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Finding Parallel Functional Pearls: Automatic Parallel Recursion Scheme Detection in Haskell Functions *via* Anti-Unification

Adam D. Barwell, Christopher Brown, and Kevin Hammond

School of Computer Science, University of St Andrews, St Andrews, United Kingdom {adb23,cmb21,kh8}@st-andrews.ac.uk

Abstract

This paper describes a new technique for identifying potentially parallelisable code structures in functional programs. *Higher-order functions* enable simple and easily understood abstractions that can be used to implement a variety of common recursion schemes, such as maps and folds over traversable data structures. Many of these recursion schemes have natural parallel implementations in the form of algorithmic skeletons. This paper presents a technique that detects instances of potentially parallelisable recursion schemes in Haskell 98 functions. Unusually, we exploit *anti-unification* to expose these recursion schemes from source-level definitions whose structures match a recursion scheme, but which are not necessarily written directly in terms of maps, folds, etc. This allows us to automatically introduce parallelism, without requiring the programmer to structure their code a priori in terms of specific higher-order functions. We have implemented our approach in the Haskell refactoring tool, HaRe, and demonstrated its use on a range of common benchmarking examples. Using our technique, we show that recursion schemes can be easily detected, that parallel implementations can be easily introduced, and that we can achieve real parallel speedups (up to $23.79 \times$ the sequential performance on 28 physical cores, or $32.93 \times$ the sequential performance with hyper-threading enabled).

1. Introduction

Structured parallelism techniques, such as *algorithmic skeletons*, expose parallelism through suitable language-level constructs [1]. By exploiting a range of high-level, composable patterns that have natural parallel implementations, such as *maps* or *folds*, parallelism can be introduced quickly, easily and automatically. Unlike common lower-level approaches, such as OpenCL, the programmer does not need to explicitly deal with implementation details such as communication, task creation, scheduling, etc. [2]. Structured parallel code also avoids common and difficult-to-debug problems, such as race conditions and deadlocks, and allows the definition of simple, but effective, *cost models* that can accurately

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predict parallel runtimes and speedups. There is a close, and long-observed, correspondence between structured approaches and functional programming. In particular, algorithmic skeletons can be seen as parallel implementations of higher-order functions (e.g. [3, 4]). There are further advantages to structured parallel approaches when writing parallel software for heterogeneous [5] or high performance parallel systems [6], which may involve massive numbers of processors, of possibly different types. Given a set of skeletons for the desired target architecture(s), the programmer need only be concerned with which skeletons to call, where they should be called, and what parameters should be given to them so that they give the best performance on the given architecture. Unfortunately, determining precisely where to introduce skeletons can be a non-trivial exercise, requiring significant programmer effort [2, 7, 8]. It follows that if instances of higher-order functions can be discovered automatically, and if cost information can be used to direct the choices of skeletons, parameters, etc., then parallel functional programs can be produced quickly, easily and efficiently. Such patterns occur frequently in real applications. For example, the *spectral* part of the NoFib suite of Haskell benchmarks [9] comprises a total of 48 programs. Manual inspection shows that at least 19 (39.6%) of these have one or more functions that could be rewritten in terms of map or fold patterns. This gives a potentially large corpus of easily parallelisable programs.

1.1. Novel Contributions

This paper introduces a new technique for discovering instances of generalised higher-order functions (recursion schemes), using anti-unification [10]. As part of our technique, we define a new, specialised anti-unification algorithm, and demonstrate how pattern arguments can be derived from the substitutions that are inferred by the anti-unification process. Our anti-unification analysis and corresponding refactorings are implemented for Haskell in the HaRe refactoring tool. Our implementation allows the automatic discovery and refactoring of parallelisable recursion schemes, detecting instances that could otherwise be difficult to detect. Moreover, manually checking a large code base for possible instances of parallelisable patterns would be a time-consuming and error-prone task, even for an expert programmer. We have tested our prototype on four standard benchmark programs: matrix multiplication, n-body, n-queens, and sumeuler, demonstrating that it can successfully expose the underlying recursion schemes. The corresponding parallel implementations achieve maximum speedups of 32.93 (21.66 using physical cores) for matrix multiplication, 27.08 (23.46 using physical cores) for n-body, 22.65 (20.48 using physical cores) for n-queens, and 30.50 (22.24 using physical cores) for sumeuler, on a 28-core hyper-threaded experimental testbed machine. Our technique is not restricted to Haskell, or to a specific set of recursion schemes, but is, in principle, completely general. It can be applied to a wide range of recursion schemes/skeletons, and to a wide range of programming languages, including both functional languages, such as Erlang, Scala, CAML, or Clojure, and other languages such as C, C++, or Java. The only requirement is that the code structure matches the recursion schemes of interest, and that there are no unwanted side effects.



Figure 1: *sumeuler*, speedups on *corryvreckan*, a 28-core hyper-threaded Intel Xeon server, dashed line shows extent of physical cores.

1.2. Illustrative Example

We illustrate our approach using an example from the *NoFib* suite, sumeuler, that calculates Euler's totient function for a list of integers and sums the results.

```
sumeuler :: [Int] -> Int
sumeuler xs = sum (map euler xs)
```

Here, euler is Euler's totient function, and sum sums the values in its argument, a list of integers. sumeuler can be easily parallelised using, e.g., the Haskell Strategies library:

```
sumeuler :: [[Int]] -> Int
sumeuler xs =
sum (map (sum . map euler) xs `using` parList rdeepseq)
```

Here, parList, from the standard Control.Parallel.Strategies library, parallelises the map over xs without changing the original definition. It is parameterised on rdeepseq, a nested strategy that forces each element of the result to be fully evaluated. By cleanly separating the functional definition of the program from its evaluation strategy, it is easy to introduce alternative parallelisations, while ensuring that the parallel version is functionally equivalent to the original definition. Fig. 1 shows the raw speedups that we obtain for this program for varying sizes on n on our 28-core hyper-threaded experimental machine, *corryvreckan*. We obtain a maximum speedup of 30.50 for n = 50000on 48 hyper-threaded cores. Other parallelisations are possible. Eden [11], the Par Monad [12], GPUs via the Accelerate library [13], etc., can all substitute for parList to introduce parallelism over the *map* operation. It is also possible to rewrite sumeuler to use a direct *fold* (or *reduce*) operation. This applies an operation, g between each pair of elements in an input list, returning q, when the end of the input list is reached. So, for example, foldr (+) 0 [1,2,3] is expanded to foldr (+) 0 ((:) 1 ((:) 2 ((:) 3 [])), where (:) is the binary list constructor function of type $a \rightarrow [a] \rightarrow [a]$ and [] is the empty list. This is reduced to (+) 1 ((+) 2 ((+) 3 0)), the result of which is 6, the sum of the elements in the input list.

```
sumeuler [Int] -> Int
sumeuler xs = foldr ((+) . euler) 0 xs
where foldr g q [] = q
foldr g q (w:ws) = g w (foldr g q ws)
```

This new definition of sumeuler can then be parallelised in an alternative way to the parallel map above, using e.g. a parallel implementation of foldr. Taking advantage of parallelism in sumeuler is therefore both *simple* and potentially *automatic* [14]. Moreover, alternative parallelisation approaches are possible. However, if sumeuler was defined without using either an explicit map or foldr, e.g. as shown below using direct recursion, then parallelisation would be less straightforward, and lower, or no, speedups might result.

```
1 sumeuler :: [Int] -> Int
2 sumeuler [] = 0
3 sumeuler (x:xs) = ((+) . euler) x (sumeuler xs)
```

Since this version of sumeuler is *implicitly* an instance of foldr, it is best to first restructure the definition so that it *explicitly* calls **foldr** before attempting parallelisation. In order to rewrite the inlined sumeuler as an explicit foldr, however, we need to derive concrete arguments to foldr that will yield a functionally equivalent definition. These arguments can be derived by inspecting the inlined definition of sumeuler, as shown in Fig. 2. To *automatically* derive the arguments to foldr, we inspect the definitions of sumeuler and foldr using anti-unification, which aims to find the least general generalisation between two terms. In Plotkin and Reynolds' original work [10, 15], anti-unification was defined for totally ordered terms, where terms consisted of variables, literals, and function application. More recent approaches to anti-unification have applied the technique to programming languages such as Haskell [16, 17], primarily for clone detection and elimination [18]. In these approaches, anti-unification compares two terms (expressions) to find their shared structure, producing an anti-unifier term (representing the shared structure), plus two sets of substitutions that enable the original term to be reconstructed from the anti-unifier. For example, anti-unifying the [] clause of sumeuler with foldr compares the unequal 0 and z expressions, producing the anti-unifier, h:

h g z [] = z

Where the anti-unified structures diverge, a variable is introduced in the antiunifier that can be substituted for the original term. Substitutions, σ , have the



Figure 2: Rewriting sumeuler as an instance of foldr



Figure 3: ASTs for the (:) branch of sumeuler (left) and foldr (right); shared structure in black, differing structure in dark green and red.

form $(v \mapsto t_i)$, where v is a variable and t_i is some term. Substitutions are applied to terms, written using postfix notation, e.g. $t \sigma$, and can be composed like functions. An applied substitution replaces all instances of v with t_i in t. For example, the substitutions for the [] clause of sumeuler are:

$$\sigma_1 = (\mathbf{z} \mapsto \mathbf{0})$$

and the substitutions for the [] clause of foldr are ε , i.e. the identity substitution. Anti-unifying the respective (:) clauses of sumeuler and foldr produces the anti-unifier:

h g z (x:xs) = g x (
$$\alpha$$
 xs)

where α is a free variable. As shown in Fig. 3, the structures of the two clauses are very similar, consisting primarily of application expressions. Differences can be found at the leaves of foldr, highlighted in both dark green and red in Fig. 3. Since x and xs are in the same relative positions, they feature in h. As ((+) . euler) \neq g (highlighted in dark green), g is used to represent

the function applied to two arguments in h. Finally, since the recursive call prefixes sumeuler \neq (foldr g z) (highlighted in red), the variable α is used in their place in h. This produces the substitutions: (g \mapsto ((+) . euler)) and ($\alpha \mapsto$ sumeuler) for sumeuler, and: ($\alpha \mapsto$ (foldr g z)) for foldr. Since it is possible to reconstruct the original terms by applying the relevant substitutions to an anti-unifier, when given the anti-unifier h:

```
\begin{array}{cccc} 1 & h & g & z & [] & = & z \\ 2 & h & g & z & (x:xs) & = & g & x & (a & xs) \end{array}
```

we can, in principle, rewrite sumeuler and foldr in terms of h using substitutions as arguments:

```
sumeuler xs = h ((+) . euler) 0 xs
```

```
2
```

```
foldr g z xs = h g z xs
```

Furthermore, because foldr is *equivalent* to h, we conclude that sumeuler must be an instance of foldr. As both sumeuler and foldr can be rewritten in terms of h, it must be the case that sumeuler can be rewritten in terms of foldr. The substitutions for sumeuler inferred as part of anti-unification are valid as arguments to foldr, allowing sumeuler to be rewritten:

```
sumeuler xs = foldr ((+) . euler) 0 xs
```

Parallelism can now be introduced using a parallel implementation foldr, either manually or using e.g. a refactoring/rewriting tool (e.g. [19, 8]). Alternatively, the foldr operation can be split into its map and foldr (sum) components, perhaps by using the laws of hylomorphisms as in [4], producing:

```
sumeuler xs = foldr (+) 0 (map euler xs)
```

which is equivalent to our original definition of sumeuler:

```
sumeuler xs = sum (map euler xs)
```

This can then be parallelised using the parallel map that we originally showed.

