Shimple: An Investigation of Static Single Assignment Form

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Abstract

In designing compiler analyses and optimisations, the choice of intermediate representation (IR) is a crucial one. *Static Single Assignment* (SSA) form in particular is an IR with interesting properties, enabling certain analyses and optimisations to be more easily implemented while also being more powerful. Our goal has been to research and implement various SSA-based IRs for the Soot compiler framework for Java.

We present three new IRs based on our Shimple framework for Soot. Simple Shimple is an implementation of the basic SSA form. We explore extensions of SSA form (Extended SSA form and Static Single Information form) and unify them in our implementation of Extended Shimple. Thirdly, we explore the possibility of applying the concepts of SSA form to array element analysis in the context of Java and present Array Shimple.

We have designed Shimple to be extensible and reusable, facilitating further research into variants and improvements of SSA form. We have also implemented several analyses and optimisations to demonstrate the utility of Shimple.

Résumé

En concevant des analyses et des optimisations, le choix de la représentation intermédiaire (RI) utilisée par le compilateur est crucial. La forme *Static Single Assi*gnment (SSA) est en particulier une RI avec des propriétés intéressantes, facilitant la conception d'analyses et d'optimisations plus puissantes. Notre but a été de rechercher et d'executer l'implementation d'une RI basée sur la forme SSA pour le cadre de compilateur Soot pour Java.

Nous présentons trois nouvelles RIs fondées sur notre cadre Shimple pour Soot. Simple Shimple est une implementation de base de la forme SSA. Nous explorons des extensions de la forme SSA (eSSA et SSI) et nous les unifions dans notre implementation de Extended Shimple. Troisièmement, nous explorons la possibilité d'appliquer les concepts de la forme SSA à l'analyse des éléments de tableaux et présentons Array Shimple.

Nous avons conçu Shimple pour être extensible et réutilisable, facilitant davantage la recherche dans des variantes ou des améliorations de la forme SSA. Nous avons également implementé plusieurs analyses et optimisations pour démontrer la valeur de Shimple.

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Chapter 1 Introduction

High-level programmers regularly employ high-level language features, abstractions and tools such as automated code generators to facilitate the programming task, often with the expectation that the resulting code will prove reasonably efficient.

Given a particular language with particular features and abstractions, it is reasonable to expect a compiler to eliminate inherent inefficiencies of that language. However, analysing a complex language for the purpose of optimisation is by no means a trivial task, and when confronted with the optimisation of programs in different languages of various complexities and idiosyncrasies, the task for compiler writers becomes monumental. Analyses must be rewritten for each language and each target architecture, and there is little or no sharing of code involved.

The problem of redesigning individual optimisations for different high-level languages is often side-stepped by compiling these languages to a common lower-level language, also known as an *intermediate representation* (IR), and instead applying analyses and optimisations directly at this shared IR level. The advantage of using a shared intermediate representation is clear: optimisations and analyses at that level benefit *all* the higher-level languages that target the IR.

Unfortunately, optimisations on a low-level IR often suffer from a loss of higherlevel information potentially applicable to optimisation. For instance, succinct highlevel iterations may translate to verbose loop constructs implemented with simple conditionals, goto statements and additional supporting statements as shown in Figure 1.1. Hence, at the IR level, it becomes harder, a priori, to identify usage patterns, and therefore useful information for potential optimisations, that may have been obvious in the original language.

```
i = 0
j = array.length
foreach value in array:
print value
i = i + 1
if (i < j) goto label
```

Figure 1.1: A high-level loop construct, when translated to a lower level IR.

A potential approach to the problem of lost information is to construct higherlevel IRs, in effect re-computing some of the information, based on an analysis of the original shared IR. Often, a specific IR has specific properties that make it suitable for a certain class of optimisations.

Static Single Assignment (SSA) form [CFR⁺91], in particular, denotes an intermediate representation with certain guaranteed properties that make the IR suitable for many analyses and optimisations. This thesis is an investigation of SSA form, its properties and use, as well as an application of fundamental ideas to yield useful new intermediate representations suitable for other classes of analyses. As part of this thesis we have also implemented the Shimple SSA framework to aid in our investigations.

1.1 Context and Motivation

Java [GJS05] is a general-use high-level programming language that is most often compiled to Java bytecode, a well-defined, low-level and portable executable format. Java bytecode is executed by a Java Virtual Machine [LY99]. Java is not the only language that can be compiled to Java bytecode. Python, Scheme, Prolog, Smalltalk, ML and Eiffel are just a few the many languages that can and have been compiled to this shared bytecode format [Tol06].

Hence, Java bytecode is a natural starting point for devising analyses and optimisations that can benefit the many higher-level languages that target the Java Virtual Machine. Often, in fact, such as in the case of third-party optimisation tools, *only* the bytecode may be available for analysis and optimisation.

Soot is a Java bytecode analysis and optimisation framework, devised by the Sable Research Group, which provides several intermediate representations of Java bytecode that are suitable for different levels of optimisations [VR00, VRHS⁺99].

In particular, Soot provides the Jimple IR, a typed and compact 3-address code representation of the bytecode that is suitable for general optimisation. To illustrate the idea behind Static Single Assignment form, it will be instructive to take a brief look at what Jimple code looks like at various stages.

Consider Figure 1.2, depicting Java code and the corresponding 'naive' Jimple representation. For the sake of clarity, the Jimple output has been simplified but the salient features have been preserved.

From the Java code, it is obvious to the reader that the first print statement, if reached, will output 0 and the second print statement, if reached, will output 5. The definition and use of the variable i in different *contexts* causes the print(i) statement to yield different results depending on that particular context. It is also clear that the two chunks of Java code are not particularly dependent on each other, although they do happen to share and use common variables.

The naive Jimple code seems a little more confusing and requires closer examination. The main reason for the added complexity is the lower-level nature of the subroutine implementation. Here we are confronted with explicit subroutine labels as well as if/goto statements. Other than this, the Jimple code represents a fairly direct mapping to the Java code.

Since we already know that the first print statement outputs 0 and the second print statement outputs 5, it is reasonable to expect a compiler to optimise the code by eliminating the unnecessary variable i and propagating the appropriate constants.

```
i = 0;
                                      i = 0;
                                      if bool == 0 goto label0;
if(bool) return;
print(i);
                                      return;
i = 5;
if (bool) return;
                                    label0:
print(i);
                                      print(i);
                                      i = 5;
                                      if bool = 0 goto label1;
                                      return;
                                    label1:
                                      print(i);
                                      return;
```

Figure 1.2: Simple code fragment in Java and naive Jimple form.

However, a compiler must first analyse the definitions and uses of the variable i and then attempt to determine, based on the particular context, whether the use of i is really the use of a known constant. This task is complicated by the fact that i is defined multiple times in the program, and due to the control structure, it may not be immediately clear which definition of i reaches a particular use of that variable.

A key observation here is that the different definitions and corresponding uses of the variable i in the original Java code are really independent. The variable i is simply being reused in a different context.

Fortunately, Soot performs what is known as *variable splitting* [Muc97] on naive Jimple before producing the final Jimple IR. Variable splitting was originally introduced in Soot to enable the task of the type assigner analysis [GHM00], but as shown in Figure 1.3, variable splitting also tends to result in code that is easier to analyse since overlapping definitions and uses of variables can be disambiguated in the process.

What has happened in Figure 1.3 is that the variable i has been split into variables i0 and i1 which represent the independent, non-overlapping, definitions and uses of i. Since i0 and i1 are only defined once, it is now easy for an analysis to determine whether the uses of these variables are in fact uses of a constant – it is no longer necessary to analyse the particular context of a use.

Static Single Assignment form [CFR⁺91] guarantees that every variable is only ever defined once in the static view of a program. The claim is that the resulting properties of a program in SSA form make it easier to analyse e.g. by removing the need for explicit flow-sensitive analysis.

The final Jimple code in the example is already in SSA form thanks to Soot's variable splitter. Unfortunately, however, the variable splitter is not always sufficient to guarantee the SSA property.

Consider the code in Figure 1.4. The variable *i* is defined twice, and hence the code is not in SSA form, since SSA form guarantees a single definition for every variable in the static text of the program. Nor can a simple variable splitter perform a renaming of *i* since it is not clear which definition of *i* the **print** statement is using. Indeed, the **print** statement could use either definition, depending on the runtime

```
i0 = 0;
i0 = 0;
                                      if b0 = 0 goto label0;
if (b0) return;
print(i0);
                                      return;
i1 = 5;
if (b0) return;
                                    label0:
                                      print(i0);
print(i1);
                                      i1 = 5;
                                      if b0 = 0 goto label1;
                                      return;
                                    label1:
                                      print(i1);
                                      return;
```

Figure 1.3: Variable splitting applied to the example from Figure 1.2. Variable i has now been split into variables i0 and i1, which can be analysed independently.

value of bool.

if(bool)	if bool == 0 goto label0;
i = 1;	
else	i = 1;
i = 2;	goto label1;
<pre>print(i);</pre>	label0:
	i = 2;
	label1:

Figure 1.4: Code fragment in Java and Jimple form where *i* is defined twice, hence the example is *not* in SSA form, and it is not entirely obvious whether variable *i* can be split.

print(i);

The crux of the matter is that the Jimple IR is not sufficient to always represent SSA form. The initial motivation for this thesis was hence to implement *Shimple*, an SSA-version of the Jimple IR. Shimple is first produced from Jimple, analysed and optimised, and then transformed back to Jimple, eventually becoming Java bytecode, as shown in Figure 1.5.

SSA form also raises many questions. For instance, if we consider simple variable splitting as illustrated in the previous example, it is obvious that many new variables will be introduced to the IR. One might wonder at the impact and implications of this overhead. The Shimple framework was in part designed to answer such questions by enabling experimentation and hence investigation of the pros and cons of SSA form.

This thesis also goes beyond the basic SSA form. We explore several improvements over SSA form, evolving from the variable splitting in Jimple to Simple Shimple, Extended Shimple, and Array Shimple. Given the many variations of and extensions of SSA form, we have also designed Shimple to be extensible and reusable, facilitating future SSA research in Soot.

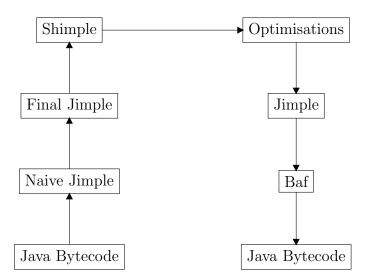


Figure 1.5: An illustration of the phases of Soot from Java bytecode through Shimple and back to optimised Java bytecode.

1.2 Contributions

The contributions of this thesis include the design and implementation of the Shimple framework in Soot, the SSA analyses and optimisations implemented on Shimple, as well as the insights gained in the process.

1.2.1 Design and Implementation

Shimple provides 3 different types of IRs, constructed in a bottom-up fashion, with each incorporating increasing amounts of analysis information.

- *Simple Shimple* was designed to investigate some of the more basic aspects of SSA. Runtime options as well as finer-grained control at the API-level are provided to allow the observation and modification of the behavior of SSA transformations.
- *Extended Shimple* is based on Simple Shimple and is an interesting example of how additional variable splitting over the basic SSA form can benefit certain analyses.

• *Array Shimple* introduces additional variable splitting for arrays, enabling the implementation of array element analysis in SSA form.

