Region-Based Memory Management for Mercury Programs

QUAN PHAN, GERDA JANSSENS
Department of Computer Science
Katholieke Universiteit Leuven
Celestijnenlaan, 200A
B-3001 Leuven, Belgium
(e-mail: quan.leuven@gmail.com, gerda.janssens@cs.kuleuven.be)

ZOLTAN SOMOGYI
National ICT Australia and
Department of Computer Science and Software Engineering
The University of Melbourne
Victoria, 3010 Australia
(e-mail: zs@unimelb.edu.au)

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Abstract

Region-based memory management (RBMM) is a form of compile time memory management, well-known from the functional programming world. In this paper we describe our work on implementing RBMM for the logic programming language Mercury. One interesting point about Mercury is that it is designed with strong type, mode, and determinism systems. These systems not only provide Mercury programmers with several direct software engineering benefits, such as self-documenting code and clear program logic, but also give language implementors a large amount of information that is useful for program analyses. In this work, we make use of this information to develop program analyses that determine the distribution of data into regions and transform Mercury programs by inserting into them the necessary region operations. We prove the correctness of our program analyses and transformation. To execute the annotated programs, we have implemented runtime support that tackles the two main challenges posed by backtracking. First, backtracking can require regions removed during forward execution to be “resurrected”; and second, any memory allocated during a computation that has been backtracked over must be recovered promptly and without waiting for the regions involved to come to the end of their life. We describe in detail our solution of both these problems. We study in detail how our RBMM system performs on a selection of benchmark programs, including some well-known difficult cases for RBMM. Even with these difficult cases, our RBMM-enabled Mercury system obtains clearly faster runtimes for 15 out of 18 benchmarks compared to the base Mercury system with its Boehm runtime garbage collector, with an average runtime speedup of 24%, and an average reduction in memory requirements of 95%. In fact, our system achieves optimal memory consumption in some programs.

KEYWORDS: region-based memory management, region analysis, runtime support, backtracking, logic programming, Mercury
1 Introduction

Memory management is an integral part of all practical programming language systems. Traditionally, memory has been left to the programmer to manage using constructs such as C’s `malloc` and `free`, but experience has shown that such manual systems require a large amount of quite tedious work from programmers, and are very hard to use correctly. More recent programming languages therefore automate memory management. The standard way to implement automatic memory management is runtime garbage collection. This provides memory safety, good memory reuse, and reasonable performance, but it does have a significant downside, which is that decisions about which parts of memory can be reused are made completely at runtime, which can incur significant overheads.

Region-based memory management or RBMM (Tofte and Talpin 1997) is a recent technique for avoiding these overheads by moving decisions from runtime to compile time, thus shifting most of the responsibility to the compiler. RBMM is based on the idea of putting each group of heap objects that have the same lifetime into their own regions, the motive being that reclaiming entire regions at the end of their lifetime makes collection very fast. A typical scenario is a function storing its intermediate results in a region that is freed once the final result of the function has been computed. All the decisions about which objects are allocated into which regions and when each region should be created and removed are made at compile time.

Since the fundamental work on RBMM for functional programming (Tofte and Talpin 1997), there have been several improvements and new developments in that context (Aiken et al. 1995; Birkedal et al. 1996; Henglein et al. 2001). RBMM has also been adapted to other programming paradigms, such as imperative programming (Gay and Aiken 1998; Grossman et al. 2002), object-oriented programming (Cherem and Rugina 2004; Chin et al. 2004), and logic programming (Makholm 2000a; Makholm 2000b; Makholm and Sagonas 2002).

The initial work on RBMM for logic programming languages applied RBMM to Prolog. However, the first attempt (Makholm 2000a; Makholm 2000b) was developed for a non-standard implementation of Prolog which would require substantial changes before it could be applied in any standard implementation. The authors of (Makholm and Sagonas 2002) fixed this problem by implementing RBMM in the context of the standard technology for implementing Prolog, the Warren Abstract Machine (WAM). Nevertheless, this work mainly concentrated on the runtime extensions needed to run Prolog programs with RBMM. As its analysis algorithm, it used an adapted version of a type-based region analysis originally developed for the strongly typed functional language SML (Henglein et al. 2001). Since Prolog has no static type system and more importantly no static mode system, the region inference has to get the information it needs from type and mode inferences, which often yield imprecise results. Moreover, a Prolog implementation’s lack of knowledge about the determinism of a program’s predicates generally requires them to be treated as nondeterministic. These limitations prevent the application of most of the optimizations that would improve the performance of RBMM, making it hard
for it to become a practical alternative to native runtime collectors in Prolog systems. The logic programming language Mercury has none of these limitations; the Mercury compiler knows the type of every variable and the mode and determinism of every goal in the program. This fact, the pure nature of Mercury (the absence of side-effects), and the limited research on RBMM in logic programming motivated us to investigate whether region-based memory management could be developed and implemented efficiently for Mercury.

In this paper we describe the first automated RBMM system for Mercury. Given a Mercury program,

- our system determines the set of regions the program should use;
- it decides, for each allocation site in the program, which region the allocation should happen in;
- it inserts instructions into the program to create each region just before it is first needed; and
- it inserts instructions into the program to remove each region as soon as it is safe to do so.

The main contributions of our work are as follows.

1. We develop the static program analyses needed for generating region-annotated programs. These include a region points-to analysis to divide Mercury terms into regions, a liveness analysis that assigns lifetimes to the regions, and a program transformation to annotate the original programs with the derived region information.
2. We prove several safety properties for memory accesses and region operations in the resulting annotated programs.
3. Our runtime support system handles the interaction of RBMM with backtracking correctly and without incurring excessive overheads.
4. Our RBMM-enabled system achieves faster execution times and much lower memory requirements for most of our benchmark programs than the standard Mercury system, which uses the Boehm-Demers-Weiser garbage collector (Boehm and Weiser 1988) for memory management. The region system actually achieves optimal memory consumption on some benchmarks.
5. We make a detailed analysis of the RBMM behavior of a selection of programs, including some well-known difficult cases. This study reveals the impact of sharing on memory reuse in RBMM systems.

A previous version of our region analysis and transformation was published in (Phan and Janssens 2007). In (Phan et al. 2008) we described the runtime support for RBMM. They all have been reformulated, extended and/or refined in this paper.

The structure of the paper is as follows. In Section 2 we introduce Mercury and the compiler’s internal representation of Mercury programs. Section 3 describes intuitively how RBMM can be realized for Mercury, and explains our decisions on how to support backtracking. Section 4 explains how we decide which terms should be stored in which regions, taking into account sharing among terms. Based on this region model, we develop the static analyses of our system: Sections 5, 6,
and 7 contain respectively our region points-to analysis, our region liveness analysis, and our program transformation, together with theorems about their correctness. Section 8 shows the basic extensions to the Mercury runtime system needed to support RBMM in deterministic code, while Section 9 describes the extensions needed to support backtracking (nondeterminism). Section 10 presents a detailed evaluation of our RBMM system, as well as a discussion of the relation between sharing and memory reuse in region-based systems. We discuss related research in Section 11, present our ideas for future work in Section 12, and conclude in Section 13.

2 Background

2.1 Mercury

Mercury is a pure logic programming language intended for the creation of large, fast, reliable programs (Somogyi et al. 1996). While the syntax of Mercury is based on the syntax of Prolog, semantically the two languages are very different due to Mercury’s purity, its type, mode, determinism and module systems, and its support for evaluable functions. (Mercury treats functions as predicates with the return value as an extra argument, so in the rest of the paper we will talk only about predicates.)

Mercury has a strong Hindley-Milner type system very similar to Haskell’s. Some types are built into the language (e.g. `int`), but users can also introduce new types using type definitions such as the one in Example 1.

Example 1

The declaration of the type `list_int`.

```mercury
:- type list_int --> []; [int | list_int].
```

This defines the type of lists of integers.

Mercury programs are statically typed; the compiler knows the type of every argument of every predicate (from declarations or inference) and every local variable (from inference).

The mode system classifies each argument of each predicate as either input or output; there are exceptions, but they are not relevant to this paper. If input, the argument passed by the caller must be a ground term. If output, the argument passed by the caller must be a distinct free variable, which the callee will instantiate to a ground term. It is possible for a predicate to have more than one mode; the usual example is `append`, which has two principal modes: `append(in,in,out)` and `append(out,out,in)`. We call each mode of a predicate a procedure. The Mercury compiler generates separate code for each procedure.

Each procedure has a determinism, which puts limits on the number of its possible solutions. Procedures with determinism `det` succeed exactly once; `semidet` procedures succeed at most once; `multi` procedures succeed at least once; while `nondet` procedures may succeed any number of times.
Example 2

Figure 1 shows the quicksort program written in Mercury, including declarations of the types, modes, and determinisms for its two essential predicates, qsort and split. We include the code of main for completeness, but it is of no relevance to the topic of the paper. The notation !IO represents two variables, which in this case stand for the initial and final states of the world, i.e. the state before the program writes out its result with io.write, and the state after. (The io.write predicate is defined in the io module of the Mercury standard library.)

We support a very large subset of Mercury: unifications, first order calls, conjunctions, disjunctions, switches, if-then-elses, negations, and quantification. The only parts we do not support are higher order calls (including typeclass method calls), calls to foreign language code, and multi-module programs. A complete description of Mercury can be found in (Mercury team 2009).

2.2 Mercury Code inside the Compiler

The compiler converts all predicate definitions into an internal form. For our subset of Mercury, this internal form is given by the following abstract syntax:

\[
\begin{align*}
\text{predicate } P & : \quad p(x_1, \ldots, x_n) \leftarrow G \\
\text{goal } G & : \quad x = y | x = f(y_1, \ldots, y_n) | p(x_1, \ldots, x_n) | G_1, \ldots, G_n | \neg G | (\text{if } G_2 \text{ then } G_3 \text{ else } G_4) | \text{some}[x_1, \ldots, x_n] G
\end{align*}
\]

We call the first three kinds of goals (unifications and calls) atomic goals or just atoms. The rest are called compound goals, in which a sequence of goals separated by commas is a conjunction, while a sequence of goals separated by semicolons is a disjunction.

As this implies, the Mercury compiler internally converts any predicate definition with two or more clauses into a single clause with an explicit disjunction. The clauses themselves are transformed into superhomogeneous form, in which each atom (including clause heads) must be one of the forms \( p(X_1, \ldots, X_n) \), \( Y = X \), or \( Y = f(X_1, \ldots, X_n) \), where all of the \( X_i \) are distinct.

Inside the compiler, every goal (compound as well as atomic) is annotated with
mode and determinism information. For unifications, we show the mode information by writing \( \leq \) for construction unifications, \( \Rightarrow \) for deconstruction unifications, \( \equiv \) for equality tests, and \( :\) for assignments. The compiler reorders conjunctions as needed to ensure that goals that consume the value of a variable always come after the goal that produces its value. We show the quicksort program in this abstract syntax in Figure 2. For readability, we have chosen meaningful names for some additional variables that are added automatically by the Mercury compiler. We also replace the sequence of unifications needed to construct a single ground term with a single goal. For example, the list construction at (1) in \texttt{main} in Figure 2, actually stands for

\[
V_0 \leq [], \\
V_1 \leq 3, V_2 \leq [V_1 | V_0], \\
V_3 \leq 1, V_4 \leq [V_3 | V_2], \\
V_5 \leq 2, L \leq [V_5 | V_4]
\]

These extra details are of no interest in this paper.

In the rest of the paper, we will ignore negation, since \texttt{not G} can be implemented as \texttt{if G then fail else true}, where \texttt{fail} and \texttt{true} are two builtin goals, with \texttt{fail} always failing and \texttt{true} always succeeding. Note that in Mercury (unlike in Prolog), the condition of an if-then-else is allowed to succeed several times. Whether the condition of a particular if-then-else can do so will be recorded in its determinism annotation, and many parts of the compiler, including the RBMM implementation, handle conditions of different determinisms differently.

Another situation in which determinism information is important is existential quantification. (Mercury also supports universal quantification, but the compiler internally converts all \([x_1, \ldots, x_n]G\) to \texttt{not some}[\texttt{not} \ G], so we do not have to deal with it.) If \texttt{some}[\ldots] \ G quantifies away all the output variables of \ G, then different solutions of \ G would be indistinguishable, so even if \ G can have more than one solution, \texttt{some}[\ldots] \ G will not. We call such a quantification a \texttt{commit}, and we handle commits differently from other quantifications.

\begin{verbatim}
main(!IO) :-
    (1) L =\leq [2, 1, 3],
    (2) A =\leq [],
    (3) qsort(L, A, S),
    (4) io.write(S, !IO),

split(X, L, L1, L2) :-
    (1) L =\leq [],
    (2) L1 =\leq [],
    (3) L2 =\leq [],
    qsort(L, A, S) :-
        (1) L =\leq [],
        (2) A =\leq [],
        (3) qsort(L, A, S),
        (4) L1 =\leq [],
        (5) L2 =\leq [],

qsort(L, A, S) :-
    (2) S =:= A

Fig. 2: quicksort program in superhomogeneous form.
\end{verbatim}
3 Overview of Region-Based Memory Management for Mercury

We divide the task of realizing RBMM for Mercury into two parts: (a) two static analyses and a program transformation, which work entirely at compile time, and (b) dynamic runtime support, which executes at runtime code added to the program by the compiler at compile time.

The goal of the static analyses and transformation is to annotate Mercury programs with information about regions. An annotated program contains information about the regions in which terms are constructed and when regions are created and freed. To obtain this information, we first use a region points-to analysis to detect the regions used by a program, and then we compute the lifetimes of these regions using a region liveness analysis. The program transformation then uses these pieces of information to convert the program into a region-annotated program.

The runtime support for RBMM has two main tasks. First, it has to implement the necessary operations on regions: the creation of regions, allocation into regions, and the removal of regions (Section 8). Second, it has to provide support for the interaction of backtracking with RBMM. There are two main forms of interaction: instant reclaiming and backward liveness (Section 9).

The memory allocated by computations that have been backtrack over will never be accessed again, since backtracking effectively “erases” such computations. To prevent memory leaks, this memory should be recovered immediately when forward execution resumes again; we call this instant reclaiming. This obviously has to be done at runtime, so in our system, the compiler inserts the code required to do this into the program at both resume points (points in the program where forward execution can resume after backtracking, such as the starts of second and later disjuncts in a disjunction) and at program points that establish resume points (such as just before entry into a disjunction).

In logic programming languages, the presence of backtracking requires the notion of liveness to be divided into two parts. A variable, memory location or region is forward live at a program point if it can be accessed during forward execution from that program point, and it is backward live at a program point if it can be accessed during backward execution (i.e. after backtracking to a choice point established before that program point). The two notions of liveness are independent: all four combinations of forward and backward liveness and deadness are possible. Regions can be reclaimed only when they are both forward dead and backward dead.

Our region liveness analysis takes into account only forward liveness, and we ensure safety with respect to backward liveness through runtime support. Our reasons for why we handle backward liveness this way are that

- handling it purely at compile time is not possible, since runtime support will still be needed in some cases, as we will point out in Section 12, and a purely-runtime solution is simpler than a solution that mixes compile time and runtime aspects; and
- we can implement a large part of this runtime support using the machinery we need anyway for instant reclaiming.
However, handling backward liveness at least partially at compile time may turn out to be more efficient, which is why we intend to explore it in future work.

### 3.1 Region Variables

We use region variables to refer to regions, just as we use program variables to refer to values. To allocate a new region, we use the instruction `create(R)`, which creates a region and binds the region variable R to it. To free a region we use the instruction `remove(R)`, which frees the memory of the region to which R is currently bound. Our regions can and actually do live across procedure boundaries, and thus we pass region variables as extra arguments to procedure calls. Figure 3 shows the region-annotated quicksort program after our region transformation. Our source-to-source transform represents these instructions, and the instructions we introduce later, as calls to builtin predicates. We describe the implementation of these predicates in Section 8.

```
main(!IO) :-
  create(R20), create(R21),
  (1) L <= [2, 1, 3] in R20,
  create(R22),
  (2) A <= {} in R22,
  qsort(L@R6, A@R8, S@R8),
  (3) L1 <= {} in R3,
  create(R4),
  remove(R21), remove(R22).

 split(X, L@R1, L1@R3, L2@R4) :-
  (1) L => [],
  create(R3),
  (2) L1 <= {} in R3,
  create(R4),
  (3) L2 <= {} in R4
  ;
  (4) L => [L | Ls],
  if X >= Le then
  (1) L => [],
  (6) split(X, Ls@R1, L11@R3, L2@R4),
  remove(R6),
  (7) L1 <= [Le | L11] in R3
  else
  (8) split(X, Ls@R1, L10@R3, L20@R4),
  (9) L2 <= [Le | L21] in R4
  ;
  (5) A1 <= [Le | S2] in R8,
  (7) qsort(L10@R9, A1@R8, S2@R8).
```

Fig. 3: Region-annotated quicksort program.

In the region-annotated code, we use the postfix @Ri to annotate both actual and formal arguments with their region variables. We also annotate each unification that constructs a new memory cell with the region in which the cell will be allocated. For example, in `main`, the skeleton of the list L is in the region (bound to) R20, while that of the accumulator A is in R22. The elements of the lists are in R21 (but see below). In the call to `qsort`, R20 and R22 are passed as actual region arguments, corresponding to the formal arguments R6 and R8 in the definition of `qsort`. We do not need to pass the region of the elements because `qsort` and `split` just read from it. The region R20 is passed to `qsort` from `main` and is removed in the base case branch of `split` in the call to `split` at (4) in `qsort`. The two new lists L1 and L2 are allocated in two separate regions referred to by R9 and R10. These regions are created by the base case branch of `split`, and removed (indirectly) by the
recursive calls to `qsort` at (5) and (7). If \( L_1 \) and \( L_2 \) are empty lists, the removals will happen in the base case branch of `qsort`; otherwise, they will happen in the base case branch of `split`. The region \( R_{22} \) of the resulting list is the region of the accumulator, which is created in `main`.

4 Region Modelling

4.1 Storing Terms in Regions Based on Their Types

As we want to distribute terms over different regions, we first discuss the representation of terms when the heap memory is divided into regions.

We assume that a term that does not fit into one word will be represented by a pointer to a memory cell on the heap. We also assume that a term that can be represented by a single memory word does not need storage on the heap in its own right. When those terms are on their own, they will be stored in registers or in stack slots. When they are arguments of a larger term, they will be stored in a word on the heap, but this word will be counted as belonging to the memory cell representing the larger term.

Our assumptions are compatible with the implementation of Mercury in the Melbourne Mercury Compiler (MMC). The MMC knows the types of all variables, and these types give us information about the storage size of terms. Terms of primitive types such as `int` and `char` are stored in one word, and the same is true of enumeration types (types in which all functors have arity zero). The principal functor of a term that needs heap space is represented by a possibly-tagged pointer to a block of memory words on the heap. The compiler knows all the functors in the type of the term. It also knows that all words in the Mercury heap are aligned, so pointers to them have two free bits on 32 bit machines, and three free bits on 64 bit machines. Therefore if a type has at most four function symbols (eight on 64 bit machines), the principal functor can be represented by what Mercury calls a “primary tag” on the lowest bits of the pointer. When a type has only one functor, even this is not needed. When a type has more than four or eight functors (on 32 and 64 bit machines respectively) the compiler will use one primary tag value to represent several function symbols, and will use the first word of the pointed-to memory block as a secondary tag to distinguish between them. (The usual implementations of Prolog have a similar word in every heap cell other than those storing lists, increasing their memory footprint.)

Example 3

Consider the following types.
```prolog
:- type elem ---> f; g(int); h(list_int, int).
:- type list_elem ---> []; [elem | list_elem].
```

Figure 4 shows MMC’s representation of the term \([f, g(1), h([1, 2], 2)]\) bound to the variable \( L \), which is of type `list_elem`. Boxes with slim border are locations on the stack or in registers, while boxes with bold borders are locations on the heap. Note the representation of the term \( h([1, 2], 2) \) in the last element of the
list: we need a two-word block for h’s arguments, but the functor itself is stored implicitly in the tagged pointer.

\[ L = [f, g(1), h([1, 2], 2)] \]

Fig. 4: Term representation of \( L = [f, g(1), h([1, 2], 2)] \).