2. Algorithmic Skeletons and Structured Parallelism

Algorithmic skeletons [3] abstract commonly-used patterns of parallel computation, communication and interaction into parameterised templates that can be used to implement high-level patterns of parallelism. There is a long-standing connection between the skeletons community and the functional programming community [20]. In the functional world, skeletons are implemented as *higher*order functions that can be instantiated with specific user code to give some concrete parallel behaviour [2]. Using a skeleton approach allows the programmer to adopt a top-down structured approach to parallel programming, where skeletons are composed to give the overall parallel structure of the program. This gives a flexible approach, where parallelism is exposed to the programmer only through the choice of skeleton and perhaps through some specific behavioural parameters (e.g., the number of parallel processes to be created, or how elements of a parallel list are to be grouped to reduce communication costs). Details of communication, task creation, task or data migration, scheduling, etc. are embedded within the skeleton implementation, which may be tailored to a specific parallel architecture or class of architectures [5].

Common skeletons include: farm, reduce, feedback, and pipeline [1]. A farm applies some function, $(f :: a \rightarrow b)$, to each element in a stream of inputs in parallel, where the number of processes created may be controlled by some parameter. The farm skeleton corresponds to the sequential map operation. The *reduce* skeleton takes an associative function, $(f :: a \rightarrow a \rightarrow a)$, and repeatedly applies it to a collection of two or more elements in parallel until one element, i.e. the result, remains. This corresponds to a *fold* when the fold fixpoint function is associative. The *feedback* skeleton enables looping operations for skeletons, taking some skeleton, \mathbf{s} , to which inputs are applied repeatedly until some function, $(f :: a \rightarrow Bool)$, is satisfied. This corresponds to an *iterate*, where the result is the n^{th} element of the produced list. Finally, the pipeline skeleton enables the composition of two or more skeletons, where parallelism arises from executing multiple pipeline stages in parallel. This corresponds to function composition (.). Many implementations of these skeletons exist, e.g. [1, 7, 8, 12, 11, 19, 21, 22, 23], with some for multiple architectures such as GPUs, e.g. [5, 13, 24]. The choice of implementation is beyond the scope of this paper. In principle, however, this correspondence allows skeletons to be introduced over their sequential equivalents, introducing parallelism for free [4]. This paper takes full advantage of these correspondences. Our antiunification technique therefore focuses on finding instances of recursion scheme implementations such as map, foldr, and iterN.

```
iterate :: (a \rightarrow a) \rightarrow a \rightarrow [a]
1
    iterate f x = x : iterate f (f x)
2
    -- iterate ((+) 1) 1 = [1, 1+1, 1+2, 1+3,...]
3
4
    iterN :: Int \rightarrow (a \rightarrow a) \rightarrow a \rightarrow a
\mathbf{5}
    iterN n f x = iterate f x !! n
6
      - iterN 1 ((+) 1) 1 = 2
7
8
    map :: (a -> b) -> [a] -> [b]
9
    map f [] = []
10
    map f (x:xs) = f x : map f xs
11
     -- map ((+) 1) [1,2,3,..] = [1+1,1+2,1+3,..]
12
13
    foldr :: (a \rightarrow a \rightarrow a) \rightarrow a \rightarrow [a] \rightarrow a
14
    foldr f z []
                        = z
15
    foldr f z (x:xs) = f x (foldr f z xs)
16
  -- foldr (+) 0 [1,2,3,4,5] = 1 + (2 + (3 + (4 + (5 + 0))))
17
```

- \mathcal{P} = Set of pattern implementations
- f = Function to be transformed
- p = A function in \mathcal{P}
- h = Result of the anti-unification of f and p
- f' = f rewritten as an instance of p
- a_i = Argument of f or p being recursed over
- v_{a_i} = Variable declared in a_i
- t, t_i = Terms for anti-unification
- σ, σ_i = Substitutions for anti-unification
 - ε = The identity substitution
 - α = Hedge variables, found only in h

Figure 4: Key to terms and notation.

These higher-order functions can be automatically introduced for suitable Haskell 98 data types, using e.g. the Haskell type class Functor for map pattern instances. We assume they are supplied in a library such as the Haskell standard Prelude. Here, iterate creates an n-element list where the first item is calculated by applying the function f to the second argument, x; the second item is computed by applying **f** to the previous result and so on. The example on Line 3 shows how iterate can be used to generate an infinite list of incrementing integers. For convenience, we also define iterN, which takes the n^{th} element from the result of iterate. Line 7 illustrates this. map applies a given function (((+) 1), which defines the function that adds 1 to its argument) to a list of elements. Line 12 shows how map can be used to apply the function ((+) 1) to a list of integers, [1,2,3,..], to produce a list of integers where each element is incremented by one. As discussed above, foldr reduces a list by replacing (:) with a binary right associative function. The result reduces the elements of the list by applying the binary function, f, pair-wise to the elements of the list. Line 17 shows how foldr can be used to sum a list. For the rest of this paper, we refer to these higher-order functions as *patterns*, and we denote these patterns collectively by \mathcal{P} (see Figure 4).

While some of the above parallel patterns have additional requirements, such as associativity of operation for fold, proving these requirements is beyond the scope of this paper. Through the discovery of these recursion schemes, we are able to expose *components* available for potential parallelisation. Further property checking, such as those for associativity, and further transformations that adjust the configuration, such as chunking [25], may be necessary or worth-while to produce the best speedups. By exposing components for parallelism, we expose properties, such as how a data structure is traversed, and are able to effectively guide the programmer to where any extra property checking is required and where parallelism might be introduced.

3. Detecting and Refactoring Pattern Instances for Haskell

This section gives an overview of our technique. Fig. 4 provides a key to our notation. Some function **f** that matches a pattern, **p**, in the set of patterns, \mathcal{P} , is rewritten to a new function, **f**', that uses **p** and that is functionally equivalent to **f**. Our approach determines whether **f** is an instance of **p**, and if so, derives the arguments that are needed to make **f** and **p** functionally equivalent. This is achieved in two main stages: *i*) anti-unifying **f** and **p** to derive argument candidates; and *ii*) using the result of the anti-unification to determine if **f** is an instance of **p**, and if so, validating the argument candidates. Our approach compares two functions, and given a finite set of \mathcal{P} , can be applied repeatedly to discover a set of potential rewrites. When multiple rewrites are valid, we take the first, although other selection methods are also valid. We give our assumptions in Sec. 3.1; we define a specialised anti-unification algorithm in Sec. 3.2; and define the properties to check whether **f** can be rewritten as a call to **p**. We give definitions for all our concepts, and provide a proof of soundness for our anti-unification algorithm.

3.1. Preliminaries and Assumptions

We illustrate our approach using the Programatica AST representation of the Haskell 98 standard [26]. Programatica is used as a basis for the Haskell refactorer, HaRe, which we extend to implement our approach. Our prototype implementation is available at https://adb23.host.cs.st-andrews.ac.uk/hare. html. Programatica represents a Haskell AST using a collection of 20 Haskell data types, comprising a total of 110 constructors [16]. These types are parameterised by the location of the syntactic elements they represent, and in the case of variables and constructors, where they are declared. For example, the expression 42 is represented in Programatica by (Exp (HsLit loc (HsInt 42))). Here, HsLit indicates that 42 is an expression; HsInt indicates that it is an integer; and loc represents its location in the source code. We refer to the representation of Haskell *expressions* in Programatica as terms; i.e. given some expression eand some term t, we say that t is the Programatica representation of e (denoted $\llbracket e \rrbracket = t$). In the above example, $\llbracket 42 \rrbracket = (\text{Exp (HsLit loc (HsInt 42))}); i.e.$ (Exp (HsLit loc (HsInt 42))) is the term that represents the expression 42. There is a one-to-one mapping between expressions and terms. In the Programatica tool set, terms are values of the HsExpI type. We make the distinction between expressions and terms since terms can be easily generalised over using their constructors, as we do in Sec. 3.2.

Although in principle we anti-unify the Haskell functions f and p, we do so in a structured way that we define in Sec. 3.2. Beyond this, we do not need to consider (the representations of) arbitrary Haskell declarations or modules since we only anti-unify terms that form the right-hand side of like-equations in f and p. Since location information will mean that any two compared terms will *always* be unequal, we *discard location information*. Finally, and to simplify our presentation, we will omit the outer Exp constructor, the common Hs prefix of constructors, and any explicit specification of literal terms (e.g. integer or string). For example, (Exp (HsLit loc (HsInt 42))) is instead recorded as (Lit 42). Where variables and constructors are both represented as identifiers, we will instead record these using Var and Con, respectively. For example, the term of the variable expression x is (Var x), and the term of the cons operator expression, (:), is (Con (:)). While our approach works for all terms, here we only need to explicitly refer to: Var (representing variables), Lit (representing literals), and App (representing application expressions). All other terms fall under the generic constructor C, and lists of terms (e.g. $[t_1, \ldots, t_n]$) that used as arguments to C to represent lists and tuples. Other common expressions found in functional languages, such as lambdas, fall under the general case since they are both unlikely to appear in recursion scheme implementations, and are otherwise safe to anti-unify due to the assumptions below.

Assumptions. In order to determine whether f is an instance of p, we will assume:

- 1. that variables are unique in f and p;
- 2. that no variables are free in p, and that all variables in p are declared as arguments to p;
- 3. that f and p recurse on the same type (denoted τ);
- 4. that **f** and **p** only pattern match on the argument that is being traversed (denoted a_i , where a_i is the *i*th argument to **f** and/or **p**);
- 5. that for every clause that matches a constructor of τ in p, there is a corresponding clause that matches the same constructor in f, and vice versa; and
- 6. that no parameter to a_i (denoted v_{a_i}) occurs as part of any binding in an as-pattern, let-expression, or where-block.

Clauses in **p** are always anti-unified with like clauses in **f**, according to the constructor of τ matched for that clause, C. a_i then acts as 'common ground' between **f** and **p**; i.e. both functions declare the parameters of the constructor in a_i as arguments. This enables a check to ensure that the inferred substitutions are *sensible* (Def. 14). It is therefore useful to know which v_{a_i} in **f** correspond to which v_{a_i} in **p**. Despite this, since variables in **f** and **p** are assumed to be unique, it follows that no variable term in **f** will ever be syntactically equal to any variable term in **p**. We instead consider v_{a_i} with the same position in **f** and **p** to be equivalent.

Definition 1 (Shared Argument Equivalence). Given the argument of type τ that is pattern matched in both **f** and **p**, a_i , where in **f**, $a_i = (D v_1 \ldots v_n)$, and in **p**, $a_i = (D w_1 \ldots w_n)$, we say that each v_i is equivalent to each w_i (denoted $v_i \equiv w_i$); i.e. $\forall i \in [1, n], v_i \equiv w_i$. Given two arbitrary v_{a_i} in **f** and **p**, $v_{a_i}^{\mathbf{f}}$ and $v_{a_i}^{\mathbf{p}}$, we denote their equivalence by $v_{a_i}^{\mathbf{f}} \equiv v_{a_i}^{\mathbf{p}}$.

