Modular Decompilation of Low-Level Code by Partial Evaluation*

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Abstract

Decompiling low-level code to a high-level intermediate representation facilitates the development of analyzers, model checkers, etc. which reason about properties of the low-level code (e.g., bytecode, .NET). Interpretive decompilation consists in partially evaluating an interpreter for the low-level language (written in the high-level language) w.r.t. the code to be decompiled. There have been proofs-ofconcept that interpretive decompilation is feasible, but there remain important open issues when it comes to decompile a real language: does the approach scale up? is the quality of decompiled programs comparable to that obtained by ad-hoc decompilers? do decompiled programs preserve the structure of the original programs? This paper addresses these issues by presenting, to the best of our knowledge, the first modular scheme to enable interpretive decompilation of low-level code to a high-level representation, namely, we decompile bytecode into Prolog. We introduce two notions of optimality. The first one requires that each method/block is decompiled just once. The second one requires that each program point is traversed at most once during decompilation. We demonstrate the impact of our modular approach and optimality issues on a series of realistic benchmarks. Decompilation times and decompiled program sizes are linear with the size of the input bytecode program. This demostrates empirically the scalability of modular decompilation of low-level code by partial evaluation.

1. Introduction

Decompilation of low-level code (e.g., bytecode) to an intermediate representation has become a usual practice nowadays within the development of analyzers, verifiers, model checkers, etc. For instance, in the context of *mo*-

bile code, as the source code is not available, decompilation facilitates the reuse of existing analysis and model checking tools. In general, high-level intermediate representations allow abstracting away the particular language features and developing the tools on simpler representations. As a representative example, Java bytecode is decompiled to a rule-based representation in [1], to clause-based programs in [18], to a three-address code view of bytecodes in Soot [20] and to the typed procedural language BoogiePL in [5]. Also, PIC programs are transformed to logic programs in [10]. Rule-based representations used in declarative programming in general-and in Prolog in particular-provide a convenient formalism to define such intermediate representations. E.g., as it can be seen in [1, 18, 20, 10], the operand stack used in a low-level language can be represented by means of explicit logic variables and that its unstructured control flow can be transformed into recursion.

All above cited approaches (except [10]) develop adhoc decompilers to carry out the particular decompilations. An appealing alternative to the development of dedicated decompilers is the so-called interpretive decompilation by partial evaluation (PE) [11]. PE is an automatic program transformation technique which specializes a program w.r.t. part of its known input data. Interpretive compilation was proposed in Futamura's seminal work [6], whereby compilation of a program P written in a (source) programming language L_S into another (*object*) programming language L_O is achieved by specializing an interpreter for L_S written in L_O w.r.t. P. The advantages of interpretive (de-)compilation w.r.t. dedicated (de-)compilers are wellknown and discussed in the PE literature (see, e.g., [3]). Very briefly, they include: *flexibility*, it is easier to modify the interpreter in order to tune the decompilation (e.g., observe new properties of interest); easier to trust, it is more difficult to prove that ad-hoc decompilers preserve the program semantics; easier to maintain, new changes in the language semantics can be easily reflected in the interpreter.

There have been several proofs-of-concept of interpretive (de-)compilation (e.g., [3, 10, 13]), but there remain interesting open issues when it comes to assess its power and/or limitations to decompile a real language: (a) does the

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approach scale? (b) do (de-)compiled programs preserve the structure of the original ones? (c) is the "quality" of decompiled programs comparable to that obtained by dedicated decompilers? This paper answers these questions positively by proposing a modular decompilation scheme which can be steered to control the structure of decompiled code and ensure quality decompilations which preserve the original program's structure. Our main contributions are summarized as follows:

- 1. We present the problems of *non-modular* decompilation and identify the components needed to enable a modular scheme. This includes how to write an interpreter and how to control an *online* partial evaluator in order to preserve the structure of the original program w.r.t. method invocations.
- 2. We present a modular decompilation scheme which is correct and complete for the proposed big-step interpreter. The *modular-optimality* of the scheme allows addressing issue (*a*) by avoiding decompiling the same method more than once, and (*b*) by ensuring that the structure of the original program can be preserved.
- 3. We introduce an interpretive decompilation scheme for low-level languages which answers issue (c) by producing decompiled programs whose *quality* is similar to that of dedicated decompilers. This requires a *block-level* decompilation scheme which avoids code duplication and code re-evaluation.
- 4. We report on a prototype implementation which incorporates the above techniques and demonstrate it on an set of realistic Java bytecode programs.

For the sake of concreteness, our decompilation scheme is formalized in the context of logic programming but the techniques to enable modularity can also be applied to compilation for any instantiation of languages (not necessarily low-level languages).