We now consider the storage of terms when the heap is split into regions. The idea is to use different regions to store different parts of a term so that we can reclaim the memory of a part by destroying its region as soon as that part becomes dead. Many programs (including quicksort) create temporary lists in which the elements have much longer lifetimes. Therefore storing the elements and the list skeletons in different regions will allow us to recover the memory of the list skeletons much earlier. Generalizing from this, we divide each term into regions based on the type of each of its subterms. We will develop this idea in the next subsection.

In Figure 4, the regions used to store our example term are shown by the dashed lines. We put the two-word memory blocks making up the skeleton of the list \( L \) into one region because they have the type \texttt{list_elem}. We also put all the elements, which have the type \texttt{elem}, into another region. Finally, the first subterm of the third element, which is of type \texttt{list_int} rather than \texttt{list_elem}, is stored in yet another region.

The representation of the list of integers here seems inconsistent with what we said in Section 3, where we have an extra, separate region for the integers. The reason for this is because in this section we want to give a region model as close as possible to the implementation of Mercury in the MMC, in which integers do not need their own memory cells on the heap. Here we have two different viewpoints: a theoretical one that wants to treat all types the same way, and a practical one that wants to accurately reflect how the implementation handles values of each type. For convenience, we take the liberty of switching between the two viewpoints at will. When talking about theoretical topics such as static analyses and transformation for convenience we generally assume that all types (including \texttt{int}) require heap storage; when talking about the actual implementation, we will assume that the implementation does not create regions without having anything to put into them. We will be more specific only if the context is not clear.
4.2 Modelling Regions of a Type

Our system needs a storage scheme that specifies how the terms of a type are stored. Consider a type \( t \) declared as follows.

\[
:- \text{type } t \longrightarrow \ldots; f(t_1, \ldots, t_i, \ldots, t_n); \ldots
\]

We associate a region variable \( R^t \) with the type. The block of memory words corresponding to a principal functor, such as \( f \), of a term of type \( t \) is stored in the region bound to \( R^t \). In the rest of the paper we abbreviate this by simply saying that a principal functor is stored in \( R^t \). The principal functor of an argument of \( f \) that has type \( t_i \) is stored in the region bound to \( R^{t_i} \), which is associated to \( t_i \).

If a type \( t \) is recursive or mutually recursive, we still use only one region variable \( R^t \). This implies that any term of a recursive type is modelled by a finite number of regions.

We model the storage scheme using a type-based region graph, \( TG(N, E) \) with \( N \) being a set of nodes and \( E \) being a set of directed edges. A node stands for a region variable. A directed edge from one node to another represents the fact that the region bound to the region variable represented by the source node of the edge contains references into (points-to) the region bound to the region variable represented by the target node of the edge. The reference relation represented by the edges is actually defined by the type.

Consider the type-based region graph of the type \( t \), \( TG_t \), with the region variables \( R^t \), \( R^{t_1} \), \( R^{t_2} \) and so on. If \( R^t \) is represented by the node \( n \), then for each node \( m \) representing \( R^{t_i} \), we have exactly one edge \((n, (f, i), m)\) with the label \((f, i)\). We refer to \( n \) as the principal node of \( TG_t \).

Example 4

The type-based region graph for the type \( \text{list} \cdot \text{elem} \) in Example 3 is shown in Figure 5. The \([1]\) principal functor is stored in \( R^{\text{list} \cdot \text{elem}} \). It has two arguments, the first having the type \( \text{elem} \) and the second having the same type \( \text{list} \cdot \text{elem} \). Thus we have two edges from \( R^{\text{list} \cdot \text{elem}} \), the first pointing to \( R^{\text{elem}} \) where the principal functors of \( \text{elem} \) (\( g/1 \) and \( h/2 \)) are stored, and the second being a self-edge. The edge labelled \((h, 1)\) is due to the first argument of the functor \( h/2 \). The reader may want to compare this type-based region graph with Figure 4, which shows the memory representation of a term of this type. \( \square \)

![Fig. 5: The type-based region graph of the type list\cdot elem.]

Example 5

Consider the following types \( t_1 \) and \( t_2 \), which are mutually recursive.
The type-based region graph for these types is shown in Figure 6.

![Type-based region graph](image)

**Fig. 6: Type-based region graph of mutually recursive types.**

### 4.3 Region Points-To Graph

Now that we have the region model for types, our next goal is to model the memory used by a Mercury program in terms of regions. A program consists of a set of procedures, each having its own set of program variables that, at runtime, are instantiated with relevant terms. Therefore we define the notion of a region points-to graph that models the memory used by a set of variables. The memory used by a procedure is modelled by a region points-to graph for its variables. Finally, the memory model for the whole program is expressed through the region points-to graphs of its procedures.

In Mercury, variables are instantiated by unifications. A construction unification \( X \leftarrow f(\ldots, Y, \ldots) \) allocates new memory for storing the functor \( f \) (actually the block of memory words storing \( f \)'s arguments, and, if the tag on the pointer to the block is not enough for this, \( f \)'s identity), and creates sharing between \( X \) and each \( Y \). In a deconstruction unification \( X \Rightarrow f(\ldots, Y, \ldots) \) or an assignment unification \( Y := X \), \( Y \) is instantiated and shares a subterm or the whole term with \( X \), respectively. Hence the region points-to graphs should capture the memory locations of the variables and the sharing among them.

A **region points-to graph**, \( G(N, E) \), for a set of variables \( V \), consists of a set of nodes \( N \), representing region variables and a set of directed edges, \( E \), representing references between the regions bound to these region variables. The edges here serve exactly the same purpose as those in a \( TG \) graph. However, each node \( n \) in the region points-to graph has an associated set of program variables, \( \text{vars}(n) \), whose principal functors are stored in the region that is bound to the region variable that is represented by \( n \). The \( \text{vars} \) sets of the various nodes must represent a partition of the set of variables of interest (e.g. the set of variables in a procedure): each variable in the set must appear in the \( \text{vars} \) set of exactly one node. (Note that the \( \text{vars} \) set of a node may be empty; this can happen when a variable's value has some subterms that the code in question does not access.) We have \( V = \bigcup_{n \in N} \text{vars}(n) \).

The notation \( n_X \) denotes the node where \( X \in \text{vars}(n_X) \) and we refer to \( n_X \) as the **location** of \( X \), since this node represents the region where the principal functor of the term that \( X \) is bound to is stored. The function \( \text{node}(n_X, (f, i)) \) returns the node \( m \) if \( (n_X, (f, i), m) \in E \), otherwise its result is undefined.

**Sharing** is represented in a region points-to graph in two ways. First, directed
edges represent the sharing of subterms, and second, a node whose \textit{vars} set contains more than one variable represents the fact that these variables may be bound to the same term. An example of the latter is given by the variables of an assignment unification: they are bound to the same term and therefore they should be in the \textit{vars} set of the same node. A region points-to graph represents sharing at the level of the regions.

\textit{Definition 1 (Region-sharing in a region points-to graph)}

Two variables $X$ and $Y$ \textbf{region-share} in a region points-to graph if there exists a node that can be reached from both $n_X$ and $n_Y$.

For convenience, we also say \textbf{a node represents a region}, by which we mean the region to which the region variable represented by the node is bound at runtime. Then we can say \textbf{a functor is stored in a node} meaning that the functor (i.e. the memory block corresponding to it) is stored in the region represented by the node.

For a procedure $p$, we denote its region points-to graph by $G_p(N_p, E_p)$. $G_p$ should represent the locations and sharing among all the variables in $p$. It is possible to form a region points-to graph for a procedure exactly from the type-based region graphs of all of its variables (whose types are known to the compiler). Although this region points-to graph adequately models the locations of the procedure’s relevant terms, it does not represent the sharing among them. Actually, as we will see in Section 5, we use that region points-to graph as the starting point in our region points-to analysis of a procedure, with the ultimate aim of producing a region points-to graph that also represents all the possible sharing among the procedure’s variables.

\textit{Example 6}

Consider the following sequence of code to construct the term that $L$ in Example 3 is bound to. The type of $K$ is of no importance.

```
..., 
X <= [1, 2], 
Y := X, 
Z <= h(Y, 2), 
L <= [f, g(1), Z], 
K <= k(Z), 
...  
```

The region points-to graph that represents the memory manipulated by this sequence is shown in Figure 7. $X$ and $Y$ are in the \textit{vars} set of the same node because the assignment makes $Y$ point to the term to which $X$ is bound. The direct sharing between $Z$ and $Y$, and between $L$ and $Z$, is represented by the edges between their corresponding nodes. The indirect sharing between $L$ and $Y$ is modelled by the fact that $n_Y$ is reachable from $n_L$ through the directed edges. The sharing between $L$ and $K$ is represented by the fact that $n_Z$ is reachable from both $n_L$ and $n_K$. \hfill $\square$
5 Region Points-To Analysis

The region points-to analysis aims at computing for each procedure in a Mercury program a region points-to graph that represents the locations of its variables and the sharing among them.

The region points-to analysis is unification-based and flow-insensitive, i.e. the execution order of the atomic goals in a procedure does not matter, and consists of an intraprocedural analysis and an interprocedural analysis. Both analyses make use of the unify operation shown in Algorithm 1, whose task is capture sharing between two nodes in a region points-to graph. This algorithm should be invoked when the analyses learn that two variables whose nodes are $n$ and $m$ respectively can refer to the same storage; it will update the points-to graph by unifying the two nodes, i.e. merging them into one. To ensure that there is only one out-edge with a specific label from any given node, unifying two nodes will cause their corresponding child nodes to be unified as well, unless they are the same node already.

**Algorithm 1** $\text{unify}(n, m)$

**Require:** $G(N, E)$, $n, m \in N$.

**Ensure:** $G(N, E)$ with $n$ representing the unified node.

$N = N \setminus \{m\}$
vars($n$) = vars($n$) $\cup$ vars($m$)
for all $(m, (f, i), k) \in E$ do
  $E = E \setminus \{(m, (f, i), k)\}$
  if $(n, (f, i), k) \not\in E$ then
    $E = E \cup \{(n, (f, i), k)\}$
  end if
end for
for all $(k, (f, i), m) \in E$ do
  $E = E \setminus \{(k, (f, i), m)\}$
  if $(k, (f, i), n) \not\in E$ then
    $E = E \cup \{(k, (f, i), n)\}$
  end if
end for
for all $l, l' \in N$ do
  if $(n, (g, j), l) \in E \land (n, (g, j), l') \in E \land l \neq l'$ then
    $\text{unify}(l, l')$
  end if
end for
We will describe the analyses in turn with the assumption that we are analyzing a procedure \( p \).

Recall that, when describing the static region analysis and transformation, for convenience, we make the assumption that all terms are stored on the heap and therefore we need regions for them. In a concrete implementation, such as ours inside the MMC (Sections 8 and 9), if certain terms do not need heap storage, their corresponding regions can just be ignored.

### 5.1 Intraprocedural Analysis of a Procedure

The intraprocedural analysis initializes \( G_p \) and then captures the sharing created by the explicit unifications. Its definition is in Algorithm 2. (See section 2.2 for the definition of superhomogeneous form.)

**Algorithm 2** \( \text{intraproc}(p) \): intraprocedural analysis of a procedure \( p \)

**Require:** \( p \) is in superhomogeneous form.

**Ensure:** The sharing created by explicit unifications is represented in \( G_p \).

\[
G_p = (\emptyset, \emptyset)
\]

for all \( X \in p \) do

\[
G_p = G_p \sqcup \text{init}\_\text{ptg}(X)
\]

end for

for all \( \text{unif} \in p \) do

if \( \text{unif} \equiv (X := Y) \) then

\[
\text{unify}(n_X, n_Y)
\]

else if \( \text{unif} \equiv (X \Rightarrow f(Y_1, \ldots, Y_n) \text{ or } X \Leftarrow f(Y_1, \ldots, Y_n)) \) then

for \( i = 1 \) to \( n \) do

\[
\text{unify}(\text{node}(n_X, (f, i)), n_Y)
\]

end for

end if

end for

As we know the type of each variable in \( p \), we initialize \( G_p \) by using the \( TG \) graphs of the variables. In Algorithm 2, we use a function \( \text{init}\_\text{ptg}(X) \) that

- generates a region points-to graph for \( X \) from the type-based region graph of the type of \( X \), \( TG_{\text{type}}(X) \),
- sets the \text{vars} set of the node corresponding to the principal node in \( TG_{\text{type}}(X) \) to \( \{X\} \) and the \text{vars} set of all others nodes to the empty set,
- and generates a fresh region variable for each node in the region points-to graph.

The intraprocedural analysis then adds to \( G_p \) all the sharing created by the unifications in the procedure. For assignment, construction and deconstruction unifications we unify the nodes corresponding with the sharing created by them. We ignore test unifications because they do not create any sharing.
5.2 Interprocedural Analysis

The interprocedural analysis, Algorithm 3, updates $G_p$ by integrating into it the relevant region-sharing information from the region points-to graphs of the called procedures.

Algorithm 3 \textit{interproc}(p): interprocedural analysis of a procedure $p$

\textbf{Require:} $p$ is in superhomogeneous form.
\textbf{Ensure:} The sharing created by procedure calls is represented in $G_p(N_p, E_p)$.

\begin{verbatim}
repeat
  for all call sites in $p$ do
    Assume that the call is $q(Y_1, \ldots, Y_n)$, with $X_1, \ldots, X_n$ being the corresponding formal arguments, and that $G_q$ is available.
    \% Build an $\alpha$ relation.
    for $k = 1$ to $n$ do
      $\alpha(n_{X_k}) = n_{Y_k}$
    end for
    \% Ensure $\alpha$ is a function.
    for all $X_i, X_j$ do
      if $\alpha(n_{X_i}) = n_{Y_i}$ \& $\alpha(n_{X_j}) = n_{Y_j}$ \& $n_{X_i} = n_{X_j}$ \& $n_{Y_i} = n_{Y_j}$ then
        unify($n_{Y_i}, n_{Y_j}$)
      end if
    end for
    \% Integrate sharing in $G_q$ into $G_p$.
    In the graph $G_q$, do a depth-first traversal starting from each $n_{X_i}$, visiting each node only once and applying the rules P1 and P2 in Figure 8 when applicable.
  end for
until There is no change in either $G_p$ or in any of the $\alpha$ functions.
\end{verbatim}

Consider a call $q(Y_1, \ldots, Y_n)$ in the body of $p$, with the head of the called procedure being $q(X_1, \ldots, X_n)$. Any region-sharing among the $X_i$ in $G_q$ may not currently be present in $G_p$ as region-sharing among the $Y_i$. The interprocedural analysis makes sure that any such sharing in $G_q$ will be copied to $G_p$. First, it builds the function $\alpha: N_q \rightarrow N_p$ that maps the nodes of the formal arguments ($X_i$'s) to the nodes of the corresponding actual arguments ($Y_i$'s). Then these nodes are the starting points for the integration of the remaining region-sharing. This is done by following the relevant edges in $G_q$ to extend the $\alpha$ function to all the relevant nodes in $G_q$ (rule P2) and to unify the relevant nodes in $G_p$ (rule P1).

For a whole program, we start by performing the intraprocedural analysis for every procedure. Since our interprocedural analysis propagates information only upwards, from the graphs of callees to those of callers, we compute the strongly connected components of the call-dependency graph and analyze the components in bottom-up order. Algorithm 4 illustrates this approach.

The points-to graphs of the \texttt{split} and \texttt{qsort} procedures in the \texttt{quicksort} program in Example 2 are shown in Figure 9. For \texttt{split}, the region points-to analysis detects
Algorithm 4 Region points-to analysis of a program

Require: A Mercury program $P$ with its procedures in superhomogeneous form.
Ensure: Region points-to graphs for all procedures.

for all procedure $p$ in $P$ do
  intraproc($p$)
end for

Compute the strongly connected components (SCCs) of $P$’s call-dependency graph.

for all SCCs in bottom-up order do
  repeat
    for all $p$ in SCC do
      interproc($p$)
    end for
  until we have reached a fixpoint
end for

5.3 Correctness of the Region Points-To Graphs

In this section we have a number of theorems about the correctness of our algorithms. The proofs of these theorems can be found in the corresponding section in a longer version of this paper that is available in the CoRR repository as http://arxiv.org/abs/1203.1392.

Theorem 1
The region points-to analysis of a program terminates.

**Theorem 2**
The graphs that result from the region points-to analysis of a program represent all the locations of the terms that can possibly be constructed during the execution of the program, and the possible sharing among the terms.

In the rest of the paper, when we mention region points-to graphs, we mean the ones obtained by the region points-to analysis of the program.

### 5.4 Regions that a Procedure Allocates Into

During the region points-to analysis of a procedure, we can track the regions that are possibly allocated into in the procedure. A construction unification is the only construct in Mercury that allocates memory. When processing a construction unification $X \leftarrow f(\ldots)$ we mark the node $n_X$ as *allocated*. When two nodes are unified, if one node is marked as allocated then the unified node is also marked as allocated. At a call site, if a node $n$ reachable from a formal parameter in the callee is marked as allocated, and $\alpha(n) = m$, then we mark $m$ in the caller as allocated as well. We call the set of nodes in procedure $p$ marked in this way $\text{allocation}(p)$. In the quicksort example of Figure 2 and Figure 9, $\text{allocation}(\text{split}) = \{R_3, R_4\}$, and $\text{allocation}(\text{qsort}) = \{R_8, R_9, R_{10}\}$.

### 6 Region Liveness Analysis

After the region points-to analysis, we know the region variables of each procedure and how the program variables are distributed over the regions to which these region variables are bound.

In this section, we construct a region liveness analysis that approximates the lifetimes of the region variables, i.e. their liveness, to decide when a region needs to be created and when it can safely be reclaimed. We make a distinction between local liveness and global liveness. Local liveness concerns the lifetime of the region variable inside the procedure itself, namely when we consider the procedure alone. Global liveness concerns liveness with respect to the whole program, namely when we take into account the call sites that call the procedure. We show how we compute local liveness in Section 6.2, while Section 6.3 shows how we compute global liveness.

#### 6.1 Technical Background

A region variable being live means that (a) it should be bound to a region, and (b) that region may possibly be used in future (forward) execution. During its lifetime, the region bound to a region variable may be allocated into by procedures other than the one that created the region, so we often need to pass region variables as arguments of procedures.

Consider a procedure $p$. We associate a *program point* with every atomic goal in
the body of $p$. An execution path in $p$ is a sequence of program points, such that at runtime the atomic goals associated with these program points are executed in sequence. We denote an execution path by $\langle \text{atom}_1, \ldots, \text{atom}_n \rangle$, in which the $\text{atom}_i$’s are the atomic goals involved, and the indexes $i$’s are a dense sequence giving the order among the atomic goals in this execution path. The function $pp(\text{atom})$ returns the program point associated with $\text{atom}$. We use the notions before and after a program point. Before a program point means right before the associated atomic goal is going to be executed; while after a program point means its atomic goal has just been completed. The set of live region variables at a program point is computed via the set of live variables at the program point. We also use two functions, $in\_args(\text{atom})$ and $out\_args(\text{atom})$, that respectively return the sets of input and output arguments of $\text{atom}$. For specialized unifications they are defined in Table 1. If $\text{atom}$ is a procedure’s head, they return formal parameters, whereas if $\text{atom}$ is a call they return actual parameters. Those sets can be computed from the mode information of Mercury procedures.

<table>
<thead>
<tr>
<th></th>
<th>$in_args$</th>
<th>$out_args$</th>
</tr>
</thead>
<tbody>
<tr>
<td>construction</td>
<td>$X \leftarrow f(X_1, \ldots, X_n)$</td>
<td>${X_1, \ldots, X_n}$</td>
</tr>
<tr>
<td>deconstruction</td>
<td>$X \Rightarrow f(X_1, \ldots, X_n)$</td>
<td>${X}$</td>
</tr>
<tr>
<td>test</td>
<td>$X \leftarrow Y$</td>
<td>${X, Y}$</td>
</tr>
<tr>
<td>assign</td>
<td>$X := Y$</td>
<td>${Y}$</td>
</tr>
</tbody>
</table>

### 6.2 Live Region Variables at a Program Point

In this subsection we specify the analysis that computes the local liveness of region variables in a procedure. We express local liveness by the sets of region variables that are live before and after every program point in a procedure. The liveness of a region variable at a program point is determined by the liveness of the variables that are stored in the corresponding region.

**Live variables.** A variable is live before a program point if it has been instantiated before the point and may be used in the goal associated with the program point or after it. A variable is live after a program point if it has been instantiated before or at the point and may be used after the point.