In sumeuler and foldr from Sec. 1.2, for example, x and xs are v_{a_i} in sumeuler, and w and ws are similarly v_{a_i} in foldr. Here, both $x \equiv w$ and $xs \equiv ws$ hold, since x and w are the first arguments to their respective *cons* operations, and xs and ws are the second arguments.

Traditional anti-unification algorithms use syntactic equality to relate two terms, t_1 and t_2 , their anti-unifier, t, and the substitutions σ_1 and σ_2 . To take advantage of argument equivalence we must use a weaker form of syntactic equality. First, we define the binary relation \sim over terms, a form of alpha equivalence.

Definition 2 (Alpha Equivalent Terms). Given two variables, $v_{a_i}^{f}$ and $v_{a_i}^{p}$, we say that the term representations of $v_{a_i}^{f}$ and $v_{a_i}^{p}$ are equivalent (denoted $[v_{a_i}^{f}] \sim [v_{a_i}^{p}]$) when:

$$v_{a_i}^{\mathtt{f}} \equiv v_{a_i}^{\mathtt{p}}$$
Var $v_{a_i}^{\mathtt{f}} \sim \mathtt{Var} \; v_{a_i}^{\mathtt{p}}$

We then replace syntactic equality with a weaker form of equivalence using Def. 2.

Definition 3 (Syntactic Equivalence). We define syntactic equivalence to be a binary relation over two terms, t_1 and t_2 , denoted $t_1 \cong t_2$, where \cong is the reflexive structural closure of \sim .

For example, (Lit 42) \cong (Lit 42) holds; $t_1 \cong t_2$ holds for all $t_1 = (C t_{11} \ldots t_n)$ and $t_2 = (C t_{21} \ldots t_{2n})$ when $\forall i \in [1, n], t_{1i} \cong t_{2i}$; and finally, (Var $v_{a_i}^{\mathbf{f}}) \cong$ (Var $v_{a_i}^{\mathbf{p}}$) holds only when $v_{a_i}^{\mathbf{f}} \equiv v_{a_i}^{\mathbf{p}}$ is true. We note that for all other variables, v, w, (Var v) \cong (Var w) does not hold.

Where τ is a product type, e.g. as in **zipWith**, we permit pattern matching on all arguments that represent the data structure(s) being traversed. All other arguments must be declared as simple variables. For example, given:

3 f2 0 (b',b'') c [] = []

f1 is permitted, but f2 is not.

As-patterns, let-expressions, and where-blocks all enable occurrences of some v_{a_i} to be aliased, or *obfuscated*, and thereby potentially obstruct argument derivation. Since our analysis inspects the syntax of **f** and **p**, such patterns and expressions can obfuscate the exact structure of **f**; i.e. lift potentially important information out of the main body of **f** and **p**. For example, given the definition,

2

$$_{2}$$
 f3 (x:xs) = g1 x : f3 xs where g1 = g2 xs

the fact that xs is passed to g2 is obfuscated by the where-block, and could lead to an incorrect rewriting of f3 as, e.g., a map.

We do not restrict the type of recursion; general recursive forms are allowed, for example. Partial definitions are also allowed. For example, both zipWith

and $\mathtt{zipWith1}$ are valid as implementations of the general $\mathit{zipWith}$ recursion scheme:

```
zipWith g []
                     [] = []
1
   zipWith g (x:xs) (y:ys) = g x y : zipWith xs ys
2
3
   zipWith1 g1 g2 g3 []
                             [] = []
4
   zipWith1 g1 g2 g3 (x:xs) [] = g1 x : zipWith1 xs []
\mathbf{5}
   zipWith1 g1 g2 g3 []
                           (y:ys) = g2 y : zipWith1 [] ys
6
   zipWith1 g1 g2 g3 (x:xs) (y:ys) = g3 x y : zipWith1 xs ys
7
```

This permits more implementations of a given scheme, and so increases the likelihood of discovering an instance of a scheme in **f**.

Finally, and since our approach effectively requires that f has a similar syntactic structure to p as a result of the anti-unification, we permit any valid and finite normalisation procedure that rewrites some arbitrary f0 to f, such that f0 is functionally equivalent to f. Normalisation can be used, e.g., to ensure that any of the above assumptions are met. For example, consider assumption 6: as-patterns and definitions in let-expressions and where-blocks can be inlined (or unfolded in the transformational sense). For example, the where-block definition of g1 in f3 can be unfolded to produce:

```
1 f3 [] = []
```

```
2 f3 (x:xs) = g2 xs x : f3 xs
```

Alternatively, normalisation procedures can be used to *reshape* [2] functions into a form to allow, or simplify, the discovery of recursion schemes. For example, the definition,

1 f4 xs0 = case xs0 of 2 [] -> 0 3 (x:xs) -> x + f xs

can be rewritten to lift the case-split into a pattern match, allowing the function to be anti-unified against a foldr:

```
1 f4 [] = 0
2 f4 (x:xs) = x + f xs
3
4 foldr g z [] = z
5 foldr g z (x:xs) = g x (foldr g z xs)
```

More ambitious normalisation procedures may split a function, e.g. f5:

1 **f5 []** z = g2 z 2 **f5** (x:xs) z = f5 xs (g1 x z)

which can be can considered a *near fold*. Here, we can lift out the application of g2 in the base case, and therefore expose a foldl instance:

1 f5 xs z = g2 (f5' xs z)
2
3 f5' [] z = z
4 f5' (x:xs) z = f5' xs (g1 x z)

Splitting of functions in this way is allowed provided that any such splitting is *finite* and *reduces the size of the split function*. For example, some f0 cannot be split into the composition of f0' and f0'', where f0' or f0'' is functionally equivalent to the identity operation. In line with the intention for f to be antiunified against a set of recursion schemes, we do *not* require the normalisation procedure to be confluent.

3.2. Argument Derivation via Anti-Unification

In order to derive arguments to express f as an instance of p, we must anti-unify f and p. We define anti-unification for two levels: *i*) at the term level for two arbitrary terms; and *ii*) at the function level where we anti-unify f and p. At the term level, the anti-unification of two terms, t_1 and t_2 , (denoted $t_1 \triangleq t_2$) we obtain the triple $(t \times \sigma_1 \times \sigma_2)$, where t is the anti-unifier with respect to the substitutions σ_1 and σ_2 . At the function level, the anti-unification of f and p produces a list of triples that represents the results of anti-unification for each pair of clauses in f and p.

We define a substitution to be a function that takes a variable, x, and returns a term, t_2 . When applied to a term, t_1 , a substitution returns t_1 but with all occurrences of x replaced by t_2 .

Definition 4 (Substitution). Given two terms, t_1, t_2 , and a variable name, x, that occurs in t_1 , substituting x for t_2 in t_1 replaces all occurrences of (Var x) in t_1 with t_2 . This substitution is denoted $(x \mapsto t_2)$; and the application of substitutions to terms is denoted $t_1 (x \mapsto t_2)$, where $(x \mapsto t_2)$ is applied to t_1 . We use σ to refer to substitutions; e.g. $\sigma = (x \mapsto t_2)$.

Here, we update the classical definition of substitution [15] to account for the Programatica representation. Substitutions are denoted as substituting a variable name for some term. When applied to a term, a substitution replaces all variable terms that are parameterised by the given variable name with another term. For example, the result of the substitution σ applied the term t, t σ , is:

$$\begin{split} t &= \texttt{Var g} \\ \sigma &= (\texttt{g} \mapsto (\texttt{App} \; (\texttt{App} \; (\texttt{Var } \; .) \; (\texttt{Var +})) \; (\texttt{Var euler}))) \\ t \; \sigma &= (\texttt{App} \; (\texttt{App} \; (\texttt{Var } \; .) \; (\texttt{Var +})) \; (\texttt{Var euler})) \end{split}$$

Here, g is substituted for a term representing the expression ((+) . euler), from sumeuler. While the *name* of the variable being substituted is given in σ , it is the variable term parameterised by g, i.e. (Var g), in t that is substituted. We call the variables in substitutions that are substituted for terms *hedge variables*. Uniqueness of variables means that all occurrences of a hedge variable in a term can be substituted safely.

Definition 5 (Identity Substitution). The identity substitution, denoted ε , is defined such that for all terms t, $t = t \varepsilon$.

Substitutions can be composed using an n-ary relation such that composition forms a rose tree of substitutions.

Definition 6 (Substitution Composition). For all terms t_0, \ldots, t_n , and for all substitutions $\sigma_1, \ldots, \sigma_n$, where $t_0 = (C t_1 \ldots t_n)$, there exists some substitution $\sigma_0 = (\sigma_1, \ldots, \sigma_n)$ such that $t_0 \sigma_0 = (C (t_1 \sigma_1) \ldots (t_n \sigma_n))$. Similarly, where $t_0 = [t_1, \ldots, t_n]$, there exists some substitution $\sigma_0 = (\sigma_1, \ldots, \sigma_n)$ such that $t_0 \sigma_0 = [(t_1 \sigma_1), \ldots, (t_n \sigma_n)]$.

For the function f, defined

$$f v_{11} \dots v_{1n} = e_1$$
$$\vdots$$
$$f v_{21} \dots v_{2n} = e_n$$

and for the substitutions $\sigma_1, \ldots, \sigma_n$, there exists some substitution

$$\sigma_0 = (\!(\sigma_1, \ldots, \sigma_n)\!)$$

such that $\llbracket f \rrbracket \sigma_0$ is defined:

$$f v_{11} \dots v_{1n} = e'_1$$

$$\vdots$$

$$f v_{21} \dots v_{2n} = e'_n$$

$$i \in [1, n] \quad \llbracket e' \rrbracket = \llbracket e : \rrbracket =$$

where

Composed substitutions are *ordered*, such that when applied to some term
$$t_0$$
, the first composed substitution is applied to the first direct subterm to t_0 , the second composed substitution is applied to the second direct subterm, and so on.
Each composed substitution is only applied to its corresponding direct subterm in t_0 , and does not interfere with any other subterm of t_0 . For example, the expression ($\mathbf{g} \times \mathbf{x}$) is represented by the term t_0 .

$$t_0 = \text{App}(\text{App}(\text{Var g})(\text{Var x}))(\text{Var x})$$

and given the substitution, σ_0 ,

$$\sigma_0 = (((\mathbf{g} \mapsto \mathsf{Var euler}), \varepsilon), (\mathbf{x} \mapsto \mathsf{Var p}))$$

when we apply σ_0 to t_0 , the resulting term is:

$$t_0 \sigma_0 = \text{App} (\text{App} (\text{Var euler}) (\text{Var x})) (\text{Var p})$$

A composition of substitutions may be applied to either f or p, where each substitution applies to the right-hand side of each equation in order. We will refer to such compositions by σ_f and σ_p . For example, given the function sum,

 $1 \quad sum [] = []$

 $_2$ sum (x:xs) = x + (sum xs)

and the substitution $\sigma_{\mathbf{f}} = (\sigma_1, \sigma_2)$, applying $\sigma_{\mathbf{f}}$ to sum, applies σ_1 to the term representation of [], i.e. the right-hand side of the first equation, and σ_2 to the term representation of $(\mathbf{x} + (\mathbf{sum xs}))$, i.e. the right-hand side of the second equation.