Array Shimple incorporates select may-alias information into the IR to facilitate the task of analyses. The amount and precision of the information that is available to Shimple can be controlled by the user e.g. by enabling or disabling interprocedural points-to analysis. Analyses on Array Shimple automatically benefit from any improved precision gains.

1.2.2 Shimple Analyses

Analyses we have implemented on Shimple include:

- A simple intraprocedural points-to analysis.
- Powerful conditional constant propagation and folder optimisations.
- Global value numbering and definitely-same information.
- Value range analysis.

Furthermore, existing Jimple analyses, such as Soot's Spark interprocedural pointsto analysis [Lho02], are shown to improve in precision simply by being applied to Shimple.

1.3 Thesis Organisation

The rest of this thesis is organised as follows. Chapter 2 provides a detailed background of basic SSA form including details of the algorithm involved in its construction and deconstruction. Chapter 3 introduces Shimple, details some of the challenges involved in its implementation, and describes a selection of analyses we have implemented on Simple Shimple. Chapter 4 provides an overview of the design and use of Extended Shimple. Chapter 5 provides a detailed description of Array Shimple. Each of the above chapters also includes a section on related work where appropriate. Finally, Chapter 6 concludes the thesis and summarises the main insights gained through this work.

Chapter 2 SSA Background

This chapter presents some of the background on Static Single Assignment form in its more basic incarnation [CFR⁺91]. In Sections 2.1 and 2.2 we pick up from where we left off in Chapter 1 and give a more detailed overview of basic SSA form. Sections 2.3 and 2.4 respectively detail the construction and deconstruction of SSA form. Finally, Section 2.5 gives a brief overview of some of the related work on SSA form.

2.1 Overview

In the grand scheme of things, SSA is often computed from a non-SSA IR, analysed, optimised and transformed, and then converted back to the syntax of the original IR [CFR⁺91]. However, due to the difficulties of defining transformations on more complex SSA-variants, another approach sometimes used is to generate the SSA IR from the non-SSA IR, analyse it, and then use the information gained to apply optimisations and transformations directly on the *original* IR [KS98, FKS00]. Both approaches are illustrated in Figure 2.1.

It is interesting to note that it may also be useful to first transform a program to SSA form, then directly out of SSA form, and subsequently analyse and optimise the resulting output. This is sometimes useful because the direct transformation to SSA

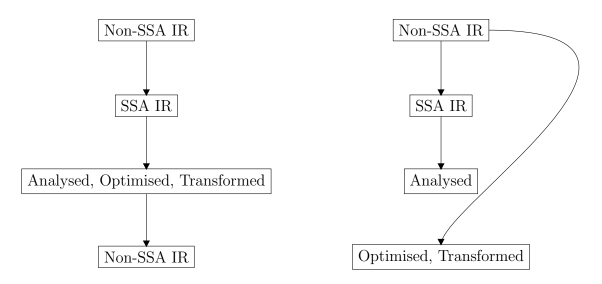


Figure 2.1: Two possible approaches towards using SSA form in a compiler.

and back, without any further optimisation, results in variables being split. However, any advantage dependent on SSA structures in particular may be lost.

2.2 Definition

In *Static Single Assignment* form, every variable is guaranteed to have a *single definition* point in the *static view* of the program text [CFR⁺91].

2.2.1 Example 1

The example in Figure 2.2 illustrates the *single definition* aspect of SSA form. The code is first shown in non-SSA form; we can see this is the case since **i** is defined twice.

We can transform the code to SSA form by splitting i into the variables i1 and i2 at the definition points. It is clear that the first two print statements should also be updated to refer the appropriate new variable. However, it is not so clear what needs to be done for the third print statement, since it may involve a use of either i1 or i2.

As shown in the figure, SSA form solves the problem by introducing a construct

if(bool)	if(bool)
i = 1	i1 = 1
print(i)	print(i1)
else	else
i = 2	i2 = 2
<pre>print(i)</pre>	print(i2)
print(i)	$i3 = \phi(i1, i2)$
	print(i3)

Figure 2.2: Simple example in non-SSA and SSA forms.

known as a ϕ -function. ϕ -functions are sometimes referred to as merge operators [BP03] and can be simplistically viewed as a mechanism for remerging a split-variable. With i1 and i2 remerged as variable i3, the third print statement can now be updated to use the new variable.

Intuitively, we can see from this example that by splitting i into i1 and i2 we gain the context-sensitivity benefits mentioned in Chapter 1 e.g. we can easily tell that the first two print statements use constants simply by looking at the definition for the variable being used.

If we view the ϕ -function as a simple merge, we do not notice any particular gains in precision for the third print statement. We will later see that it is perhaps more meaningful to refer to a ϕ -function as a *choice* operation rather than a merge. We will also note situations (e.g. Section 3.3.2) where we can gain more precision by analysing ϕ -functions as performing a choice rather than simply merging all its arguments.

Before we move on, we should note that a ϕ -function generally has as many arguments as it has control-flow predecessors. Each argument corresponds to the name of the relevant variable along a particular control-flow path. In this example, the ϕ -function can be reached from the **if** block or the **else** block, hence it has two arguments corresponding to each control-flow path.

2.2.2 Example 2

This next example illustrates the emphasis on *static* in Static Single Assignment form.

SSA form is sometimes described as exhibiting "same name, same value" behaviour i.e. two different references of a variable might be assumed to be uses of the same value [LH96]. In a sense this is true, however, the "same name, same value" statement requires clarification.

Consider the code in Figure 2.3. Generally, the first step in converting a program to SSA form is to split multiply-defined variables, in this case i, at the definition points. The next step is to remerge the split-variables where necessary i.e. adding ϕ -functions where appropriate.

i = 0

while(i != 10)
print(i)
i = random()

print(i)

Figure 2.3: High-level loop with a variable assignment.

As a precondition, however, we generally assume that we are working with a control flow graph or at least lower-level control-flow primitives. In this example, the while condition can refer to either of the two definitions of i; hence, in SSA form, we will need an appropriate position to place a merge function – this cannot be conveniently done with the high-level while loop syntax.

Figure 2.4 shows the same example with lower-level Jimple control-flow constructs instead of a while loop, both in non-SSA and SSA forms.

As shown, i has been split into i0 and i1 at the definition points. A ϕ -function has also been strategically inserted to remerge i0 and i1 into i2. Notice that the lower-level control-flow primitives allow the loop entry point to be separated from the

```
i0 = 0;
  i = 0;
loop:
                                   loop:
  if (i = 10) goto exit;
                                     i2 = \phi(i0, i1);
                                     if (i2 = 10) goto exit;
  print(i);
  i = random();
                                     print(i2);
                                     i1 = random();
  goto loop;
                                     goto loop;
exit:
                                   exit:
  print(i);
                                     print(i2);
```

Figure 2.4: Example from Figure 2.3 shown with lower-level loop constructs in both non-SSA and SSA forms.

main loop conditional, such that a merge operation can now be placed between the two.

The interesting thing to note is that even though i1 and i2 are defined once in the static view of the program, they are inside a loop and hence may be assigned to many times during the execution. It is also interesting to note that the two print statements, although identical, *never* print the same value. We will take advantage of this observation later, in Chapter 4.

The "same name, same value" property is violated in this example because although i2 is defined once in the static view of the program, during the dynamic execution, i2 is redefined many times. In particular, the two uses of i2 in the print statements reference distinctly different dynamic assignments.

The "same name, same value" property can however be guaranteed to hold if the uses of a variable occur within the scope of the same *dynamic* assignment. In particular, if the variable uses can be shown to be present within the same (possibly nested) loop iterations and same context invocation, then we can assume that they reference the same value.

2.2.3 ϕ -functions

Definition

Having illustrated the basic SSA definition, it would be instructive to further examine the meaning of ϕ -functions. Our description of ϕ -functions as merge operators is somewhat over-simplistic and insufficient when one needs to perform operations on them.

A ϕ -function is generally placed at the beginning of a join node in the control-flow graph and has an argument for each of its corresponding control-flow predecessors. Each argument corresponds to the appropriate name of the split variable defined on the path from the given control-flow predecessor, as indicated in Figure 2.5.

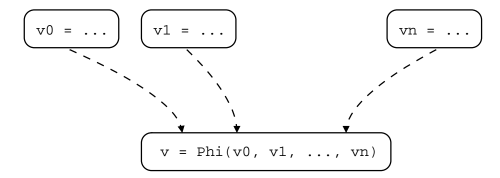


Figure 2.5: A ϕ -function over an *n* split-variable.

Traditionally, $\phi(v0, v1, ..., vn)$ is defined as a function that evaluates to the value of the argument corresponding to the flow of control at *runtime* [CFR⁺91]. That is, $\phi(v0, v1, ..., vn)$ really denotes a *choice* or selection of a single argument rather than a merging of all the arguments.

Powerful static optimisations that take control-flow into account can be devised by analysing which choice will be made at runtime. In the cases where it is not possible to determine that a single choice is guaranteed, conservative approximations can be made. In particular, in the worst case $\phi(v0, v1, ..., vn)$ can be estimated as resulting in the merge of all the arguments. Even in the worst case, by analysing the arguments themselves, useful information can potentially be gained.

 ϕ -functions hence provide a powerful means of analysing the results of control flow. Examples of this will later be demonstrated in Section 3.3.

Caveats

The traditional definition of ϕ -functions as given often elicits mystified reactions. There is a good reason for this confusion, since the notation for ϕ -functions as presented is fundamentally incomplete.

Consider that function $\phi(v1, v2)$ may evaluate to v1 at runtime, and yet the very same function may evaluate to v2. ϕ -functions certainly do not seem to behave as pure functions which are expected to evaluate to the same value when given the same input.

The reason for this difficulty is the assertion that each argument in a ϕ -function is associated with a particular control-flow predecessor at runtime. This information is not explicitly present in the function notation itself, but could be expressed with one or more additional arguments to the ϕ -function.

It is fair to say that without this additional information, additional ancillary computation would be required in order for a runtime to execute ϕ -functions.

There have been efforts to express the full information in the IR to permit runtime execution and analysis [OBM90, KS98]. However, the resulting IR often tends to be significantly more verbose since the additional arguments now have to be explicitly computed and updated.

For the purposes of static analysis and for algorithmic exposition, it is often more practical to use the abbreviated ϕ -functions with the understanding that additional runtime information may be required to evaluate a ϕ -function. In Section 3.2 we will see the approach that Shimple takes when handling ϕ -functions.

2.3 Construction

SSA form was known at least as far back as 1969 [SS70] and many SSA-based variants and analyses have since been formulated. It was only in 1989 however, when an efficient and practical algorithm for computing SSA was first formulated by Cytron *et al.* [CFR⁺91], that SSA form really took off. Since that time, there have been valiant attempts to improve the efficiency of basic SSA computation [BP03] but the Cytron *et al.* [CFR⁺91] algorithm has remained one of the most useful and efficient in practice.

SSA form is typically constructed in two stages:

- 1. Insertion of trivial ϕ -functions at appropriate places in the control-flow graph (see Figure 2.6).
- 2. Renaming of variable at definition points as well as corresponding uses such that the program is in SSA form while preserving the original semantics.

The key step is the first one and is where most of the variation in the different algorithms tends to occur. For the purposes of this thesis we will focus on the Cytron *et al.* [CFR⁺91] algorithm. However, Shimple was designed such that alternative algorithms [BP03] could be accommodated for experimental purposes in Soot.