The live variable analysis for a procedure $p$ is defined in Algorithm 5. It traverses each execution path ($ep$) backwards, starting with the last program point, computing sets of live variables along the way. At each program point, we update its $LV_{after}$ and $LV_{before}$ sets. The $LV_{after}$ of the last program point(s) is defined to be $out\_args(p)$, while the $LV_{before}$ of the first program point(s) will be $in\_args(p)$. This assumes that every procedure uses all its arguments, but since we run this
Algorithm 5 \texttt{lva}(p): live variable analysis of a procedure \texttt{p}.

\textbf{Require}: \texttt{p} in superhomogeneous form.
\textbf{Ensure}: The sets of live variables before (\texttt{LV}_{\text{before}}) and after (\texttt{LV}_{\text{after}}) all program points in \texttt{p}.

\begin{algorithmic}
\ForAll{program points \texttt{i} in \texttt{p}}
    \State \texttt{LV}_{\text{before}}(i) = \texttt{LV}_{\text{after}}(i) = \emptyset \\
\EndFor
\ForAll{\texttt{ep} \equiv \langle \texttt{atom}_1, \ldots, \texttt{atom}_n \rangle \text{ in } \texttt{p}}
    \For{$j = n$ \textbf{down to} $1$}
        \State \texttt{i} = \texttt{pp} (\texttt{atom}_j)
        \If{$j = n$}
            \State \texttt{LV}_{\text{after}}(i) = \texttt{out} \texttt{args}(\texttt{p})
        \Else
            \State \texttt{LV}_{\text{after}}(i) = \texttt{LV}_{\text{after}}(i) \cup \texttt{LV}_{\text{before}}(\texttt{pp} (\texttt{atom}_{j+1}))
        \EndIf
    \If{$j = 1$}
        \State \texttt{LV}_{\text{before}}(i) = \texttt{in} \texttt{args}(\texttt{p})
    \Else
        \State \texttt{LV}_{\text{before}}(i) = (\texttt{LV}_{\text{after}}(i) \setminus \texttt{out} \texttt{args}(\texttt{atom}_j)) \cup \texttt{in} \texttt{args}(\texttt{atom}_j)
    \EndIf
\EndFor
\EndFor
\end{algorithmic}

Analysis after a Mercury compiler pass that removes unused arguments, this is a justified assumption.

\textbf{Live region variables}. A region variable is live before (after) a program point if its node is reachable from a variable that is live before (after) the program point.

The set of nodes that are reachable from a variable \texttt{X} is defined as follows:

\[ \text{Reach}(X) = \{n_X\} \cup \{m \mid \exists(n_X, label_0, n_1), \ldots, (n_{i-1}, label_{i-1}, n_i) \in E \land m = n_i\}. \]

The live region variable analysis of a procedure is specified in Algorithm 6. This algorithm computes the sets of live region variables before (\texttt{LR}_{\text{before}}) and after (\texttt{LR}_{\text{after}}) each program point as the unions of the \texttt{Reach} sets of all variables in the \texttt{LV}_{\text{before}} and in \texttt{LV}_{\text{after}} sets of the program point, respectively.

Algorithm 6 \texttt{tra}(p): live region variable analysis of a procedure \texttt{p}

\textbf{Require}: \texttt{LV}_{\text{before}} and \texttt{LV}_{\text{after}} of all program points in \texttt{p}.
\textbf{Ensure}: The sets of live region variables before (\texttt{LR}_{\text{before}}) and after (\texttt{LR}_{\text{after}}) all program points in \texttt{p}.

\begin{algorithmic}
\ForAll{program points \texttt{i} in \texttt{p}}
    \State \texttt{LR}_{\text{before}}(i) = \texttt{LR}_{\text{after}}(i) = \emptyset \\
    \ForAll{\texttt{X} \in \texttt{LV}_{\text{before}}(i)}
        \State \texttt{LR}_{\text{before}}(i) = \texttt{LR}_{\text{before}}(i) \cup \text{Reach}(\texttt{X})
    \EndFor
    \ForAll{\texttt{X} \in \texttt{LV}_{\text{after}}(i)}
        \State \texttt{LR}_{\text{after}}(i) = \texttt{LR}_{\text{after}}(i) \cup \text{Reach}(\texttt{X})
    \EndFor
\EndFor
\end{algorithmic}
6.3 Lifetime of Regions across Procedure Boundary

Sometimes we have to pass region variables between procedures. For a procedure, the region variables reachable from its arguments are all candidates to be region arguments. But as we will see later, not all of them may actually need to be arguments. This subsection introduces an analysis that, by looking at the calling contexts of a procedure in the whole program, decides which region variables become live or become dead inside the procedure. With this global liveness information, we can give regions shorter lifetimes, achieving better memory reuse.

Consider a procedure \( q \) that is called by some procedure \( p \). We define:

- \( \text{bornR}(q) \) is the set of region variables of \( q \) that are mapped (by the \( \alpha \) function at the call site) to region variables of \( p \) that definitely become live inside \( q \), i.e. in the code of \( q \) or in one of the procedures \( q \) calls.
- \( \text{deadR}(q) \) is the set of region variables of \( q \) that are mapped to region variables of \( p \) that definitely cease to be live (i.e. they become dead) inside \( q \).
- \( \text{outlivedR}(q) \) is the set of region variables of \( q \) that are mapped to region variables of \( p \) that outlive the call to \( q \). They are live before the call and are still live after the call.

The idea is that, in the transformed program, the region variables in \( \text{bornR}(q) \) will get bound to a region inside \( q \) and \( q \) will return the bound region variable to \( p \), while the region variables corresponding to \( \text{deadR}(q) \) are passed by \( p \) to \( q \) and have their regions safely removed during the call to \( q \). The alternative would be that \( p \) creates the regions corresponding to \( \text{bornR}(q) \) just before the call to \( q \), and removes the regions corresponding to \( \text{deadR}(q) \) right after the call. With that approach, many regions would have a longer lifetime, which is why we prefer to create regions as late as possible and remove them as soon as possible.

For a procedure \( q \), we initially set \( \text{bornR}(q) = \text{outputR}(q) \setminus \text{inputR}(q) \) and \( \text{deadR}(q) = \text{inputR}(q) \setminus \text{outputR}(q) \), where \( \text{inputR}(q) \) and \( \text{outputR}(q) \) are the sets of region variables reachable from the variables in \( \text{inargs}(q) \) and \( \text{outargs}(q) \), respectively. This is an overestimate in which all the region variables that contain input terms but are not involved with output terms are assumed to become dead in \( q \), while all the region variables where output terms are stored but are not yet bound at the entry of \( q \) are assumed to become live in \( q \). We use \( \text{localR}(q) \) to denote the set of the region variables that are local to \( q \) (not reachable from input or output variables); it is computed by \( N_q \setminus (\text{inputR}(q) \cup \text{outputR}(q)) \). Initially, \( \text{outlivedR}(q) = \text{inputR}(q) \cap \text{outputR}(q) \). It is clear that \( \text{localR}(q), \text{bornR}(q), \text{deadR}(q), \text{outlivedR}(q) \) form a partition of \( N_q \).

The calling contexts of a procedure influence what it can do to its non-local region variables. Therefore when analyzing a procedure \( p \), the analysis applies the rules in Figure 10 to any \( \text{atom} \) in \( p \) that is a call to \( q \). These rules update the \( \text{deadR} \) and \( \text{bornR} \) sets of \( q \) according to the calling context. Rule L1 requires a region variable to be moved from \( \text{deadR}(q) \) to \( \text{outlivedR}(q) \) if its region needs to be live in \( p \) after the call to \( q \). Rule L2 is there to avoid the problems that would arise if we let a region that is referred to by more than one region variable in \( q \) be
removed when one of those region variables becomes dead. Either that region can still be referred to through the other region variables, in which case we would have removed it too early, or the other region variables are also in \( \text{deadR}(q) \), in which case the region would be removed again. Repeated application of L2 will ensure that our system never removes aliased regions during the call to \( q \) through any of the region variables referring to them. Rule L3 is analogous to L1; it moves a region variable from \( \text{bornR}(q) \) to \( \text{outlivedR}(q) \) if it is already live before the call to \( q \). Rule L4 is analogous to L2 in the same way; just as we do not want to remove a region twice, we do not want to create it twice. Rules L2 and L4 together ensure that region variables that are involved in a region alias never belong to either \( \text{bornR} \) or \( \text{deadR} \) sets.

When there is a change to any of the sets of \( q \), \( q \) must be analyzed to propagate the change to the procedures it calls. Therefore, this analysis requires a fixpoint computation. After a fixpoint is reached, each procedure has exactly one \( \text{bornR} \) set and one \( \text{deadR} \) set, and these will be suited for its most restrictive calling context. For calls in a less restrictive context, some regions will be created or removed outside the call, which will mean that some regions will be created earlier than needed and/or some other regions will be removed later than needed. For call sites that are sufficiently heavily used, we could avoid the inefficiency inherent in that by creating a specialized copy of the callee that exactly matches the caller’s context, but this could be fairly expensive, since it may (and generally will) require specialized copies of many of the specialized callee’s descendants as well.

In the \texttt{quicksort} program from Figure 1, \texttt{split} has three execution paths: \(( (1), (2), (3)) \), \(( (4), (5), (6), (7)) \), and \(( (4), (8), (9)) \), while \texttt{qsort} has two paths: \(( (1), (2)) \) and \(( (3), (4), (5), (6), (7)) \).\footnote{Note that the third execution path of \texttt{split} does not contain the test at (5) because of the semantics of if-then-else. The \( \text{LV} \) and \( \text{LR} \) sets of \texttt{split} are in Table 2(a), while the sets of \texttt{qsort} are in Table 2(b) (see also Figure 2 and Figure 9). In this example, the sets after one program point are always equal to the corresponding sets before the next point in the execution path. However, this is not true in all cases. Consider the last program point before a disjunction. The set of live variables after this point contains the region variables.}
Table 2: Live variable and live region variable sets in the *quicksort* program.

(a) split

<table>
<thead>
<tr>
<th>pp</th>
<th>LV</th>
<th>LR</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1_a)</td>
<td>{X, L}</td>
<td>{R5, R1, R2}</td>
</tr>
<tr>
<td>(1_a, 2_a)</td>
<td>{}</td>
<td>{}</td>
</tr>
<tr>
<td>(2_a, 3_a)</td>
<td>{L1}</td>
<td>{R3, R2}</td>
</tr>
<tr>
<td>(3_a)</td>
<td>{L1, L2}</td>
<td>{R3, R2, R4}</td>
</tr>
<tr>
<td>(4_a)</td>
<td>{X, L}</td>
<td>{R5, R1, R2}</td>
</tr>
<tr>
<td>(4_a, 5_a)</td>
<td>{X, Le, Ls}</td>
<td>{R5, R2, R1}</td>
</tr>
<tr>
<td>(5_a, 6_a)</td>
<td>{X, Le, Ls}</td>
<td>{R5, R2, R1}</td>
</tr>
<tr>
<td>(6_a, 7_a)</td>
<td>{L2, Le, L11}</td>
<td>{R4, R2, R3}</td>
</tr>
<tr>
<td>(7_a)</td>
<td>{L1, L2}</td>
<td>{R3, R2, R4}</td>
</tr>
<tr>
<td>(4_a, 8_a)</td>
<td>{X, Le, Ls}</td>
<td>{R5, R2, R1}</td>
</tr>
<tr>
<td>(8_a, 9_a)</td>
<td>{L1, Le, L21}</td>
<td>{R3, R2, R4}</td>
</tr>
<tr>
<td>(9_a)</td>
<td>{L1, L2}</td>
<td>{R3, R2, R4}</td>
</tr>
</tbody>
</table>

(b) qsort

<table>
<thead>
<tr>
<th>pp</th>
<th>LV</th>
<th>LR</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1_a)</td>
<td>{L, A}</td>
<td>{R6, R7, R8}</td>
</tr>
<tr>
<td>(1_a, 2_a)</td>
<td>{A}</td>
<td>{R8, R7}</td>
</tr>
<tr>
<td>(2_a)</td>
<td>{}</td>
<td>{R8, R7}</td>
</tr>
<tr>
<td>(3_a)</td>
<td>{L, A}</td>
<td>{R6, R7, R8}</td>
</tr>
<tr>
<td>(4_a, 4_b)</td>
<td>{A, Le, Ls}</td>
<td>{R8, R7, R6}</td>
</tr>
<tr>
<td>(4_a, 5_b)</td>
<td>{A, Le, L1, L2}</td>
<td>{R8, R7, R9, R10}</td>
</tr>
<tr>
<td>(5_a, 6_a)</td>
<td>{Le, L1, S2}</td>
<td>{R9, R7, R8}</td>
</tr>
<tr>
<td>(6_a, 7_b)</td>
<td>{L1, A1}</td>
<td>{R9, R7, R8}</td>
</tr>
<tr>
<td>(7_b)</td>
<td>{}</td>
<td>{R8, R7}</td>
</tr>
</tbody>
</table>

Table 3: Partition of the set of region variables.

<table>
<thead>
<tr>
<th></th>
<th>localR</th>
<th>bornR</th>
<th>deadR</th>
<th>outlivedR</th>
</tr>
</thead>
<tbody>
<tr>
<td>split</td>
<td>{}</td>
<td>{R3, R4}</td>
<td>{R1}</td>
<td>{R2, R5}</td>
</tr>
<tr>
<td>qsort</td>
<td>{R9, R10}</td>
<td>{}</td>
<td>{R6}</td>
<td>{R7, R8}</td>
</tr>
</tbody>
</table>

that are live in any of the disjuncts; in general, some of these variables will be live in only some of the disjuncts, not all.

When computing the *deadR* and *bornR* sets of these procedures, the initial partition is changed only once, when \(R5\) is removed from *deadR*(split) by an application of rule L1 to the call to *split* inside *qsort*. The final result is as in Table 3.

### 6.4 Correctness

Algorithm 6, the algorithm that detects live region variables locally at each program point is an extension of live variable analysis, which is a standard, well-known program analysis (Nielsen et al. 1999). Theorem 2 guarantees that the locations of variables and their possible sharing are represented in the region points-to graphs. Therefore Algorithm 6 computes all the live region variables by starting from the live variables and collecting all the reachable region variables using the region points-to graphs.

The analysis in Section 6.3 aims to compute a shortest possible lifetime for a
region. Its termination follows from the facts that each procedure uses a finite set of region variables (which guarantees that the initial bornR and deadR sets are finite), and that the analysis only ever reduces the sizes of these sets. The rules in Figure 10 enforce all the cases where a caller of a procedure needs to restrict what the callee can do to its region variables. The eager application of the rules therefore ensures that after a fixpoint has been reached, the bornR and deadR sets obtained for a procedure will respectively contain exactly the region variables that the procedure will safely create and remove.

7 Program Transformation

The purpose of the program transformation is to annotate all the procedures in the program with the information the code generator needs about regions. For each procedure, the tasks of the transformation are:

- extend the procedure definition with the formal region arguments;
- extend its procedure calls with the corresponding actual region arguments;
- annotate each construction unification with the region variable representing the region into which the new memory cell should be put;
- insert instructions to create and remove regions at suitable points.

The third task is straightforward because the new cell is always put into the region associated with the variable on the left hand side of the construction unification, and the map from variables to the region variables representing their regions is available after the region points-to analysis.

We elaborate the other tasks in the next three subsections.

7.1 Region Arguments

The region variables in bornR and deadR must be arguments because their regions will be created and removed inside the procedure. Besides these region variables, we also need to pass as arguments the region variables that are reachable from the input and output variables and are allocated into in the procedure. This set of arguments, which we call allocR, is therefore computed by \( allocR = (inputR \cup outputR) \cap allocation \) (Section 5.4). Note that allocR is not necessarily disjoint with any of bornR, deadR and outlivedR.

So all in all, the set of formal region arguments of a procedure is deadR \( \cup \) bornR \( \cup \) allocR. In the quicksort program, \( allocR(split) = \{R1, R2, R3, R4\} \cap \{R3, R4\} = \{R3, R4\} \), \( allocR(qsort) = \{R6, R8\} \cap \{R8\} = \{R8\} \), and the region arguments are \( \{R1\} \cup \{R3, R4\} \cup \{R3, R4\} = \{R1, R3, R4\} \) for split and \( \{R6\} \cup \emptyset \cup \{R8\} = \{R6, R8\} \) for qsort.

The actual region arguments of a procedure call are computed simply by looking up the formal region arguments of the called procedure and applying the \( \alpha \) function of the call site.
Region-Based Memory Management for Mercury Programs

7.2 Insertion of create and remove Instructions

Regions are created and removed only by the create and remove instructions respectively. When a region is created, the region variable in the create instruction is bound to it. Removing a region consists of calling remove on the region variable bound to the region. We implement create and remove as builtin Mercury procedures. Calls to other procedures may also create and remove regions, but only if those procedures directly or indirectly invoke create or remove. Unifications can never either create or remove regions.

7.2.1 Transformation Rules

The transformation rules in Figure 11 make use of the local and global liveness of region variables to introduce create and remove instructions when necessary.

**Creation rules T1 and T2.** A region variable will never become locally live between atomic goals; a region cannot be not live after a program point but live before the immediately next program point in some execution path. A region variable can become locally live only within atomic goals. Let this be the atomic goal atom at program point \( i \) in procedure \( p \). T1’s first condition says that this rule covers the case where atom is a call, for example to \( q \). The second condition is true for a region \( r \) that is not live before atom but is live after atom. The third condition checks whether \( p \) itself is allowed to create the region. It is intuitively clear that \( p \) needs to create regions bound to region variables in born\( R(p) \) and local\( R(p) \). The reason why we also allow \( p \) to create regions in dead\( R(p) \) is that it is OK for \( p \) to remove the region bound to \( r \) at some point before atom, if that is safe, and then recreate \( r \) right before atom. The new region will be removed later because \( r \) is in dead\( R(p) \). Such deletion-followed-by-recreation is not allowed for regions in outlived\( R(p) \) because the caller needs their contents. The fourth condition checks whether the call will create the region; if it will, then \( p \) itself need not do so. Overall, if the third condition is false, then \( p \)’s caller will have created the region; if the third condition is true, but the fourth condition is false, then \( q \) will create the region; if

<table>
<thead>
<tr>
<th>Rule</th>
<th>Description</th>
</tr>
</thead>
</table>
| (T1) | \( r \in LR_{after}(pp(\text{atom})) \setminus LR_{before}(pp(\text{atom})) \) \( r \in \text{local}(p) \cup \text{born}(p) \cup \text{dead}(p) \)
| \( r = \alpha(r') \rightarrow r' \notin \text{dead}(q) \) |
| | add “create(r)” before atom |
| (T2) | \( atom \equiv \{ \ldots \} \) \( r \in LR_{after}(pp(\text{atom})) \setminus LR_{before}(pp(\text{atom})) \)
| \( r \in \text{local}(p) \cup \text{born}(p) \cup \text{dead}(p) \) |
| | add “create(r)” before atom |
| (T3) | \( atom \equiv \{ \ldots \} \) \( r \in LR_{before}(pp(\text{atom})) \setminus LR_{after}(pp(\text{atom})) \) \( r \in \alpha(r') \rightarrow r' \notin \text{dead}(q) \) |
| | add “remove(r)” after atom |
| (T4) | \( atom \equiv \text{unif} \) \( r \in LR_{before}(pp(\text{atom})) \setminus LR_{after}(pp(\text{atom})) \) \( r \in \text{local}(p) \cup \text{dead}(p) \cup \text{born}(p) \) |
| | add “remove(r)” after atom |
| (T5) | \( atom \equiv \{ \ldots \} \) \( r \in LR_{after}(pp(\text{atom}')) \setminus LR_{before}(pp(\text{atom}')) \) \( r \in \text{local}(p) \cup \text{dead}(p) \cup \text{born}(p) \) |
| | add “remove(r)” before atom |
| (T6) | \( atom \equiv \{ \ldots \} \) \( r \in \text{local}(p) \cup \text{dead}(p) \cup \text{born}(p) \) |
| | add “remove(r)” after atom |

Fig. 11: Transformation rules.
both the third and fourth conditions are true, then the instruction that T1 inserts before the call will create the region.