2 Basics of Partial Deduction

We assume familiarity with basic notions of logic programming [16]. Executing a program P for a call A consists in building an *SLD tree* for $P \cup \{A\}$ and then extracting the *computed answers* from every non-failing branch of the tree. PE in logic programming (see e.g. [7]) builds upon the SLD trees mentioned above. We now introduce a generic function PE, which is parametric w.r.t. the *unfolding rule*, unfold, and the *abstraction operator*, abstract and captures the essence of most algorithms for PE of logic programs:

```
1: function PE (P, \mathcal{A}, S_0)
```

```
2: repeat
```

```
3: T^{pe} := unfold(S_i, P, \mathcal{A});
```

```
4: S_{i+1} := \mathsf{abstract}(S_i, \mathit{leaves}(T^{pe}), \mathcal{A});
```

```
5: i := i + 1;
```

- 6: **until** $S_i = S_{i-1}$ % (modulo renaming)
- 7: **return** codegen $(T^{pe}, unfold);$

Function PE differs from standard ones in the use of the set of annotations A, whose role is described below. PE starts from a program P, a (possibly empty) set of annotations \mathcal{A} and an initial set of calls S_0 . At each iteration, the so-called *local control* is performed by the unfolding rule unfold (L3), which takes the current set of terms S_i , the program and the annotations and constructs a partial SLD tree for each call in S_i . Trees are partial in the sense that, in order to guarantee termination of the unfolding process, it must be possible to choose not to further unfold a goal, and rather allow leaves in the tree with a non-empty, possibly non-failing, goal. The particular unfold operator determines which call to select from each goal and when to stop unfolding. The partial evaluator may have to build several SLD-trees to ensure that all calls left in the leaves are "covered" by the root of some tree. This is known as the closedness condition of PE [17]. In the global control, those calls in the leaves which are not covered are added to the new set of terms to be partially evaluated, by the operator abstract (L4). At the next iteration, an SLD-tree is built for such call. Thus, basically, the algorithm iteratively (L2-6) constructs partial SLD trees until all their leaves are covered by the root nodes. An essential point of the operator abstract is that it has to perform "generalizations" on the calls that have to be partially evaluated in order to avoid computing partial SLD trees for an infinite number of calls. A partial evaluation of P w.r.t. S is then systematically extracted from the resulting set of calls T^{pe} in the final phase, codegen in L7. The notion of *resultant* is used to generate a program rule associated to each root-to-leaf derivation of the SLD-trees for the final set of terms T^{pe} . Given an SLD derivation of $P \cup \{A\}$ with $A \in T^{pe}$ ending in B and θ be the composition of the mgu's in the derivation steps, the rule $\theta(A) : -B$ is called the *resultant* of the derivation. A PE is defined as the set of resultants associated to the derivations of the constructed partial SLD trees for all $P \cup T^{pe}$.

The notions of *completeness* and *correctness* of PE [7] ensure that the specialized program produces no less resp. no more answers than the original program. A sufficient condition to ensure completeness is that the specialized program is *closed* by the resulting set of terms T^{pe} . Intuitively, the closedness condition ensures that all calls which may arise during the computation of $P \cup S$ are instances of T^{pe} and hence there is a matching resultant for them (solutions are not lost). The abstraction operator is encharged of ensuring that the closedness condition is met by means of a proper generalization of calls. Correctness is achieved when the resulting set T^{pe} is independent, i.e., there are no two calls in T^{pe} which unify. Independence can be easily recovered by a post-processing of renaming, which often does argument filtering [7]. Finally, the role of the annotations \mathcal{A} (often manually provided) in *offline* PE is to give information to the control operators to decide when to stop derivations in the local control and how to perform generalizations in the global control to ensure termination. In *online* PE, all control decisions are taken during the specialization phase, without the use of annotations. We trivially turn function *PE* into online by just ignoring the annotations.

3 Non-Modular Interpretive Decompilation

This section describes the state of the art in interpretive decompilation of low-level languages to Prolog, including recent work in [10, 2, 9, 3]. We do so by formulating nonmodular decompilation in a generic way and identifying its limitations. The low-level language we consider, denoted as \mathcal{L}_{bc} , is a simple imperative bytecode language in the spirit of Java bytecode but, to simplify the presentation, without object-oriented features (our implementation supports full Java bytecode). It uses an operand stack to perform computations. It has an unstructured control flow with explicit conditional and unconditional goto instructions and manipulates only integer numbers. A bytecode program P_{bc} is organized in a set of methods which are the basic (de-) compilation units of \mathcal{L}_{bc} . The code of a method m, denoted code(m), consists of a sequence of bytecode instructions $BC_m = \langle pc_0 : bc_0, \dots, pc_{n_m} : bc_{n_m} \rangle$ with pc_0, \dots, pc_{n_m} being consecutive natural numbers. The \mathcal{L}_{bc} instruction set is:

```
\begin{array}{l} \textit{BcInst} ::= \texttt{push}\left(\texttt{x}\right) \mid \texttt{load}\left(\texttt{v}\right) \mid \texttt{store}\left(\texttt{v}\right) \mid \texttt{add} \mid \texttt{sub} \mid \texttt{mul} \mid \texttt{div} \mid \texttt{rem} \mid \\ \mid \texttt{neg} \mid \texttt{if} \diamond \left(\texttt{pc}\right) \mid \texttt{if0} \diamond \left(\texttt{pc}\right) \mid \texttt{goto}\left(\texttt{pc}\right) \mid \texttt{return} \mid \texttt{call}\left(\texttt{mn}\right) \end{array}
```

where \diamond is a comparison operator (eq, le, gt, etc.), v a local variable, x an integer, pc an instruction index and mn a method name. Instructions push, load and store transfer values or constants from the local variables to the stack (and viceversa); add, sub, mul, div, rem and neg perform the usual arithmetic operations, being rem the division remainder and neg the arithmetic negation; if and if 0 are conditional branching instructions (with the special case of comparisons with 0); goto is an unconditional branching; return marks the end of methods and call invokes a method. A method m is uniquely determined by its name. We write calls(m) to denote the set of all method names invoked within the code of m. We use $defs(P_{bc})$ to denote the set of *internal* method names defined in P_{bc} . The remaining methods are *external*. We say that P_{bc} is self-contained if $\forall m \in P_{bc}, calls(m) \subseteq defs(P_{bc}), \text{ i.e., } P_{bc}$ does not include calls to external methods.