2.3.1 Step 1: Insertion of ϕ -functions

Overview

The first question is, where should ϕ -functions be inserted? In Shimple we focus on minimal SSA form [CFR⁺91]. The two conditions for minimal SSA form to hold are:

1. If a join node in a CFG has several reaching definitions (2 or more) of a variable v, a trivial *n*-argument ϕ -function of the form $v = \phi(v, v, ...)$ is inserted at the beginning of the join node, where *n* is the number of control-flow predecessors of the join node in the CFG (e.g. Figure 2.6).

2. The number of ϕ -functions inserted is as small as possible, subject to the previous condition.

if (boolean)	if (boolean)
v = 1	v = 1
else	else
v = 2	v = 2
print(v)	$\mathrm{v}~=~\phi(\mathrm{v},~\mathrm{v})$
	print(v)

Figure 2.6: A trivial ϕ -function is added in the first step of computing SSA form.

Note that the insertion of a new ϕ -function definition statement may induce the need for other ϕ -functions to be inserted, since the new definition may also reach a join node reachable by other definitions. It is also worth noting that condition 1 implies that the number of ϕ -functions inserted is not necessarily the *minimum* required to support SSA form. For example, consider Figure 2.7. Since **i3** is not used, the ϕ -function is unnecessary. However, minimal SSA form demands that it be inserted.

if(bool)	if(bool)
i = 1	i1 = 1
else	else
i = 2	i2 = 2
i = 3	$i3 = \phi(i1, i2)$
	i4 = 3
<pre>print(i)</pre>	<pre>print(i4)</pre>

Figure 2.7: Example of a dead ϕ -function in minimal SSA form.

In pruned SSA form and some other variants of SSA, the dead ϕ -function might be omitted or eliminated in a subsequent step. It is interesting to note that even these dead assignments may occasionally become useful for analysis purposes [CFR+91] (e.g. detecting program equivalence) and, even if they are not used, they will eventually be eliminated when translating out of SSA form.

Condition 1 assumes that all variables have an initial definition in the start node and hence are guaranteed to be defined along all paths to a join node. This is not necessarily the case in Java or Jimple, and therefore we strengthen the condition such that we insert a ϕ -function for a variable V only if V is guaranteed to be defined along all paths to the join node. This is a safe modification since Java guarantees that a variable which is not defined on one or more control flow paths may not be used subsequently [LY99].

Next we summarise the key concept of dominance frontiers per Cytron *et al.* [CFR⁺91] which will subsequently enable us to efficiently locate exactly those positions where ϕ -functions need to be inserted.

Dominance

Definition 1 A node X in a CFG is said to dominate node Y if every path from the start node to Y must pass through X $[CFR^+91]$.

Definition 2 X is a strict dominator of Y if X dominates Y and X is not equal to $Y [CFR^+91]$.

Definition 3 Cytron et al define the dominance frontier of a node X as being the set of all nodes Y such that X dominates a predecessor of Y but does not strictly dominate $Y [CFR^+91]^{.1}$

Cytron *et al.* prove that if a variable V has a definition V_x in node X, then ϕ -functions are required in every node Y of the dominance frontier of X [CFR⁺91].

Intuitively, we can see that this is the case, since by the definition of the dominance frontier, V_x reaches a predecessor of Y (of its dominance frontier) and hence reaches

¹In particular, from this definition, it is possible for X to be in its own dominance frontier.

Y itself. However, since V_x does not strictly dominate Y, other definitions of V reach the join node and hence the insertion of a ϕ -function is required.

Given the dominance frontier, it is simple to express the algorithm for inserting trivial ϕ -functions, as shown in Figure 2.8 [CFR⁺91].

```
for each variable V do
for each node X that defines V:
add X to worklist W
for each node X in worklist W:
for each node Y in dominance frontier of X:
if node Y does not already have a \phi-function for V:
prepend ''V = \phi(V, ..., V)'' to Y
if Y has never been added to worklist W:
add Y to worklist W
```

Figure 2.8: Algorithm for inserting ϕ -functions [CFR⁺91].

Although the above algorithm represents $O(n^2)$ time complexity for each variable V with respect to the size of the CFG, in practice the performance is found to be quite acceptable and competitive [BP03]. Cytron *et al.* focus on optimising the algorithm for computing the dominance frontiers.

The dominance relation itself can be computed with a simple flow analysis [ASU86] as shown in Figure 2.9. If the forward flow analysis proceeds in topological order, the dominator sets will shrink in each iteration for a maximum of n times. Depending on the efficiency s of the set operations, the time cost would be $O(n^2 * s)$. More efficient, linear-time, algorithms are known for computing dominance [AL96].²

Given the dominance sets and the definition of dominance frontier, we can compute the latter straightforwardly though inefficiently by a brute-force search of the CFG

²A faster dominance algorithm [CHK01] was implemented for Shimple by Michael Batchelder, although no significant overall speedup was experienced.

```
initialise dominance sets:

start node dominates itself

every other node is assumed to be dominated by all nodes

until no more changes in dominance sets:

for each node N in CFG:

DomSet(N) = \{N\} \cup

{intersection of DomSet(P) for all predecessors P of N}
```

Figure 2.9: Simple flow analysis to compute dominance sets [ASU86].

and dominance sets. We will instead outline the more efficient approach by Cytron $et \ al. \ [CFR+91].$

Definition 4 X is an immediate dominator of Y if X strictly dominates Y and X is the closest dominator of Y in the CFG $[CFR^+91]$.

Definition 5 The dominator tree of a CFG is defined [CFR⁺91] as follows:

- 1. The root of the tree is the start node.
- 2. The children of a node X in the dominator tree are all the nodes immediately dominated by X in the CFG.

The dominator tree can be computed from the dominance sets or incrementally from scratch using a more efficient algorithm [BP03]. We are interested in the dominator tree because of the observation that if we compute the dominance frontiers for each node in a bottom-up fashion on the dominator tree, we can compute the dominance frontier mapping in time linear to the size of the sets in the mapping, by reusing the information computed for previous nodes.

To see this, we present the proof per Cytron *et al.* [CFR⁺91] that the dominance frontier of a node X, or DF(X), can be computed in terms of the sets DF_{local} and DF_{up} :

$$DF(X) = DF_{local}(X) \cup \bigcup_{\forall Z \in children(X)} DF_{up}(Z)$$
 (2.1)

where Z are the children of X in the dominator tree.

The sets DF_{local} and DF_{up} are defined as follows:

Definition 6 The set $DF_{local}(X)$ is defined as containing all Y such that Y is a successor of X in the CFG and X does not strictly dominate Y [CFR⁺91].

Definition 7 The set $DF_{up}(Z)$, where is Z is a child of X in the dominator tree, is defined as containing all Y such that Y is in the dominance frontier of Z and X does not strictly dominate Y [CFR⁺91].

From the definitions, DF(X) is computed by observing all the CFG successors as well as the nodes immediately dominated by X – hence the requirement that we traverse the dominator tree in bottom-up order for efficiency. We need to prove that equation 2.1 is indeed correct.

Proof:

 \leftarrow First we prove that elements in $DF_{local}(X)$ and $DF_{up}(Z)$ are indeed elements of DF(X) [CFR⁺91].

- 1. Since X self-dominates and hence dominates a predecessor of Y while not strictly dominating Y itself, in accordance with the definitions of $DF_{local}(X)$ and DF(X), all elements Y of $DF_{local}(X)$ are indeed in the dominance frontier of X.
- 2. Since Z is a child of X in the dominator tree, in accordance with the definition of DF_{up}(Z), all nodes dominated by Z are also dominated by X. Hence, any node Y that is in the dominance frontier of Z has a predecessor that is dominated by X. If Z is not strictly dominated by X, then Z is in the dominance frontier of X according to the definition of DF(X).

 \Rightarrow Next we prove that if an element is in DF(X), it is in either $DF_{local}(X)$ or $DF_{up}(Z)$ [CFR⁺91].

Consider any Y that is in DF(X). Y must have a predecessor P that is dominated by X.

- 1. If P is X, then Y is a successor of X and is hence in $DF_{local}(X)$.
- 2. If P is not X, then X must have a child Z in the dominator tree that dominates P but does not strictly dominate Y since Y is in the dominance frontier of X. Hence, Y is in $DF_{up}(Z)$.

The next steps are to reformulate the definitions of $DF_{local}(X)$ and $DF_{up}(Z)$ so that they are easier to compute.

Lemma 1 The set $DF_{local}(X)$ can be defined as containing all Y such that Y is a successor of X in the CFG and X is not an immediate dominator of Y [CFR⁺91].

Proof:

Assuming that Y is successor of X in the CFG, we have to prove that the statement X is not an immediate dominator of Y is equivalent to the statement in the original definition that X does not strictly dominate Y [CFR⁺91].

It suffices to prove that the statement X is an immediate dominator of Y is equivalent to the statement that X strictly dominates Y.

 \Rightarrow By definition, if X is an immediate dominator of Y, then X strictly dominates Y.

 \Leftarrow If X strictly dominates Y, then X is on every path from the start node to Y. Furthermore, since X is a predecessor of Y in the CFG, X is already the closest strict dominator of Y and hence its immediate dominator. \Box

Lemma 2 The set $DF_{up}(Z)$, where Z is a child of X in the dominator tree, can be defined as containing all Y such that Y is in the dominance frontier of Z and X is not an immediate dominator of Y [CFR⁺91].

Proof: Assuming that Y is in the dominance frontier of Z, we have to prove that the statement X is not an immediate dominator of Y is equivalent to the statement in the original definition that X does not strictly dominate Y [CFR+91].

It suffices to prove that the statement X is an immediate dominator of Y is equivalent to the statement that X strictly dominates Y.

 \Rightarrow By definition, if X is an immediate dominator of Y, then X strictly dominates Y.

 \Leftarrow If X strictly dominates Y, then X has a child C in the dominator tree that dominates Y since strict domination is a stronger relation than simple domination.

Since Y is in the dominance frontier of Z, let P be a predecessor of Y that is dominated by Z.

Since C dominates Y, C must appear on any path from the start node to P to Y via the $P \to Y$ edge. Hence C either dominates P or C is Y.

If C is Y, then Y is immediately dominated by X since C is a child of X in the dominator tree and we are done.

If C is not Y then C dominates P. Since Z also dominates P, and both C and Z are children of X in the dominator tree, since there can be only one child of X in the dominator tree that dominates P, C must be equal to Z. Hence Z dominates Y which contradicts our assumption that P is in the dominance frontier of Z [CFR⁺91]. \Box

As shown in Figure 2.10, we can now formulate the algorithm [CFR⁺91] for computing the dominance frontiers of every node in the CFG.

As we have shown at the beginning of this section, once we have computed the dominance frontiers for each node, we may proceed to insert our trivial ϕ -function assignments.

2.3.2 Step 2: Variable Renaming

Two structures are maintained during the renaming process [CFR⁺91].

• C[V] is an array that holds an integer for each variable. C[V] is initialised to 0 and incremented to generate unique subscripts for the new variables split from for each node X in a bottom-up traversal of the dominator tree: $DF(X) = \{\}$

compute $DF_{local}(X)$: for each node Y that is a successor of X in the CFG: if the immediate dominator of Y is NOT X: $DF(X) = DF(X) \cup \{Y\}$

compute $DF_{up}(Z)$: for each node Z that is a child of X in the dominator tree: for each node Y that is in the dominance frontier of Z: if the immediate dominator of Y is NOT X: $DF(X) = DF(X) \cup \{Y\}$

Figure 2.10: Algorithm for efficiently computing dominance frontiers [CFR⁺91].