Rule T2 covers the case where a region becomes live in a unification. The first condition looks only for construction unifications because for all other kinds of unifications, the second condition always fails. T2 is analogous to T1, the main difference being that unifications can never create regions.

Removal Rules T3, T4, and T5. Removal rule T3 is analogous to creation rule T1. If a region variable locally ceases to be live during a call, the situation described by the first and second conditions, what happens is governed by the third and fourth conditions. If the third condition is false, then p’s caller or one of its ancestors will (eventually) remove the region; if the third condition is true, but the fourth condition is false, then q will remove the region; if both the third and fourth conditions are true, then the instruction that T3 inserts after the call will remove the region. Note that it is OK for p to remove a region in bornR(p), a region it must have previously created; since the region will be live at the end of p, p will later create it again, and that is all that p’s caller expects.

Removal rule T4 is likewise analogous to creation rule T2, but a region can become dead in any kind of unification, not just constructions.

While a region cannot be not live after one program point and then magically become live before an immediately following program point, it is possible for a region to be live after one program point (atom in T5) and dead before an immediately following program point (atom’). This can happen e.g. when the following program point is the first goal of a disjunct in a disjunction or switch, and the region is live in other disjuncts of the disjunction or switch. In that case, the region is live after atom because it is live in some execution paths that do not include atom’ . In such cases, rule T5 removes the region before atom’, provided as usual that p is allowed to do so.

Handling instantly-dead variables: rule T6. In some cases, a variable may be instantiated at some point but then never used after that. We call them instantly-dead variables. In logic programming in general and in Mercury in particular, they can be void or singleton variables. A void variable’s name starts with the underscore (see e.g. the first clause of split in Figure 1) to explicitly tell the compiler that we do not care about its value. A singleton variable is a variable that occurs exactly once in a clause whose name does not start with an underscore. Singleton variables often represent mistakes, so the Mercury compiler issues a warning for them; programmers who believe the code to be correct can avoid the warning by adding a leading underscore, turning the singleton into a void variable.

Because it is useless to do a construction unification that binds the new term to an instantly-dead variable, we assume that such unifications are eliminated before our region analysis and transformation; the Mercury compiler has an optimization that does this. However, this is not a full solution. A procedure can return several output arguments, and it may be that the caller ignores some and pays attention only to the others. The ignored arguments pose a problem for our analysis. Being instantiated means that we need regions to store their terms, and of course we want those regions to eventually be removed. However, the fact that the ignored arguments are not
% p(in, out). % q(in, out).
\begin{verbatim}
p(A, B) :- q(X, Y) :- length(L) = N :-
  if ( if L == [], A == 1 then then ; B := C (3) V := X L => [_ | T],
(5) Y := Z + length(V).
\end{verbatim}

Fig. 12: Effect of re-creation of regions.

\begin{verbatim}
p(A, B@R1) :- q(X@R2, Y) :- create(R1), (1) Z := length(X),
  if ( if in R1, (if (2) Z == 1 then then (3) V := X else
    else remove(R2), remove(R1), create(R2),
    create(R1), (4) V <= [1] in R2 (4) Y <= [1] in R2,
    (5) Y := Z + length(V), remove(R2)).
\end{verbatim}

Fig. 13: Effect of re-creation of regions: region-annotated version.

used in the future makes them never live according to our concept of live variables (Section 6). Therefore we may not rely on the change of their liveness from live to dead (the basis of rules T3-T5) to remove the regions storing their terms. That is why we have rule T6, which tries to remove regions reachable from void variables right after the point where the void variables get instantiated. We assume that at each program point \( i \), we have available the set of such instantly-dead variables, \( VV(i) \) (\( i \) is the point at which they get instantiated). We then compute \( VR(i) \), the set of region variables that are reachable from the variables, by \( \bigcup_{V \in VV(i)} Reach(V) \).

The basic idea of T6 is to remove the region of a region variable reachable from an instantly-dead variable right after the point where the variable gets instantiated, provided of course that the region variable is not reachable from any of the live variables after the point.

**Example of re-creation and re-removal.** We illustrate (a) creating, removing and recreating a region on the one hand and (b) removing, creating, re-removing a region on the other hand using the two procedures in Figure 12 and their region-annotated counterparts in Figure 13. For completeness, we include the definition of the function \( length \), which returns the number of elements of the input list, though its code is not important in this case. We also assume that there is no region for integers. Therefore the focus is only on the variables \( B \) and \( C \) in the procedure \( p \) and \( V \) and \( X \) in \( q \), which are of the type \texttt{list_int} (see Example 1). Each pair of them is assigned to the same region variables, \( R1 \) in \( p \) and \( R2 \) in \( q \) due to the assignments
at the program points (3) in both procedures. \( p \) and \( q \) are unrelated; we use them to demonstrate different situations.

Assume that \( p \) can create \( R_1 \), i.e. no calling context forces it otherwise. So \( R_1 \) is in \( \text{bornR}(p) \). In Figure 13, the \textit{create} instructions added for it before (1) and (4) are due to the rule T2. The \textit{remove} instruction added before (4) is due to rule T5. If execution reaches the else branch, the \( R_1 \) that was live after (1) is no longer live before (4), and we can reclaim the memory occupied by \( \lbrack 1 \rbrack \) by removing this incarnation of \( R_1 \), before creating a new incarnation of it and putting \( \lbrack 2 \rbrack \) into it.

For \( q \), assume that \( R_2 \) is in \( \text{deadR}(q) \). \( R_2 \) is not live before the program point (4), and the \textit{remove} instruction there is added by rule T5. As \( R_2 \) is live after (4), T2 adds the \textit{create} instruction there as well. The \textit{remove} instruction after (5) is added by rule T4. If execution reaches the else branch, we reclaim the memory of the input list \( X \) by removing \( R_2 \) before recreating it to construct \( V \).

In both cases, we need to make sure that the two operations before program point (4) are done in the right order. This is ensured by the following algorithm.

### 7.3 Insertion Algorithm

The insertion of the instructions is specified by Algorithm 7, which says how the transformation rules in Figure 11 should be applied to the atomic goal at each program point.

**Algorithm 7** Insertion of region instructions in a procedure \( p \).

**Require:** \( p \) in superhomogeneous form; all points-to graphs and region liveness sets are available.

```
for all program points \( i \) in \( p \) do
  \( \text{atom = atom}_{-\text{at}(i)} \)
  apply rule T6 to \( \text{atom} \)
  if \( \text{atom} \equiv \text{unif} \) then
    apply rule T4 to \( \text{atom} \)
    if \( \text{atom} \equiv X <\equiv f(\ldots) \) then
      apply rule T2 to \( \text{atom} \)
    end if
  else
    apply rules T1 and T3 to \( \text{atom} \)
  end if
end for
for all \( \text{ep} \equiv (\text{atom}_1, \ldots, \text{atom}_n) \) in \( p \) do
  for \( j = 1 \) to \( n - 1 \) do
    apply rule T5 to \( \text{atom}_j \), with \( \text{atom}' \equiv \text{atom}_{j+1} \)
  end for
end for
```

Each program point is associated with three sets of region instructions: a set of \textit{remove} instructions added before it, a set of \textit{create} instructions added before it, and a set of \textit{remove} instructions added after it. The instructions in the first set will be executed before the instructions in the second set.
The first loop in Algorithm 7 applies all the transformation rules except T5 to the atomic goals at all the program points in a procedure. We use the function $atom@i$ to refer to the atomic goal at program point $i$. While rule T6 can be applied to any atomic goal, T4 needs to be tried only when the atom at a program point is a unification, T2 only when the atom is a construction unification, and T1 and T3 only when the atom is a procedure call. The second loop follows every execution path to try rule T5, which needs to consult information at two consecutive program points at the same time.

The result of the program transformation of the quicksort program in Example 2 was shown in Figure 3. The additions of the remove instructions after the first program points in both qsort and split result from the applications of T4. The two create instructions in split were added by T2.

7.4 Correctness of Region-Annotated Programs

Region-annotating a program does not change its computational behavior; it changes only the locations of terms in memory. In this section we have a number of theorems about this. The proofs of these theorems can be found in the corresponding section in a longer version of this paper that is available in the CoRR repository as http://arxiv.org/abs/1203.1392.

Theorem 3
Consider a procedure $p$ in a program $P$. We call $P'$ the region-annotated program that is produced by applying the analyses and transformation in Sections 5, 6, and 7 to $P$, in which $p'$ is the region-annotated version of $p$. If a region variable is live before (after) a program point $i$ in $p'$, then in $P'$ it is bound to a region before (after) $i$.

Theorem 4
In region-annotated programs, allocations of memory, and the associated memory write accesses, are safe.

Theorem 5
When a variable appears as an input argument to an atomic goal at a program point, we say that the variable is read at that point. In region-annotated programs, when a variable is read at a program point, the term it is bound to is available.

8 Runtime Support for Regions During Forward Execution

We now describe the runtime support needed to execute region-annotated programs. In this section, we cover the support needed for forward execution, while in the next section we will look at the support needed for backward execution, i.e. backtracking. The latter is much more extensive, partly because our analyses in Section 6 determine liveness only with respect to forward execution.

Let us look at the lifespan of a region during forward execution. A region comes into existence with the execution of a $\text{create}(R)$ instruction that assigns memory
to the region and binds the region variable $R$ to a so-called region handle, which refers to the assigned memory. From then on, terms are allocated into the region by construction unifications annotated with $R$. When the memory referred to by the region handle bound to $R$ is no longer needed, the program will end the lifetime of $R$ by executing $\text{remove}(R)$, which reclaims that memory.

This aspect of our implementation is generally similar to the “standard” RBMM implementations for SML and Prolog, which are described in detail in (Makholm 2000a; Makholm 2000b). In our system, a region is a singly-linked list of fixed-size region pages. Each region page has a data area, an array of words that can be used to store program data, and a pointer to the next region page to form the singly-linked list. The handle of the region, which is how the rest of the system refers to it, is the address of the region header. Besides some other fields that we will introduce later, the header structure includes a region size record: a pointer to the newest region page, and a pointer to the next available word in the newest region page. Since region pages have a fixed size, these two values implicitly also specify the amount of free space in the newest region page. As is usual in RBMM systems, we store each region header at the start of the data area of its region’s first region page. Figure 14 shows a region with two region pages; the shaded areas represent memory allocated to user data.

There is no bound on the sizes of regions. When a region is created it will contain only one region page, but it can be extended by adding more region pages when necessary. The program maintains a global list of free region pages. If the free list runs out, the program requests a big chunk of memory from the operating system, divides it into region pages, and adds them to the free list. When a region needs to be extended, we take a region page from the free list and add it to the region as its new last region page, and then update the region’s size record. When a region is reclaimed, we return all its region pages to the free list. An allocation into a region always happens in its newest region page simply by incrementing the pointer to the next available word. When the amount of free memory in this region page is not enough for the allocation, we extend the region before allocating.

---

2 Storing the headers separately from the region pages would require the system that now keeps track of which region pages are free to also keep a separate free list for header-sized blocks. This would cause fragmentation that would not occur with the standard header-in-first-region-page design.
The advantage of this implementation is that the basic region management actions are bounded in time; even freeing all the region pages in a region can be done in constant time (we can destructively append the region’s list of pages to the free list in constant time because we maintain pointers to the tails as well as the heads of the lists). Disadvantages are that there is no natural size for the region pages (Tofte et al. 2004), and that if the remaining space of a region page is not enough for an allocation, that space will be wasted when a new region page is added.

Like most RBMM systems, we do not do garbage collection inside regions.

9 Runtime Support for Backtracking

Backtracking introduces two issues that need to be handled: reclaiming the memory allocated by the computations backtracked over, and ensuring that regions are reclaimed only when they are dead with respect to both forward and backward execution. The first issue obviously has to be handled at runtime. For our initial implementation, we have chosen to deal with the second issue, backward liveness, in the runtime system too. We expect this to give us the insights we will need later to redesign the program analysis in Section 6 to handle backward liveness both safely and precisely. Moreover, our current system can serve as a reference for that work.

In Mercury, disjunctions are the main source of backtracking because they provide alternatives. However, backtracking is also possible in if-then-elses, since they are just a special kind of disjunction: 

\[
(if \ C \ then \ T \ else \ E) \ \text{is semantically equivalent to} \ (C, T; not \ some \ C, E).
\]

Operationally, Mercury will try \(C\). If \(C\) succeeds, Mercury executes \(T\); if \(C\) fails, it executes \(E\) as if \(C\) had never been tried. The handling of commit (Section 2.2) is related to the handling of backtracking because committing to a solution may prune some alternatives of relevant disjunctions. Therefore, we need to provide runtime support for backtracking in the context of these three language constructs.

The region-annotated program in Figure 15 illustrates our two tasks.

We constructed this program, which unfortunately has no intuitive meaning, to illustrate the interaction between regions and backtracking; we will use it as our running example when describing the runtime support. (We could find no equally useful real code of manageable size. Also, we include the definitions of \texttt{member} and \texttt{length} only for completeness; their behavior is of no importance in this example.)

Regarding the lifetime of the regions, \texttt{main} creates \(R1\) and \(R2\) before the constructions of the lists \(X\) and \(A\). \texttt{main} creates \(R3\) before the call to \texttt{p} at (3), and \texttt{p} will use this region to store the skeleton of \(Y\). All the \texttt{remove} instructions for regions are added after the last \texttt{forward} uses of the terms stored in them. \texttt{member} and \texttt{length} only read their input variables, so they need no region arguments. For \(p\), \texttt{deadR}(\(p\)) = \{\(R4\)\}, \texttt{bornR}(\(p\)) = \emptyset, \texttt{outlivedR}(\(p\)) = \{\(R5\), \(R6\)\}, and \texttt{allocation}(\(p\)) = \{\(R5\), \(R6\)\}.

**Task 1:** Preventing the reclamation of backward live regions. The condition of the if-then-else in \texttt{main} is the call to the semidet procedure \(p\). The RBMM transformation marks the region \(R1\) for removal in the call because it is forward dead (it is not used in the then part) even though it is backward live (it is used in
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main(!IO) :- :- pred p(list_int, list_int, list_int, list_int).
create(R1), :- node p(in, in, out, out) is semidet.
(1) X <= [1, 3, -1, 3] in R1, create(R2),
(2) A <= [-2] in R2, ( if
create(R3), (3) p(X@R4, A@R2, B@R2, Y@R3) :-
(2) H <= 0
then
(1) X => [H | T],
(4) A <= [-2] in R2, 
(5) then Y := U
(3) Y <= [H | T] in R5,
(6) else V := U
(4) io.write(B, !IO),
(7) io.write(A, !IO),
(8) if member(H, U)
(5) io.write(Y, !IO),
(9) else V <= [H | U] in R5
(6) remove(R4),
(7) remove(R2),
(10) else Y <= [H | Y1] in R6
(8) remove(R3)
(9) remove(R2)
).
).

% node(in, in), semidet
member(X, L) :- length(L) = N :-
L => [H | T],
( L => [], N := 0
H => X
; L => [H | T],
; member(X, T)
).

Fig. 15: Illustrating the interaction of regions and backtracking.

the else part). We must make sure that R1 is not actually removed while it is backward live. In this case, that means we need to delay the reclamation of R1 until we reach the then part, since it is not safe to destroy R1 if the condition fails. We therefore distinguish reclaiming a region, which makes the memory of the region available for other uses and thus potentially destroys its contents, from the operation of removing a region, which causes the region to be reclaimed only when it is safe to do so. Basically, a region is removed when it is forward dead, and it is reclaimed when it is both forward and backward dead.

Task 2: Reclaiming the memory used by backtracked-over computations. The call to p has two output arguments, B and Y. main tells p to put any cells for B in R2, and creates R3 so that p can put Y into it. If the condition succeeds, we must leave both regions alone. If the condition fails, we should restore R2 to its size before the condition, and we should reclaim R3 in its entirety.

We now define several runtime concepts that we will use in the rest of the paper.

Old vs new regions. A region is old with respect to a point during the execution of a program if it was created before that point, otherwise it is new with respect to that point. We also refer to old regions as the existing regions. To allow efficient checks whether a region is old or new, we maintain a global region sequence number counter (starting at one) and include a sequence number field in region headers. When we create a region, we timestamp it by setting its sequence number from the global counter, and increment the counter. When execution reaches a point in the program that sets up later backtracking, such as the entry point of a disjunction, we
save the current sequence number. Then all the regions which are created before that point, i.e. the old regions with respect to the point, will have their sequence numbers smaller than the saved value; the regions which are created after that point, i.e. the new regions with respect to the point, will have their sequence numbers greater than or equal to the saved value. When the program backtracks to that point, we can use the saved value to check whether a region has been created before or after the point. In the context of RBMM, the memory that we want to reclaim at a resumption point will be new allocations into existing regions, and new regions in their entirety (since they have been created by the computation we have just backtracked over).

**Region list.** To do instant reclaiming of new regions, knowing the sequence numbers of the new regions is not enough; we also need to reach them. We therefore link all the live regions into a doubly-linked region list (using two additional pointers in the region header). We maintain a global pointer to the head of the list, which will be the newest live region. When a region is created, we add it to the head of the region list; when a region is reclaimed, we remove it from the list. We maintain the invariant that the region list is ordered by regions’ creation time, newest first. To reclaim new regions, we can traverse the region list from its head and reclaim each region until we meet an old one.

**Region size snapshots.** To do instant reclaiming of new allocations into an existing region, we need the old size of the region. When we need to remember the size of a region at a point, we can save its region size record at that point.

**Protection.** We will prevent the destruction of backward live regions by protecting them so that when a removal happens to the region during forward execution, the removal will be ignored.

**Changes to live regions by a goal.** When providing support for backtracking, sometimes we want to know about the changes which may be caused by a goal to the set of regions the goal may refer to. This means we need to know about any new regions the goal creates, any live regions the goal removes, and any live regions in which the goal performs allocations. We refer to these sets of regions as the goal’s created, removed, and allocated sets, respectively. We have computed several sets of region variables for procedures, such as inputR, bornR, deadR, and allocation. The created, removed, and allocated sets of goals can be computed from these in a fairly straightforward manner, as shown by the following paragraphs.

**Changes to live regions by a goal: creation.** Only create instructions and procedure calls may create regions. A create instruction always creates the region in its argument. A procedure call will create the regions that are the actual region arguments corresponding to the formal arguments in the bornR set of the called procedure. For a compound goal, its created set is the set of all regions created inside it, either directly or through a procedure call, even if the region is also removed later, because at compile time we cannot know whether a removed region is actually reclaimed.

**Changes to live regions by a goal: removal.** We can similarly use remove instructions and the deadR sets of procedures to compute the removed set of each goal. Some of these regions may be removed, created and removed again. Since we
only care about the old regions which are removed inside a goal, we exclude regions created inside the goal (i.e. the goals created set) from its removed set.

**Changes to live regions by a goal: allocation.** A region is allocated into by construction unifications and by procedure calls. A construction unification will allocate into the region with which it is annotated. A procedure call possibly allocates into the regions of region variables that are mapped to by those in the procedure’s allocation set. Because we are only interested in allocations in old regions (allocations into new regions being reclaimed by reclaiming the whole region), we restrict the allocated set to the regions in $\text{inputR} \cap \text{allocation}$.

**Changes to live regions by a goal: an example.** Take the condition of the if-then-else in the procedure $p$ in Figure 15 as an example goal. We say that the region $R_4$ is removed in the condition because $R_4$ is live before the condition and $\text{remove}(R_4)$ has been added to the condition. Or take the condition of the if-then-else in main. We say region $R_3$ is created in the condition because $\text{create}(R_3)$ has been inserted into the condition, while region $R_1$ is removed in the condition because it is live before the condition and is removed in the call to $p$. We have $\text{allocation}(p) = \{R_5, R_6\}$, but while $R_5$ is an input argument of $p$, $R_6$ is not, so the only old region $p$ allocates into is $R_5$. So the allocation set of the condition in main is $R_2$, since $R_2 = \alpha(R_5)$.

We provide the runtime support for backtracking for a program by generating extra supporting code at the right places to achieve our goals. In the next three subsections we will describe in detail the support for disjunctions, if-then-elses, and commits.

### 9.1 Support for Disjunctions

The Mercury compiler supports only one search strategy: depth-first search with chronological backtracking, so that the disjuncts of each disjunction are tried in order. Given a disjunction $(g_1; \ldots; g_i; \ldots; g_n)$, we refer to $g_1$ as the first disjunct, to the $g_i$s for all $1 < i < n$ as middle disjuncts, and to $g_n$ as the last disjunct of the disjunction. We will also use “later disjunct” to refer to any $g_i$ for $i > 1$.