Anti-unification computes the *anti-unifier* of two terms. In the literature, all hedge variables are *fresh*. Conversely, and since we aim to rewrite **f** in terms of **p**, we can use the variables declared in **p** as hedge variables. This allows us to easily derive expressions that can be passed to **p** in **f**. We refer to the set of hedge variables from **p** by \mathcal{V}_p . All hedge variables not in \mathcal{V}_p are fresh hedge variables, which we denote by α .

Another difference to traditional anti-unification algorithms can be found the concept of *unobstructive recursive prefixes*. As we are anti-unifying two different recursive functions, by definition they will diverge in at least one respect: their recursive calls. We extend this notion to *recursive prefixes*, where a recursive prefix is a variable expression comprising the name of the function, i.e. **f** or **p**, or an application expression that applies **f** or **p** to one or more expressions. For example, given the definition of **foldr**,

```
<sup>1</sup> foldr g z [] = z
```

² foldr g z (x:xs) = g x (foldr g z xs)

the recursive call (foldr g z xs), has four recursive prefixes: *i*) (foldr g z xs); *ii*) (foldr g z); *iii*) (foldr g); and *iv*) foldr. It is useful to further extend the notion of recursive prefixes with the concept of *unobstructiveness*. We define an *unobstructive recursive prefix* to be a recursive prefix for which the values of its arguments *do not change*. In the above example, all recursive prefixes, with the exception of (foldr g z xs), are *unobstructive*.

Definition 7 (Unobstructive Recursive Prefix). Given a recursive function f,

 $f v_1 \ldots v_n = e$

for all recursive calls in e,

$$f e_1 \ldots e_n$$

we say that a recursive prefix is either:

- 1. the direct subexpression $f e_1 \ldots e_m$, where $1 \le m \le n$; or
- 2. the expression f itself.

We say that a recursive prefix is unobstructive when $\forall i \in [1, m]$, $e_i = v_i$. An expression, e' is denoted as being an unobstructive recursive prefix by membership of the set of unobstructive recursive prefixes for f; i.e. $e' \in \mathcal{R}_f$.

To help ensure that as many instances of some scheme are found, we will also assume the existence of the identity operator, which is defined in Haskell as id. **Definition 8** (The Identity Operation). We define id to be the lambda expression $(\langle \mathbf{x} \rangle \mathbf{x})$ such that for all terms, t, $(App (Var id)(t)) \cong t$.

Our anti-unification algorithm calculates the anti-unifier t and substitutions σ_1 and σ_2 for the terms t_1 and t_2 , such that $t_1 \cong t \sigma_1$ and $t_2 \cong t \sigma_2$ hold, denoted:

$$t_1 \triangleq t_2 = t \times \sigma_1 \times \sigma_2$$

Our rules in Fig 5 define how to infer t, σ_1 , and σ_2 from t_1 and t_2 . Rules have the form,

$$\frac{p}{\left[\begin{array}{c}t_1\\t_2\end{array}\right] \cong (t)\left[\begin{array}{c}\sigma_1\\\sigma_2\end{array}\right]}$$

where p denotes any additional assumptions necessary for $t_1 \cong t \sigma_1$ and $t_2 \cong t \sigma_2$ to hold. We use matrix notation as shorthand for the two equations; i.e.

$$\left[\begin{array}{c}t_1\\t_2\end{array}\right]\cong(t)\left[\begin{array}{c}\sigma_1\\\sigma_2\end{array}\right]=t_1\cong t\,\sigma_1\wedge t_2\cong t\,\sigma_2$$

When t_1 and t_2 are the same, i.e. $t_1 \cong t_2$, EQ holds. For all other cases when t_2 is a variable term, and is not (Var p), t_2 is used as the hedge variable in t (VAR). When t_2 is an application term applying some function u to a v_{a_i} , and t_2 is a variable term that is equivalent to the v_{a_i} in t_2 , then the application term is used in t, and u is substituted for id in σ_1 (ID). When t_2 is an unobstructive recursive prefix of p, RP holds. CONST states that when t_1 and t_2 are terms with the same constructor, C, then t is a term with constructor C and its direct subterms are the anti-unifiers of each of the respective direct subterms in t_1 and t_2 . LIST is similar to CONST, but satisfies when the t_1 and t_2 are lists of terms, where t_1 and t_2 are the same length. In all other cases, OTHERWISE holds. For example, the terms, t_1 and t_2 ,

$$t_1 = \operatorname{App}(\operatorname{Var} e) (\operatorname{Lit} 42)$$

 $t_2 = \operatorname{App}(\operatorname{Var} g) (\operatorname{Lit} 108)$

both apply some function to some literal, and have the anti-unifier and substitutions:

$$t = (\operatorname{App} (\operatorname{Var} g) (\operatorname{Var} \alpha))$$

$$\sigma_1 = ((g \mapsto (\operatorname{Var} e)), (\alpha \mapsto (\operatorname{Lit} 42)))$$

$$\sigma_2 = (\varepsilon, (\alpha \mapsto (\operatorname{Lit} 108)))$$

Here, σ_1 and σ_2 are compositions reflecting the two arguments to App (CONST). Since the first argument in both t_1 and t_2 are variable terms, but with different names, t_2 is chosen as the anti-unifier with the substitution for σ_1 replacing g with e, and the identity substitution for σ_2 (VAR). Conversely, the second argument differs in t_1 and t_2 , and since (Lit 108) is not an identifier, a fresh hedge variable, α , is used for t, with the corresponding substitutions replacing (Var α) in (Lit 42) and (Lit 108) for t_1 and t_2 , respectively (OTHERWISE).

Figure 5: Inference rules to calculate the anti-unifier t for the terms t_1 and t_2 .

As we are interested in finding instances of a pattern in a function, it is convenient to relax the *least general* property found in traditional anti-unification algorithms. Specifically, the RC rule in Fig. 5 can result in an anti-unifier that is not the least general generalisation of two terms. To illustrate this, consider the example of elem anti-unified against a foldr, where

```
1 elem a [] = False
2 elem a (x:xs) =
3 (\y ys \rightarrow if a == y then True else ys) x (elem a xs)
4 5 foldr g z [] = z
6 foldr g z (x:xs) = g x (foldr g z xs)
```

Here, the anti-unification of the recursive calls in the *cons*-clauses produces the term:

App (Var α) (Var xs)

since (foldr g z) is an unobstructive recursive prefix. In this case, the RC rule applies instead of CONST. CONST would otherwise apply since both (foldr g z) and (elem a) are application expressions, and would produce a less general generalisation than the result of applying the RC rule. Intuitively, the least general generalisation is the term t that shares the closest structure to t_1 and t_2 , and requires the least number of substitutions in σ_1 and σ_2 . In the above example, the least general generalisation for the recursive call is:

App
$$(\text{App} (\text{Var } \alpha) (\text{Var } z)) (\text{Var } xs)$$

The 'shortcuts' produced by application of the RC rule are *useful*. Unobstructive recursive prefixes are uninteresting, aside from their relative location, since they produce no valid *argument candidates* in σ_{f} (Sec. 3.3). Moreover, without the RC rule, extra work would be necessary to derive and/or choose between argument candidates.

Theorem 1 (Soundness of Anti-Unification Algorithm). Given the terms t_1 and t_2 , we can find t, σ_1 , and σ_2 , such that $t_1 \cong t \sigma_1$ and $t_2 \cong t \sigma_2$ hold.

We give the proof for the above soundness property in Appendix A.

Having defined anti-unification for terms, we can now define it for f and p. Given the functions f and p,

$$\begin{array}{cccc} \mathbf{f} \ v_{11} \ \dots \ v_{1m} = e_1^{\mathbf{f}} & & \mathbf{p} \ w_{11} \ \dots \ w_{1l} = e_1^{\mathbf{p}} \\ & & \vdots & & \vdots \\ \mathbf{f} \ v_{n1} \ \dots \ v_{nm} = e_n^{\mathbf{f}} & & & \mathbf{p} \ w_{n1} \ \dots \ w_{nl} = e_n^{\mathbf{p}} \end{array}$$

where v_{ij} and w_{ij} are arguments to **f** and **p**, respectively, and we define the anti-unification of **f** and **p**, denoted **h**, to be the list:

$$\mathbf{h} = \begin{bmatrix} [e_1^{\mathtt{f}}]] \triangleq [[e_1^{\mathtt{p}}]], \dots, [[e_n^{\mathtt{f}}]] \triangleq [[e_n^{\mathtt{p}}]] \end{bmatrix}$$

where each element, $\forall i \in [1, n], e_i^{h}$, in **h** corresponds to the anti-unification of the *i*th clauses in **f** and **p**; and elements are the triple $(t_i \times \sigma_{1i} \times \sigma_{2i})$. The substitutions for each element in **h** can be composed, and we refer to the composition of substitutions for **f** as $\sigma_{\mathbf{f}}$, and the composition of substitutions for **p** as $\sigma_{\mathbf{p}}$; i.e.

$$\sigma_{\mathbf{f}} = (\!(\sigma_{11}, \dots, \sigma_{1n})\!)$$
$$\sigma_{\mathbf{p}} = (\!(\sigma_{21}, \dots, \sigma_{2n})\!)$$

h is, in principle, equivalent to a function with n equations and where: i) the anti-unifier term in each element represents the right-hand side of its respective equation; and ii) the set of all hedge variables are the parameters to the function. For example, the functions **f** and **p**,

when anti-unified, produce:

$$\begin{split} \mathbf{h} &= \left[(\operatorname{App} \left(\operatorname{App} \left(\operatorname{Var} \, \mathbf{y} \right) \right) \left(\operatorname{Var} \, \mathbf{z} \right) \right) \\ &\times \left(\left(\varepsilon, (\mathbf{y} \mapsto (\operatorname{Var} \, \mathbf{x})) \right), (\mathbf{z} \mapsto (\operatorname{Id} 42)) \right) \times \left(\left(\varepsilon, \varepsilon \right), (\mathbf{z} \mapsto (\operatorname{Id} 108)) \right) \right] \end{split}$$

Here, h can be represented as:

where, h is the anti-unifier, and the substitutions are represented using standard function application. To simplify our presentation, in examples we confuse h for its equivalent function definition.

3.3. Deriving Pattern Arguments

Given the anti-unifier **h** of **f** and **p**, with substitutions $\sigma_{\mathbf{f}}$ and $\sigma_{\mathbf{p}}$ respectively, we next derive the arguments to **p** that are needed to rewrite **f** as an instance of **p**. Our anti-unification algorithm is designed such that σ_1 provides *candidate* arguments. These candidates are considered *valid* when they adhere to three properties:

- 1. that **p** and **h** are equivalent (Def. 10);
- 2. that there does not exist a (sub-)substitution in $\sigma_{\mathbf{f}}$ or $\sigma_{\mathbf{p}}$ where a v_{a_i} occurs as either a hedge variable or as a term (Def. 12); and
- 3. that all substitutions in σ_{f} substituting for the same hedge variable that is derived from p must substitute for the same term (Def. 14).

Equivalence of Pattern and Anti-Unifier. Recall that we aim to rewrite f in terms of p. As stated in Sec. 3.2, the syntactic equivalence properties of the produced anti-unifier allow f to be rewritten as a call to the function representation of h. Furthermore, by equational reasoning [27], if p and h are equivalent, then f can be rewritten to replace the call to h with a call to p; i.e. f is an instance of p. Since anti-unification will always produce an anti-unifier and substitutions between any two arbitrary terms, the production of an anti-unifier cannot be used as a test of equivalence. We instead define an equivalence relation between p, h, σ_f , and σ_p . In order to do this, we first define equivalence between two terms t_2 and t and substitutions σ_1 and σ_2 with respect to t_1 .