V.

• S[V] is an array of integer stacks that are used to keep track of the scope of the new variable definitions so that the uses can be renamed appropriately.

The variables are initialised at the beginning of the renaming process as shown in Figure 2.11. The renaming process [CFR⁺91] proceeds in a top-down fashion beginning from the entry-node and processing each node by following a depth-first search path in the dominator tree. We shall consider the **rename(node)** function in four sequential steps, each consisting of a **for** loop.

Step 1 of rename(node) traverses each statement S in the node as shown in Figure 2.12. Since the dominator tree is traversed in a top down fashion and the statements in the node are processed sequentially, the if statement never sees a use of a variable before its definition is processed in the for statement.

In effect the for loop generates a unique name V_i for each definition of V and remembers the newest subscript *i* by pushing it onto S[V], since if this node dominates for each variable V: C[V] = 0S[V] = []

Figure 2.11: Initialisation phase for the renaming process [CFR⁺91].

```
for each statement S in node:

if S is not a \phi-assignment:

for each use (not def) of variable V in S:

i = S[V].top()

replace use of V by use of V_i

for each definition of V in S:

i = C[V]

replace V by new variable V_i

S[V].push(i)

C[V] = i + 1
```

Figure 2.12: Step 1 of the renaming process [CFR⁺91].

another node with a use of V we are interested in this latest definition of V_i , provided it is not killed by another definition. The **if** block subsequently ensures that uses of V that are dominated by the new V_i definition are renamed appropriately.

Uses of V in a ϕ -function are handled next in step 2 of rename(node) as shown in Figure 2.13. We can see that the above code takes care of uses of V that are in the local dominance frontier (DF_{local}) of node and that need to be renamed to V_i . Furthermore, given that we are traversing the dominator tree in top down fashion and storing the most recent dominant definition of V in S[V], we are also taking care of uses of V that are in the relevant DF_{up} sets. Hence we have renamed all uses of the new definition V_i since we have renamed uses of V that are dominated by and are in the dominance frontier of the new definition.

for each successor succ of node in the CFG: for each ϕ -function on variable V in succ: $j = \text{index of argument in } \phi$ -function that corresponds to the predecessor node of succ $i = S[V] \cdot \text{top}()$ replace jth argument V in ϕ -function with the use V_i

Figure 2.13: Step 2 of the renaming process [CFR+91].

The next code segment makes the traversal of the dominator tree explicit, as shown in Figure 2.14. The code ensures that nodes in the dominator tree are traversed in a top-down fashion.

```
for each child of node in the dominator tree:
    rename(child)
```

Figure 2.14: Step 3 of the renaming process [CFR⁺91].

Finally in Figure 2.15, we see that as the traversal has reached the end of the dominator tree and needs to backtrack to other branches (in typical depth-first search

fashion), the no longer relevant re-definition of V (all its uses have been renamed appropriately) is popped from the name stack.

```
for each definition statement S in node:
for each original definition of V in S:
S[V].pop()
```

Figure 2.15: Step 4 of the renaming process $[CFR^+91]$.

The full code for the renaming algorithm $[CFR^+91]$ is shown in its entirety in Figure 2.16. The running-time of the above code is dependent on the number of nodes N in the dominator tree (and hence the number of nodes in the CFG), the number of edges E in the CFG (since the successors of each node must be analysed) and T the total number of variable uses and definitions that need to be processed.

2.3.3 Summary

The ϕ -function insertion algorithm coupled with the renaming algorithm results in code that is in minimal SSA form. We have given an outline of the algorithms as well as an intuition into their workings. A full proof of the validity of the approach as well as a detailed analysis of the running-time can be found in the work of Cytron *et al.* [CFR⁺91].

2.4 Deconstruction

To translate out of SSA form, it suffices to remove the ϕ -functions and replace them with equivalent statements. A statement of the form $v = \phi(v_1, v_2, ..., v_n)$ can be replaced by n assignment statements of the form $v = v_x$. Since $\phi(v_1, v_2, ..., v_n)$ evaluates to v_x based on the control-flow at run-time, the n assignments can each be placed on the appropriate CFG edge leading into the join node as shown in Figure 2.17.

To further illustrate this, consider Figure 2.18 where the ϕ -assignment is replaced by two ordinary assignments. There are a couple of points worth noting here.

```
initialisation()
rename (Entry)
rename(node):
   for each statement S in node:
      if S is not a \phi-assignment:
          for each use (not def) of variable V in S:
             i = S[V] \cdot top()
             replace use of V by use of V_i
      for each definition of V in S:
         i = C[V]
          replace V by new variable V_i
          S[V]. push (i)
         C[V] = i + 1
   for each successor succ of node in the CFG:
      for each \phi-function on variable V in succ:
         j = index of argument in \phi-function that corresponds to
                 the predecessor node of succ
         i = S[V] \cdot top()
          replace jth argument V in \phi-function with the use V_i
   for each child of node in the dominator tree:
      rename(child)
   for each definition statement S in node:
      for each original definition of V in S:
          S[V].pop()
      Figure 2.16: Algorithm for the variable renaming process [CFR<sup>+</sup>91].
```

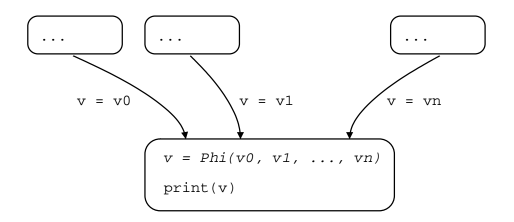


Figure 2.17: ϕ -function shown with equivalent copy statements.

```
if (bool)
                                    if (bool)
   i1 = 1
                                       i1 = 1
                                       print(i1)
   print(i1)
                                       i3 = i1
else
   i2 = 2
                                    else
   print(i2)
                                       i2 = 2
                                       print(i2)
i3 = \phi(i1, i2)
                                       i3 = i2
print(i3)
                                    print(i3)
```

Figure 2.18: Example of naive ϕ -function elimination.

First, since the code is represented in a flat text format, the ordinary assignment statements are placed at the end of the predecessor blocks instead of on the control flow edges. In this particular case, this is not an issue given that there is no effective difference between whether the statements are placed at the end of the predecessor blocks or on the actual control flow edges.

The second more important point is that the resulting code after ϕ -function elimination does *not* look like the original code before ϕ -function insertion. This can be seen in Figure 2.19.

if(bool)	if(bool)
i = 1	i1 = 1
print(i)	print(i1)
else	i3 = i1
i = 2	else
1 = 2	erse
print(i)	i2 = 2
	i2 = 2

print(i3)

Figure 2.19: Comparison of code before ϕ -function insertion and after ϕ -function elimination.

The original variable i remains split into the 3 variables i1, i2, i3. Due to this fact, code that has been converted to SSA form and straight back out results in code that may still expose useful control-flow sensitive information. For instance, we can see that i1 and i2 are both constants in Figure 2.18 and hence their uses can be replaced by constants.

The code in Figure 2.18 could be considered wasteful and inefficient. It is longer than the original code, uses significantly more variables and often may contain ineffective statements.

As detailed by Cytron *et al.* [CFR+91], we can obtain efficient code by apply-

ing simple analyses such as dead code elimination [Muc97] before replacing the ϕ -functions and, after ϕ -function elimination, applying a variable packing [Muc97] algorithm that optimises storage allocation e.g. by a graph colouring algorithm.

By applying dead code elimination before removing ϕ -functions, we can get rid of those ϕ -functions that have been added to satisfy the minimal SSA requirement but aren't otherwise used. For example, if there had been no further use of *i*3 in Figure 2.18, applying dead code elimination would have simply removed the ϕ -function. Subsequently applying variable packing will usually undo the variable splitting that results from SSA form.

2.5 Related Work

The origins of SSA form can be traced back to the work of Shapiro and Saint [SS70]. The form was later popularised through the work of Cytron *et al.* [CFR⁺91], for the first time introducing an algorithm for computing SSA form that was efficient enough in practice that it could be readily adopted by compiler writers. The algorithms and proofs in this chapter are largely based on this work of Cytron *et al.*

There have been several attempts to improve on the theoretical complexity of the algorithm introduced by Cytron *et al.*, including work by Johnson and Pilardi who proposed using a structure called the dependence flow graph [JP93], a generalisation of SSA form and def-use chains useful for sparse forwards and backwards data flow analysis of a program, in order to compute SSA form. However, many of the algorithms with better theoretical complexity have been found to fare worse than the Cytron *et al.* algorithm in practice [BP03].

Bilardi and Pingali [BP03] propose a framework for comparing the various SSA algorithms and their relative performances and provide an overview of the various algorithms for computing SSA form. Strategies for optimising the computation of SSA form have tended to focus on the optimisation of the computation of dominance frontier sets by balancing various techniques such as lazy or on-demand computation, precomputation and caching of the dominance frontier sets, often resulting in the development of new data structures to represent and compute the relevant information.

One such algorithm was described by Sreedhar and Gao [SG95]. The algorithm used linear preprocessing time in building a structure called the DJ-graph (a dominator tree augmented by what the authors called join edges) and could use the structure in order to efficiently compute the dominance frontier sets on demand. Although the algorithm had a better theoretical complexity than the Cytron *et al.* algorithm, it was later acknowledged to have worse performance in practice [BP03].

The Cytron *et al.* [CFR⁺91] approach can be seen at one end of the spectrum since it completely precomputes the dominance frontier sets, storing the structure in memory, and subsequently computing SSA form, whereas the Sreedhar and Gao algorithm, where dominance frontier sets are computed on demand, can be seen at the other end. Bilardi and Pingali devised an approach using a structure called the augmented dominator tree [PB95] that subsumed both the Cytron *et al.* and Sreedhar and Gao algorithms since it could be tuned to behave as either of those algorithms (i.e. full precomputation versus on-demand computation of the dominance frontier sets). More notably, the augmented dominator tree could be tuned such that dominator frontier sets could be computed with better theoretical complexity while actually outperforming the Cytron *et al.* algorithm in practice.

Chapter 3 Shimple

In this chapter we introduce Shimple, our implementation of an SSA framework for Soot. In Section 3.1 we overview the overall goals and design of Shimple. Section 3.2 further details some of the challenges we faced while implementing Shimple. Section 3.3 presents a selection of analyses that we have implemented on Simple Shimple and which will also help illustrate the use of SSA form. Finally, Section 3.4 gives a brief overview of the relevant related work.

3.1 Overview and Design

There has long been a demand for an SSA-based IR in Soot. Shimple was implemented partly to fulfill this demand as well as to facilitate research of SSA form, including variants and extensions of the latter. As such, Shimple has been designed to be flexible and extensible, while fitting naturally into the existing Soot framework.

3.1.1 Shimple from the Command Line

As with Jimple, the Soot command-line user can create Shimple output for inspection and program understanding from Java class or source files using a variety of configuration and optimisation options. SSA-based optimisations on Shimple form can also be applied directly to class files for program optimisation. Further details of using Shimple from the command-line can be found in the phase option documentation for Soot and the Shimple user guide [Dev06].

3.1.2 Shimple for Development

At the basic API level, Shimple follows the design of Jimple [VR00], and as such a Soot developer will quickly be at ease. The Soot framework user can create Shimple bodies from Jimple and can apply or implement a variety of SSA analyses and transformations. Since the Shimple design is based very closely on Jimple, existing analyses can be easily ported to Shimple, automatically gaining from the flow sensitive benefits of SSA form.