A disjunction can have any determinism. The most general determinism is of course nondet, but if one of the disjuncts always has at least one solution, then the disjunction as a whole does too, so a disjunction can also be multi. And if the disjunction has no outputs (which happens frequently for disjunctions in the conditions of if-then-elses), then the disjunction as a whole cannot have more than one solution, which means that it will be either det or semidet, depending on whether it has an always-succeeding disjunct. (Typical programs do not contain det disjunctions, since they are equivalent to true.)

For our purposes, the important distinction is between nondet and multi disjunctions on the one hand, in which backtracking may reach a later disjunct from code executed outside the disjunction, after the success of a previous disjunct, and semidet and det disjunctions on the other hand, in which backtracking to a
later disjunct is possible only from code within an earlier disjunct. Since we do not care about the minimum number of solutions of each disjunction, our support treats multi disjunctions the same as nondet ones and det disjuncts the same as semidet ones. In the following, we will therefore talk only about nondet and semidet disjunctions. We consider nondet disjunctions first, since they are more general.

Figure 16 shows in pseudo-code form the supporting code we add to a nondet disjunction. We insert code at the following points: (d1) which is the start of the first disjunct, (d2) which represents the start of every middle disjunct, and (d3) which is the start of the last disjunct. These code fragments communicate using shared data in what we call a disj frame. Each entry to a disjunction creates a new disj frame. Since multiple nested disjunctions can be active at the same time, we link these frames together to form the disj stack (this is possible due to chronological backtracking). The disj stack is not a separate stack; we reserve space for its frames in the usual stacks used by the Mercury language implementation. We maintain a global pointer to the top disj frame on the disj stack.

A disj frame has a fixed part and a nonfixed part. In Figure 17, the fixed part is the 4-slot box separated by a thick line from the nonfixed part. The four slots in the fixed part are:

- The prev_disj_frame slot holds the pointer to the previous disj frame, or null if there is none.
- The saved_seq_num slot holds the value of the global region sequence number at the time when the disjunction was entered.
- The num_prot_region field gives the number of regions which are protected by

---

Fig. 16: RBMM runtime support for nondet disjunctions.

---

3 Semidet code in Mercury never does deep backtracking; it only ever does local, shallow backtracking. Semidet procedures return a success/failure indication, which is then tested by the caller. An arm of a semidet disjunction can call nondet code, but only if that nondet code is wrapped in a commit (see later); the commit will convert any deep backtracks done by the code inside it to shallow backtracking for the code outside it.
a semidet disjunction (which we will discuss later). For a nondet disjunction, this slot will contain zero.

- The num_size_rec field gives the number of region size records saved in the nonfixed part.

```plaintext
| prev_disj_frame     | (previous disj frame) |
| saved_seq_num       | (saved sequence number) |
| num_prot_region     | (number of protected regions) |
| num_size_rec        | (number of saved region snapshots) |
| prot_region_id      | (handle of a protected region) |
| ...                |                           |
| snapshot_region_id  | (handle of a region in a snapshot) |
| snapshot_size_record| (snapshot size record of that region) |
| ...                |                           |
```

Fig. 17: The structure of a disj frame.

**Disj-protecting backward live regions.** Consider a region which was created before the execution of a disjunction. Assume that this region is removed during forward execution, either by the code of a disjunct, or after the success of that disjunct by code following and outside the disjunction, but that this region is backward live with respect to a later disjunct of the disjunction. In this case, we need to make sure that if the region is removed during forward execution, it will not be actually reclaimed. Of course, the instruction that removes the region may not be reached because forward execution may fail before it gets there. But in general, we have to assume that the remove instruction will be executed, and that if the region may be needed after backtracking, we will need to prevent it from being reclaimed during the forward execution. We achieve this by disj-protecting such regions as follows. At the start of the disjunction, at (d1), we push a disj frame on the disj stack and save the current global sequence number into the saved_seq_num slot of the disj frame. A region is disj-protected by a disj frame if its sequence number is smaller than the sequence number saved in that disj frame. The remove instruction will only reclaim a region if the region is not disj-protected. Due to chronological backtracking, the order of the frames on the disj stack always corresponds to the order of the creation of those frames. Together with the fact that the global region sequence number is monotonically increasing, this implies that if a region is protected by a disj frame, it is also protected by all the later frames on the disj stack. This invariant means that to check if a region is disj-protected or not, we only need to check if it is protected by the top disj frame.

The program will no longer backtrack into a disjunction after starting the execution of its last disjunct. This means that no regions need to be protected any more by this disjunction. Therefore, at the start of the last disjunct, at (d3), we disj-unprotect them by popping the disj frame. The regions which had previously been protected only by this disj frame will be reclaimed when execution reaches their remove instructions.
**Instant reclaiming of new regions.** When the program backtracks to a later disjunct, we want to reclaim all the regions that have been created during the computation that has just been backtracked over, i.e. all the regions that were created after entry to the disjunction. At (d1), we saved the global sequence number in the disj frame. Therefore at the start of a later disjunct of the disjunction, at (d2) and (d3), we just need to traverse the region list, and reclaim all the regions we see until we encounter a region whose sequence number indicates that it was created before the disj frame.

**Instant reclaiming of new allocations in old regions.** When arriving at a later disjunct, we want to restore all the regions that existed before the disjunction to the sizes they had when entering the disjunction, recovering any memory that has been allocated in them. For each old region, we need to save the region’s size record in the nonfixed part of the disjunction’s disj frame at (d1), so that we can restore the region’s size at (d2) and (d3). We need three slots for each region: one for the region handle so that we know to which region the saved record belongs, and the other two for the record itself (see Figure 17). To be able to loop through the saved records and restore the regions at (d2) and (d3), we store the number of saved records in the fixed \texttt{num\_size\_rec} slot. The first saved record can be located by taking the address of the frame, and adding both the size of the fixed part and the number of slots for protected regions (which is zero for nondet disjunctions).

The set of regions that existed before the disjunction and that may be allocated into by code following the disjunction is not available to the compiler. In theory, we could implement a global analysis to make it available, but such an analysis would be very complicated, especially for multi-module programs. Even if such an analysis existed, we would still have a big problem, which is that the number of regions in this set is not bounded, and in many cases the set would contain tens, hundreds or even thousands of regions. Saving and then restoring the sizes of that many regions can take a significant amount of both memory and time. We do not want this overhead to outweigh the benefits of instant reclaiming.

In our implementation, we have chosen to save and restore the sizes of only the regions that are \textit{locally} forward live at the start of the disjunction; this means the regions that are forward live before the disjunction \textit{and} whose region variables are visible at that point. (This information is readily available inside the Mercury compiler.) This means that we do not recover memory in regions that are forward live before the disjunction but whose identity was not passed to the current procedure, and are visible only in its ancestors. Since nondet disjunctions are quite rare in most Mercury programs (most programs that do serious searching tend to program their own searches instead of relying on chronological backtracking), we do not expect this to be too much of a problem. We will see below that we do not miss memory recovery opportunities for semidet disjunctions.

We save and restore the sizes of \textit{all} regions that are locally forward live at the start of the disjunction (the number of these regions governs how much space we reserve for the nonfixed part of the disj frame). We save and restore the sizes even of regions that are never allocated into before backtracking, since (in the absence of the analysis mentioned above) we do not know which ones of those are. This may
lead to some unnecessary saving and restoring, but in typical programs, the number of regions whose size we save and restore at a disjunction is usually relatively small, and in that case the memory or runtime cost of these unnecessary saves and restores is negligible. In some cases, however, the cost can be significant, and an optimization that eliminates saves/restores with a poor cost/benefit ratio would be useful. Such an optimization would probably need access to profiling information about region reclamation. We do not yet generate such information.

**Specialized treatment of semidet disjunctions.** Because at most one disjunct of a semidet disjunction may succeed, when one of its disjuncts is reached, it means that all the previous disjuncts have failed and that therefore (more importantly for us) execution has not passed outside the disjunction’s scope. Therefore, we only need to provide runtime support for a semidet disjunction if in its scope there is some change with respect to the set of existing regions. This basically means that the runtime support for nondet disjunctions described above will only be applied to semidet disjunctions whose created, removed and allocated sets are not all empty. In our practical experience with Mercury, most semidet disjunctions contain only tests, and rarely make changes to the heap. Therefore the support we describe below is needed only by a relatively small fraction of semidet disjunctions.

For a semidet disjunction, the Mercury compiler generates code such that when one of its non-last disjunct succeeds, the execution will commit to it and not go back to try any later disjuncts. This means the code we add at (d3) may not be reached after the success of a non-last disjunct, causing two problems. First, the disj frame will not be popped. Second, the regions which are removed by this disjunction but are protected against reclamation while later disjuncts exist will not be first unprotected at the start of the execution of the last disjunct and then reclaimed in the body of the last disjunct, as in the case of nondet disjunctions. Our solution is to do these two tasks at the end of any non-last disjuncts, i.e. after their success at (e1) and (e2) as in Figure 18.

To solve the first problem, we pop the frame at (e1.b) and (e2.b). To solve the second problem, at (d1) we loop through the regions in the disjunction’s removed set. If a region is already protected, we do not want it to be reclaimed in the disjunction and its `remove` instructions inside the disjunction will be ineffective anyway, so we do not need to do anything. If a region is not already protected, we save its handle in the nonfixed part of the disj frame. At the end, we store the number of region handles we saved in the frame’s `num_prot_region` slot. The code at (e1.a) and (e2.a) will loop through the saved handles, and reclaim all the saved regions (they were logically removed during the disjunct, but the protection of this disjunction prevented their `remove` instructions from actually reclaiming them.)

At (d1.c), we save the sizes of only the regions in the disjunction’s allocated set. Since execution cannot leave a semidet disjunction, we do not miss any memory recovery opportunities by restricting ourselves to these regions.
Fig. 18: RBMM runtime support for semidet disjunction.

9.1.1 Disjunctions: Summary

To summarize Section 9.1, we review how we handle Tasks 1 and 2 for disjunctions; first nondet disjunctions, and then semidet disjunctions.

We prevent the reclamation of backward live regions (Task 1) by disj-protecting all regions whose sequence number indicates they were created before the disjunction was entered. The protection of such regions starts at the beginning of the first disjunct (d1.a and d1.b), and ends at the beginning of the last disjunct (d3.c). Such regions are no longer protected by this disjunction during the execution of the last disjunct, so that if they are removed, they can be reclaimed.

Task 2, the reclaiming of memory, consists of two parts. Instant reclaiming of new regions happens at the beginning of every nonfirst disjunct (at d2.a and d3.a); the new regions are identified as such by their sequence numbers. Instant reclaiming of new allocations in old regions also happens at the beginning of every nonfirst disjunct (at d2.b and d3.b). To allow us to restore each old region to its state before the disjunction, each disj frame contains a list of the old regions that are allocated into during the disjunction, together with the sizes of these regions at the start of the disjunction (d1.c).

Task 1 needs extra support in the case of semidet disjunctions. The disj frames of such disjunctions have a list of the disj-protected regions, namely the regions in the removed list of the disjunction which are disj-protected only by this disj frame (set at d1.d). We use this list to explicitly reclaim these regions if a nonlast disjunct succeeds (e1 and e2).
9.2 Support for If-then-elses

The condition of an if-then-else (ite) can be either semidet or nondet. In most Mercury programs, the overwhelming majority are semidet, and this is the case we will look at first. Such if-then-elses share some properties with semidet disjunctions. If the condition succeeds, the execution will never enter the else part, and if the condition fails, the failure must have occurred in the scope of the condition.

Like disjunctions, if-then-elses need to protect regions from being reclaimed while backward live. But in the case of if-then-elses, we can restrict our attention to regions removed in the condition (i.e., in the condition’s removed set), since this is the only part of the code in which the if-then-else itself can make a region backward live. When execution reaches the start of the then part, backtracking to the else part is no longer possible, which means that any regions that have been marked for removal in the condition have to be reclaimed for real, unless they are protected by a surrounding scope.

Also, if-then-elses, like disjunctions, should do instant reclaiming of memory allocated by backtracked-over computations. In the case of if-then-elses, this means that at the start of the else part, we should recover any memory allocated by the condition.

In general, we only need to provide support for changes to regions which occur inside the condition. This is good, because the conditions of if-then-elses are often very simple, containing only one or a few tests. Conditions whose created, removed and allocated sets are all empty are therefore fairly common. For such if-then-elses, the mechanisms we describe below are unnecessary, and so we optimize them away. If at least one these three sets is not empty, we add code at the starts of the condition, the then part, and the else part, i.e., at points (i1), (i2), and (i3) in Figure 19.

```plaintext
{ if
  (i1): start of the condition
    (a) push an ite frame
    (b) save the protected regions and their number
    (c) save size records and their number
    ...
  then
  (i2): start of the then part
    (a) reclaim the ite-protected regions
    (b) pop the ite frame
    ...
  else
  (i3): start of the else part
    (a) unprotect the ite-protected regions
    (b) do instant reclaiming of new regions
    (c) do instant reclaiming of allocations in old regions
    (d) pop the ite frame
    ...
}
```

Fig. 19: RBMM runtime support for if-then-else with semidet condition.

For each if-then-else, we use a data structure called an ite frame to store the information used for its runtime support. As with disj frames, we embed ite frames...
in the ordinary stacks used by the Mercury implementation, and link them together into the  
*ite stack*, with a global variable pointing to its top. The structure of an  
ite frame is exactly analogous to that of a disj frame, the only difference being that  
the first slot of the fixed part, $\text{prev_{ite\_frame}}$, holds a pointer to the previous  
te frame, or null if there is none.

**Ite-protecting backward live regions.** Since the compiler knows the regions in  
the removed set of the condition (in our example in Figure 15, $R_1$ is such a region),  
we will stop them from being reclaimed by *ite-protecting* them at the entry to the  
if-then-else. To allow us to ite-protect regions, we add to the region header a pointer  
field, $\text{ite\_protected}$, which is set to null when a region is created. A region is ite-  
protected if its $\text{ite\_protected}$ field is not null. The *remove* instruction will now  
only reclaim a region if its $\text{ite\_protected}$ field is null and it is not disj-protected.  
(We do not use the same protection mechanism as in the case of disjunctions. We  
will explain the reason for this when we describe how we handle if-then-elses with  
nondet conditions.) Before entering the condition, i.e. at (i1), we push an ite frame,  
and then iterate over the to-be-protected regions. If one of these regions is already  
protected by a surrounding disjunction or if-then-else, we ignore it. Otherwise, we  
protect it by setting its $\text{ite\_protected}$ field, which must be currently null, to point  
to the ite frame. For such a protected region, we add its handle to a  
$\text{region\_id}$ slot in the nonfixed part of the ite frame. Then we also put the final number of regions  
we protect in this way into the frame's $\text{num\_prot\_region}$ slot. We do this so that we  
can loop over all the regions protected by this ite frame in two places: at the start  
of the then part (i2.a), where we reclaim all these regions (giving delayed effect to  
the *remove* instructions in the condition), and at the start of the else part (i3.a),  
where we undo their protection by resetting their $\text{ite\_protected}$ fields to null.

**Instant reclaiming.** When the condition fails, we want to reclaim both the new  
regions created inside it and any new allocations into old regions. In our example  
in Figure 15 we want to reclaim all of $R_3$ and some of $R_2$.

To reclaim new regions, at (i1.a) we save the current sequence number into the  
ew frame's $\text{saved\_seq\_num}$ slot, and at (i3.b), we add code that traverses the region  
list and reclaims all the regions until it meets an old region.

To reclaim new allocations into an old region, at (i1.c) we save its size record  
into the nonfixed part of the ite frame. Although it is reasonable to do this for the  
regions in the allocated set of the condition, it would be wasteful to reclaim new  
allocations into the regions which will be reclaimed right at the start of the else part.  
Unfortunately, while the compiler knows which old regions have *remove* instructions  
at the start of the else part, it does not know which of these will actually reclaim  
their regions, since it does not know which regions are protected by surrounding  
code. We handle this uncertainty as follows. We generate code at (i1.c) for every  
old region which is live at that point. For those that are not removed at the start of  
the else branch, this code always saves their size records unconditionally. For those  
that are removed at the start of the else branch, this code checks whether they are  
protected before this if-then-else, and saves their size records only if they are. This  
is an optimization because the test to see if a region is protected takes less time  
than saving its size record, and restoring it if the condition fails. We record the
number of size records we saved in the `num_size_record` slot, so that code at (i3.c) can restore them all.

The final action of the support code for an if-then-else with a semidet condition is to pop the ite frame at either (i2.b) or (i3.d).

**If-then-else with nondet condition.** Unlike Prolog, Mercury allows the condition of an if-then-else to have more than one solution. If the condition is nondet, then execution can backtrack into the condition from the then part or later code. This poses two problems we need to solve.

First, since the condition can succeed more than once, the code we add at the start of the then part (i2) can also be executed more than once. Because we need the ite frame every one of these times, we cannot let the code pop it at (i2.b); we must keep it until after the last time it may be used, i.e., after the last success of the condition. We arrange for this to happen by modifying the way the code generator handles the failure of the condition.

Normally, the code generator arranges for failures of the condition before the condition succeeds for the first time to cause a branch to the start of the else part, while a failure of the condition after it has succeeded represents a failure of the if-then-else as a whole, and will be handled accordingly, in whatever way the surrounding context demands. For example, if the if-then-else is one disjunct of a disjunction, its failure will cause execution to resume at the start of the next disjunct. We call the place to branch to on failure of the whole if-then-else the **failure continuation**.

We modified the code generator so that if the nondet condition needs support for region operations, i.e., it has a nonempty created set, removed set or allocated set, we branch to the failure continuation only after we execute code to pop the ite frame, the same code that for semidet conditions we would execute at (i2.b).

Second, the condition being nondet means that it must include, directly or indirectly, a nondet disjunction (since this is the only Mercury construct that can introduce nondeterminism). Therefore we must ensure that the supporting code fragments we generate for the if-then-else and the disjunction inside it do not step on each other’s toes.

Our support for if-then-elses with semidet conditions provides ite-protection for regions in the condition’s removed set that are not yet protected before the if-then-else. For such a region in a nondet condition, there are two cases. The first case is when the region is removed before the first nondet disjunction inside the condition. That means that when the `remove` instruction is executed, the region is ite-protected but not disj-protected. The `remove` instruction will (correctly) not reclaim it. Later on, the region will be reclaimed when the condition succeeds for the first time by the supporting code added at (i2). Because the program may backtrack into the condition and may reach the then part again, when the region is reclaimed at (i2.a), we need to nullify its entry in the ite frame so that it will not be wrongly reclaimed again the next time execution reaches (i2.a). This explains our saving of the pointer to the ite frame in the `ite_protected` field in the region header of a protected region.

In the second case, the region is removed after the start of the first disjunction in
the condition, either in the disjunction itself or at some point after it. In an execution containing a non-last disjunct, when the \texttt{remove} instruction is encountered the region is not reclaimed because it is both ite- and disj-protected. We need to ensure that if the condition succeeds and execution reaches the then part, the region should not be reclaimed at (i2) because it may be needed when execution backtracks into the condition. We therefore put different code at (i2.a) if the condition is nondet; this code will reclaim a region only if it is not currently disj-protected (Figure 20). The region will remain both ite- and disj-protected until the execution enters the last disjunct, at that time it will lose its disj-protection (Section 9.1). When the \texttt{remove} instruction in the condition is executed after this, it will not reclaim the region because it is still ite-protected, but the code at (i2.a) will reclaim it.

When the nondet condition fails, in both cases above, the region is only ite-protected, not disj-protected. It is because in the first case, the region is never disj-protected and in the second case, the failure happens only after all the disjuncts of the nondet code have been tried and failed, and the region has been disj-unprotected at the start of the last disjunct. This situation is exactly the same as when a semidet condition fails. Therefore the code at (i3) is exactly the same for nondet conditions as for semidet conditions.

\textit{9.2.1 If-then-elses: Summary}

To summarize Section 9.2, we review how we handle Tasks 1 and 2 for if-then-elses; first if-then-elses with semidet conditions, and then those with nondet conditions.

