally, the parallel behaviour of the FFT implementation is shown in Figure 9. This program scales well on both machines, achieving a relative speedup of 4.4 on with 7 threads on the Xeon and 12.7 on 14 threads on the SPARC. Compared to the Laplace

³ Yes, those really are the results in the first graph of the figure.

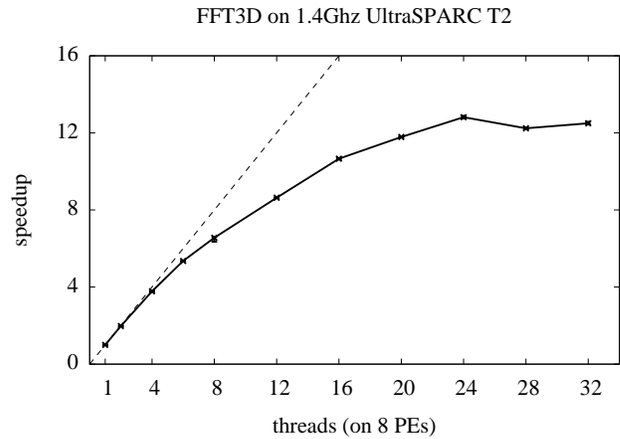
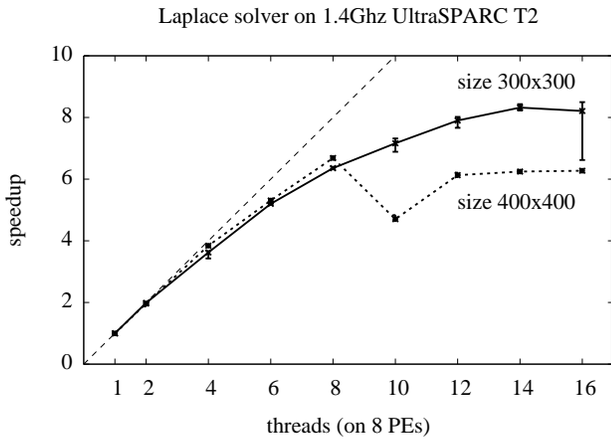
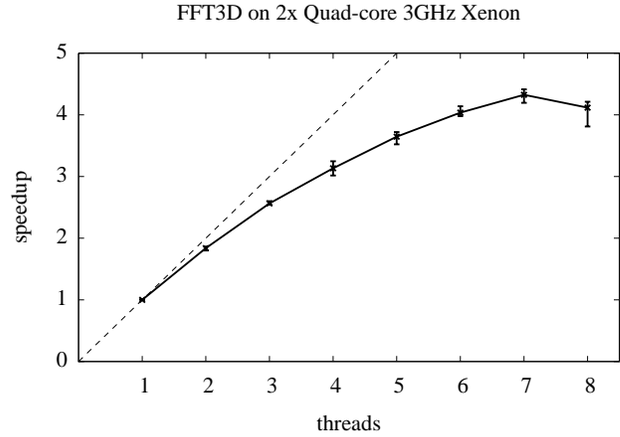
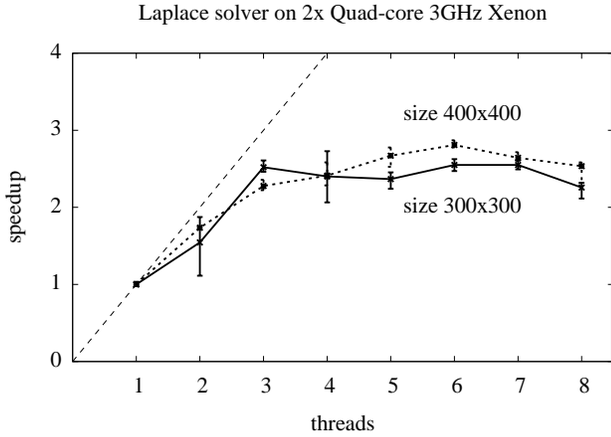


Figure 8. Laplace solver, 1000 iterations.