Definition 9 (Equivalence of Terms with Substitutions). Given the terms, t, t_1 , t_2 , and the substitutions σ_1 and σ_2 , such that

$$\left[\begin{array}{c} t_1 \\ t_2 \end{array}\right] \cong (t) \left[\begin{array}{c} \sigma_1 \\ \sigma_2 \end{array}\right]$$

we say that t_2 and t are equivalent with respect to σ_1 , σ_2 , and t_1 , (denoted $t_2 \equiv_{t_1} t \times \sigma_1 \times \sigma_2$) when:

$$\begin{array}{c} \underbrace{t_2 \cong t}_{\text{EQ} \quad t_2 \equiv_{t_1} t \times \varepsilon \times \varepsilon} \quad \underset{\text{HEDGE} \quad v \neq \texttt{p} \quad v \in \mathcal{V}_\texttt{p}}{(\text{Var } v) \equiv_{t_1} (\text{Var } v) \times (v \mapsto t_1) \times \varepsilon} \\ \\ \underbrace{t_1 \in \mathcal{R}_\texttt{f} \quad t_2 \in \mathcal{R}_\texttt{p} \quad \alpha \notin \mathcal{V}_\texttt{p}}_{\text{FRESH} \quad \underline{t_2 \equiv_{t_1} (\text{Var } \alpha) \times (\alpha \mapsto t_1) \times (\alpha \mapsto t_2)}} \end{array}$$

$$\underbrace{\forall i \in [1, n], t_{2i} \equiv_{t_1} t_i \times \sigma_{1i} \times \sigma_{2i}}_{(C \ t_{21} \ \dots \ t_{2n}) \equiv_{t_1} (C \ t_1 \ \dots \ t_n) \times (\sigma_{11}, \dots, \sigma_{1n}) \times (\sigma_{21}, \dots, \sigma_{2n}) }$$

$$\underbrace{\forall i \in [1, n], t_{2i} \equiv_{t_1} t_i \times \sigma_{1i} \times \sigma_{2i}}_{[t_{21}, \dots, t_{2n}] \equiv_{t_1} [t_1, \dots, t_n] \times (\sigma_{11}, \dots, \sigma_{1n}) \times (\sigma_{21}, \dots, \sigma_{2n})}$$

Here, EQ states that when t_2 and t are the same, and when σ_1 and σ_2 are both ε , then t_2 and t are equivalent with respect to σ_1 and σ_2 . When t_2 and t are both variable terms, t_2 and t are equivalent when the variables have the same name and when the the variable is a hedge variable derived from p (HEDGE). Similarly, when t_2 and t are both fresh hedge variables, α , then both terms substituted for α must be unobstructive recursive prefixes in their respective functions (FRESH). Finally, CONST and LIST state that terms consisting only of subterms are equivalent when all respective subterms are equivalent. The definition of equivalence for **p** and **h** then follows naturally.

Definition 10 (Equivalence of Pattern and Anti-Unifier). Given two functions \mathbf{f} and \mathbf{p} , and the result of their anti-unification, \mathbf{h} , containing the substitutions $\sigma_{\mathbf{f}}$ and $\sigma_{\mathbf{p}}$, we say that \mathbf{p} and \mathbf{h} are equivalent when for all clauses in \mathbf{p} , $e_i^{\mathbf{p}}$, and elements in \mathbf{h} , $e_i^{\mathbf{h}} = (t_i \times \sigma_{1i} \times \sigma_{2i})$, $[\![e_i^{\mathbf{p}}]\!] \equiv_{t_{1i}} e_i^{\mathbf{h}}$.

Absence of Shared Arguments in Substitutions. As stated in Sec. 3.1, the variables, v_{a_i} , that are declared in a_i are shared between **f** and **p**. As such, if **f** is an instance of **p**, v_{a_i} must be used in **f** in an equivalent way to their use in **p**; i.e. their relative locations in the structure of **f** and **p** must be the same. For example, in sumeuler and foldr,

```
sumeuler [] = 0
sumeuler (x:xs) = ((+) . euler) x (sumeuler xs)
foldr g z [] = z
foldr g z (w:ws) = g w (foldr g z ws)
```



the heads of the matched lists, **x** in sumeuler and **w** in foldr, are both passed as the first argument to some function, **g**; and the tails of the matched lists, **xs** and **ws** respectively, are passed as the last argument to the recursive calls in **f** and **p**. While the equivalence of pattern and anti-unifier (Def. 10) ensures the structural equivalence of **p** and **h**, it does not inspect hedge variables derived from **p**. Our anti-unification rules in Fig. 5 means that no v_{a_i} will occur as either a hedge variable or as a subterm in either $\sigma_{\mathbf{f}}$ or $\sigma_{\mathbf{p}}$ when all equivalent v_{a_i} have the same relative positions in **f** and **p**. Conversely, the occurrence of some v_{a_i} in $\sigma_{\mathbf{f}}$ or $\sigma_{\mathbf{p}}$ indicates that not all equivalent v_{a_i} have the same relative positions in **f** and **p**.

Definition 11 (Variable Occurrence in Terms). For all variables v, and for all terms, t, we say that v occurs in t (denoted $v \ll t$) when t = (Var v) or when there exists a subterm, t_i , in t such that $t_i = (Var v)$.

Definition 12 (Shared Argument Absence). Given the functions \mathbf{f} and \mathbf{p} , and their anti-unification \mathbf{h} , containing the substitutions $\sigma_{\mathbf{f}}$ and $\sigma_{\mathbf{p}}$, we say that no v_{a_i} occurs in $\sigma_{\mathbf{f}}$ or $\sigma_{\mathbf{p}}$ (denoted $v_{a_i} \notin (\sigma_{\mathbf{f}}, \sigma_{\mathbf{p}})$) when, for all v_{a_i} :

$$\begin{array}{c|c} v \not\cong v_{a_i} & (\operatorname{Var} v_{a_i}) \not\ll t \\ \hline v_{a_i} \not\in (v \mapsto t) & v_{a_i} \not\in (\sigma_1, \dots, \sigma_n) \end{array}$$

Substitution Uniqueness. Substitution composition (Def. 6) ensures that the same hedge variable can be used to substitute for different terms without the problem of substitution interference. For example, consider some **f** that is anti-unified against **scanl**:

```
1 f a [] = [a]
2 f a (x:xs) = (+) a x : f ((-) a x) xs
3
4 scanl g z [] = [z]
5 scanl g z (x:xs) = g z x : scanl g (g z x) xs
```

Here, the result of the anti-unification of the two *cons*-clauses will include substitutions where g is substituted for both (+) and (-) in σ_{f} . In scanl, however, g is the same in both instances. We therefore also require that all substitutions that substitute for the same hedge variable in σ_{f} are substituted for the same term.

Definition 13 (Substitution Flattening). For all substitutions, σ , we say the flattening of σ (denoted $\lfloor \sigma \rfloor$) is the list of substitutions, where:

$$\lfloor (v \mapsto t) \rfloor = [(v \mapsto t)] \qquad - \lfloor (\sigma_1, \dots, \sigma_n) \rfloor = \lfloor \sigma_1 \rfloor + \dots + \lfloor \sigma_n \rfloor$$

Here, + appends two lists, and [a] denotes the singleton list.

A list of substitutions cannot be applied to a term, allowing us to safely expose any potential substitution interference across f. **Definition 14** (Substitution Uniqueness). Given the functions f and p, and their anti-unification h. Given that σ_{f} is the composition of substitutions in h for the clauses of f. We say that substitutions are unique in σ_{f} when for all hedge variables, $v \in \mathcal{V}_{p}$, and for all pairs of substitutions in $\lfloor \sigma_{f} \rfloor$, $(v \mapsto t_{1})$ and $(v \mapsto t_{2})$, $t_{1} = t_{2}$ holds.

Sufficiency of Validity Properties. Given f, p, h, σ_{f} and σ_{p} , the above three properties are sufficient to determine whether the candidate arguments in σ_{f} are valid. Equivalence of p and h (Def. 10) ensures that for all clauses in p and **h**, the terms for that clause, t_2 and t, are syntactically equivalent (Def. 3). Hedge variable terms have additional information in their respective substitutions, σ_1 and σ_2 . There are two cases: fresh hedge variables, α , and hedge variables derived from \mathbf{p} , v. For α , Def. 10 requires that both terms, t_1 and t_2 , substituted for α are unobstructive recursive prefixes (Def. 7). No other terms are allowed. For v, two cases are possible: $v = v_{a_i}$ and otherwise. All equivalent v_{a_i} in f and p must have the same relative locations to ensure that a_i is traversed in the same way, and that no free variables occur in any argument candidate expression. The rules in Fig. 5 mean that the identity substitution is derived for all equivalent v_{a_i} in the same relative position in f and p. For the case when a v_{a_i} is not anti-unified with its equivalent, Shared Argument Absence (Def. 12) requires that no v_{a_i} occurs in any substitution and so will fail in this case. For all other v, Def. 10 requires that v will be in scope, due to the assumption that all variables that occur in p are in scope and are declared as parameters of p. Any updates to the value of v must be reflected in p, e.g. such as in fold1 and scan1. Finally, and since each argument may be instantiated only once for each call to p, Substitution Uniqueness (Def. 14) ensures that for all substitutions that substitute for the same hedge variable, the substituted terms are the same.

3.4. Deriving Arguments from Substitutions

Given that f, p, and their anti-unification, h, adhere to Def. 10 (Equivalence of Pattern and Anti-Unifier), Def. 12 (Shared Argument Absence), and Def. 14 (Substitution Uniqueness), the arguments for p can be directly obtained from σ_f . For sumeuler, given σ_1 for the [] clause:

$$\sigma_1 = (\mathsf{z} \mapsto (\mathsf{Lit} \ 0))$$

and σ_1 for the (:) clause:

$$\begin{split} \sigma_1 = ((g \mapsto (\operatorname{App} (\operatorname{App} (\operatorname{Var} \, . \,) (\operatorname{Var} \, +)) (\operatorname{Var} \, \operatorname{euler}))), \varepsilon) , \\ ((\alpha \mapsto (\operatorname{Var} \, \operatorname{sumeuler})), \varepsilon)) \end{split}$$

((+). euler) is passed as g; 0 is passed as z; and a_i (i.e. (x:xs)) is passed as itself due to Shared Argument Absence and Argument Equivalence properties. Substitutions with fresh hedge variables are discarded. We can now refactor f in terms of p using the derived arguments above, to give:

sumeuler xs = foldr ((+) . euler) 0 xs

Which, as before, can be rewritten to give our original definition:

```
sumeuler xs = foldr (+) 0 (map euler xs)
```

 $i/\partial Maximum$ Segment Sum. Where sumeuler is a simple foldr, consider the less obvious example of maximum segment sum, as defined by Kannan [7]:

```
1 mss xs0 = mss2 0 0 xs0
2
3 mss2 y z [] = z
4 mss2 y z (x:xs) = mss2 (max y (x+z)) (max x (x+z)) xs
```