Shimple method bodies as well as Shimple body elements (such as ϕ -functions) can be created from the Shimple constructor class, with ShimpleBody providing a high-level interface that allows one to reconstruct SSA form or exit out of SSA form as desired.

Developers have convenient and efficient access to variable definition-use chains and use-definition chains respectively through the ShimpleLocalUses and Shimple-LocalDefs classes. The optimisations currently implemented on Shimple (Section 3.3) are also available for developer use.

The various components implemented for the purposes of constructing Shimple have been, where possible, added as general analyses to the Soot framework. Particularly, the dominator analysis classes have proven popular for independent use from Shimple e.g. for constructing control dependence graphs [CFR⁺91].

3.1.3 Improving and Extending Shimple

Although ShimpleBody provides an interface for constructing and interacting with a Shimple method body, it does not contain any logic pertaining to the construction of the body. A Shimple body is constructed and deconstructed through an implementation of the ShimpleBodyBuilder interface.

Shimple provides a default implementation of ShimpleBodyBuilder; Default-ShimpleBodyBuilder is the central location for invoking other classes involved in the computation of Shimple, such as the various NodeManagers which have specific functionality related to the insertion and removal of SSA-related nodes¹ and the ShimpleRenamer for SSA variable renaming.

Shimple implements the factory pattern [GHJV95] in order to centralise the location where key components of Shimple, such as the default ShimpleBodyBuilder implementation itself as well as the default implementations of NodeManager and the ShimpleRenamer can be instantiated from. Other useful analyses such as those used to compute dominance frontiers are also instantiated from the Shimple factory.²

Hence, developers interested in improving or extending Shimple can do so by supplying new algorithms that implement the required interfaces, or by simply selecting other pre-existing algorithms in Soot, and providing an updated factory in order to load the new classes. ShimpleFactory is the basic interface through which Shimple expects to obtain its components, with a default implementation of the factory being provided in DefaultShimpleFactory. Developers can either extend Default-ShimpleFactory or provide another implementation altogether of ShimpleFactory.

Since the code for computing Shimple is generally well-modularised (e.g. the default algorithms for inserting and eliminating ϕ -functions can be found in the Phi-NodeManager class), a developer can easily access existing code and algorithms while tweaking specific algorithms.

3.2 Implementation

The default implementation of Shimple closely follows the Cytron *et al.* algorithms described in Chapter 2. We have found the Cytron *et al.* approach to be reasonably efficient while being well understood and relatively easy to implement.

Shimple is constructed by computing the dominance frontiers for ϕ -function insertion, followed by the SSA renaming algorithm as described in the previous chapter. It is deconstructed by naively eliminating ϕ -functions preceded and followed by stan-

¹Such as ϕ -functions, or as we will later see π -functions, as well as array-related functions.

 $^{^2 {\}rm Indeed},$ Michael Batchelder has recently provided a new efficient drop-in replacement for the dominators algorithm used by Shimple.

dard Soot optimisations such as LocalPacker which implements a variable packing algorithm to optimise storage allocation, DeadAssignmentEliminator which prunes unused assignment statements, and various other optimisations, if desired.

The approach we generally take in Soot for Shimple-based analyses is to first create Shimple from Jimple, perform analysis and optimisations as desired, and to exit Shimple by transformation to Jimple. A Soot user is of course free to create Shimple and exit immediately to Jimple in order to take advantage of the additional variable splitting introduced without needing to handle ϕ -functions, as previously mentioned. From Jimple, Soot can subsequently output bytecode or Java source as desired.

In the remainder of this section we first present some relevant background on Jimple, followed by an overview of some of the challengers encountered when implementing Shimple in Soot.

3.2.1 Jimple Background

Soot's SSA implementation, Shimple, is based on the Jimple IR, a typed and compact 3-address code representation of Java bytecode. Internally, Jimple is represented as a chain of instructions, sequentially listed from the first to the last as shown in Figure 3.1.

Although the printed Jimple fragment includes labelled blocks with the branching statements referencing those labels, it can be seen in the figure that the internal chain representation does not in fact include this abstraction. Any instruction that may branch to another out of the normal sequential ordering simply contains a pointer to the target instruction (i.e. the first instruction in the target block) – all labels in the printed representation are computed at print time.

The chain representation of Jimple allows simple operations such as iterating sequentially though the instructions of a method body, and also allows modifications to the chain such as instruction insertions or removals.

Consider the instruction i3 = 1 in Figure 3.1. This instruction could be removed, for example, by an analysis that has determined that it is a dead assignment. Since

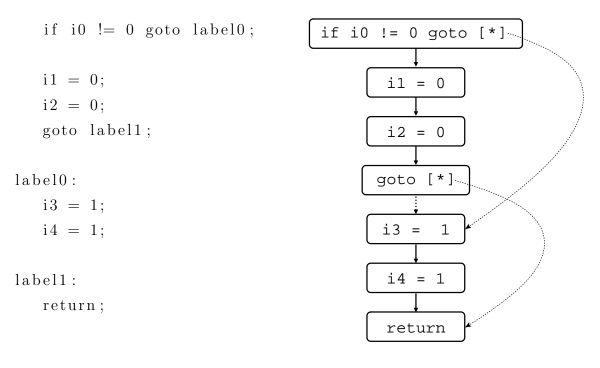


Figure 3.1: Printed Jimple fragment with its Soot internal chain representation.

the instruction is at the beginning of a block, removing it from the chain requires updating any pointers that used to point to that instruction. In particular, any pointers pointing to i3 = 1 must now point to i4 = 1.

Similarly, consider an instruction insertion before instruction i3 = 1. Since the intent is to insert the new instruction at the beginning of the block, any pointers formerly pointing to i3 = 1 must now be updated to point to the new instruction.

The implication of Jimple's internal chain representation is that any instruction insertion or removal may require internal pointer patching. Fortunately, Jimple provides a PatchingChain interface by default which attempts to patch all internal pointers using a few simple rules. In complex control-flow manipulations on the chain, user-intervention may be necessary.

Jimple's chain representation may not be so convenient when analyses may require control flow or block information. For this reason, Soot provides a variety of convenient graph structures that can be built from Jimple.

These structures are *secondary* in the sense that they are built, used, and ulti-

mately discarded. Changes to the structures may or may not result in updates to the Jimple chain, structural changes to the Jimple chain will not result in any secondary structures being updated, and changes to one secondary structure will also not affect other secondary structures.

Given this background, we will now take a look at the implementation of ϕ -functions in Shimple.

3.2.2 ϕ -functions

As we have previously noted, each argument to a ϕ -function is associated with a particular control-flow predecessor. Although this is often not made explicit in typical SSA notation, the importance of this association is crucial to the semantics of the representation.

Given that we first create a control-flow graph structure from Jimple before computing SSA form, each argument to a ϕ -function could implicitly be considered to correspond to the appropriate control-flow predecessor in the graph. Unfortunately, as we have noted in Section 3.2.1, our control-flow graph structure is only a secondary structure and will ultimately be discarded. Furthermore, there is no guarantee that other secondary structures subsequently built from the Shimple form will maintain the same implicit correspondence between ϕ -function arguments and control-flow predecessors.

It is hence necessary for us to explicitly encode the correspondence between ϕ -function arguments and control-flow predecessors in the internal Shimple representation.³ An example of Shimple with its printed and internal representation is shown in Figure 3.2.

As shown in the figure, each ϕ -function argument is associated with a pointer to the relevant control flow predecessor i.e. the last statement at the end of the relevant control flow block.

 $^{^{3}}$ We decided that it would be beneficial to base Shimple closely on the Jimple design rather than creating a radical new Soot representation. The main benefit of this approach is that existing Soot developers will feel instantly at ease with Shimple, as well as the fact that Jimple analyses can be more readily ported to Shimple.

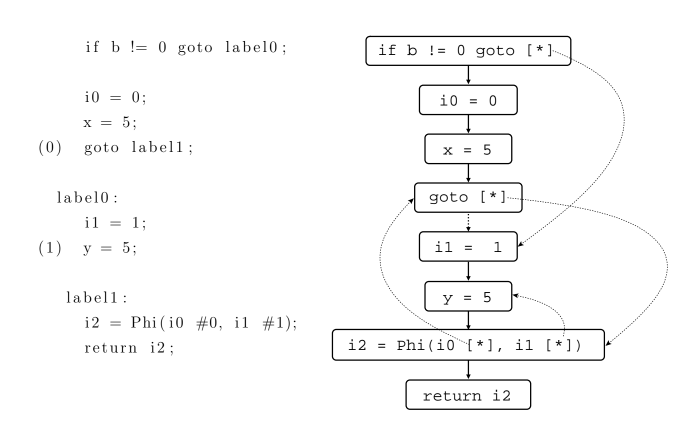


Figure 3.2: Printed Shimple fragment with its Soot internal chain representation.

It is worth noting that pointers in the ϕ -function context tend to track the end of control flow blocks ('backward pointers'), as opposed to pointers in branching statements (e.g. **goto** or **if** statements) which tend to track the beginning of control flow blocks ('forward pointers').

Among other things, tracking the end of control flow blocks aids us when transforming out of Shimple form. In Figure 3.3, the #0 and #1 pointers are used to determine where to place the copy statements when eliminating ϕ -functions. In the case of the #0 pointer, we place the copy statement before the statement pointed at since it happens to be a goto; whereas in the case of the #1 pointer, we place the copy statement at the end of the statement being pointed at since it is a non-branching statement.

;

	if b $!= 0$ goto label0;	if b $!= 0$ goto label0
	i0 - 0	i0 - 0.
	i0 = 0;	i0 = 0;
	$\mathbf{x} = 5;$	$\mathbf{x} = 5;$
(0)	goto label1;	i2 = i0;
		goto label1;
la	bel0:	
	i1 = 1;	label0:
(1)	y = 5;	i1 = 1;
		y = 5;
1	abel1:	i2 = i1;
	$i2 = Phi(i0 \ \#0, \ i1 \ \#1);$	
	return i2;	label1:
		return i2;

Figure 3.3: Shimple fragment when naively transformed to Jimple.

We track and patch backward pointers in the same manner that Jimple tracks and patches forward pointers by providing an extended patching chain implementation. The patching chain handles the more simple and frequent cases; a developer should keep these limitations in mind when complex or arbitrary control flow manipulations are desired.

For instance, in Figure 3.2, an instruction insertion after the statement y = 5 would result in the #1 pointer being moved down to the newly inserted statement. Similarly, if y = 5 were to be removed, the #1 would have to be moved so that it is at the end of the new statement at the end of the block. On the other hand, if a statement were to be added after the goto label1 statement, no action would have to be taken for the #0 pointer since the new statement would be in a different control-flow block.

By tracking the end of the block, we provide a high degree of flexibility for codemotion transformations enabled by SSA form. For instance, as long as other requirements⁴ are respected, the i0 = 0 statement could be moved higher up in the control flow graph to any block that dominates its originating block or as far down as the last statement of its originating block without affecting the final semantics of the program when translated out of SSA form.

3.2.3 Exceptional Control Flow

Exceptions in Java have the potential to affect control flow in a program and hence may have implications for any analysis that depends on correct control flow information. Soot has a variety of options for representing exceptional control flow in CFGs, from allowing one to ignore exceptions altogether to allowing one to create the most conservative graph possible.