3.1 Non-modular, Online Decompilation

We rely on the so-called "interpretive approach" to compilation by PE described in Sect. 1, also known as first

```
main(Method, InArgs, Top) :-
   build_s0(Method, InArgs, S0), execute(S0,Sf),
   Sf = st(fr(_, [Top]], ), )).
execute(S,S) :
   S = st(fr(M, PC, [_Top|_],_),[]),
   bytecode (M, PC, return, )
execute(S1,Sf) :
   S1 = st(fr(M, PC, _, _), _), bytecode(M, PC, Inst, _),
   step(Inst,S1,S2), execute(S2,Sf).
step(goto(PC),S1,S2) :-
   S1 = st(fr(M,_,S,LV),FrS)
   S2 = st(fr(M, PC, S, LV), FrS).
step(push(X), S1, S2) :
   S1 = st(fr(M, PC, S, L), FrS), next(M, PC, PC2),
   S2 = st(fr(M, PC2, [X|S], L), FrS).
step(call(M2),S1,S2) :-
   S1 = st(fr(M, PC, OS, LV), FrS), split_OS(M2, OS, Args, OS3),
   build_s0(M2,Args,st(fr(M2,PC2,OS2,LV2),_)),
   S2 = st(fr(M2, PC2, OS2, LV2), [fr(M, PC, OS3, LV) | FrS]).
step(return,S1,S2) :
   p(leculi,fr(_, [RV|], _), [fr(M,PC,OS,LV)|FrS]),
next(M,PC,PC2), S2 = st(fr(M,PC2, [RV|OS],LV),FrS).
```

Figure 1. Fragment of (small-step) \mathcal{L}_{bc} interpreter

Futamura projection [6]. In particular, the decompilation of a \mathcal{L}_{bc} -bytecode program P_{bc} to LP (for short LPdecompilation) might be obtained by specializing (with an LP partial evaluator) a \mathcal{L}_{bc} -interpreter written in LP w.r.t. P_{bc} . In Fig. 1 we show a fragment of a (small-step) \mathcal{L}_{bc} interpreter implemented in Prolog, named $Int_{\mathcal{L}_{bc}}$. We assume that the code for every method in the bytecode program P_{bc} is represented as a set of facts bytecode/3 such that, for every pair $pc_i : bc_i$ in the code for method m, we have a fact bytecode (m, pc_i, bc_i) . The state carried around by the interpreter is of the form st (Fr, FrStack) where Fr represents the current frame (environment) and FrStack the stack of frames (call stack) implemented as a list. Frames are of the form fr(M, PC, OStack, LocalV), where M represents the current method, PC the program counter, OStack the operand stack and LocalV the list of local variables. Predicate main/3, given the method to be interpreted Method and its input method arguments InArgs, first builds the initial state by means of predicate $build_{0}/3$ and then calls predicate execute/2. In turn, execute/2 calls predicate step/3, which produces S2, the state after executing the bytecode, and then calls predicate execute/2 recursively with S2 until we reach a return instruction with the empty stack. For brevity, we only show the definition of step/3 for a selected set of instructions and omit the code of $build_s0/3$ and localVar_update/4. The latter simply updates the value of a local variable. By using this interpreter, in a purely online setting, we define a non-modular decompilation scheme in terms of the generic function PE as follows.

Definition [DECOMP_{\mathcal{L}_{bc}}] Given a self-contained \mathcal{L}_{bc} -bytecode program P_{bc} , the (non-modular) LP-decompilation of P_{bc} can be obtained as:

 $DECOMP_{\mathcal{L}_{bc}}(P_{bc}) = PE(Int_{\mathcal{L}_{bc}} \cup P_{bc}, \emptyset, S)$ where S is the set of calls {main(m, _, _) | m \in defs(P_{bc})}.

<pre>int gcd(int x, int y) { int res; while (y != 0) { res = x%y; x = y; y = res; } return abs(x); }</pre>		<pre>int lcm(int x, int y) { int gcd = gcd(x,y); if (gcd == 0) return 0; else return x*y/gcd; int fact(int x) { if (x == 0)</pre>	
int abs(int x){		return 1;	
if $(x < 0)$ return $-x;$		else	
else return x; }		return x*fact(x-1);}	
Method gcd/2 0:load(1) 1:if0eq(11) 2:load(0) 3:load(1) 4:rem 5:store(2) 6:load(1) 7:store(0) 8:load(2) 9:store(1) 10:goto 0 11:load(0) 12:call(abs) 13:return	Method abs/1 0:load(0) 1:if0ge(5) 2:load(0) 3:neg 4:return 5:load(0) 6:return	Method lcm/2 0:load(0) 1:load(1) 2:call(gcd) 3:store(2) 4:load(2) 5:if0ne 8 6:push(0) 7:return 8:load(0) 9:load(1) 10:mul 11:load(2) 12:div 13:return	<pre>Method fact/1 0:load(0) 1:ifOne(4) 2:push(1) 3:return 4:load(0) 5:load(0) 6:push(1) 7:sub 8:call(fact) 9:mul 10:return</pre>

Figure 2. Source code and \mathcal{L}_{bc} -bytecode for working example

Recent work in interpretive, online decompilation has focused on ensuring that the layer of interpretation is completely removed from decompiled programs, i.e., *effective* decompilations are obtained. This requires the use of advanced control strategies as explained in [2] and [9]. Our starting point is thus a state-of-the-art partial evaluator which incorporates such advanced techniques and which is able to remove the layer of interpretation.