Figure 9. 3D Fast Fourier Transform, size 128x128x128

solver, this time the scalability is much better on the Xenon but practically unchanged on the SPARC. Note that FFT is less memory intensive than Laplace. The fact that Laplace with a 300×300 matrix does not scale as well on the Xenon as it does on the SPARC supports our conclusion that this benchmark suffers from lack of memory bandwidth. For Laplace with a 400×400 matrix on the SPARC, we suspect the sharp drop off after 8 threads is due to the added threads contending for cache. As written, the implementation from Section 5.4 operates on a row-by-row basis. We expect that changing to a block-wise algorithm would improve cache-usage and reduce the bandwidth needed.

9. Related Work

Array programming is a highly active research area so the amount of related work is quite significant. In this section, we have to restrict ourselves to discussing only a few most closely related approaches.

9.1 Haskell array libraries

Haskell 98 already defines an array type as part of its prelude which, in fact, even provides a certain degree of shape polymorphism. These arrays can be indexed by arbitrary types as long as they are instances of `Ix`, a type class which plays a similar role to our `Shape`. This allows for fully shape-polymorphic functions such as `map`. However, standard Haskell arrays do not support at-least constraints and rank generalisation which are crucial for imple-

menting highly expressive operations such as `sum` from Section 4.3. This inflexibility precludes many advanced uses of shape polymorphism described in this paper and makes even unboxed arrays based on the same interface a bad choice for a parallel implementation.

Partly motivated by the shortcomings of standard arrays, numerous Haskell array libraries have been proposed in recent years. These range from highly specialised ones such as `ByteString Coutts et al. [2007b]` to full-fledged DSLs for programming GPUs `Lee et al. [2009]`. However, these libraries do not provide the same degree of flexibility and efficiency for manipulating regular arrays if they support them at all. Our own work on `Data Parallel Haskell` is of particular relevance in this context as the work presented in this paper shares many of its ideas and large parts of its implementation with that project. Indeed, `Repa` can be seen as complementary to `DPH`. Both provide a way of writing high-performance parallel programs but `DPH` supports irregular, arbitrarily nested parallelism which requires it to sacrifice performance when it comes to purely regular computations. One of the goals of this paper is to plug that hole. Eventually, we intend to integrate `Repa` into `DPH`, providing efficient support for both regular and irregular arrays in one framework.

9.2 C++ Array Libraries

Due to its powerful type system and its wide-spread use in high-performance computing, C++ has a significant number of array libraries that are both fast and generic. In particular, `Blitz++` `Veld-`

huizen [1998] and `Boost.MultiArray` `boo` [2010] feature multi-dimensional arrays with a restricted form of shape polymorphism. However, our library is much more flexible in this regard and also has the advantage of a natural parallel implementation which neither of the two C++ libraries provide. Moreover, these approaches are inherently imperative while we provide a purely functional interface which allows programs to be written at a higher level of abstraction.

9.3 Array Languages

In addition to libraries, there exist a number of special-purpose array programming languages. Of these, Single Assignment C (SAC) Scholz [2003] has exerted the most influence on our work and is the closest in spirit as it is purely functional and strongly typed. SAC provides many of the same benefits as `Repa`: high-performance arrays with shape polymorphism, expressive collective operations and extensive optimisation based on *with-loops*, a special-purpose language construct for creating, traversing and reducing arrays. It also comes with a rich library of standard array and matrix operations which `Repa` has not yet acquired.

However, `Repa` has the advantage of being integrated into a mainstream functional language and not requiring specific compiler support. This allows `Repa` programs to utilise the entire Haskell infrastructure and to drop down to a very low level of abstraction if required in specific cases. This, along with strong typing and purity, are also the advantages `Repa` has over other array languages such as APL, J and Matlab The International Standards Organisation [1989], Burke [1996], Gilat [2004].

Acknowledgements. We are grateful to Arvind for explaining the importance of delaying index space transformations and thank Simon Winwood for comments on a draft. We also thank the anonymous ICFP'10 reviewers for their helpful feedback on the paper. This research was funded in part by the Australian Research Council under grant number LP0989507.

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