We prevent the reclamation of backward live regions (Task 1) by ite-protecting any regions that are removed in the condition, but are backward live, and are not protected by any other mechanism. The mechanism we use for ite-protection takes the form of \texttt{ite\_protected} fields in region headers: if this field is not null, the region is ite-protected. At the beginning of the condition (i1.a and i1.b), we set this field to point to the ite frame of the if-then-else for all the regions that meet the conditions listed above. If the condition succeeds, then execution enters the then part, and the code at (i2.a) reclaims these regions (since backtracking to the else case is no longer possible, and the regions are therefore no longer backward live). If the condition fails, code at (i3.a) unprotects these regions.

Task 2 consists of two parts. Instant reclaiming of new regions happens at the beginning of the else part (at i3.b): as with disjunctions, new regions are identified as such by their sequence numbers. Instant reclaiming of new allocations in old regions also happens at the beginning of the else part (at i3.c). To allow us to
Fig. 21: RBMM runtime support for commit.

9.3 Support for Commit

When the goal inside a commit succeeds for the first time, we commit to that solution by discarding the inner goal’s outstanding alternatives. We call the point in the code where this happens the commit point. If the inner goal is nondet (rather than multi), it may also fail. When it fails, the compiler’s failure-handling mechanism causes execution to pass through a failure point before the program resumes forward execution at the resumption point of the next surrounding goal. The failure point is there to allow the execution of some cleanup code. We add code to support region operations at two or three points in Figure 21: the entry point of the commit (c1), the commit point (c2), and the failure point (c3). If the inside goal has determinism multi, there is no (c3) to modify as execution would never reach there.

Consider a region that is in the removed set of a commit goal. If it is already protected by a disjunction or if-then-else when execution arrives at (c1), then the region should not be reclaimed by any code inside the commit, and the mechanisms we have described so far are sufficient to ensure this. If the region is not already protected at (c1), then the region should be reclaimed before execution reaches (c2). Ensuring this needs a new mechanism because the goal inside a commit will

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<th>Original Script</th>
<th>Parsed Data</th>
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<tbody>
<tr>
<td>some [...]</td>
<td>(c1): entry to the commit</td>
</tr>
<tr>
<td></td>
<td>(a) push a commit frame</td>
</tr>
<tr>
<td></td>
<td>(b) save the sequence number</td>
</tr>
<tr>
<td></td>
<td>(c) save the pointer to the top disj frame</td>
</tr>
<tr>
<td></td>
<td>(d) save the pointer to the top ite frame</td>
</tr>
<tr>
<td></td>
<td>(e) save the to-be-reclaimed old regions and their number</td>
</tr>
<tr>
<td></td>
<td>(the inner goal)</td>
</tr>
<tr>
<td></td>
<td>(c2): commit point</td>
</tr>
<tr>
<td></td>
<td>(a) reclaim the saved old regions</td>
</tr>
<tr>
<td></td>
<td>(b) reclaim the new regions</td>
</tr>
<tr>
<td></td>
<td>(c) restore the state of the disj stack</td>
</tr>
<tr>
<td></td>
<td>(d) restore the state of the ite stack</td>
</tr>
<tr>
<td></td>
<td>(e) pop the commit frame</td>
</tr>
<tr>
<td></td>
<td>(c3): failure point</td>
</tr>
<tr>
<td></td>
<td>(a) restore status of the saved regions</td>
</tr>
<tr>
<td></td>
<td>(b) pop the commit frame</td>
</tr>
</tbody>
</table>
contain, directly or indirectly, at least one disjunction that can succeed more than once (if it did not, it would have at most one solution, and there would be no commit operation), and the runtime support for this disjunction will protect the region from being reclaimed during the execution of its non-last disjuncts. On the other hand, we cannot simply insert code at (c2) to reclaim the region, since it can already be reclaimed by its remove instruction in the execution of the last disjunct before reaching (c2). We do not need to worry about the case when regions are protected only by semidet disjunctions or by if-then-elses with semidet conditions inside a commit, since these constructs, if they occur, protect regions only temporarily, and ensure that any regions that are removed inside them and are not protected when execution enters them will be reclaimed before execution exits them. If-then-elses with nondet conditions cannot protect regions either, though the nondet disjunctions inside their conditions can.

As before, our solution involves a new embedded stack, the commit stack. We push a new commit frame at (c1), and fill in its fixed fields, which we will discuss shortly. Following this will be the code that, for each region in the removed set of the commit goal, checks whether the region is already protected. If it is, that region is left alone. If it is not, we add the handle of the region to the commit frame’s nonfixed part, and record the address where this handle is stored in the commit frame in the region’s own header, in a new field called commit_slot. This way, when a region that should be reclaimed inside the commit actually survives to (c2) due to the protection of an inner disjunction, code at (c2) can iterate through all the region handles in the commit frame and reclaim those regions. However, we cannot do this for regions that are actually reclaimed inside the commit (whose remove instructions were executed in the last disjuncts). That is why, when we reclaim a region, we check whether its header’s commit_slot field is null. If not, then it will contain the address of a pointer to the region header, an address that will be in a commit frame, and the reclaim operation will replace that pointer in the commit frame with a null. Making the loop at (c2.a) ignore such nulled-out region handle pointers ensures that each region recorded in the commit frame’s list is reclaimed exactly once, and that this will happen as soon as possible.

If the goal inside the commit fails, we need to undo the update of the saved regions’ commit_slot fields, so at (c3.a) we reset them all to their original values. To make this possible, we save each original value in the commit frame next to the pointer to the region header from which it is taken. This effectively chains together all the entries referring to a given region in the commit stack. The reclaim operation will set to null not just the first slot in this chain, but all of them.

This mechanism is sufficient to correctly handle any old regions that are in the commit goal’s removed set. To handle any new regions (regions created inside the commit) that are also removed inside the commit, we record the current region sequence number in the commit frame at (c1). When a new region is removed in the commit, if it is not protected, it is reclaimed. If it is protected, we mark it so that at the commit point we can reclaim it. We add a field destroy_at_commit to the region header, and we augment the remove instruction again so that when a protected, new region is removed in a commit, the remove instruction will set the
region’s `destroy_at_commit` field to true (it is always initialized to false). At the (c2.b) part of the commit point, we traverse the region list until meeting an old region, and reclaim the new regions whose `destroy_at_commit` field is true.

We do not need to worry about instant reclaiming of new regions in the created set and of new allocations into regions in the allocated set of the commit, since that will be done by the goals surrounding the commit.

At the commit point, the Mercury execution algorithm throws away all the remaining alternatives of the goal inside the commit. To reflect this, at (c2) we need to restore the embedded disj stack to the state it had at (c1). This is why at (c1.c), we save the current disj stack pointer in a fixed slot in the new commit frame, and at (c2.c), we restore the disj stack pointer from there. The regions protected by the disj frames thrown away by this action will be exactly the ones removed by the code at (c2.b).

In some rare cases, the thrown-away disj frames will be from disjunctions inside if-then-elses with nondet conditions. Such if-then-elses cannot protect any regions in any code outside their conditions, but we do still need to ensure that we leave the embedded ite stack in the same state as we found it. This is why at (c1.d) and (c2.d), we save and restore its stack pointer. (The ite frames of if-then-elses with semidet conditions will have been popped by the time we get to c2, but the ite frames of if-then-elses with nondet conditions may still be there.)

The layout of commit frames is shown in Figure 22, with the fixed and nonfixed parts are separated by a thick line.

| prev_commit_frame | (previous commit frame) |
| saved_seq_num | (saved sequence number) |
| saved_disj_sp | (saved disj stack pointer) |
| saved_ite_sp | (saved if-then-else stack pointer) |
| num_saved_regions | (number of saved regions) |
| region_id | (handle of a saved region) |
| prev_commit_slot | (original commit slot of the saved region) |
| ... |

Fig. 22: The structure of a commit frame.

The meaning of the first two fields should be clear. The third and fourth fields contain the values of the disj and ite stack pointers respectively at the time when the commit was entered. The last fixed field gives the number of region handles and saved `commit_slot` fields actually stored by the code at (c1.d) in the nonfixed part.

### 9.3.1 Commits: Summary

To summarize Section 9.3, we review how we handle commits.

A commit does not need to protect any regions against reclamation, as it does not make any regions backward live. When the commit goal succeeds, it cuts away any backtrack points set up inside it, so we need to take away all the protections
associated with those backtrack points, and if this leaves a region (old or new) unprotected, we need to reclaim it.

We keep in each commit frame a list of the old regions (existing before the commit) that may be subject to such reclamation. We store this list at (c1.e), and we reclaim the regions in it at (c2.a), provided they have not been reclaimed within the commit goal itself, by code executing within or after a last disjunct. We set the \texttt{commitSlot} of each of these regions' headers to point to their entry in the commit frame; if and when the region is reclaimed within the commit goal, we delete this entry to prevent double reclamation.

Since commits may be nested, a given to-be-reclaimed region may be listed in several commit frames. We keep its entries in these frames in a chain, and when a region is actually reclaimed, we delete its entries in \textit{all} these frames.

To reclaim new regions, we store a snapshot of the sequence region number in the commit frame at (c1.b). When the commit goal succeeds, we reclaim all regions younger than this whose \texttt{destroyAtCommit} field has been set to true by a \texttt{remove} instruction.

If the commit goal fails, all the protections set up by any disjunctions or if-then-elsees inside it must have expired already, so we need do no more than simply restore the commit stack to its original state.

### 9.4 Compatibility with Tabling

Mercury supports three forms of tabling: loop checking (which detects the simplest form of infinite loops, and aborts the program if found), memoization (caching of results), and minimal model tabling.

The mechanisms we have discussed in this section so far are compatible with loop checking because the only two changes loop checking makes to the flow of execution are to force the execution of some table lookups, which have no effect on our data structures, and (maybe) to abort the program, in which case what our mechanisms do does not matter.

Our mechanisms are also compatible with automatic caching for det and semidet procedures. This tabling method surrounds the body of the tabled procedure with code that checks whether a call with the current argument list has been seen before. If it has not been seen, it computes the answer and records it. For det procedures, the answer consists of the values of the output arguments; for semidet procedures, it includes the success/failure indication as well. If this call has been seen before, the transformed procedure just returns the recorded answer. Neither the extra code executed at the starts and ends of new calls nor the table lookup executed for previously-seen calls interfere with any of our mechanisms.

Automatic caching for nondet and multi procedures is a more complex case, because the code that adds answers to a table adds one answer at a time, and only when execution is about to backtrack out of a new call does the tabling system know that its set of answers is complete. The Mercury system handles the interaction of tabled nondet/multi procedures with commits, just as it handles the interaction of nondet/multi procedures using RBMM with commits, but it does
not handle the interaction of tabled nondet/multi procedures using RBMM with commits. There is no reason why it could not do so, we just have not implemented it yet, mainly because memoization is not as useful for nondet and multi procedures as minimal model tabling.

The current implementation of minimal model tabling in the Mercury system works by saving segments of the Mercury stacks and restoring them later, possibly several times (Somogyi and Sagonas 2006). This makes minimal model tabling fundamentally incompatible with the mechanisms we have presented earlier in this section.

10 Experimental Evaluation

10.1 The Experimental Systems

We have implemented the region analysis and transformation shown in Sections 5, 6, and 7, as well as the runtime support describe in Sections 8 and 9 by incorporating them in the Melbourne Mercury compiler. The runtime support is currently available in the backend that generates low-level C code.

We use three variants of our RBMM system in our experiments. The first one, rbmm1, is similar to the RBMM system in (Phan et al. 2008) in which we do not track which regions that are allocated into. In rbmm1, while the region operations (Section 8) are implemented as C functions, the runtime support for backtracking (Section 9) is implemented using C macros. The functionality of the second system, rbmm2, is exactly the same as rbmm1, however we consistently implement the whole runtime support in functions. The third system, rbmm3, also uses only functions in the runtime system, but differs from rbmm2 in that it does track which regions are allocated into (using the algorithms in Section 5.4), which allows us to restrict the set of old regions for which we take size snapshots for later reclaiming (see Section 9) to just the regions for which this may have an effect. We chose these three versions to evaluate because comparing rbmm1 and rbmm2 tells us which implementation technology is better, while comparing rbmm2 and rbmm3 can reveal the impact of tracing which regions are allocated into and which are not. We also compare these RBMM variants with a Mercury compiler that is identical in all aspects except that instead of RBMM, it uses the Boehm garbage collector (Boehm and Weiser 1988), which is Mercury’s standard garbage collector. We call this system boehm.

For all three RBMM systems, we use a region page size of 2,048 words, of which 2,047 are available to store program data. When needed, we request blocks of 100 region pages from the OS. The three systems use the same regions and create and remove them in exactly the same places. However, they do differ in other aspects, such as compilation time, size of object files, and runtime performance.

Next, we will present the benchmarks and give the results of our experiments, and then we will discuss the RBMM behavior of the benchmarks in more detail. The experiments were performed on a Dell Optiplex 760 PC with a 2.83 GHz Core 2 Quad Q9550 CPU, 8 GB of RAM, running Ubuntu Linux, with the kernel version being 2.6.24-25-server SMP. The Mercury programs were compiled to C with the
3 December 2009 release-of-the-day of the Mercury system (with different options for the different variants). This and other releases-of-the-day are available on the Mercury web site. The resulting C files were compiled to executables by gcc 3.4.4. Every time we report was derived by running the program eight times, discarding the lowest and highest times, and averaging the rest.

### 10.2 The Benchmark Programs

In our experiments, we used a set of relatively small benchmark programs. We selected the benchmarks carefully; they are actually more like a collection of case studies that illustrate the strong and weak points of RBMM. While we would have liked to test our system with bigger, more realistic programs, we are currently not able to do so because the region analysis and transformation do not yet support higher order code, foreign language code and multi-module programs.

#### Table 4: Information about the benchmarks.

<table>
<thead>
<tr>
<th></th>
<th># Predicates</th>
<th># LOC</th>
<th>if-then-else</th>
<th>disjunction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>semidet</td>
<td>nondet</td>
</tr>
<tr>
<td>dna</td>
<td>16</td>
<td>251</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>isort</td>
<td>6</td>
<td>101</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>nrev</td>
<td>5</td>
<td>72</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>primes</td>
<td>8</td>
<td>93</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>qsort</td>
<td>6</td>
<td>92</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>bigcatch</td>
<td>12</td>
<td>159</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>boyer</td>
<td>17</td>
<td>372</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>bsolver</td>
<td>41</td>
<td>805</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>crypt</td>
<td>15</td>
<td>219</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>filev</td>
<td>12</td>
<td>154</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>life</td>
<td>18</td>
<td>338</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>healthy</td>
<td>24</td>
<td>485</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>queens</td>
<td>9</td>
<td>128</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>sudoku</td>
<td>22</td>
<td>441</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>rdna</td>
<td>17</td>
<td>262</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>risort</td>
<td>7</td>
<td>111</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>rlife</td>
<td>19</td>
<td>343</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>rqueens</td>
<td>10</td>
<td>138</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

The benchmark programs in Table 4 are divided into three groups. The first group contains benchmarks that do not need any runtime support for backtracking. The benchmarks in the second group do need such support. The third group consists of manually modified versions of benchmarks that illustrate how programs can be made more region-friendly (hence the “r” as prefix on their names).
The programs in the first group contain only det code, and maybe some if-then-elses with semidet conditions whose created, removed and allocated sets are all empty. dna computes similarities between gene sequences, isort implements insertion sort on a list of 10000 integers, nrev reverses a list of 5000 integers, primes finds all the primes less than 20000, and qsort sorts a list of 100000 integers.

The programs in the second group need runtime support for if-then-elses and/or disjunctions. bigcatch and filrev are Mercury versions of programs used in (Aspinall et al. 2008). They manipulate lists of lists of integers and introduce sharing between the input, the temporary data and the output and as such they also present difficult cases for RBMM. bsolver is a simple solver for systems of binary linear equations and inequations over integers; boyer is a toy theorem prover; crypt finds the unique answer to a cryptoarithmetic puzzle; life implements the Game of Life (known to be a difficult case for RBMM); healthy is a nondeterministic variant of life that searches for a generation that after a certain number of reproductions (8) still has a number of live cells that is higher than a threshold (80); queens solves the 12-queens problem by first generating permutations and then checking; sudoku finds the solution for a sudokuz puzzle by doing propagation on finite domains.

The programs rlife and rdna are versions of life and dna that have been manually made region-friendly by copying some data instead of letting it be shared. rqueens is a modified form of queens; its delete predicate (called by permute) copies the list remaining after a deletion. Similarly, risort copies the remaining list when inserting an element into a sorted list. We will come back to this group of programs when discussing the benchmarks in detail.

10.3 Experimental Results

10.3.1 Compilation Times and Object File Sizes

We first compare the three RBMM systems and the Boehm system with respect to their compilation times and the sizes of their object files (the text sections). The results are given in Table 5, which contains two sets of columns, for compilation time and object file size respectively. The first four columns in each group report results for each of our four system variants, rbmm1/2/3 and boehm, while the fifth column is computed by (rbmm3 - boehm)/boehm * 100.

Compilation times for most benchmarks are so short that we get significant fluctuations due to clock granularity; times in the table that differ only by a couple of tenths of seconds are effectively indistinguishable in practice. That said, compilation is always somewhat slower for the RBMM systems than when targeting the Boehm collector, which is not surprising, given the analysis we have to do. However, the cost of including RBMM is reasonable; the average slowdown for rbmm3 is 17%, and it is only a bit higher for rbmm1 and rbmm2. Compilation with the function-based systems is usually faster than compilation for the partly macro-based rbmm1 because the runtime support functions in rbmm2 and rbmm3 are compiled just once (when the runtime system itself is built) while in rbmm1 the macros containing their functionality are expanded and compiled several times during the compilation
Table 5: Compilation time and object file size.

<table>
<thead>
<tr>
<th></th>
<th>Compilation time (s)</th>
<th>Object file size (bytes)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>boehm 1 2 3 r3/b (%)</td>
<td>boehm 1 2 3 r3/b (%)</td>
</tr>
<tr>
<td>dna</td>
<td>0.51 0.66 0.60 0.60 18</td>
<td>4.782 6.670 6.366 6.142 28</td>
</tr>
<tr>
<td>isort</td>
<td>0.41 0.47 0.43 0.45 10</td>
<td>1.048 1.800 1.512 1.512 44</td>
</tr>
<tr>
<td>nrev</td>
<td>0.38 0.43 0.42 0.42 13</td>
<td>0.976 1.728 1.408 1.408 44</td>
</tr>
<tr>
<td>primes</td>
<td>0.39 0.44 0.43 0.42 8</td>
<td>1.026 1.712 1.408 1.408 37</td>
</tr>
<tr>
<td>qsort</td>
<td>0.41 0.47 0.45 0.47 15</td>
<td>1.209 2.088 1.768 1.768 46</td>
</tr>
<tr>
<td>dna</td>
<td>0.51</td>
<td>4,782</td>
</tr>
<tr>
<td>isort</td>
<td>0.41</td>
<td>1,048</td>
</tr>
<tr>
<td>nrev</td>
<td>0.38</td>
<td>0.976</td>
</tr>
<tr>
<td>primes</td>
<td>0.39</td>
<td>1.026</td>
</tr>
<tr>
<td>qsort</td>
<td>0.41</td>
<td>1.209</td>
</tr>
<tr>
<td>bigcatch</td>
<td>0.45</td>
<td>1,601</td>
</tr>
<tr>
<td>boyer</td>
<td>0.78</td>
<td>13,748</td>
</tr>
<tr>
<td>solver</td>
<td>0.97</td>
<td>16,034</td>
</tr>
<tr>
<td>cryp</td>
<td>0.57</td>
<td>5,656</td>
</tr>
<tr>
<td>file</td>
<td>0.40</td>
<td>1,650</td>
</tr>
<tr>
<td>life</td>
<td>0.56</td>
<td>5,564</td>
</tr>
<tr>
<td>healthy</td>
<td>0.61</td>
<td>7,906</td>
</tr>
<tr>
<td>queens</td>
<td>0.42</td>
<td>1,880</td>
</tr>
<tr>
<td>sudoku</td>
<td>0.65</td>
<td>7,685</td>
</tr>
<tr>
<td>rdna</td>
<td>0.55</td>
<td>4,831</td>
</tr>
<tr>
<td>risort</td>
<td>0.40</td>
<td>1,194</td>
</tr>
<tr>
<td>rlife</td>
<td>0.55</td>
<td>5,741</td>
</tr>
<tr>
<td>rqueens</td>
<td>0.43</td>
<td>2,155</td>
</tr>
</tbody>
</table>

of each benchmark. Compared to rbmm2, tracing and making use of the allocated regions in rbmm3 sometimes helps to reduce the compilation time, but the effect is quite small. This is because the overhead of tracking is rather small, and having information about allocated regions allows the compiler to do less work: it does not need to pass as many region arguments in calls, and it can skip adding some runtime support code.