3.2 Limitations

This section illustrates by means of the bytecode example in Fig. 2 that non-modular decompilation does not ensure a satisfactory handling of issues (a) and (b). In the examples, we often depict the Java source code for clarity, but the partial evaluator works directly on the bytecode. The program consists of a set of methods that carry out arithmetic operations. Method gcd computes the greatest-common divisor, abs the absolute value, lcm the least-common multiple and fact the factorial recursively. The LP-decompilation obtained by applying Def. 3.1 is shown in Fig. 3. We identify the following limitations of non-modular decompilation:

(L1) Method invocations from lcm to gcd (index 2) and from gcd to abs (index 12) do not appear in the decompiled code. Instead, such calls have been *inlined* within their calling contexts and, as a consequence, the structure of the original code has been lost. For instance, the last two rules in the decompilation for lcm, execute_1, correspond to the while loop of gcd.

(L2) As a consequence, decompilation might become very inefficient. E.g., if n calls to the same method appear within a code, such method will be decompiled n times. This might be even worse in teh case of loops.

<pre>main(lcm,[B,0],A) :- B>0, C is B*0, A is C//B. main(lcm,[0,0],0). main(lcm,[B,0],A) :-</pre>	<pre>main(gcd,[A,0],A) :-A>=0. main(gcd,[B,0],A) :-</pre>	
B<0, D is B*0, C is -B, A is D//C.	C = 0, D is B rem C, execute $2(C, D, A)$.	
main(lcm,[B,C],A) :-		
C\=0, D is B rem C,	execute_2(A,0,A) :-	
execute_1(C,D,B,C,A).	A>=0.	
	execute_2(A,0,C) :-	
execute_1(A,0,B,C,D) :-	A<0, C is-A.	
A>0, E is $B*C$, D is $E//A$.	execute_2(A,B,G) :-	
execute_1(0,0,_,_,0).	B \=0,	
execute_1(A,0,B,C,D) :-	I is A rem B,	
A<0, E is-A,	execute_2(B,I,G).	
F is B*C, D is F//E.		
execute_1(A,B,C,D,I) :-	<pre>main(abs,[A],A) :- A>=0.</pre>	
B\=0, K is A rem B,	main(abs,[B],A) :-	
execute_1(B,K,C,D,I).	B<0, A is-B.	

Figure 3. Decompiled (non-modular) code for working example

(L3) The non-modular approach does not work incrementally, in the sense that it does not support *separate* decompilation of methods but rather has to (re)decompile all method calls. Thus, decompiling a real language becomes unfeasible, as one needs to consider system libraries. Limitation L2 together with L3 answer issue (*a*) negatively.

(L4) The decompiled program does not contain the code corresponding to recursive fact due to space limitations, as the decompiled code contains basically the whole interpreter. The problem with recursion is: assume we want to decompile method m1 whose code is $< pc_0 : bc_0, \ldots, pc_j :$ $call(m1), \ldots, pc_n : return >$. There is an initial decompilation for $A_k = \texttt{execute}(\texttt{st}(\texttt{fr}(\texttt{m1},\texttt{pc}_1,\texttt{os},\texttt{lv}),[]),\texttt{S}_f)$ in which the call stack is empty. During decompilation, a call of the form A_l its $execute(st(fr(m1, pc_i, os', lv'), [fr(m1, pc_i, os, lv)]), S_f)$ with the call stack containing the previous frame appears when we get to the recursive call. At this point, the derivation must be stopped as $A_k \leq T A_l$. In order to ensure termination, the global control generalizes the above calls into execute(st(fr(m1, pc₁, -, -), -), S_f), where _ denotes a fresh variable and thus the call-stack has become unknown. As a consequence, after evaluating the return statement, the continuation obtained from the call-stack is unknown and we produce the call $execute(st(fr(_,_,_,_),_), S_f)$ to be decompiled. Here, the fact that the method and the program counter are unknown prevents us from any chance of removing the interpretation layer, i.e., the decompiled code will potentially contain the whole interpreter. This indeed happens during the decompilation of fact. Partial solutions to the recursion problem exist and will be discussed later. Limitations L1 and L4 answer issue (b) negatively.

4 A Modular Decompilation Scheme

By *modular* decompilation, we refer to a decompilation technique whose decompilation unit is the method, i.e., we

decompile a method at a time. We show that this approach overcomes the four limitations of non-modular decompilation described in Sect. 3.2 and answers issues (a) and (b) positively. In essence, we need to: (i) give a compositional treatment to method invocations, we show that this can be achieved by considering a *big-step* interpreter; (ii) provide a mechanism to residualize calls in the decompiled program, we automatically generate program annotations for this purpose; (iii) study the conditions which ensure that *separate* decompilation of methods is sound.

4.1 Big-step Semantics Interpreter

Traditionally, two different approaches have been considered to define language semantics, *big-step* (or *natural*) semantics and *small-step* semantics (see, e.g., [12]). Essentially, in big-step semantics, transitions relate the initial and final states for each statement, while in small-step semantics transitions define the *next* step of the execution for each statement. In the context of bytecode interpreters, it turns out that most of the statements execute in a single step, hence making both approaches equivalent for such statements. This is the case for our \mathcal{L}_{bc} -bytecode interpreter for all statements except for *call*. The transition for *call* in small-step defines the next step of the computation, i.e., the current frame is pushed on the call-stack and a new environment is initialized for the execution of the invoked method. Note that, after performing this step, we do not distinguish anymore between the code of the caller method and that of the callee. This avoids modularity of decompilation.