Here, mss2 traverses over the list xs0. Since mss2 does not return a list, we instead consider a fold pattern. Two possibilities are available: a foldr or a fold1. The standard Prelude definitions of these functions take only one initial/accumulator argument, and neither will produce a rewritten mss2. To solve this, y and z in mss2 may be tupled as part of a normalisation process:

```
1 mss2 y z xs0 = fst (mss2' (y,z) xs0)
2
3 mss2' w [] = w
4 mss2' w (x:xs) =
5 mss2' ((\(y1,z1) x -> (max y1 (x1+z1), max x1 (x1+z1))) w x) xs
Will w i: if the integral of the provided of the integral of the provided of the integral of the provided of the provided
```

When anti-unified against foldr and foldl, mss2' will produce the following anti-unifiers:

```
1 hr g z [] = z

2 hr g z (x:xs) = g x \alpha

3

4 hl g z [] = z

5 hl g z (x:xs) = \alpha (g z x) xs
```

where hr is the result of anti-unifying mss2' and foldr, and hl is the result of anti-unifying mss2' and foldl. For hr, z is substituted for w, g is substituted for mss2', x is substituted for

```
1 (\(y1,z1) x1 -> (max y1 (x1+z1), max x1 (x1+z1))) w x)
```

and α is substituted for xs in mss2' and (foldr g z xs) in foldr. hr will not pass the validation properties, since: *i*) x and xs (i.e. v_{a_i}) both occur in substitutions; and *ii*) hr is not equivalent to foldr, in part because α is substituted for (foldr g z xs) which is not an unobstructive recursive prefix. Conversely, an instance of foldl can be found, and f rewritten:

1 mss2' w xs0 = foldl ($(y,z) \times \rightarrow$ (max y (x+z), max x (x+z))) w xs0

An alternative and equally valid approach can be to anti-unify against a foldl that takes two accumulative variables; e.g.:

```
foldl g1 g2 y z [] = y
1
   foldl g1 g2 y z (x:xs) = foldl g1 g2 (g1 y z x) (g2 y z x) xs
2
   When anti-unified against mss2' in the form:
   mss2' y z [] = y
1
   mss2' y z (x:xs) =
2
     mss2' ((\y1 z1 x1 -> max y1 (x1+z1)) y z x)
3
            ((\y1 z1 x1 -> max x1 (x1+z1)) y z x)
4
            xs
\mathbf{5}
   the following anti-unifier is produced, and mss2' is rewritten:
   h g1 g2 y z [] = y
1
   h g1 g2 y z (x:xs) =
2
     \alpha (g1 y z x) (g2 y z x) xs
3
4
   mss2' y z xs0 =
\mathbf{5}
     foldl (\y1 z1 x1 -> max y1 (x1+z1))
6
            (\y1 z1 x1 -> max x1 (x1+z1))
7
            xs0
8
```

where α is substituted for mss2' and (foldl g1 g2) in mss2' and foldl, respectively. This example demonstrates the increased importance of normalisation as functions become more complicated; a limitation we will explore in future work.

Termination of Approach. As an additional result, we observe that our approach will always terminate, given our assumptions, that the definitions of f and p are themselves finite, and \mathcal{P} contains a finite number of scheme implementations. Our first-order anti-unification algorithm can be represented as a fold over two finite terms, and so will both always terminate and produce a finite result. Similarly, our validation properties will always terminate, since they too can be represented as a fold over finite substitutions and function definitions. Should the validation properties *not* hold for a given **f** and **p**, then the process terminates with no rewriting. Conversely, when the validation properties hold for a given f and p, the selection of arguments to p and rewriting of f will both always terminate. The selection of arguments can be represented as a search for a particular node of a finite tree. Similarly, the rewrite always constructs a single clause function whose body is an application expression that applies p to a finite number of arguments. Should our approach be extended by a normalisation procedure, as suggested in Sec. 3.1, we require and assume that the normalisation procedure is both finite and valid. Therefore, whether our approach is extended by a normalisation procedure or not, our approach will always terminate.

n	MUT	SD	Total	SD	Sparks	Heap	Residency
1000	0.06	0.02	0.06	0.02	3	67795856	68256
5000	1.29	0.01	1.31	0.01	11	1958285432	214792
10000	5.34	0.03	5.39	0.03	21	8296937744	224584
15000	12.43	0.03	12.54	0.03	31	19280868336	498624
20000	22.82	0.08	23.03	0.07	41	35050884184	761720
25000	36.29	0.03	36.6	0.04	51	55705003000	1024160
30000	53.24	0.18	53.7	0.18	61	81319554136	1228304
35000	73.36	0.3	73.99	0.3	71	111956760792	1428304
40000	96.77	0.04	97.6	0.04	81	147666965368	1612232
45000	123.98	0.13	125.04	0.13	91	188497463480	1612232
50000	154.33	0.25	155.65	0.26	101	234487830776	1778184

Figure 6: Sequential mutator (MUT) and total (Total) times in seconds (with standard deviations), total number of sparks for parallel version, and heap allocation and residency in bytes for sumeuler on *corryvreckan*.

4. Examples

We have applied our approach to four examples from the *NoFib* suite [9] of Haskell benchmarks that are known to have *parallel* implementations. For each example we provide: performance and speedup results on our 28-core testbed machine; a brief description of the size and structure of the program; the number of explicit patterns that it contains; and a description of any other interesting functions/patterns. All four examples have been anti-unified against relevant map/fold patterns using our prototype anti-unification implementation.

Experimental Setup. All our examples have been executed on our experimental testbed machine, corryvreckan, a 2.6GHz Intel Xeon E5-2690 v4 machine with 28 physical cores and 256GB of RAM. This machine allows turbo boost up to 3.6GHz, and supports automatic dynamic frequency scaling. Using built-in hyper-threading support, it is possible to scale to 56 virtual cores, but this does not always give performance improvements. All four examples have been compiled using GHC 7.6.3 on Scientific Linux version 3.10.0. The original source programs have been taken from the repository at git.haskell.org/nofib.git (commit ce4b36b). We have not changed this code, apart from adding wrappers to aid profiling where this is necessary. Our code is available at https://adb23. host.cs.st-andrews.ac.uk/nofib-parallel.zip. We compiled the examples using -0, -feager-blackholing, -threaded, and -rtsopts flags, which we found gave the best general performance. Our measurements represent the average of three runs and are taken from the -s Haskell RTS option, which records execution statistics. Timing information is taken from two sources: mutator time (MUT) and total time (Total), both in seconds. The mutator time represents the CPU time that is spent purely on executing the program, and is thus a measure of pure parallel performance. The total time is the elapsed or



Figure 7: Speedups for sumeuler on *corryvreckan* using reported mutator (MUT, reduplicated from Sec. 1.2) times. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

real-time for the program as a whole, and also includes the additional system runtime overheads of initialisation, finalisation and garbage collection time.

4.1. Sumeuler

As described in Sec. 1.2, sumeuler calculates Euler's totient function for a list of integers and sums the results. In the NoFib suite, sumeuler consists of three Haskell files: ListAux (41 lines), SumEulerPrimes (36 lines), and SumEuler (290 lines). Collectively, these introduce 31 functions, 12 of which have explicit maps or folds as part of their definitions. Our prototype implementation rediscovers all the implicit maps and folds. One interesting case is primesIn:

primesIn is a foldr over an *infinite* list when the standard Prelude definition of foldr does not define behaviour for the empty list.

We executed sumeuler for n = 1000, 5000 and between 10000 and 50000 at intervals of 5000, with a chunk size of 500. Fig. 6 gives average sequential times (with standard deviations), the number of sparks, heap allocation, and residency for all n. All sparks are converted for all n. Fig. 7 and Fig. 8 give speedups for sumeuler using mutator and total time, achieving maximum speedups of 30.50 for n = 50000 on 48 virtual cores (mutator) and 20.92 for n = 50000 on virtual 48 cores (total). This clearly shows that sumeuler scales both with n and with



Figure 8: Speedups for **sumeuler** on *corryvreckan* using reported total (Total) times. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

n	MUT	SD	Total	SD	Sparks	Heap	Residency
11	0.1	0.03	0.1	0.03	101	95722624	64088
12	0.42	0.08	0.43	0.08	122	569715376	1693112
13	2.13	0.02	2.19	0.03	145	3590068664	9854376
14	13.65	0	14.1	0.01	170	24108663032	59041312
15	96.36	0.17	103.11	0.18	197	171560553832	479706512
16	717.66	0.81	821.03	0.35	226	1293635766936	3097146864

Figure 9: Sequential mutator (MUT) and total (Total) times in seconds (with standard deviations), total number of sparks for parallel version, and heap allocation and residency in bytes, for queens on *corryvreckan*.

the number of cores. Total time speedups are more likely to *reduce* as further cores are added for a given n, since garbage collection time *increases* with more cores. The irregularity of the tasks in **sumeuler** is also likely to have contributed to this result. For this example, all of the potential parallelism (Haskell *sparks*) was converted into actual parallelism (Haskell ultra-lightweight *threads*). In general, however, the sophisticated *evaluate-and-die* mechanism will throttle excess parallelism, giving a better load-balance, and avoiding system overload when massive amounts of parallelism are present.

4.2. N-Queens

The n-queens problem (queens) calculates the positions of n queens on a $n \times n$ chess board such that no two queens threaten each other. In the NoFib suite, queens consists of one Haskell file, Main, that is 42 lines long and defines 6



Figure 10: Speedups for queens on *corryvreckan* using reported mutator (MUT) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

functions. Two of these 6 functions have explicit maps and folds (including list comprehensions), and an explicit call to iterate. One other function, safe,

```
1 safe :: Int -> Int -> [Int] -> Bool
2 safe x d [] = True
3 safe x d (q:1) = x /= q && x /= q+d && x /= q-d && safe x (d+1) l
is an implicit foldr over lists:
1 safe :: Int -> Int -> [Int] -> Bool
2 safe x d qs = fst $ foldr f (True,d) qs
```

```
f q (l,d) = (x /= q && x /= q+d && x /= q-d && l, d+1)
```

where

з

4

Since d is updated between recursive calls, our prototype implementation will not currently detect this foldr¹. By rewriting safe as a composition of an unfold and an (implicit) foldr, our prototype can detect the foldr over the tree generated by the unfold. safe may alternatively be considered an instance of a *zygomorphism* [28], it may therefore be worth investigating this pattern as part of our future work. Another function, pargen, enumerates the different possible locations of the queens on the board.

¹This is because in the original definition of **safe**, **d** is updated between recursive calls meaning a standard **foldr** is not equivalent (Def. 10).