The object files of the RBMM systems are, as expected, larger than those of the Boehm system. The use of macros in rbmm1 can double the size compared to boehm, as shown by bigcatch and healthy, with average increase being 74%. Replacing macros with calls reduces the overhead significantly; the object size ratio between rbmm2 and boehm ranges from 27% to 66%, averaging 43%. Rbmm3 yields even smaller object files, since keeping track of allocated-into regions allows the compiler to reduce the number of region arguments passed and the amount of support code generated; the object size ratio between rbmm3 and boehm ranges from 18% to 47%, averaging only 35%. This shows that for larger programs, rbmm3 is likely to be preferable.

10.3.2 Memory Usage

We measured the memory consumption of the regions for the RBMM systems. Note also that the runtime support consumes some memory as will be discussed later. Here we focus on the storage of program data. The results in Table 6 are the same in all three RBMM systems. For each benchmark, we give the total number of regions created during its execution, and the maximum number of regions coexisting during
Table 6: Memory use in rbmm systems.

<table>
<thead>
<tr>
<th>Region</th>
<th>Total</th>
<th>Max</th>
<th>Words used</th>
<th>Total</th>
<th>Max</th>
<th>SLR</th>
<th>S (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dna</td>
<td>2,082,006</td>
<td>8</td>
<td>18,926,797</td>
<td>4,590,797</td>
<td>4,096,000</td>
<td>75.7</td>
<td></td>
</tr>
<tr>
<td>isort</td>
<td>3</td>
<td>1</td>
<td>67,029,222</td>
<td>67,009,222</td>
<td>67,009,222</td>
<td>0.0</td>
<td></td>
</tr>
<tr>
<td>nrev</td>
<td>5,003</td>
<td>2</td>
<td>25,015,000</td>
<td>10,000</td>
<td>10,000</td>
<td>99.9</td>
<td></td>
</tr>
<tr>
<td>primes</td>
<td>2,265</td>
<td>1</td>
<td>5,221,386</td>
<td>39,998</td>
<td>39,998</td>
<td>99.2</td>
<td></td>
</tr>
<tr>
<td>qsort</td>
<td>200,003</td>
<td>21</td>
<td>5,865,744</td>
<td>200,000</td>
<td>200,000</td>
<td>96.6</td>
<td></td>
</tr>
<tr>
<td>bigcatch</td>
<td>3</td>
<td>2</td>
<td>25,015,000</td>
<td>25,015,000</td>
<td>25,005,000</td>
<td>0.0</td>
<td></td>
</tr>
<tr>
<td>boyer</td>
<td>5</td>
<td>3</td>
<td>143,561</td>
<td>143,561</td>
<td>143,505</td>
<td>0.0</td>
<td></td>
</tr>
<tr>
<td>bsolver</td>
<td>78</td>
<td>7</td>
<td>2,914,444</td>
<td>2,911,528</td>
<td>2,908,442</td>
<td>0.1</td>
<td></td>
</tr>
<tr>
<td>crypt</td>
<td>417</td>
<td>3</td>
<td>3,442</td>
<td>94</td>
<td>64</td>
<td>97.3</td>
<td></td>
</tr>
<tr>
<td>filev</td>
<td>6</td>
<td>3</td>
<td>25,023,004</td>
<td>25,019,000</td>
<td>25,009,000</td>
<td>0.0</td>
<td></td>
</tr>
<tr>
<td>life</td>
<td>50,304</td>
<td>102</td>
<td>894,336</td>
<td>8,208</td>
<td>6,486</td>
<td>99.1</td>
<td></td>
</tr>
<tr>
<td>healthy</td>
<td>3,917,124</td>
<td>82</td>
<td>62,639,310</td>
<td>2,794</td>
<td>2,054</td>
<td>99.9</td>
<td></td>
</tr>
<tr>
<td>queens</td>
<td>4,545,703</td>
<td>2</td>
<td>121,453,230</td>
<td>114</td>
<td>90</td>
<td>99.9</td>
<td></td>
</tr>
<tr>
<td>sudoku</td>
<td>6,651</td>
<td>88</td>
<td>84,080</td>
<td>16,678</td>
<td>10,916</td>
<td>80.1</td>
<td></td>
</tr>
<tr>
<td>rdna</td>
<td>2,083,006</td>
<td>9</td>
<td>18,930,797</td>
<td>501,752</td>
<td>428,733</td>
<td>97.3</td>
<td></td>
</tr>
<tr>
<td>risort</td>
<td>373,214</td>
<td>1</td>
<td>289,968,666</td>
<td>2,000</td>
<td>2,000</td>
<td>99.9</td>
<td></td>
</tr>
<tr>
<td>rlife</td>
<td>50,356</td>
<td>102</td>
<td>894,594</td>
<td>2,056</td>
<td>1,722</td>
<td>99.8</td>
<td></td>
</tr>
<tr>
<td>rqueens</td>
<td>23,080,416</td>
<td>13</td>
<td>142,047,288</td>
<td>156</td>
<td>24</td>
<td>99.9</td>
<td></td>
</tr>
</tbody>
</table>

its run. We also include the total number of words allocated and the maximum number of words that coexist. SLR is the Size of the Largest Region and S (%) is the saving, calculated by 1 - Max words/Total words.

RBMM achieves optimum memory management in nrev, in primes, and in qsort. For the nondeterministic programs crypt, healthy, queens, and sudoku, the memory savings are also high. The impact of instant reclaiming on memory reuse differs among these programs (Table 7): in crypt and queens, instant reclaiming collects most of the words, while in healthy it collects only a small fraction and it reclames none at all in sudoku.

For cases such as isort, bigcatch, bsolver and filrev, we see that most of the memory goes to the biggest region. Typically, this biggest region contains some garbage data, but as it also holds some live data it cannot be reclaimed.

The boehm version of our system uses the Boehm-Demers-Weiser garbage collector (Boehm and Weiser 1988) for memory management. In our experiments, we just use the default configuration of this collector as it is in the Mercury compiler distribution. It is a stop-the-world, sequential mark-and-sweep collector that uses 1024-word pages. It starts with a heap of 64k words and heuristically carries out collections of garbage or expands the heap on demand.

Data about memory use in the boehm system is shown in Table 8. The second
Table 7: Words reclaimed by runtime support. (Other words are reclaimed by `remove` instructions.) Only programs with some nontrivial numbers are shown.

<table>
<thead>
<tr>
<th>New allocations</th>
<th>New regions</th>
<th>Start of then</th>
<th>Commit point</th>
</tr>
</thead>
<tbody>
<tr>
<td>Words</td>
<td>%</td>
<td>Words</td>
<td>%</td>
</tr>
<tr>
<td>bigcatch</td>
<td>0</td>
<td>0.00</td>
<td>0</td>
</tr>
<tr>
<td>crypt</td>
<td>0</td>
<td>0.00</td>
<td>3,270</td>
</tr>
<tr>
<td>queens</td>
<td>12,356,378</td>
<td>10.17</td>
<td>109,096,776</td>
</tr>
<tr>
<td>rqueens</td>
<td>81,862</td>
<td>0.13</td>
<td>3,314</td>
</tr>
<tr>
<td>sudoku</td>
<td>0</td>
<td>0.00</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 8: Memory use in one iteration.

<table>
<thead>
<tr>
<th>dna</th>
<th># gc</th>
<th># expans</th>
<th>boehm max size</th>
<th>rbmm words requested</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>kB</td>
<td>max size</td>
</tr>
<tr>
<td>dna</td>
<td>7</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>isort</td>
<td>20</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>nrev</td>
<td>9</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>primes</td>
<td>3</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>qsort</td>
<td>3</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>bigcatch</td>
<td>5</td>
<td>10</td>
<td>119,804</td>
<td>30,669,824</td>
</tr>
<tr>
<td>isort</td>
<td>2</td>
<td>2</td>
<td>17,168</td>
<td>4,395,008</td>
</tr>
<tr>
<td>nrev</td>
<td>2</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>crypt</td>
<td>1</td>
<td>2</td>
<td>17,168</td>
<td>4,395,008</td>
</tr>
<tr>
<td>rlife</td>
<td>5</td>
<td>10</td>
<td>119,804</td>
<td>30,669,824</td>
</tr>
<tr>
<td>healthy</td>
<td>19</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>queens</td>
<td>36</td>
<td>4</td>
<td>30,524</td>
<td>7,814,144</td>
</tr>
<tr>
<td>sudoku</td>
<td>1</td>
<td>2</td>
<td>17,168</td>
<td>4,395,008</td>
</tr>
</tbody>
</table>

The numbers show that, in almost all of the benchmarks, the RBMM systems can work within spaces that are smaller than those requested by the Boehm collector. RBMM systems often need to request only the minimum, which in our system is 100 * 2048 words. The worst case for RBMM is isort in which RBMM is not able to reuse memory efficiently. The Boehm system can work with only a bit more than one tenth the memory in this case.
### 10.3.3 Runtime Performance

We also studied the runtime performance of our benchmark programs because this is probably the most important criterion for the practicality of RBMM. To control the uncertainty involved in measuring small times, we ran each program many times in a loop. Each benchmark has a row in Table 9 that gives the number of iterations, the actual execution times with boehm (boehm), the boehm system’s gc time (gc), and the boehm system’s runtime minus the gc time (nogc), and then the runtime with the three RBMM systems (all in seconds, all for user mode only). Each row also includes the number of collections executed by the Boehm collector, and the savings achieved by using our preferred RBMM system, rbmm3, instead of the boehm system. The savings are given by 1 - rbmm3 runtime / boehm runtime.

The rbmm3 system gets clearly better runtimes than the boehm system for 15 out of our 18 benchmark programs, including both deterministic and nondeterministic programs. The speedups range from around 8% to more than 60%. (We do not count the 2.3% speedup as “clearly better”.) The overall average speedup, even including the two programs with slowdowns, is about 24%. We get this promising result because with RBMM, we avoid the burden of runtime garbage collection, and because the overhead of supporting regions is reasonably modest. Moreover, the runtimes of 10 of these 15 programs are smaller than the corresponding runtimes in the boehm system even excluding garbage collection times, which strongly suggests that RBMM also improves data locality. In bigcatch and filrev, two difficult cases for RBMM, their memory-use pattern actually has even more adverse effects on the operation of the Boehm collector. These programs all build very large lists that are live data before producing any garbage, so during their initial phase, the traversal

<table>
<thead>
<tr>
<th>Program</th>
<th># Iter</th>
<th>boehm runtime</th>
<th>gc</th>
<th>nogc</th>
<th># gc</th>
<th>RBMM runtime</th>
<th>Saving</th>
</tr>
</thead>
<tbody>
<tr>
<td>dna</td>
<td>100</td>
<td>25.27</td>
<td>8.80</td>
<td>16.47</td>
<td>549</td>
<td>20.81</td>
<td>16.1%</td>
</tr>
<tr>
<td>isort</td>
<td>60</td>
<td>53.47</td>
<td>17.90</td>
<td>35.57</td>
<td>1141</td>
<td>21.43</td>
<td>59.5%</td>
</tr>
<tr>
<td>nrev</td>
<td>160</td>
<td>50.09</td>
<td>17.58</td>
<td>32.51</td>
<td>1134</td>
<td>20.39</td>
<td>57.8%</td>
</tr>
<tr>
<td>primes</td>
<td>400</td>
<td>49.94</td>
<td>9.46</td>
<td>31.48</td>
<td>597</td>
<td>24.86</td>
<td>39.9%</td>
</tr>
<tr>
<td>qsort</td>
<td>400</td>
<td>41.41</td>
<td>12.65</td>
<td>28.76</td>
<td>701</td>
<td>20.62</td>
<td>48.9%</td>
</tr>
<tr>
<td>bigcatch</td>
<td>30</td>
<td>28.31</td>
<td>5.70</td>
<td>22.61</td>
<td>20</td>
<td>20.39</td>
<td>28.0%</td>
</tr>
<tr>
<td>boyer</td>
<td>8,000</td>
<td>25.69</td>
<td>5.60</td>
<td>20.09</td>
<td>357</td>
<td>22.59</td>
<td>34.3%</td>
</tr>
<tr>
<td>boolver</td>
<td>1500</td>
<td>55.00</td>
<td>19.44</td>
<td>35.56</td>
<td>1242</td>
<td>22.92</td>
<td>58.3%</td>
</tr>
<tr>
<td>crypt</td>
<td>300,000</td>
<td>21.19</td>
<td>4.53</td>
<td>16.66</td>
<td>293</td>
<td>18.85</td>
<td>2.3%</td>
</tr>
<tr>
<td>filrev</td>
<td>50</td>
<td>34.80</td>
<td>11.03</td>
<td>27.37</td>
<td>54</td>
<td>24.09</td>
<td>37.9%</td>
</tr>
<tr>
<td>life</td>
<td>700</td>
<td>27.18</td>
<td>2.77</td>
<td>24.41</td>
<td>179</td>
<td>26.16</td>
<td>12.8%</td>
</tr>
<tr>
<td>healthy</td>
<td>30</td>
<td>37.65</td>
<td>8.34</td>
<td>29.31</td>
<td>533</td>
<td>41.63</td>
<td>21.3%</td>
</tr>
<tr>
<td>queens</td>
<td>15</td>
<td>32.90</td>
<td>7.97</td>
<td>24.93</td>
<td>517</td>
<td>22.34</td>
<td>8.7%</td>
</tr>
<tr>
<td>sudoku</td>
<td>20,000</td>
<td>23.02</td>
<td>6.45</td>
<td>16.58</td>
<td>413</td>
<td>17.65</td>
<td>23.7%</td>
</tr>
<tr>
<td>rdna</td>
<td>120</td>
<td>30.41</td>
<td>10.52</td>
<td>19.89</td>
<td>657</td>
<td>24.38</td>
<td>22.2%</td>
</tr>
<tr>
<td>risort</td>
<td>25</td>
<td>89.81</td>
<td>31.84</td>
<td>57.98</td>
<td>2051</td>
<td>35.28</td>
<td>60.3%</td>
</tr>
<tr>
<td>rlife</td>
<td>700</td>
<td>27.02</td>
<td>2.74</td>
<td>24.28</td>
<td>179</td>
<td>26.04</td>
<td>12.9%</td>
</tr>
<tr>
<td>rqueens</td>
<td>15</td>
<td>35.65</td>
<td>9.57</td>
<td>26.08</td>
<td>604</td>
<td>43.06</td>
<td>23.7%</td>
</tr>
</tbody>
</table>
The five columns related to disj frames are as follows: Total is the total number of disj frames used in one iteration; M is the maximal number of disj frames coexisting at some point; \# Words is the total number of words used for all the disj frames; Mw is the maximal number of words used at some point; and Sr is the total number of size records saved. No regions are protected at semidet disjunctions in these benchmarks. For ite frames, the first five columns have meanings analogous to those for disj frames, while the last column gives the total number of regions that are protected by the ite frames by having their handles saved in these frames. The Mw columns show that the memory used by both these kinds of embedded frames is negligible in all benchmarks. We do not show information about commit frames.
The rbmm3 system is only a little faster than boehm on crypt. Despite being a nondeterministic program, the runtime support for backtracking it needs is rather cheap (see Table 11). However, the program handles a large number of small regions, more than 125 million regions in total (417 regions in each of 300,000 iterations), with an average of just over eight words per region, and the largest region being 64 words. The cost of creating and destroying the region has to be amortized over the words stored in the region. In large regions, the proportion of this overhead falling on any one word is negligible, but in small regions, it can be substantial. So rbmm3’s gain due to avoiding runtime garbage collection is almost exactly counterbalanced by the overhead of handling many small regions, resulting in just a small overall speedup.

This problem also manifests itself to various extents in the other programs that handle many small-to-medium size regions (more than ten million of them). This can be seen in programs such as dna, life, healthy, sudoku, rdna, and rlife, where we still have clear speedups but they are not as good as the speedups for programs with fewer, larger regions. The memory results in Table 6 show that with rbmm3, rdna indeed needs much less memory than dna, since it can reuse memory better with the help of its copying predicate. Unfortunately, the overhead of copying still causes rdna to be about 12% slower than dna, though the slowdown for rbmm3 is less than for boehm (where it is 20%). However, compared to crypt, queens has many more nondet disjunctions so it has to pay the cost of supporting backtracking within them many times (see Tables 10 and 11), and it has to pay for handling many small regions (68M regions with an average of about 27 words each), and yet rbmm3 gets a speedup of 8.7% over boehm on this benchmark.

The two worst cases for rbmm3 are rqueens and boyer. rqueens uses about five times as many regions as queens, which makes the average region much smaller than the already too small regions in queens. This is the negative side-effect of copying terms to new regions to allow their old ones to be freed earlier. That copying does achieve its objective; we can see in Table 7 that the memory queens recovers from within regions is recovered by rqueens in the form of whole regions. rqueens actually never recovers memory from within regions, which means that overhead it pays for trying to do that (saving size records at disj frames) is useless while being quite expensive. The slowdown in boyer is mainly due to the cost of saving size records (more than 306 million of them) at ite frames, which are also all in vain. A closer look at boyer reveals that it contains some semidet procedures that allocate into their input regions, and the conditions of some if-then-elses call these procedures. So the compiler needs to save the size records of those regions if it wants to have instant reclaiming. However, for the specific input used in our benchmark, the calls to these semidet predicates all succeed, so instant reclaiming has no words to reclaim. See Section 12 for an idea that would allow us to eliminate such unprofitable overhead.

Comparing the runtime results for rbmm2 and rbmm3 gives us an idea about the usefulness of tracking allocated regions. While the reduction in the number of region arguments does not have a strong impact in these benchmarks, having less
supporting code for backtracking shows marked speedups for life, healthy and rlife. This enhanced performance corresponds with the reductions in Table 11 compared to Table 10. We can see that the main impact is on the ite frames. For filrev and life, we can get rid of them completely, except for one needed by the benchmarking mechanism itself. For some others, we no longer have to save any size records to ite frames. This is very important because while nondet disjunctions are rare in Mercury programs, if-then-elses are very common. Ensuring their efficiency is therefore vital to the efficiency of Mercury programs as a whole. However, tracking of allocated regions cannot help in all cases, such as in the case of boyer. For the programs for which rbmm3 seems slower than rbmm2, this is purely a chance cache effect. We have examined the C files generated by the Mercury compiler, and for each such benchmark, the only difference between the two versions is that the rbmm2 version executes some statements that the rbmm3 version does not, while using larger stack frames.

Comparing runtimes for rbmm1 and rbmm2, we see that in the programs that use runtime support for backtracking, using macros to implement that support may improve performance. Table 9 shows that boyer, life, healthy, queens, rlife and rqueens are all at least 5% faster in rbmm1 than they are in rbmm2. This is because using macros avoids the cost of calling functions, and because these programs are so small that the increase in code size does not adversely affect instruction cache behavior. However, we expect that for larger programs, the slowdown due to the reduction in the effectiveness of the instruction cache will outweigh the cost of the calls. However, in multi-module programs, it should be possible to compile most modules with function calls while compiling with macros the modules in which the program spends most of its time, thus getting the best of both worlds.

10.4 The Impact of Sharing on Reusing Regions

One can argue that sharing is the most basic and natural form of memory reuse. However, sharing can conflict with RBMM, because in RBMM we want terms with different lifetimes to be stored in different regions, and a subterm shared between two terms of different lifetimes obviously cannot be stored in two different regions at once. In this section we study in detail some benchmark programs that we selected specifically for insights about the impact of sharing on RBMM. Some of them are known difficult cases for RBMM such as dna and life. Some others create sharing that make it hard for in-place updating such as isort, bigcatch, and filrev (Aspinall et al. 2008).

In our region points-to analysis, we essentially put two program variables into the same region in two cases: when there is an assignment between them, or they are bound to a term and its same-type subterm in a recursive data structure (Section 5). When the variables in a region have different lifetimes, we will have a sort of memory leak, because the memory of the variables with shorter lifetimes will not be reclaimed until the longest lived variable dies.