In the context of interpretive (de-)compilation of imperative languages, small-step interpreters are commonly used (see e.g. [19, 10, 3]). We argue that the use of a big-step interpreter is a necessity to enable modular decompilation which scales to realistic languages. In Fig. 4, we depict the relevant part of the big-step interpreter for \mathcal{L}_{bc} -bytecode, named $Int_{\mathcal{L}_{bc}}^{bs}$. We can see that the *call* statement, after extracting the method parameters from the operand stack, calls recursively predicate main/3 in order to execute the callee. Upon return from the method execution, the return value is pushed on the operand stack of the new state and execution proceeds normally. Also, we do not need to carry the call-stack explicitly within the state, but only the information for the current environment. I.e., states are of the form st (M, PC, OStack, LocalV). This is because the call-stack is already available by means of the calls for predicate main/3.

The compositional treatment of methods in $Int_{\mathcal{L}bc}^{bs}$ is not only essential to enable modular decompilation (overcome L1, L2 and L3) but also to solve the recursion problem in a simple and elegant way. Indeed, the decompilation based on the big-step interpreter $Int_{\mathcal{L}bc}^{bs}$ does not present L4. E.g., the decompilation of a recursive method m1 starts from the

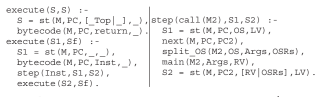


Figure 4. Fragment of big-step \mathcal{L}_{bc} interpreter $Int_{\mathcal{L}_{bc}}^{bs}$

call $main(m1, _, _)$ and then reaches a call $main(m1, args, _)$ where args represents the particular arguments in the recursive call. This call is flagged as dangerous by local control and the derivation is stopped. The important points are that, unlike before, no recomputation is needed as the second call is necessarily an instance of the first one and, besides, there is no information loss associated to the generalization of the call-stack, as there is no stack. The recursion problem was first detected in [8] and a solution based on computing regular approximations during PE was proposed. Although feasible in theory, such technique might be too inefficient in practice and problematic to scale it up to realistic applications due to the overhead introduced by the underlying analysis. Another solution is proposed in [10], where a simpler control-flow analysis is performed before PE in order to collect all possible instructions which might follow the return. Such information may then be used to recover information lost by the generalization. This solution turns out to be also impractical for our purposes when considering realistic programs that make intensive use of library code (e.g. Java bytecode) as many continuations can follow. Our solution does not require the use of static analysis and, as our experiments show, does not pose scalability problems.

4.2 Guiding Online PE with Annotations

We now present the annotations we use to provide additional control information to PE. They are instrumental for obtaining the quality decompilation we aim at. We use the annotation schema: " $[Precond] \Rightarrow Ann Pred$ " where *Precond* is an optional precondition defined as a logic formula, *Ann* is the kind of annotation (*Ann* \in {**memo**, **rescall**}) and *Pred* is a predicate descriptor, i.e., a predicate function and distinct free variables. Such annotations are used by local control when a call for *Pred* is found as follows:

- **memo**: The current call is not further unfolded. Therefore, the call is later transferred to the global control to carry out its specialization separately.
- **rescall**: The current call is not further unfolded. Unlike calls marked **memo**, the current call is not transferred to the global control.

In the following, we denote by $unfold_{\leq T}^{\mathcal{A}}$ the unfolding operator of Sect. 2 enhanced to use the above annotations. We adopt the same names for the annotations as in offline PE

[15]. However, in offline PE they are the *only* means to control termination and **rescall** annotations are in principle only used for builtins.

4.3 Modular Decompilation

In order to achieve modular decompilation, it is instrumental to allow performing separate decompilation. In the interpretive approach this requires being able to perform separate PE, i.e., to be able to specialize parts of the program independently and then join the specializations together to form the residual program. For instance, consider a self-contained logic program P partitioned in a set $\{P_1, \ldots, P_n\}$ of mutually disjoint subprograms which preserve predicate boundaries, i.e., for any predicate pred in P we have that all rules for pred are in the same partition P_j , for some $j \in \{1, \ldots, n\}$. Consider also the sets of terms S_1, \ldots, S_n such that all calls in S_i correspond to predicates defined in P_i , i = 1, ..., n. We can now define $S = S_1 \cup \cdots \cup S_n$ and the usual notions of closedness and independence are applicable. A separate partial evaluation for P and S is obtained as the union of the individual specializations w.r.t. each corresponding set of calls, i.e., $\bigcup_{P_i \in P} PE(P_i, \emptyset, S_i)$. One additional difficulty for separate PE is related to the use of renaming for guaranteeing independence, since renaming requires a global table which is not available when generating code for the individual subprograms. A simple strategy which we will use in our modular decompilation is to allow polyvariant specialization for calls to predicates locally defined in the subprogram P_i being partially evaluated but to resort to monovariant specialization for predicates used across subprogram boundaries. Then, the renaming can use a local renaming table, which must guarantee that there will be no name clash with renamed calls from other subprograms.

We present now a modular decompilation scheme which, by combining the big-step interpreter with the use of **rescall** annotations, enables separate decompilation and ensures *soundness* (i.e., it is correct and complete w.r.t. internal methods).