Shimple can use any of the various CFGs provided by Soot, however, correctness is only guaranteed if the the CFG fully represents exceptional control flow in the same manner it represents regular control flow.

In CompleteBlockGraph, one of the CFG implementations provided by Soot, exceptional control flow is denoted in the most conservative manner possible i.e. it is assumed that any statement in a try block can throw an exception. Figure 3.4 shows some Jimple code with a try and catch block and Figure 3.5 shows its corresponding

⁴Such as, for instance, a variable must be defined before it is used.

CompleteBlockGraph.

Since i0 is defined multiple times in Figure 3.4, SSA form requires that it be split into several variables. Furthermore, since the catch block may subsequently use i0, a ϕ -function is required to merge the split variable.

If SSA form is constructed from the graph in Figure 3.5, the ϕ -function placed in the catch block would require 7 arguments since the block has 7 control-flow predecessors. The resulting catch block is shown in Figure 3.6.

This proves to be especially inconvenient as try blocks increase in size, since a ϕ function in a join node merging exceptional control-flow would need at least as many
arguments as there are statements in the try block. Furthermore, any operation on
the try block such as a statement removal or insertion would require a corresponding
update to the ϕ -function. Finally, eliminating the ϕ -function would require replacing
it with as many assignment statements as there are arguments, doubling the number
of statements in the try block in the best case – although it might be possible to
subsequently optimise this inefficiency.

For all these reasons, we decided on a practical compromise in Shimple. If we observe the ϕ -function in Figure 3.6, it is apparent that there are repeated arguments; i0_1 and i0_2 are repeated 3 times each.

The repetitions are mainly due to the overly-conservative edges from the try block to the catch block, and we observe that it suffices to consider only the dominant control-flow predecessors for repeated arguments. For example, if the ϕ -function has an argument for the block that defines i0_1, an argument for the same variable is not required for any other block (forcibly dominated by the defining block) since at ϕ -function elimination time, the argument will be replaced by an assignment statement that still dominates all other relevant blocks. Figure 3.7 illustrates the results of trimming ϕ -functions in this manner – in particular we see that the ϕ -function formerly with 7 arguments now has only 3.

This optimisation opportunity arises mainly as consequence of the artificial situation created by the conservative graph for exceptional control flow. In general, it does not apply to ϕ -functions for regular control flow and such optimisations would generally not be expected in SSA form.

```
i0 = 1;
   i1 = r2.nextInt(2); // i1 may assigned 0 or 1
   i2 = 3;
trystart:
   i0 = i0 / i1;
   i3 = i2;
   i4 = i3;
   i0 = i0 + i4;
   r3 = java. lang. System. out;
   $r3.println(i0);
tryend:
   goto exit;
catchblock:
   r4 := @caughtexception;
   r1 = \$r4;
   r5 = java.lang.System.out;
   $r5.println(i0);
exit:
    return;
```

catch java.lang.Exception from trystart to tryend with catchblock;

Figure 3.4: Jimple code with example try and catch blocks. Jimple denotes all exceptional control flow with catch statements at the end – in this case, any Exception thrown between trystart and tryend will be caught by catchblock.

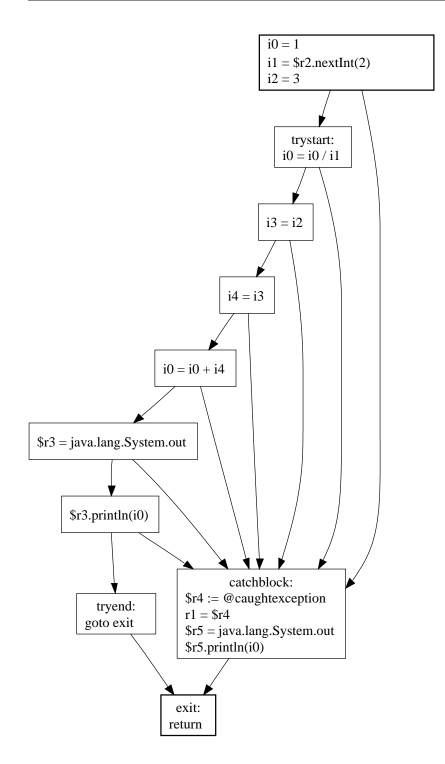


Figure 3.5: CompleteBlockGraph for code in Figure 3.4. As shown, it is assumed that any statement in the try block can throw an Exception – hence all the edges to the catch block.

catchblock: \$r4 := @caughtexception; i0_3 = Phi(i0_1, i0_1, i0_2, i0, i0_1, i0_2, i0_2); r1 = \$r4; \$r5 = java.lang.System.out; \$r5.println(i0_3);

Figure 3.6: Catch block from Figure 3.4 in SSA form. The ϕ -function has 7 arguments corresponding to the 7 control-flow predecessors in Figure 3.5.

When Shimple was first designed, Soot's CompleteBlockGraph implementation proved the most convenient option available, and the ϕ -function trimming approach was developed as a matter of necessity. Subsequently, efforts were made to improve exception analysis in Soot [Jor03] resulting in the availability of an Exceptional-BlockGraph which is now able to trim many of the excess exceptional edges from the control flow graph.

Figure 3.8 shows the results of computing Shimple with an optimised Exceptional-BlockGraph, but no further trimming of the ϕ -function. The resulting ϕ -function takes 3 arguments, although not the same ones as with the previous approach. In particular, it is interesting to note that one of the variables is repeated and had we applied our previous argument trimming optimisation, the result would be a ϕ -function with only two arguments.

3.3 Shimple Analyses

In this section we describe three analyses we have implemented that demonstrate the usefulness of Shimple and SSA form.

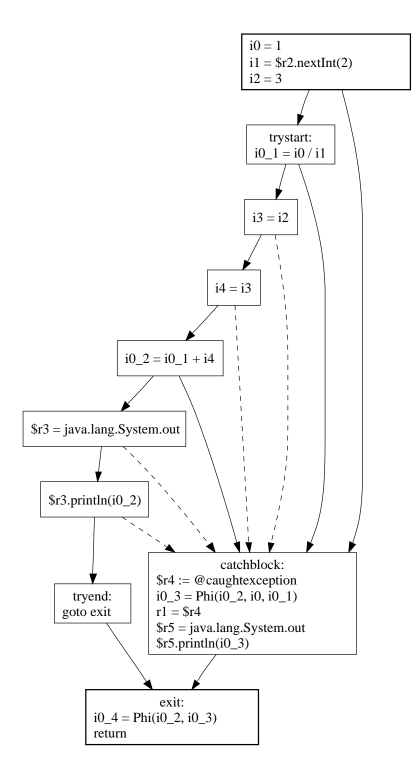


Figure 3.7: Only the blocks containing the dominating definitions of i0, i0_1 and i0_2 (non-dotted outgoing edges) are considered when trimming the ϕ -function.

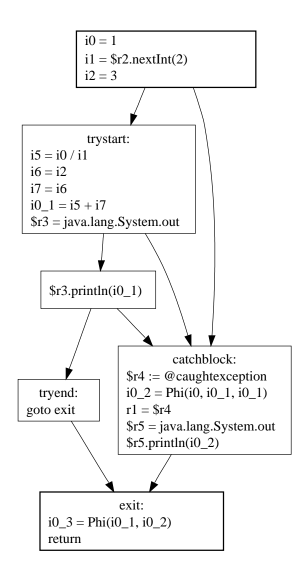


Figure 3.8: The optimised ExceptionalBlockGraph has far fewer edges resulting from exceptional control flow, and consequently the ϕ -function in the catch block has fewer arguments.

3.3.1 Points-to Analysis

We have modified Soot's Spark interprocedural points-to analysis [Lho02] to operate on Shimple and we have also implemented a simple conservative intraprocedural points-to analysis based on Shimple. Although the latter analysis is much cheaper to compute in Soot, the former analysis is evidently much more powerful.

While a full description of the intricacies of points-to analysis [Lho02] is beyond the scope of this thesis, it will suffice here to give a brief overview of the general idea so as to demonstrate how SSA form may be of use in this context.

In points to analysis, the goal is to conservatively determine all objects a variable may point to. For instance, in the code shown in Figure 3.9, variable o may point to objects allocated at allocation sites [1] and [2].

if(bool)		
o = new A()	[1]	A
<pre>print(o.type())</pre>		
else		
o = new B()	[2]	В
<pre>print(o.type())</pre>		
$\mathbf{x} = 0$	[3]	
<pre>print(x.type())</pre>		

Figure 3.9: Points-to example, code and pointer assignment graph. o may point to objects A and B allocated at sites [1] and [2], and so may x.

Points-to information has many uses and applications. As a basic example, if one knows which objects variable o may point to, one might be able to determine the effect of a method call on o.

Points-to information can be determined by constructing what is known as the pointer assignment graph and flowing information until a fixed point is reached. The nodes of the graph are variables and allocated objects. The edges of the graph are determined by the statements in the program, particularly assignment and copy statements.⁵

The pointer assignment graph in Figure 3.9 has nodes for o, x, an object (of type) A assigned in [1], and an object (of type) B assigned in [2]. The edges of the graph are determined by the three statements [1], [2] and [3] as shown. By flowing the information in the pointer assignment graph, the analysis would deduce here that o and x can point to either of objects A and B that are assigned in statements [1] and [2] respectively.

An analysis using this information could determine that the three print statements will print either 'A' or 'B' for the type of the object currently being pointed-to by o.⁶

It becomes clear how SSA form may be of use in points-to analysis if we consider the SSA version of the program as shown in Figure 3.10. The variable o has been split into variables o1, o2 and o3, and consequently, interesting flow information has been exposed, particularly in the contexts of o1 and o2, since a points-to analysis can now determine more precise information about the objects they can point to i.e. o1can only point to A and o2 can only point to B.⁷

However, a slight difficulty is now posed by the introduction of the ϕ -function given that the original points-to analysis may not know how to handle the statement. The situation can be handled trivially by adding a new rule for the construction of the pointer assignment graph. Statements of the form ' $v = \phi(v_1, v_2, ...)$ ' simply require that edges be added from all the nodes determined by the arguments to the ϕ -function to v, the variable that is the target of the assignment.

After flowing information in the pointer assignment graph (Figure 3.10), the points-to analysis can determine that o1 can point to A, o2 can point to B, and o3 and x can point to either A or B.

Hence, an analysis using this information can now determine more precise results for the first two print statements in this example. The first print statement will print

⁵This approach is flow insensitive since the ordering of the statements is not being taken into consideration.

⁶We assume here for the sake of simplicity that the variables will never be null-assigned.

⁷One can also point out that while the points-to analysis itself has remained flow insensitive, SSA form has exposed flow sensitive information.

if (bool) o1 = new A() [1] print (o1.type()) else o2 = new B() [2] print (o2.type()) o3 = $\phi(o1, o2)$ x = o3 print (x.type())



'A', the second will print 'B' and the third will print either 'A' or 'B' for the type of the object.

It is worth noting here that the ϕ -functions are being handled in the most conservative way possible. In fact if we had first generated SSA then eliminated the ϕ -functions with the naive (non-optimised) approach and subsequently applied points-to analysis on the resulting code, we would obtain results with the same improved precision as points-to analysis on SSA form without having to explicitly handle ϕ -functions.

As we will show in the next section, however, it is possible to make good use of ϕ -functions to improve the precision of analyses even further. Such techniques could also potentially be applied to improve the results of points-to analysis on SSA form.