Figure 11: Speedups for queens on *corryvreckan* using reported total (Total) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

```
1 pargen strat n b
2 n >= t = iterate gen [b] !! (nq - n)
3 otherwise =
4 concat (map (pargen strat (n+1)) (gen [b]) `using` strat)
```

This is defined as a sub-function of the top-level queens function. Here, nq denotes the number of queens that are to be searched for, and t is a threshold for controlling parallelism. The strat parameter is an evaluation strategy: either evalList or parList depending on whether sequential or parallel operation is chosen. n is used as a measure for determining whether the threshold, t, is reached; b accumulates results; and gen generates an individual result. We can unfold both iterate and map expressions as implicit iterN and map functions that are called by pargen:

```
1
   pargen n b
      n \ge t = pgI([b],(nq-n))
2
      otherwise = concat $ pgM (gen [b])
з
       where
4
         pgI (rs, 0) = rs
5
         pgI (rs,i) = pgI ((gen rs),i-1)
6
7
         pgM [] = []
8
         pgM (r:rs) = pargen' (n+1) r : pgM rs
9
```

Our prototype correctly (re)discovered instances of both iterN and map within pgI and pgM. We executed queens for n ranging from 11 to 16, with t = 2. Fig. 9



Figure 12: Garbage collection times for **queens** on *corryvreckan*. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

gives mutator and total times (with standard deviations) for sequential runs, and number of sparks, heap allocation size, and residency for each n. An average of 30% of sparks were converted on 4 cores, 65% on 8 cores, and 85% for all other cores. Fig. 10 and 11 shows speedups for **queens** in terms of mutator and total time. We achieve maximum speedups of 22.65 for n = 16 on 48 hyper-threaded cores and 10.00 for n = 12 on 36 hyper-threaded cores. When n = 11, the mutator time plateaus quickly and sharply, possibly due to lack of work. When n = 12, speedups reduce as the number of cores is increased, probably because there is insufficient work to offset increased overheads. Otherwise, mutator times plateau above 28 cores, probably because hyper-threading is not effective in this example. Unlike our other examples, total time speedups reveal an overall halving of performance gains as a consequence of increased garbage collection time. Fig. 12 shows that when n = 15 and n = 16, garbage collection times are heavily increased. This is probably due to repeated generation of lists.

4.3. N-Body

nbody calculates the movement of bodies in *m*-dimensional space according to the physical forces between them. The classical representation of this problem is between celestial bodies, modelling the effects of gravity on their trajectories. In the NoFib suite, **nbody** consists of two Haskell files: Future (16 lines) and **nbody** (135 lines) which collectively define 8 functions. Two of these 8 functions have an explicit map or fold, and one, **loop**, is an *implicit* iterN. Our prototype discovers the maps and folds in both functions where they are explicitly used, from their inlined definitions. The **loop** sub-function of **compute** is defined as:

1 loop i ax ay az 2 i == end = (# ax,ay,az #) 3 otherwise = loop (i+1) (ax+px) (ay+py) (az+pz) 4 where

n	MUT	SD	Total	SD	Sparks	Heap	Residency
10000	2.06	0.06	2.07	0.06	40	6310152	1197488
25000	11.61	0.1	11.63	0.11	40	15670360	5216880
50000	46.93	0.13	46.98	0.13	40	31270520	12311984
75000	104.9	0.35	104.96	0.34	40	46870848	12547600
100000	187.47	0.1	187.57	0.11	40	62471232	25082544
250000	1164.34	0.68	1164.52	0.65	40	156226496	40215360

Figure 13: Sequential mutator (MUT) and total (Total) times in seconds (with standard deviations), total number of sparks for parallel version, and heap allocation and residency in bytes, for nbody on *corryvreckan*.



Figure 14: Speedups for **nbody** on *corryvreckan* using reported mutator (MUT) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

5	(x,y,z) = vector
6	<pre>(x',y',z') = vecList Array.! i</pre>
7	
8	(# dx,dy,dz #) = (# x'-x, y'-y, z'-z #)
9	eps = 0.005
10	$distanceSq = dx^2 + dy^2 + dz^2 + eps$
11	$factor = 1/sqrt(distanceSq ^ 3)$
12	(# px,py,pz #) =
13	(# factor * dx, factor * dy, factor *dz #)

where i and n control the number of iterations performed; ax, ay, az are the three-dimensional cartesian coordinates of the body being updated; and, ultimately, px, py, and pz are updates to those initial coordinates. Applying our prototype to loop, with guards converted to pattern matched clauses and i and



Figure 15: Speedups for nbody on *corryvreckan* using reported total (Total) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

end inverted such that i decreases to 0, our prototype discovers the iterN.

```
1 loop ((ax,ay,az),i) = iterN g ((ax,ay,az),i)
2 where
3 g (ax,ay,az) = let ... in ((ax+px),(ay+py),(az+pz))
```

We have elided the local definitions for clarity. We executed nbody for $n = \{10000, 250000\}$ and between 0 and 100000 at intervals of 25000. Fig. 13 gives the corresponding mutator and total times (with standard deviations), and the number of sparks (potential parallelism), heap allocation size, and residency for each n. An average of 97.5% of sparks were converted on 4 cores, and 100% of sparks were converted on all other numbers of cores. Fig. 14 and 15 give speedups for nbody using mutator and total time, achieving maximum speedups of 27.08 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores and 26.82 for n = 250000 on 52 hyper-threaded cores. There is little difference between mutator and total reported speedups for nbody, showing that garbage collection is not significant for this example. The example also demonstrates good scalability for both varying n and number of cores. The growth in speedup does diminish after 28 cores, and the use of hyper-threading is again the most likely cause.

4.4. Matrix Multiplication

Matrix multiplication (matmult) multiplies two matrices. The *NoFib* suite definition consists of two Haskell files: ListAux (41 lines) and MatMult (246 lines). These collectively define 42 functions, 11 of which feature explicit *maps* and *folds* (not including the specialised versions of foldr, e.g. filter or sum, that

n	MUT	SD	Total	SD	Sparks	Heap	Residency
1000	13.15	1.59	13.37	1.58	18	317362136	44416
1500	56.09	5.12	56.33	5.14	27.5	713751520	44416
2000	131.72	14.07	131.96	14.07	38	1268624520	44416
2500	226.58	7.49	226.83	7.51	49.5	1835831694	44416
3000	402.13	2.71	402.36	2.69	62	1801405160	44416
3500	633.18	9.43	633.44	9.43	67	2451609160	44416
4000	946.42	12.1	946.67	12.08	80.5	3201845928	44416
4500	1344.39	16.09	1344.59	16.08	85.5	4052049928	44416
5000	1823.05	0.96	1823.31	0.96	100	5002253928	44416

Figure 16: Sequential mutator (MUT) and total (Total) times in seconds (with standard deviations), total number of sparks for parallel version, and heap allocation and residency in bytes, for matmult on *corryvreckan*.

are used extensively in this example). All the instances of explicit maps and folds are discovered by our prototype from their inlined definitions. Three functions, shiftRight and both addProd definitions, are implicit folds. shiftRight is defined as:

```
shiftRight c [] = []
shiftRight c (xs:xss) = (xs2++xs1):shiftRight (c-1) xss
where (xs1,xs2) = splitAt c xs
and is an implicit foldr:
```

```
1 shiftRight c xs = fst $ foldr f ([],c) xs
2 where
3 g xs (xss,c) = let (xs1,xs2) = splitAt c xs
4 in ((xs2++xs1):xss, c-1)
```

As with safe from queens (Sec. 4.2), our prototype cannot discover the foldr in shiftRight because c is updated in the recursive call. The second implicit fold, addProd, is defined as:

```
1 addProd :: Vector -> Vector -> Int -> Int
2 addProd (v:vs) (w:ws) acc = addProd vs ws (acc + v*w)
3 addProd [] [] n = n
```

addProd can be viewed as a composition of a zipWith and a fold1, or simply as a fold1. Here, both zipWith and fold1 implementations need not be total definitions, as mentioned in Sec. 3.1. We executed matmult for n between 1000 and 5000 at intervals of 500, with chunk size set to 20. Two parallel versions are possible: *i*. performs the multiplication of each row in parallel, and *ii*. divides each matrix into blocks to be computed in parallel and joins their result. We have used both modes here. Whilst the original *NoFib* suite definition provides its own matrix generation function, we have adjusted this so that matrices are generated using:



Figure 17: Speedups for matmult on *corryvreckan* using the row-wise parallelisation option and reported mutator (MUT) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

replicate n [1..n]

1

Fig. 16 reports sequential times (with standard deviations), and the number of sparks, heap allocation size, and residency for each n. The average number of sparks converted varies with n, instead of the number of cores seen in previous examples. Converted sparks fluctuate between 100% and not less than 93% (seen on 4 cores), and fluctuations grow smaller with larger n. Figs. 17, 19, 18, and 20 give speedups for matmult using mutator and total time for both parallel modes. Row-wise parallelisation achieves maximum speedups of 28.26 for n = 2500 on 56 hyper-threaded cores for both mutator and total values. Block-wise parallelisation achieves maximum speedups of 32.93 for n = 1500 on 52 hyper-threaded cores for both mutator and total values. As with nbody, there is little difference between mutator and total reported speedups for matmult, and mode 2 proves generally only slightly better than mode 1. The example demonstrates good scalability for both varying n and number of cores. Once again, speedup growth slows after 28 cores due to hyper-threading.

5. Related Work

Anti-unification was first described by Plotkin [10] and Reynolds [15] who described the approach using lattices of totally ordered terms. The technique has been explored and expanded upon since, enabling its operation on unranked terms [29], and used in a range of application areas; including generating code from analogy [30], generating invariants for program verification [17], termination checking [31], symbolic mathematical computing [32], and clone detec-



Figure 18: Speedups for matmult on *corryvreckan* using the block-wise parallelisation option and reported mutator (MUT) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

tion [33, 18]. Whilst our technique can be considered a form of clone detection, traditional anti-unification approaches to clone detection have focussed on comparing functions within the same program. Instead, we compare functions against potentially arbitrary higher-order functions that can either be from the same program, the Haskell Prelude, or a library.

One closely related technique, higher-order matching, was introduced and developed by Huet in 1976 [34] and later by Statman in 1982 [35]. Originally defined for the simply typed lambda calculus, higher-order matching has been adapted to work with Haskell-like languages and as part of program transformation tools [36]. Higher-order matching is a NP-hard problem [36]. Conversely, and by targeting the functions to which it is applied, anti-unification algorithms can be made efficient [17]. One important aspect of our technique is determining whether or not a function is an instance of a chosen pattern, a decision that is made by inspecting the structure of the anti-unifier. Higher-order matching algorithms result in a set of possible substitutions from the *language* itself [36], meaning that a decision of whether the examined function is an instance of a given pattern remains unanswered, and potentially unanswerable from the result alone. This collection of inferred substitutions presents another problem: a substitution must be chosen from the overall set, and the substitution must maintain functional equivalence between the examined function and its reconstructed form. Anti-unification derives its substitutions from both examined function and pattern, requiring no selection of inferred substitutions to ultimately derive pattern arguments. Lastly, we envisage that when comparing functions of increasingly different structures reasoning about the anti-unifier and inferred substitutions will play a greater roll in argument inference.



Figure 19: Speedups for matmult on *corryvreckan* using the row-wise parallelisation option and reported total (Total) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

Although the refactoring and clone detection fields are not traditionally concerned with the discovery of patterns, it is often a stage in the parallelisation process [8] as patterns as a simplification of parallelisation have been shown to be an effective [4, 5, 37, 38]. Pattern discovery for parallelism has primarily focused on approaches for imperative programs by analysing dependencies and machine learning techniques [39, 40, 8, 41]. Alternative automatic approaches focus on deriving parallel implementations from small programs or specifications, commonly employing the Bird Meertens formalism to calculate parts of the program [42, 43, 7, 44]. Whilst these approaches do not explicitly look for patterns, they use algorithmic structures such as list homomorphisms [45] and hylomorphisms [4] which are amenable to divide-and-conquer skeletons. These approaches have thus far focussed on translating operations into specific types, e.g. cons- or join-lists, which can then be easily parallelised [44, 7]. The approach by Geser [42] is particularly notable here, as it uses a form of anti-unification (referred to simply as *generalization*). These approaches are all limited by their type translations, and by the specific parallelism mechanics they introduce. In [46], Hu demonstrates how hylomorphisms can be found in arbitrary code, but the approach is limited to hylomorphisms. Our approach and prototype implementation works for Haskell 98, and can in principle discover arbitrary patterns in existing code. These patterns can be parallelised by, e.g., the approaches in [14, 4], but also used as part of code maintenance, and ensuring code reuse. Used for parallelism or performance reasons, our approach also allows the use of a variety of parallelisation or optimisation approaches.