Definition [MOD-DECOMP $_{\mathcal{L}_{bc}}$] Given a \mathcal{L}_{bc} -bytecode program P_{bc} , a modular LP-decompilation of P_{bc} can be obtained as:

$$\text{MOD-DECOMP}_{\mathcal{L}_{bc}}(P_{bc}) = \bigcup_{\substack{\forall m \in defs(P_{bc})}} PE(Int^{bs}_{\mathcal{L}_{bc}} \cup code(m), \mathcal{A}_{mod}, S_m)$$

where the set of annotations $\mathcal{A}_{mod} = \{(m \in calls(P_{bc})) \Rightarrow$ rescall $main(m, ..., ...)\}$ and the initial sets of calls $S_m = \{main(m, ..., ...)\}$ for each $m \in defs(P_{bc})$.

Let us briefly explain the above definition. Now the function PE is executed once per method defined in P_{bc} , starting each time from a set of calls, S_m , which contains a call of the form main(m, -, -) for method m. The set \mathcal{A}_{mod} contains a **rescall** annotation which affects all methods invoked (but not necessarily internal) inside P_{bc} . When a method invocation is to be decompiled, the call step (call (m'), -, -) occurs during unfolding. We can see that, by using the bigstep interpreter in Fig. 4, a subsequent call main (m', -, -) will be generated. As there is a **rescall** annotation which affects all methods invoked in the program, such call is not unfolded but rather remains residual. If m' is internal, a corresponding decompilation from the call main (m', -, -) will be, or has already been, performed since function PE is executed for every method in P_{bc} . Thus, completeness is ensured for internal predicates.

Example 1 By applying function MOD-DECOMP_{$\mathcal{L}bc}$ to the \mathcal{L}_{bc} -bytecode program in Fig. 2 we execute PE once for each of the four methods in the program. In each execution we specialize the interpreter w.r.t. the calls main(fact,_,_), main(gcd,_,), main(lcm,_,_), and main(abs,_,). We obtain the following LP-decompilation:</sub>

The structure of the original program w.r.t. method calls is preserved, as the residual predicate for lcm contains an invocation to the definition of gcd, which in turn invokes abs, as it happens in the original bytecode. Moreover, we now obtain an effective decompilation for the recursive method fact where the interpretive layer is completely removed without the need of any analysis. Thus, L1 and L4 have been successfully solved.

The following theorem ensures the soundness of modular decompilation for the big-step bytecode interpreter. Completeness can be ensured by excluding calls to external methods not defined in the bytecode. It is independent of the way the interpreter is defined, as the closedness condition for the internal methods is enforced by our definitions of \mathcal{A}_{mod} and S_m . Correctness holds in the case of our interpreter, because the only calls which are transferred to the global control are instances of main/3 and execute/2 and their first argument is the method's name, which makes them mutually exclusive. A post-processing of renaming is thus optional, but it can be necessary to ensure that the independence condition is met for other interpreters.

Theorem 1 (soundness) Consider a \mathcal{L}_{bc} -bytecode program P_{bc} and a concrete input I. Let P'_{bc} be the result of MOD-DECOMP $_{\mathcal{L}_{bc}}(P_{bc})$ and I' the LP representation of I. Then, A' is an answer for $P'_{bc} \cup \{I'\}$ iff A is the result of executing P_{bc} for the input I, where A' is the LP representation of A.

We now characterize the notion of *modular-optimality* in decompilation which ensures that (1) only the code associated to internal methods is decompiled, thus, we can have external calls (e.g., to libraries) which are not decompiled and overcome L3; (2) and each method is decompiled only once and thus we overcome L2.

Proposition 1 (modular-optimality) Given a \mathcal{L}_{bc} bytecode program P_{bc} , function MOD-DECOMP $_{\mathcal{L}_{bc}}$ only decompiles the code corresponding to internal methods defined in P_{bc} , and the code of each method is decompiled once.

Note that modular decompilation gives a monovariant treatment to methods in the sense that it does not allow creating specialized versions of method definitions. This is against the usual spirit in PE, where polyvariance is a main goal to achieve further specialization. However, in the context of decompilation, we have shown that a monovariant treatment is necessary to enable scalability and to preserve program structure. It naturally raises the question whether a polyvariant treatment could achieve, even if at the cost of efficiency and loss of structure, a better quality decompilation. Note that enabling polyvariant specialization in the modular setting can be trivially done by not introducing rescall annotations for certain selected methods which should be treated in a polyvariant manner. Our experience indicates that there is often a small quality gain at the price of a highly inefficient decompilation.

5 Decompilation of Low-Level Languages

Applying the interpretive approach on a low-level language introduces new challenges. The main issue is whether it is possible to obtain, by means of interpretive decompilation, programs whose *quality* is equivalent to that obtained by dedicated decompilers, issue (c) in Sect. 1. We will show now that, using the most effective unfolding strategies of PE, code for the same program point can be emitted (i.e. it can be decompiled) several times, which degrades both efficiency and quality of decompilation. In order to obtain results which are comparable to that of dedicated decompilers, it makes sense to use similar heuristics. Since decompilers first build a *control flow graph* (CFG) for the method, which guides the decompilation process, we now study how a similar notion can be used for controlling PE of the interpreter. Let us explain *block-level* decompilation by means of an example. Consider the method m_{bl} to the left of Fig. 5, where we only show the relevant bytecode instructions, and its CFG in the center. A *divergence point* (D point) is a program point (bytecode index) from which more than one branch originates; likewise, a *convergence point* (C point) is a program point where two or more branches merge. In the CFG of m_{bl} , the only divergence (resp. convergence) point is pc_i (resp. pc_k).