Finally we note that Hasti and Horwitz [HH98] have done related work on the use of SSA form to improve points-to analysis in C. The authors point out that while flow sensitive points-to analysis can be more precise than flow insensitive points-to analysis, it is usually more expensive in terms of time and space. Hasti and Horwitz present an iterative algorithm based on using flow insensitive points-to analysis with SSA form that can produce results of increasing precision ranging from flow insensitive to flow sensitive analysis, allowing a compromise to be made in terms of efficiency versus precision of the points-to analysis.

3.3.2 Constant Propagation

If a variable can be proven to be a constant, its uses can be replaced by uses of a constant, the variable definition could be eliminated altogether, and further optimisation opportunities might arise. Constant propagation algorithms vary from the very basic to the truly sophisticated and may be combined and intertwined with other analyses for an overall greater effect [LSG00].

One particularly simple constant propagation algorithm is based on using the results from reaching definitions analysis. With a reaching definitions analysis, one can determine all definitions that may reach an arbitrary point in the program. Hence, if one can determine that a single constant definition reaches a particular use, one can perform constant propagation (e.g. Figure 3.11).

x = 4 x = 5if (bool) y = x + 5x = 6

Figure 3.11: With reaching definitions analysis, an analysis can determine that the use of x is of the constant 5 and not of 4 nor 6.

In SSA form, full-blown reaching definitions analysis is unnecessary given that each variable use already corresponds to a single definition. Although SSA form makes the analysis easier to implement and potentially faster, in this case there is no particular gain in the accuracy of the results. We can, however, formulate a more powerful constant propagation algorithm that takes greater advantage of SSA form by exploiting the control-flow information exposed by ϕ -functions.

Consider the code in Figure 3.12. By inspection, one will note that only two assignments of x can ever be reached and both of these assign 100 to x. Hence x

is a constant and the program may be reduced to an empty loop and a constant return statement. Reaching definitions analysis will however conclude that x might have 3 definitions reaching the entry point of the **if** statement and hence constant propagation would fail to find any opportunities for optimisation.

return x

Figure 3.12: Harder constant propagation problem, shown with optimised version.

The code is shown with explicit control-flow structure exposed and in SSA form in Figure 3.13. We wish to determine whether the use of variable x_4 is the use of a constant or not. We can see that x_4 is defined as the result of a ϕ -function on x_1, x_2 , and x_3 . The idea of the analysis is to prove that only x_1 and x_2 can in fact reach the ϕ -function and hence x_4 is a constant. Once that is ascertained, the if/else statement can be removed as well as all of the assignments. After some further simplifications, the resulting optimised code appears as shown in Figure 3.14.

During the flow analysis, two separate sets of assumptions are maintained and corrected until stabilisation is reached. The analysis is initialised with the assumption that all control-flow edges are unexecutable as well as the assumption that all variables are of unknown constant value \top , and begins by examining all nodes reachable from the start node of the control-flow graph [CFR⁺91].

Nodes are removed from the work list and each statement in a node is examined in turn. If the statement is an assignment statement, constant assumptions are updated accordingly, and all reachable nodes with affected uses are added to the worklist. If

x = 100	$x_1 = 100$
goto looptest	goto looptest
iftest:	iftest:
if $x \ge 200$ goto else	if $x_4 \ge 200$ goto else
x = 100	$x_2 = 100$
goto looptest	goto looptest
else:	else:
x = 200	$x_3 = 200$
looptest: if doIt != 0 goto iftest return x	looptest: $x_4 = \phi(x_1, x_2, x_3)$ if doIt != 0 goto iftest return x_4

Figure 3.13: Code in Figure 3.12 with control-flow explicitly exposed. In both non-SSA and SSA forms.

```
looptest:
if doIt != 0 goto looptest
return 100
```

Figure 3.14: Optimised code.

the statement is a branching statement, the branching condition is examined and if relevant constant assumptions have changed, the appropriate control flow edges are marked as reachable and the reachable nodes are added to the worklist.

The algorithm $[CFR^+91]$ is outlined in Figure 3.15.

while nodes list not empty: remove first node from list: examine each statement in node: if assignment statement: update constant assumptions if necessary if change, add reachable affected uses to node list

- if branching statement:
 - if constant assumptions have changed or first time visit: process and update edge assumptions if necessary add nodes reachable from statement to node list

Figure 3.15: Algorithm for constant propagation on SSA form.

A ϕ -function is handled as the right-hand side of an assignment statement and requires special treatment. In particular, one has to determine what a ϕ -function evaluates to in order to determine if the assignment statement affects the constant assumptions.

A ϕ -function $\phi(..., v_x, ...)$ is evaluated by merging the currently assumed constant or non-constant values of its *reachable* uses v_x as determined by the current edge assumptions.

The merge is performed as follows:

- A constant merged with itself evaluates to that constant value.
- A constant merged with a different constant (or a non-constant) evaluates to non-constant value \perp .

Since the ϕ -function merge rules only consider information along provably reachable control-flow paths, information alongst unreachable edges does not affect the results of the merge. As the analysis proceeds and assumptions are corrected, eventually all executable edges will be identified, ensuring the final correctness of the constant assumptions.

The ϕ -function represents a natural merge point for forwards control flow information in the program and is particularly useful here because it allows one to reason about control-flow and reachability specifically at the relevant points of the program. The resulting algorithm is easy to describe and easy to understand in contrast to non-SSA-based conditional constant propagation algorithms.

We have implemented this algorithm in Shimple.

In related work, Wegman and Zadeck [WZ91] describe an even more powerful constant propagation algorithm on SSA form which makes use of constant information that can potentially be gained from conditional branching statements. We will see an example of this in the next chapter.

3.3.3 Global Value Numbering

In global value numbering, we are concerned with the equivalences of variables. If variables can be determined to be equivalent, they are assigned the same global value number for the use of future analyses.

Consider the code in Figure 3.16. It is clear that, when talking about variable equivalence, it is useful to refer to a specific context or program point. For example, in the non-SSA version of the code, x and y are guaranteed to be equivalent only at the end of the if block.

By using SSA form we sidestep the issue since the variables are split and we can speak of the equivalences of variable x1 and y1 without having to say anything about x2, y2, x3 or y3. We do need a further qualification: In particular, x1 and y1 are equivalent at a point p of a program *only* if both definitions dominate that point. In other words, both x1 and y1 must be defined at p to be considered truly equivalent.

We are also interested in determining equivalences of variables that are condition-

if (boolean)	if(boolean)
x = z	x1 = z
y = z	y1 = z
else	else
x = 2	x2 = 2
y = 3	y2 = 3
	$\mathbf{x}3 = \phi(\mathbf{x}1, \mathbf{x}2)$
	$y3 = \phi(y1, y2)$

Figure 3.16: Simple example in both normal and SSA forms.

ally defined such as y3 and x3 in our example. Since ϕ -functions essentially summarise the relevant control-flow information, SSA form is of further use here.

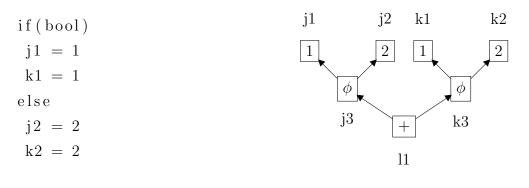
Using the approach described by Alpern et al. [AWZ88], we first build the value graph of the program. Nodes in the graph are labelled either by the values of variables they represent, in which case the nodes are leaves of the graph, or by the function that generates the value, in which case the ordered⁸ edges from the node point to the nodes that represent the arguments to the function. Trivial copy assignments such as $\mathbf{x} = \mathbf{y}$ result in node \mathbf{x} getting labelled by the label (and corresponding edges) generated for y.

An example of some code in SSA form with the corresponding value graph is shown in Figure 3.17.

Variable equivalence is computed from the value graph by determining whether the corresponding nodes for the variables are *congruent*. Congruency of nodes in this context is defined [AWZ88] as:

- The nodes have the same labels.
- The corresponding children of the nodes are congruent.

⁸In the case of commutative functions or operators such as '+' or '*', imposing an ordering may be undesireable. In such cases, one might assign a single hyperedge from the function to the operands [AWZ88].



 $j3 = \phi(j1, j2)$ $k3 = \phi(k1, k2)$ l1 = j3 + k3

Figure 3.17: Simple example in SSA form with corresponding value graph.

In our example, in Figure 3.17, $\{j1, k1\}$, $\{j2, k2\}$, $\{j3, k3\}$ and $\{l1\}$ form congruence classes and hence sets of variables that will be equivalent to each other.

Congruence is a symmetric, reflexive and transitive relation. We seek to maximise the number of congruences according to our definition and can do so using a simple optimistic partitioning algorithm.

The algorithm [AWZ88] begins with an initial partitioning of the nodes such that each partition contains nodes which may be congruent to each other; the partitioning is refined at each iteration:

Step 1: Place all nodes with same label in the same partitions.

Step i+1: Two nodes will be in the same partitioning in Step i+1 if in Step i:

- The nodes are in the same partition.
- The corresponding children of the nodes are in the same partition.

The algorithm terminates when two successive partitionings are the same. If we assume that every node has, on average, a small-constant number of children⁹, each

⁹A reasonable assumption in the case of the Shimple IR which is based on the 3 address-code Jimple. The exception of course being that ϕ -functions may have an arbitrary number of arguments in theory.

iteration can be computed in O(n) time and there are a maximum of O(n) iterations since there can be no more than n partitions, where n is the number of nodes. Hence the algorithm is $O(n^2)$. A more efficient partitioning algorithm is detailed by Alpern *et al.* [AWZ88].

It is worthwhile noting that this algorithm will find congruences even when there are loops in the graph, since it is initialised with optimistic assumptions and proceeds by refining the assumptions.

There is however a flaw in our current formulation, since we have assumed that all nodes labelled ϕ are comparable to each other. In our example in Figure 3.17 this is in fact true, since both ϕ -functions are in the same control flow block and hence they will choose the same corresponding argument at runtime i.e. if one ϕ -function chooses the nth argument, so will the other since the control flow edge into the block at runtime is the deciding factor.

It is clear however that ϕ -functions are not necessarily always comparable since they might embed differing control-flow information. One possible approach to correcting the algorithm is to associate a label with each ϕ -function e.g. each ϕ -function could be labelled with the corresponding block number. This label would allow ϕ functions within the same block to be compared while differentiating them from ϕ functions in other blocks as illustrated in Figure 3.18.

What if otherbool had been equal to bool in Figure 3.18? In that case, j3 would clearly be equivalent to k3 but since the ϕ -functions are labelled differently, our partitioning algorithm would never assume that the corresponding nodes in the value graph could be congruent.

The key here is that it is the if-predicates which determine the control-flow edges reaching the ϕ -functions. If we can identify all ϕ -functions that depend on an ifpredicate, we could simply label those ϕ -functions as ϕ_{if} and associate them with the corresponding predicate e.g. by adding an edge from the ϕ_{if} to its predicate in the value graph. With such a modification, our partitioning algorithm would determine that j3 and k3 are equivalent in Figure 3.18.