One solution for this is to copy the live data in the region to a different region,
The life benchmark encodes the Game of Life in which a new generation is generated from a previous one based on a set of production rules. From an initial generation, it uses a loop (in the life predicate) to produce several intermediate ones before reaching the final generation, which is the wanted output. We represent a generation by a list of live cells, with each cell being represented by its row and column in a 20x20 board. To store a generation, we need two regions, one for the skeleton and the other for the cells. In the program, the list skeletons of two successive generations are independent while their cells may share. In the recursive case of the predicate life, we first call next_gen to compute the next generation, whose skeleton could be in a different region, and then we call life recursively with the next generation as input. In the base case, we assign the current generation, which is the “next” generation created by the caller, to the output generation. The computation is summarized in Figure 23. Due to the assignment in the base case, which creates sharing only between the last intermediate generation and the output generation, our region points-to analysis decides that the skeletons of the input and output generations in the life predicate are in the same region, and then enforces this for all the (recursive) calls to life. This eventually means that the skeletons of all the generations are placed in one big region with a size of 6,486 words. In rlife, we replace the assignment in the base case with a call to a copying predicate that does not create any sharing, thus allowing the compiler to store the skeleton of each generation in a separate region, which then can be reclaimed in time. We see in Table 6 that the maximum amount of memory needed by rlife is 2,056 words, which is a 75% reduction compared to life’s 8,208 words. This is because in rlife, the skeletons of the old generations are reclaimed at each step.

The program dna simulates the matching of a given DNA sequence to each of the DNA sequences in a predefined set. The matching degree of two sequences is represented by a similarity, which is computed based on the similarities of their elements with respect to the spatial relation among them. The similarities between two sequences are calculated one by one and put in an ordered tree, which is a recursive data structure. To store a tree, we need two regions, one for the tree nodes and the other for the structures where the similarities are stored. Other than that, in this program, there are assignments in several predicates that establish sharing among the similarity structures in such a way that all the similarities ever computed end up in the same large region of 4M words. The maximal number of words in use during a run of the program is about 4.6M. In the so-called region-friendly version rdna, we make a fresh copy of each similarity and add the copy to the tree. This allows the region analysis to decide that the region to which the similarity is copied is the region of the nodes of the tree, and it can reclaim its previous...
region containing all the temporary similarities involving in its computation. The maximum amount of memory needed drops from 4.6M words to only 0.5M. The size of the largest region also drops from 4M words to 0.43M words; in rdna, it contains only the skeleton of the tree.

In (Phan and Janssens 2009) we proposed a more desirable solution, a more refined region analysis that, by taking into account different execution paths, can keep apart the regions of the variables in an assignment. A dedicated implementation of the improved analysis should achieve the same effect as changing life into rlife changing dna into rdna, without either requiring manual rewriting of the program or incurring the cost of copying.

Another issue that we found was that one of the Mercury compiler’s existing optimizations, common structure reuse, was reducing the effectiveness of our region analysis. This optimization looks for conjunctions in which the same term is assigned to two or more variables, and then changes the code so that the term is constructed just once, and then it is assigned to all the variables. This is always an optimization for the boehm system, but in cases where our region analysis would want to assign those variables to different regions, making them refer to the same memory cell creates unwanted sharing, requiring our region analysis to merge the two variables’ regions. In general, the unmerged regions would be reclaimed at different times. Therefore merging the two regions can delay the reclaiming of an unbounded amount of memory by an unbounded amount of time. The best way to avoid this problem is to teach the optimization about regions, and make it perform the transformation only if the variables involved are in the same region.

The problems with memory reuse in RBMM in isort and queens are typical for programs that use recursive data structures such as lists and trees, and continuously update them by adding to them and deleting from them. Because the updated structure normally shares most parts of the original, they are stored in the same regions, which prevents us from reclaiming the now-obsolete parts of the original structure. In risort and rqueens, we try to improve memory reuse by adding a predicate to copy the modified structure so that the original region can be reclaimed after the copying. In risort, the copying happens after an integer is inserted, while in rqueens, it happens after a queen is deleted. This modification obtains optimal memory management for risort (see Table 6). In rqueens, compared to queens, the peak memory usage is higher. This is due to region protection: some disj-protected regions are removed but not reclaimed, and instant reclaiming does not recover their memory until later. However, the size of largest region drops to 24 words, which is the storage needed to represent a list of 12 queens.

While memory reuse can be improved by this copying approach, its runtime overhead is very expensive. We see a 63% increase of runtime for rqueens compared to queens in Table 9, and for risort we have to reduce the input size by a factor of ten (to 1,000 integers, compared to 10,000 integers in isort) to allow the program to finish in a reasonable time. Similar problems with memory reuse in the presence of recursive data structures can also be seen in dna and rdna, which insert similarity structures into trees, and in bsolver, which reduces the domains of the integral variables, with the domains being represented as lists of integers.
The reason why bigcatch and filrev are not even faster is also related to recursive data structures. In this case the structures are not updated but only part of them is used, i.e. only a part is live data, but that still requires us to keep the whole region alive. Copying the live data out of the region would work just as well to recover memory, and at just as high an overhead, as in the previous case. We do not have an automatic solution for the problems related to the use of recursive data structures in RBMM-only systems, but then, neither does anyone else. The problem is well-known among researchers who use type systems or type inference to reason about memory structures (Baker 1990; Chase et al. 1990; Tofte and Talpin 1997; Henglein et al. 2001), who nevertheless have to accept the loss of precision as the price of having a finite model. To improve storage use in such cases, one can combine RBMM with other techniques, such as runtime or compile time garbage collection. The copying approach used by our region-friendly benchmarks can be viewed as a simulation of runtime copying garbage collection. Combining RBMM with copying garbage collection has been realized in the MLKit (Hallenberg et al. 2002).

11 Related Work

In this section, we only mention the most important and most related papers. It is not our intention to give a detailed overview of the research on RBMM for other programming paradigms. An in-depth review of RBMM research for functional programming can be found in (Tofte et al. 2004).

The research on automated region-based memory management for programming languages started with the work of Tofte and Talpin (Tofte and Talpin 1997) for functional programming, in particular for a simplified call-by-value lambda calculus. They divide program terms into regions using a technique similar to unification-based type inference in which the types have been annotated with region variables. The lifetimes of the regions are computed based on the lexical scope of the expressions and the regions themselves are forced to follow stack discipline, with the last region created always being the first one destroyed. While lexically-scoped regions and stack discipline seem natural for the evaluation of lambda expressions and they simplify the task of deciding region lifetimes, they often give regions lifetimes that are longer than needed, increasing the program’s memory requirements. Possibly even more important, the cleanup they often require after a tail call also spoils tail call optimization. (Birkedal et al. 1996) refined this system in several ways, the most important being Storage Mode Analysis, which mitigates the problems caused by the stack discipline by resetting regions to zero size when their contents are no longer needed. However, to make this region resetting possible, programmers often have to rewrite their programs in unusual ways.

While Aiken et al. also used a stack in their inference algorithm, they nevertheless thought that forcing stack discipline on the lifetimes of regions is too strict (Aiken et al. 1995), and they decoupled region creation and removal, allowing regions to have arbitrarily overlapped lifetimes. Going even further in this direction, Henglein, Makholm, and Niss in (Henglein et al. 2001) proposed an imperative sublanguage
on regions. In their system, regions are allowed not only to have arbitrary lifetimes but also to change their bindings. Their regions also contain reference counters that can give their system more flexibility in controlling their lifetimes. The most complete functional programming system with RBMM is the MLKit (Tofte et al. 2006), which manages storage solely by RBMM. This system, while still using stack discipline for the lifetimes of regions, supports both resetting regions to zero size and runtime garbage collection within regions. Its performance is competitive with other state-of-the-art SML compilers.

Our static region analysis and transformation for Mercury were inspired by the work in (Cherem and Rugina 2004), which also allowed arbitrarily-overlapped region lifetimes. The analyses in that paper take into account the data flow in a Java program in order to determine the set of needed regions and their lifetimes. Therefore the analyses had to be redefined for Mercury to deal with unification and a control flow that are fundamentally different from object manipulation and control flow in Java. Cherem and Rugina use the classes of Java to achieve a finite representation of the storage of (recursive) structures in terms of regions, but their starting assumptions are different from ours. In our analysis, we start by associating each variable with as many regions as its type requires (e.g. skeletons and elements for list\_int) whereas they start by associating each variable with only one region (the one for its class), and add the other nodes later, on demand. In the case of recursive types, we know from the start that e.g. all the list skeleton nodes of a given variable are in the same region. Given a variable \( v \) of class \( c \) whose fields include, directly or indirectly, other variables of class \( c \), they initially allocate different nodes in the region graph to \( v \) and those other variables, and merge some of those nodes only when they see a link between them. This complicates their analysis, though in some cases it allows them to keep the regions separate and thus free some memory earlier. In logic programs, recursive types are almost always processed using recursive procedures, and such cases would be vanishingly rare.

Another difference between the two systems that is likely to be more important in practice is that the liveness information we derive in Section 6 allows interprocedural creation of regions, something that was not handled in (Cherem and Rugina 2004). This can give finer lifetimes to regions, which can result in better memory reuse in certain situations. For example, for a region like \( R_1 \) in \( p \) in Figure 13, the system in (Cherem and Rugina 2004) would force \( R_1 \) to be live throughout \( p \). If we had replaced the atom at (4) with a recursive call to \( p \) (such as \( p(A - 1, B) \)) their system would build up all the temporary memory allocated at (1) in \( R_1 \).

Note that using graphs to model storage is not at all new in research about heap structures (Chase et al. 1990; Steensgaard 1996). Our graphs share many features with annotated types where the annotation on each type constructor is a location or region; see e.g. (Baker 1990; Tofte and Talpin 1997). Baker in (Baker 1990) and many others pointed out that such annotated types can also give information about sharing, very similar to the concept of region-sharing in this paper.

The first application of RBMM to logic programming was the work of Makholm for Prolog, described in (Makholm 2000b) and (Makholm 2000a). He realized that backtracking can be handled completely by runtime support, which can keep the
region inference simple. However, the Prolog system he used was not based on
the usual implementation technology for Prolog, the Warren Abstract Machine or
WAM. This shortcoming was fixed in (Makholm and Sagonas 2002) where Makholm
and Sagonas extended the WAM to enable region-based memory management. The
main differences between their work and ours are that Mercury supports if-then-
else statements with conditions that can succeed more than once, and the Mercury imple-
mentation generates specialized code for many situations that Prolog handles with
a more general mechanism. (For example, Mercury has separate implementations
for nondet disjunctions and for semidet disjunctions.) The first difference required
new algorithms, while the second posed a tough engineering challenge in keeping
overheads down, since due to Mercury’s higher speed, any given overhead would
hurt Mercury more than Prolog.

12 Future Work

Our RBMM implementation already has some support for profiling. When given
a certain option, the Mercury compiler will augment the RBMM support code it
generates with code that counts and keeps track of several things: the number
of region creations and removals, the amount of memory allocated in regions, the
maximum size of regions, the number and size of the embedded disj, ite and commit
frames, and so on. This option was the source of the information in Tables 6, 7, 10,
and 11. We would like to modify this profiling mechanism to also report, for each
region variable (both old and new) at each resume point, the number of instant
reclaiming attempts made at that point for that region variable, and the amount of
memory recovered in those attempts. We would like to then feed this information
back to the compiler, so that it can find out which attempts are too expensive for
the amount of memory they recover, so it can simply avoid generating them.

Our current system prevents the reclamation of regions that are forward dead
but backward live entirely at runtime. Such runtime protection is in fact necessary
in general. Given a procedure \( p \) and a region \( r \) with \( r \in \text{deadR}(p) \), \( p \) cannot know
whether some disjunction to the left of its caller makes \( r \) backward live or not.
We could handle this situation by generating three versions of \( p \). The first version
would assume that \( r \) is backward live and therefore never reclaim \( r \), the second
version would assume that \( r \) is backward dead, and therefore always reclaim \( r \), and
the third version would make neither assumption and would reclaim \( r \) only if it is
not protected, as in our current system. The caller would call the first version if it
itself makes the region backward live (e.g. the call may be in one disjunct, and a
later disjunct in that disjunction may need the region), or because the caller itself
is a specialized version that assumes that the region is backward live. The caller
would call the second version if it itself created the region, and if there is no nondet
construct between that creation and the call that could make the region backward
live.

Unfortunately, a procedure’s \( \text{deadR} \) set may contain several regions, and given \( n \)
regions, we may need up to \( 3^n \) copies of the procedure, which is far too many, since
that many copies would significantly degrade the effectiveness of the instruction
cache. Nevertheless, in some situations, the fraction of execution time spent in the procedure may justify creation of one or more specialized copies of the procedure. We intend eventually to implement an optimization that figures out which of the possible specialized versions can ever be called, attempts to compare their cost in lost locality to the speedup we can expect from optimizing away unnecessary remove instructions, and creates the specialized versions if and only if the comparison indicates that it is beneficial to do so. If a specialized version is not worth it, the caller can call the original version of the procedure; since this does runtime tests on all the removed regions before reclaiming them, it still works in all cases.

What we could improve without considering such complicated tradeoffs are situations where the instruction that removes a region is in a procedure that itself makes the region unconditionally protected at the removal site. In such cases, we know statically that the removal will not actually reclaim the region, and that therefore we can simply optimize it away. If such protection is only conditional, we do have to consider the tradeoff. Since we cannot guarantee optimizing away all protected removals, the mechanisms we described in Section 9 will always be needed.

The main limitation of our work is that currently, the program analysis underlying our system supports only a subset of Mercury. We intend to work on extending the analysis to handle the rest of the language. Since we already handle almost all of Mercury, “the rest of the language” covers only a few features: Mercury procedures defined in foreign languages, multi-module programs, and higher-order code. To handle them, we need to ensure two things. First, that the callers and callees involved in calls to foreign language code, cross-module calls and higher order calls all agree on the liveness of the regions involved in the call; second, that they all agree on the sharing between those regions. The first one is relatively easy to achieve by simply setting the bornR and deadR sets of those calls to empty. This will work; any creations and removals of the regions that would have been in those sets will happen around the call. The cost is that it may increase the program’s memory consumption, though only to the levels seen in some other RBMM systems. The real problem is the second issue: getting consensus between callers and callees on sharing.

Handling foreign language procedures. Always setting the bornR and deadR sets of foreign language procedures to the empty set avoids burdening programmers with the responsibility for managing the creation and removal of regions. Since most foreign language procedures do not allocate any memory, their writers do not need to know anything about regions at all. The foreign language procedures that do allocate memory need to know where the allocation of each cell should happen. In a hybrid system that combines RBMM with the Boehm collector, it is simple enough to let such foreign procedures keep doing what they do now, which is doing all their allocation on the Boehm heap. An RBMM-only system would need to make the region arguments added to each procedure by our transformation visible to the programmer, and document which of these region variables represent which part of each of the arguments originally created by the programmer, so that when he or she writes code to create a new cell that will become part of a term that will be bound to an output argument, they can allocate it in the right region. We would
also need to give programmers a mechanism that they can use to tell the compiler about any sharing they create between the regions; our Algorithm 3 could then take this information on trust. As for temporary structures that can never become part of an output argument, programmers can put them where they wish. They can put them in memory managed by malloc and free (if the foreign language is C) and their equivalents (if the foreign language is something else), or, if we expose the functions for creating and removing regions, they can put them in one or more programmer-managed regions instead.

**Handling multi-module programs.** Our current implementation actually allows cross module calls; if a program cannot call the procedures in the standard library’s I/O module, then it cannot print out its results. The reason why we cannot yet handle multi-module programs in general is that currently we do not do any region analyses across modules, and hence we never pass region variables or any other information about regions from one module to another.

The reason why implementing region analysis in multi-module programs is hard is that the fixpoint computation in Algorithm 3 is inherently incompatible with separate compilation. Mercury’s compilation system ensures that when a module changes, all other modules dependent on its interface will be recompiled before the building of the executable, but it guarantees that this will take a bounded number of steps. As it is, Algorithm 3 cannot provide a similar guarantee; the procedures in a single SCC may be in different modules, and each iteration of the search for the fixpoint must analyze code in each of those modules. We therefore need to either change the algorithm, or make the compilation system flexible enough to encompass fixpoint computations that need an unbounded number of iterations. We have looked at the second option in the past, using the ideas of (Bueno et al. 2001) as the basis, but even if it were implemented, being able to limit the number of iterations would help compile programs more quickly. There are some assumptions we can make that can help with that. For example, we can assume that all input variables of cross-module calls are in regions that the callee will not allocate in or remove; if their last use is during the call, the caller will remove them upon return. This loses some precision and therefore reduces the efficiency of memory reuse, but this is a known and fairly widespread problem: most program analysis and optimizations lose precision at module boundaries, and in almost every case this is seen as an acceptable tradeoff. The challenge will be in coming up with mechanisms for handling the regions of output variables that still allow memory to be recovered effectively enough. We have some ideas, but no solutions yet.

**Handling higher order code.** Mercury supports two forms of higher order calls: calling an ordinary higher order term (a closure), and calling a typeclass method. The challenge in both cases is that the identity of the called procedure may not be apparent when the calling module is compiled, which prevents Algorithm 3 from analyzing it. There are two avenues of possible solutions. First, the Mercury compiler already contains an analysis that attempts to find out which procedures each higher order value may call. If this analysis succeeds, an adapted version of Algorithm 3 can convey the requirements of the calling context to these procedures, and convey to the caller the worst-case demands that any of the callees may make.
(e.g. in terms of which nodes they need merged to reflect their sharing). Second, in case the analysis fails (which may happen e.g. because the caller picks up those higher order values from a data structure created elsewhere), we need an interface between caller and callee that is standard and thus does not require negotiation (which is what the fixpoint iteration in Algorithm 3 represents).

Our search for this standard interface will not be restricted to RBMM-only systems. We will also look at hybrid systems in which RBMM coexists with the Boehm general purpose garbage collector, each looking after some of the program’s memory. Hybrid system that combine RBMM with a runtime collector have proven useful in other contexts (Hallenberg et al. 2002), and they may prove useful in this one as well. We do not intend to look at hybrid schemes that integrate RBMM with Mercury’s accurate garbage collector since that collector is actually significantly slower than the Boehm collector (Henderson 2002). We do however intend to look at integrating our RBMM system with the compile time garbage collection scheme reported in (Mazur et al. 2000; Mazur et al. 2001; Mazur 2004).

13 Conclusion

We have made region-based memory management available as an alternative storage management technique for programs written in a very large subset of Mercury. This involved the design and implementation of two program analyses (region points-to analysis and region liveness analysis) and a program transformation, the modification of the Mercury code generator to use the information produced by the analyses and transformation to generate code that uses RBMM to manage its memory, and the implementation of the primitive operations used by the generated code.

We provide termination and correctness theorems for our region analyses and our transformation algorithms. These ensure the safety of memory accesses and region operations with respect to forward liveness. Our discussions in Section 9 also strongly argue that our runtime support operations guarantee the safety of memory accesses and region operations with respect to backward liveness (i.e. in the presence of backtracking). These operations also instantly reclaim the memory allocated by backtracked-over computations, which help programs to reuse memory effectively.

The main challenge for the runtime support is to support backtracking correctly without incurring significant overhead, especially in deterministic code. Our experiments show that using RBMM instead of the Boehm collector yields nontrivial speedups for 15 out of our 18 benchmark programs, these speedups ranging from near 10% to a remarkable more than 60%. We even get large speedups for some benchmarks that are known to be difficult cases for RBMM. This indicates that the runtime support we provided for backtracking incurs very modest overhead in most cases, contributing to the overall better performance.

The memory use results of the benchmarks are also positive: in some programs we obtain optimal memory consumption. On average, our benchmarks require about one-twentieth the memory with RBMM than with the Boehm collector (only 5%), and even if we exclude the region-friendly programs, the figure is about one-
eighteenth (5.4%). This even before including any of the optimizations that have been studied for RBMM, such as stack allocation of regions (Birkeda l et al. 1996; Cherem and Rugina 2004), and merging regions that are removed at the same points (Makholm 2000a).

Everything we have described is available in current releases-of-the-day from the Mercury web site. The experimental setup for this paper is available from http://www.cs.kuleuven.be/~gerda/rbmm/rbmmbenchmarks.tar; it includes the benchmark programs as well as the benchmarking script.

References