By using the decompilation scheme presented so far, we obtain the SLD-tree shown to the right of Fig. 5, in which all calls are completely unfolded as there is no termination risk (nor **rescall** annotation). The decompiled code is shown under the tree. We use { res_X } to refer to the residual code emitted for BlockX and cond_i to refer to the condition associated to the branching instruction at pc_i (cond_i denotes its negation). The quality of the decompiled code is not optimal due to:

- D. Decompiled code $\{res_A\}$ for BlockA is duplicated in both rules. During PE, this code is evaluated once but, due to the way resultants are defined (see codegen in Sect. 2), each rule contains the decompiled code associated to the whole branch of the tree. This code duplication brings in two problems: it increases considerably the size of decompiled programs and also makes their execution slower. For instance, when $cond_i$ holds, the execution goes unnecessarily through $\{res_A\}$ in the first rule, fails to prove cond_i and, then, attempts the second rule.
- C. Decompiled code of BlockD is again emitted more than once. Each rule for the decompiled code contains a (possibly different) version, $\{res_D\}$ and $\{res'_D\}$, of the code of BlockD. Unlike above, at PE time, the code of BlockD is actually evaluated in the context of $\{cond_i, \{res_B\}\}$ and then re-evaluated in the context of $\{\overline{cond_i}, \{res_C\}\}$. Convergence points thus might degrade both efficiency (and endanger scalability) and quality of decompilation (due to larger residual code).

The amount of repeated residual code grows exponentially with the number of C and D points and the amount of reevaluated code grows exponentially with the number of C points. Thus, we now aim for a *block-level* decompilation that helps overcome problems D and C above. Intuitively, a block-level decompilation must produce a residual rule for each block in the CFG. This can be achieved by building SLD-trees which correspond to each single block, rather than expanding them further.

The **memo** annotations presented in Sect. 4.2 facilitate the design of the block-level interpretive decompilation scheme. In particular, we can easily force the unfolding process to stop at D points by including a **memo** annotation for execute/2 calls whose *PC* corresponds to a D point. In

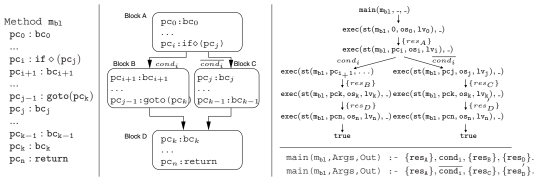


Figure 5. \mathcal{L}_{bc} -bytecode, CFG, unfolding tree and decompiled code of m_{bl} method

the example, unfolding stops at pc_i as desired. Regarding C points, an additional requirement is to partially evaluate the code on blocks starting at these points at most once. The problem is similar to the polyvariant vs monovariant treatment in the decompilation of methods in Sect. 4.3, by viewing entries to blocks as method calls. Not surprisingly, the solution can be achieved similarly in our setting by: (1) stopping the derivation at execute/2 calls whose PC corresponds to C points and (2) passing the call to the global control, and ensuring that it is evaluated in a sufficiently generalized context which covers all incoming contexts. The former point is ensured by the use of **memo** annotations and the latter by including in the initial set of terms a generalized call of the form $execute(st(m_{b1}, pc_{k}, ..., ..), ...)$ for all C points, which forces such generalization. The next definition presents the block-level decompilation scheme where $div_points(m)$ and $conv_points(m)$ denote, resp., the set of D points and C points of a method m.

Definition [BLOCK-MOD-DECOMP \mathcal{L}_{bc}] Given a \mathcal{L}_{bc} -bytecode program P_{bc} , a block-level, modular LP-decompilation of P_{bc} can be obtained as:

An important point is that, unlike annotations used in offline PE [13] which are generated by only taking the interpreter into account, our annotations for block-level decompilation are generated by taking into account the particular program to be decompiled. Importantly, both the annotations and the initial set of calls can be computed automatically by performing two passes on the bytecode (see, e.g., [1, 20]). The result of performing block-level decompilation on m_{bl} is:

Now, the residual code associated to each block appears once in the code. This ensures that block-level decompilation preserves the CFG shape as dedicated decompilers do. Thus, the quality of our decompiled code is as good as that obtained by state-of-the-art decompilers [1, 18] but with the advantages of interpretive decompilation (see Sect. 1). We formalize the quality of block-level decompilation.

Proposition 2 (block-optimality) Given a bytecode program P_{bc} , the block-level decompilation function BLOCK-MOD-DECOMP_{\mathcal{L}_{bc}} ensures that: (I) residual code for each bytecode instruction in P_{bc} is emitted once in the decompiled program, and (II) each bytecode instruction in P_{bc} is evaluated at most once during PE.