Figure 20: Speedups for matmult on *corryvreckan* using the block-wise parallelisation option and reported total (Total) time. Dashed line indicates 28 physical cores, with a total of 56 hyper-threaded cores.

6. Conclusions and Future Work

In this paper, we have presented a technique to detect instances of recursion schemes in Haskell functions using anti-unification. That is we describe how to detect recursive patterns whose behaviour is well understood. We have described how our approach can be combined with automated or semi-automated refactoring techniques to transform functions into a form that enables the detection of such recursion schemes, and subsequently to rewrite the function as an instance of the pattern that is detected. As part of our approach, we have defined a specialised anti-unification algorithm. We have prototyped our approach by adapting the HaRe refactoring tool for Haskell, and have demonstrated our technique on four benchmarks from the standard NoFib suite. We have shown that real performance improvements and speedups can be achieved using parallel versions of the patterns that we find. Whilst our approach shows promising results, our analysis does presently require code to be written in a stylised form. We envisage, however, that further work on argument derivation will allow us to detect recursion schemes in a broader range of structures. We also intend to explore how equational reasoning, in conjunction with standard reduction operations, can be applied to inferred substitutions to successfully derive pattern arguments more flexibly. Additionally, we intend to examine larger, pre-existing examples of Haskell code. Although we have explained the technique with reference to Haskell and the standard *iterN*, map and fold patterns, none of these is a fundamental limitation. The approach described in this paper is, in fact, completely general. In future, we expect to apply our work to a much broader range of Haskell higher-order functions including many of the libraries that already exist in e.g. the *hackage* repository. We also intend to apply our approach to more varied patterns, e.g. *unfolds* [47], which have been shown to be useful components of parallel programs [4].

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Appendix A. Proof for Soundness of Anti-Unification Algorithm

Theorem 1 (Soundness of Anti-Unification Algorithm). Given the terms t_1 and t_2 , we can find t, σ_1 , and σ_2 , such that $t_1 \cong t \sigma_1$ and $t_2 \cong t \sigma_2$.

Proof Sketch. Given the terms t, t_1 , t_2 , and substitutions, σ_1 and σ_2 . The proof is by case analysis on t_1 and t_2 .

Case $t_1 \cong t_2$:

By application of EQ with $t = t_2$ and $\sigma_1 = \sigma_2 = \varepsilon$, we must show that $t_1 \cong t_2 \varepsilon$ and $t_2 \cong t_2 \varepsilon$.

By Def. 5 and reflexivity, both $t_1 \cong t_2$ and $t_2 \cong t_2$ hold.

Case $t_2 = (\operatorname{Var} v) \land (v \neq p)$:

By application of VAR when $t = (\operatorname{Var} v)$, $\sigma_1 = (v \mapsto t_1)$, and $\sigma_2 = \varepsilon$, we must show that $t_1 \cong (\operatorname{Var} v) ((v \mapsto t_1))$ and $(\operatorname{Var} v) \cong (\operatorname{Var} v) \varepsilon$ hold.

By Def. 4 and reflexivity, $t_1 \cong t_1$ holds.

By Def. 5 and reflexivity, $(Var v) \cong (Var v)$ holds.

 $\mathbf{Case} \ t_1 = (\mathtt{Var} \ v_{a_i}^\mathtt{f}) \land t_2 = (\mathtt{App} \ (\mathtt{Var} \ u) \ (\mathtt{Var} \ v_{a_i}^\mathtt{p})) \land (u \neq \mathtt{p}) \land (v_{a_i}^\mathtt{f} \equiv v_{a_i}^\mathtt{p}):$

By application of ID where

$$t = (\operatorname{App} (\operatorname{Var} u) (\operatorname{Var} v_{a_i}^{\operatorname{p}}))$$

$$\sigma_1 = ((v \mapsto id), \varepsilon)$$

$$\sigma_2 = \varepsilon$$

we must show that:

$$(\operatorname{Var} v_{a_i}^{\mathbf{f}}) \cong (\operatorname{App} (\operatorname{Var} u) (\operatorname{Var} v_{a_i}^{\mathbf{p}})) ((u \mapsto id), \varepsilon)$$
(1)

$$(\operatorname{App}(\operatorname{Var} u) (\operatorname{Var} v_{a_i}^{\operatorname{p}})) \cong (\operatorname{App}(\operatorname{Var} u) (\operatorname{Var} v_{a_i}^{\operatorname{p}})) \varepsilon$$

$$(2)$$

For (1): By Def. 6, $(\operatorname{Var} v_{a_i}^{\mathbf{f}}) \cong (\operatorname{App} (\operatorname{Var} u) (u \mapsto id) (\operatorname{Var} v_{a_i}^{\mathbf{p}}) \varepsilon)$. By Def. 4 and Def. 5, $(\operatorname{Var} v_{a_i}^{\mathbf{f}}) \cong (\operatorname{App} (\operatorname{Var} id) (\operatorname{Var} v_{a_i}^{\mathbf{p}}))$. By Def. 8 and reflexivity, $(\operatorname{Var} v_{a_i}^{\mathbf{f}}) \cong (\operatorname{Var} v_{a_i}^{\mathbf{p}})$ holds.

For (2): By Def. 5 and reflexivity, $(\text{App }(\text{Var } u) (\text{Var } v_{a_i}^p)) \cong (\text{App }(\text{Var } u) (\text{Var } v_{a_i}^p))$ holds.

Case $t_2 \in \mathcal{R}_p$:

By application of RP where $t = (\operatorname{Var} \alpha)$, $\sigma_1 = (\alpha \mapsto t_1)$, and $\sigma_2 = (\alpha \mapsto t_2)$, we must show that both $t_1 \cong (\operatorname{Var} \alpha) (\alpha \mapsto t_1)$ and $t_2 \cong (\operatorname{Var} \alpha) (\alpha \mapsto t_2)$ hold.

By Def. 4 and reflexivity, both $t_1 \cong t_1$ and $t_2 \cong t_2$ hold.

Case $t_1 = (C t_{11} \dots t_{1n}) \land t_2 = (C t_{21} \dots t_{2n}) \land$ $\forall i \in [1, n], (t_{1i} \cong t_i \sigma_{1i}) \land (t_{2i} \cong t_i \sigma_{2i})$:

By application of CONST where

 $t = (C t_1 \dots t_n)$ $\sigma_1 = (\sigma_{11}, \dots, \sigma_{1n})$ $\sigma_2 = (\sigma_{21}, \dots, \sigma_{2n})$

we must show that:

$$(C t_{11} \dots t_{1n}) \cong (C t_1 \dots t_n) \left(\sigma_{1i}, \dots, \sigma_{1n} \right)$$
$$(C t_{21} \dots t_{2n}) \cong (C t_1 \dots t_n) \left(\sigma_{2i}, \dots, \sigma_{2n} \right)$$

By Def. 6, $(C t_{11} \ldots t_{1n}) \cong (C (t_1 \sigma_{11}) \ldots (t_n \sigma_{1n}))$ and $(C t_{21} \ldots t_{2n}) \cong (C (t_1 \sigma_{21}) \ldots (t_n \sigma_{2n}))$. By assumption, substitutivity and reflexivity, both $(C t_{11} \ldots t_{1n}) \cong (C t_{11} \ldots t_{1n})$ and $(C t_{21} \ldots t_{2n}) \cong (C t_{21} \ldots t_{2n})$ hold.

Case $t_1 = [t_{11}, \ldots, t_{1n}] \land t_2 = [t_{21}, \ldots, t_{2n}] \land$ $\forall i \in [1, n], t_{1i} \cong t_i \sigma_{1i} \land t_{2i} \cong t_i \sigma_{2i}$:

By application of LIST where

$$t = [t_1, \dots, t_n]$$

$$\sigma_1 = (\sigma_{11}, \dots, \sigma_{1n})$$

$$\sigma_2 = (\sigma_{21}, \dots, \sigma_{2n})$$

we must show that:

$$(\llbracket t_{11} \ldots t_{1n} \rrbracket) \cong (\llbracket t_1 \ldots t_n \rrbracket) (\llbracket \sigma_{1i}, \ldots, \sigma_{1n} \rrbracket)$$
$$(\llbracket t_{21} \ldots t_{2n} \rrbracket) \cong (\llbracket t_1 \ldots t_n \rrbracket) (\llbracket \sigma_{2i}, \ldots, \sigma_{2n} \rrbracket)$$

By Def. 6, $([t_{11} \ldots t_{1n}]) \cong ([(t_1 \sigma_{11}) \ldots (t_n \sigma_{1n})])$ and $([t_{21} \ldots t_{2n}]) \cong ([(t_1 \sigma_{21}) \ldots (t_n \sigma_{2n})])$. By assumption, substitutivity and reflexivity, both $([t_{11} \ldots t_{1n}]) \cong ([t_{11} \ldots t_{1n}])$ and $([t_{21} \ldots t_{2n}]) \cong ([t_{21} \ldots t_{2n}])$ hold.

Otherwise:

By application of OTHERWISE where $t = (\operatorname{Var} \alpha)$, $\sigma_1 = (\alpha \mapsto t_1)$, and $\sigma_2 = (\alpha \mapsto t_2)$, we must show that both $t_1 \cong (\operatorname{Var} \alpha)(\alpha \mapsto t_1)$ and $t_2 \cong (\operatorname{Var} \alpha)(\alpha \mapsto t_2)$ hold.

By Def. 4 and reflexivity, both $t_1 \cong t_1$ and $t_2 \cong t_2$ hold.

Adam Barwell is a PhD student at the University of St Andrews working in the area of static analysis and refactoring. Adam graduated from University College London in 2013 where he developed an interest in programming langauge theory, and is now studying under Kevin Hammond and Chris Brown.

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Chris Brown is a Post-Doctoral Research Fellow at the University of St Andrews working in the area of Refactoring and Parallelism. Chris has a Ph.D. from the University of Kent under the supervision of Simon Thompson, where he worked on the implementation of the HaRe refactoring tool for Haskell. Chris now works at St Andrews building radically new refactoring techniques and methodologies to exploit parallel skeleton and pattern applications in both the functional and imperative domains. Chris has published at major conferences in the field of functional programming, refactoring and parallelism.

Kevin Hammond is a Professor of Computer Science at the University of St Andrews working in the area of Parallelism. In the past, Kevin has helped implement Haskell, worked on EU Projects such as ParaPhrase, and is currently working on the RePhrase project.

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- We present a technique using anti-unification to discover recursion schemes in Haskell functions
- We present a new, specialised anti-unification algorithm.
- We have implemented our approach using the Haskell refactoring tool HaRe
- We have demonstrated our approach on a range of standard examples
- Our technique is in principle completely general, able to discover arbitrary patterns in code.