6 Experimental Evaluation

We report on our implementation of a decompiler for full (sequential) Java Bytecode into Prolog. The extensions needed to handle the features of Java Bytecode not considered in \mathcal{L}_{bc} (exception handling, dynamic memory allocation, object orientation, etc) are basically carried out by making the decompiler produce the corresponding builtins in the residual code. E.g. the bytecode instruction *putfield* will make the decompiler produce the predicate set_field/5 in the decompiled code. This naive solution might be considerably improved to increase the precision and quality of the decompilation. However this is out of the scope of this paper. For the experimental evaluation we have used the set of benchmarks in the JOlden suite [4]. Most programs make an extensive use of library methods. Hence, non-modular decompilation cannot be assessed as we run into memory problems when trying to decompile the code of library calls. The experiments have been performed on an Intel Core 2 Duo 1.86GHz with 2GB of RAM, running Linux. Figure 6 depicts four charts measuring different aspects of the decompilation. We assess the differences between the modular and the modular+block-level (just block*level* for short) approaches; as well as how the size of the programs affects the decompilation. We measure two as-

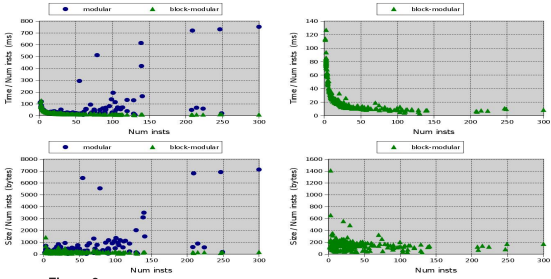


Figure 6. Evaluating modular decompilation vs. modular+block-level deompilation with the JOIden Suite

pects of the decompilation: the decompilation time (in milliseconds) per instruction and the decompiled program size (in bytes) per instruction. The decompilation time indicates the efficiency of the process and the size of decompiled programs is directly related to the decompilation quality. Each point [X, Y] in the charts corresponds to the decompilation of a single method in the JOlden suite, where X represents the number of instructions of the method and Y the measured data (time or decompiled program size). The tables in the left-hand side show the data obtained (times in the top chart and sizes in the bottom one) for both the modular and the block-level decompilation. The variations in the blocklevel decompilation cannot be appreciated when combined with modular. Thus, we include in the tables on the righthand side the figures for the block-level decompilation in isolation such that we adjust the scale on the Y-axis to the domain of the data.

From the charts, we conclude: (1) Times per instruction are notably larger for the smallest methods, as can be seen by looking at the initial curve in the charts. This is because the overhead introduced for starting a new decompilation is more noticeable when the time for decompilation itself is small, while it becomes negligible for larger methods. The same happens for the size of the decompiled programs. (2) Block-level decompilation achieves important speedups in general (for all methods with more than 40 instructions). Besides, it obtains significantly smaller decompiled programs. The speedups per package range from 3.36 in **power** to 31.4 in **bisort** for the decompilation times; and from 2.5 times smaller in **power** to 9 times smaller in **bisort** for the decompiled program sizes. Note that there is a clear correspondence between both measures, since C points introduce both inefficiency and size increase in decompilation, as explained in Sect. 5. Moreover, modular decompilation runs out of memory for some of the largest methods.

This is again related to code duplication (C and D points) and (re-)evaluation (C points), which grow exponentially. (3) The most important conclusion is that, while in modular decompilation both the times and the sizes per instruction greatly increase with the size of the benchmarks, this does not happen in the block-level scheme. In block-level decompilation, these figures are totally stable (mostly constant) for all methods with more than 40 instructions. This demonstrates that both the decompilation times and the decompiled program sizes are *linear* with the size of the input bytecode program, thus demonstrating the scalability of the block-level decompilation. One might wonder why there are still small variations in the ratio. In our experience, the following points also matter: 1) the complexity of the control flow of the methods, 2) the relative complexity of the bytecode instructions used, e.g., instructions which operate in the heap tend to produce more residual code, 3) the structure w.r.t. methods of the program, e.g., classes with methods of medium size tend to result in better decompilations than those with few large methods or many small ones.

7 Conclusions and Related Work

We argue that *declarative languages* and the technique of *partial evaluation* have nowadays a large application field within the development of analysis, verification, and model checking tools for modern programming languages. On one hand, declarative languages provide a convenient intermediate representation which allows (1) representing all iterative constructs (loops) as recursion, independently of whether they originate from iterative loops (conditional and unconditional jumps) or recursive calls, and (2) all variables in the local scope of the methods (formal parameters, local variables, fields, and stack values in low-level languages) can be represented uniformly as explicit arguments of a declar-

ative program. On the other hand, the technique of partial evaluation enables the automatic (de-)compilation of a (complicated) modern program to a simple declarative representation by just writing an interpreter for the modern language in the corresponding declarative language and using an existing partial evaluator. The resulting intermediate representation greatly simplifies the development of the above tools for modern languages and, more interestingly, existing advanced tools developed for declarative programs (already proven correct and effective) can be directly applied on it.

Previous work in interpretative (de-)compilation has mainly focused on proving that the approach is feasible for small interpreters and medium-sized programs. The focus has been on demonstrating its effectiveness, i.e., that the socalled interpretation layer can be removed from the compiled programs. To achieve effectiveness, offline [13], online [3, 10, 19] and hybrid [14] PE techniques have been assessed and novel control strategies have been proposed and proved effective [9, 2]. Our work starts off from the premise that interpretive decompilation is feasible and effective as proved by previous work and studies further issues which have not been explored before. A main objective of our work is to investigate, and provide the necessary techniques, to make interpretive decompilation scale in practice. A further goal is to ensure, and provide the techniques, that decompiled programs preserve the structure of the original programs and that its quality is comparable to that obtained by dedicated decompilers. We believe that the techniques proposed in this paper, together with their experimental evaluation, provide for the first time actual evidence that the interpretive theory proposed by Futamura in the 70s is indeed an appealing and feasible alternative to the development of ad-hoc decompilers from modern languages to intermediate representations.

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