The algorithm as described here has been implemented in Shimple. Alpern et al. [AWZ88] detail several other enhancements and refinements of the global value

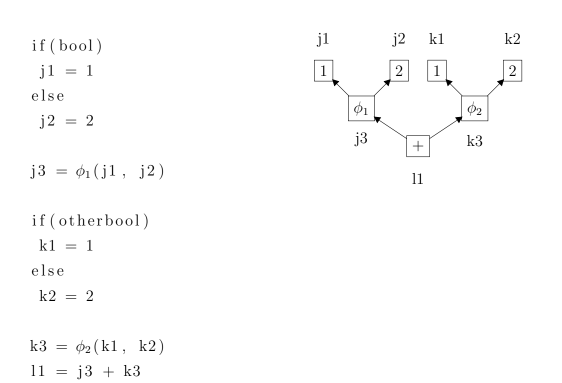


Figure 3.18: In this example, j3 and k3 are not necessarily equivalent. Since the corresponding ϕ -functions are now labelled differently, they will never be placed in the same partition.

numbering algorithm. We will not cover them here as we simply wish to draw attention to the fact that ϕ -functions do allow one to reason about control-flow dependent information in convenient ways. Here for example, ϕ -functions can be treated as any other operator or function such as – or + except that the information gained is now control-flow related and hence more variable equivalences can be determined.

3.4 Related Work

SSA form is a popular representation, and as such there have been many implementations of it for optimising compiler output for many languages and targets, including the Java bytecode. We reference three particular implementations for optimising Java through the use of SSA form below.

The JikesRVM [AAB+00] is a Java virtual machine which can create various SSA representations of Java bytecode for just-in-time optimisation in a dynamic environment. In JikesRVM, the most basic SSA representation is constructed using the classic Cytron *et al.* algorithm [CFR+91] and allows a variety of SSA-based analyses to be applied by the optimising compiler. The compiler also implements Array SSA form [KS98] and with a unification approach [FKS00] can also analyse array and object references.

The Marmot compiler $[FKR^+00]$ also implemented an SSA IR for Java bytecode analysis and optimisation. The authors specifically note the difficulties in maintaining ϕ -functions to preserve SSA variants while the control-flow graph is undergoing transformations and consider using SSA not as the primary intermediate representation but rather as a secondary source of information for the transformation of a primary (non-SSA) IR. The suggestion appears to be similar to the SSA numbering approach proposed by Lapkowski and Hendren [LH96] where instead of renaming variables with new integer subscripts corresponding to new SSA variables, the numbers are stored in secondary structures as variable annotations. Knowing the SSA number of a variable would allow an analysis to make deductions about the value and context of a variable since one would know what the corresponding SSA name of the variable would have been. However, any advantages associated with the ϕ -functions themselves would be lost in such an approach.

Another compiler tool of note with recent SSA support is the GNU Compiler Collection (GCC) [Fre]. GCC can compile Java, C, C++ and other languages to a common intermediate representation known as GIMPLE (which is similar to Jimple, given that both forms are inspired from the SIMPLE representation [HDE+93]) followed by Tree SSA [Nov03], an SSA version of GIMPLE in an abstract tree syntax representation.

Finally, it is worth noting that the Φ syntax introduced by Sarkar and Knobe [SK98] for encoding control flow information directly in the SSA nodes in order to support runtime execution of ϕ -functions is somewhat comparable to our augmented ϕ -functions in Shimple which explicitly track the control flow source to facilitate deconstruction of the form.

Chapter 4 Extended Shimple

In Chapter 1 we introduced basic variable splitting, followed in Chapter 2 by a natural extension of the concept to SSA form, where variables are split sufficiently such that every variable is only ever defined once in the static view of the program. This chapter explores the implications and possible applications of splitting variables to an even higher degree than that required by SSA form.

Section 4.1 of this chapter introduces extended SSA (eSSA) form [BGS00] and provides example applications of the representation. In Section 4.2, we introduce Static Single Information (SSI) form [Ana99], which can be considered an extension to eSSA form, as well as a sample application. Finally, in Section 4.3 we provide an overview of Extended Shimple which combines the best elements of the eSSA and SSI forms in our Soot implementation. We conclude with a brief review of related work in Section 4.4.

4.1 eSSA Form

4.1.1 Overview

Variable splitting becomes useful when it allows more precise information to be attached to a new variable name, potentially exposing context-dependent information on the value of a variable. Basic SSA form splits variables to the extent such that it is guaranteed that every variable used in a program has a single definition point. In essence, basic SSA form exposes information gained when variables are *written* i.e. multiply-defined variables are split such that the information gained at each static write point can now be associated to a new variable name. An example of this was seen in Figure 2.2 of Chapter 2, where information gained at the write points of i are exposed by splitting the variable into i1 and i2 at the relevant points.

Extended SSA form exposes additional context information gained at certain *read* points of the program, in addition to the context information exposed at the write points in basic SSA form. Any read or use of a variable that allows the inference of further information on its value may be used when generating eSSA form.¹

In the code fragment in Figure 4.1, we see a conditional branch with a comparison expression of the form $\mathbf{x} > 0$. As an observation, it is interesting to note that the first two print statements in the code will never print the same value for \mathbf{x} even though they reference the same SSA variable – the "same name, same value" property does not hold because the branch condition is implying something about the context and hence the value being printed. If one side of the branch is taken, we might deduce that \mathbf{x} is greater than 0, but if the other side of the branch is taken we would know \mathbf{x} to be less than or equal to 0 in that context. Hence, the comparison statement is exposing new context-based information on the value of \mathbf{x} .

Although in basic SSA form no splitting would occur given that the variable definition itself is not being changed, in eSSA form we split \mathbf{x} into $\mathbf{x1}$ and $\mathbf{x2}$. We do this simply by introducing π -functions as shown in the figure. Here the π -functions do nothing but denote positions where \mathbf{x} is given new names, although in an implementation additional information may be associated with a π -function for convenience, such as a pointer to the comparison statement which caused the split.

The split of x into x1 and x2 requires a new ϕ -function to be inserted, since the third print statement is not in the context of either x1 or x2.

¹Without loss of generality, we focus here on information gained at conditional statements i.e. at the branch nodes of a CFG, although it is possible to conceive of other statements such as a code assertion or an implicit array bounds check which may reveal useful information on a variable.

if $(x > 0)$	if (x > 0)
print(x)	x1 = Pi(x)
else	print(x1)
print(x)	else
	x2 = Pi(x)
print(x)	print(x2)
	x3 = Phi(x1, x2)
	print(x3)

Figure 4.1: A simple conditional branch code fragment in SSA and eSSA forms. Depending on the branch taken, we can deduce further information on the value of \mathbf{x} , hence we split \mathbf{x} in eSSA form.

We contend that eSSA form is particularly suitable for predicated data flow analyses, since it helps expose information associated with conditional branching.

4.1.2 π -functions

Conceptually, π -functions are much simpler than ϕ -functions. π -functions can be seen as glorified copy statements, the purpose of which is to assign a new name to a variable in a given context. What differentiates a π -function from an arbitrary copy statement is the reason for and location of the π -function; evidently it is up to an analysis to make good use of a π -function.

Where should π -functions be inserted when computing eSSA form?

As we have noted, conditional branch statements have the potential to expose further context information on a variable. Hence, perhaps we should insert π -functions after all conditional branch statements. The next matter to determine is which variables need to be split at a conditional branch.

In Figure 4.1 we had decided to split variable \mathbf{x} , since we could gain more contextdependent information on it. Clearly, this was because the conditional branch statement had a direct reference to \mathbf{x} in its conditional. We might deduce that it would be useful to split all variables that are referred to in a conditional.

When eSSA was first introduced [BGS00], the approach used to compute eSSA form was to first insert trivial π -functions and then compute SSA form as would be normally done. π -functions were simply added at the exits of conditional branches for any variables referenced in the conditional.

In Figure 4.2 we see an example of where it might be useful to split a variable y even though it is not directly referenced by a conditional branch statement. The split is useful because y is in fact defined in terms of x which *is* referenced by the conditional branch statement – hence we know that y1 is not the same value as y2. Another point of interest in the figure is that we have not split variable x since it would serve no purpose – the only use of x in the program is not within the context of the conditional branch.

y = x + 1	$\mathbf{y} = \mathbf{x} + 1$
if (x > 0)	if (x > 0)
print(y)	y1 = Pi(y)
else	print(y1)
print(y)	else
	y2 = Pi(y)
print(y)	print(y2)
print(x)	
	y3 = Phi(y1, y2)
	print(y3)
	print(x)

Figure 4.2: Example situation where the original eSSA algorithm would not split variable y, although this could be potentially useful since y is defined in terms of x and hence would gain context information from a split. Furthermore, the original algorithm would split x although this is not useful here.

We deduce from the previous example that the original algorithm [BGS00] may not be the optimal approach to computing eSSA. We will postpone the discussion of a more optimal placement for the π -functions till later sections.

Finally, we note that removing π -functions can be naively done by replacing them with equivalent copy statements followed by a few optimisations such as copy propagation and dead assignment elimination.

4.1.3 Improving SSA Algorithms

Constant Propation

One obvious and somewhat trivial application of eSSA form would be to improve the accuracy of the SSA constant propagation algorithm outlined in Section 3.3.2.

Where the previous algorithm obtained constant information solely from assignment statements in a program, an algorithm based on eSSA form might make use of information gained from particular comparison statements.

For example, in Figure 4.3, the SSA constant propagation algorithm would fail to detect that \mathbf{x} is a constant in the context of the if-block. In eSSA, \mathbf{x} would be further split, allowing an algorithm to identify $\mathbf{x1}$ and $\mathbf{x2}$ as constants.

In an implementation, an extended analysis would handle π -functions by considering the conditional statement associated with them as well as their context.² For instance, in the example, the π -function is found in the true branch of an $\mathbf{x} = 4$ test, hence when the assignment statement for $\mathbf{x1}$ is examined by the algorithm, the constant assumptions can be updated to reflect the fact that $\mathbf{x1}$ is 4. This will eventually lead to the discovery that $\mathbf{x2}$ is a constant.

For the purposes of constant propagation, we might also make use of information provided by an inequality or comparison statement in a conditional (Figure 4.4), although our current algorithm may not store sufficient information on the possible values of a variable once it has been determined that it is not a constant. In Section 4.1.4, we will see a value range analysis implementation which subsumes the

 $^{^2\}mathrm{As}$ we will see in Section 4.3.4, Shimple conveniently stores this information in the $\pi\text{-function}$ itself.

x = random()	x = random()
if (x == 4)	if (x == 4)
x1 = x - 1	x1 = Pi(x)
print(x1)	x2 = x1 - 1
	print(x2)
x2 = Phi(x, x1)	
print(x2)	x3 = Phi(x, x2)
	print(x3)

Figure 4.3: Example situation where constant propagation analysis might be improved. The code is shown in both SSA and eSSA forms; in eSSA form x1, and hence x2, can be identified as constants in the context of the if-block by virtue of the comparison statement associated with the π -function.

constant propagation algorithm and makes use of such information.

We note that Wegman and Zadeck [WZ91] do also track this information in their conditional constant propagation algorithm. Although they only use SSA form and not eSSA form, other means were used to track the control flow and context information in the case of comparison statements.

Points-to Analysis

In a similar manner, the SSA points-to analysis algorithm can be improved in precision when applied to eSSA form. In Figure 4.5, **b** might point to object **A** or some other unknown object. In the context of the if-test however, **b** can only point to **A**, a fact that is not recognised by our previous algorithm.

The code from Figure 4.5 is shown in eSSA form in Figure 4.6. b has been split in the context of the if-test as b1; we might conceive of a new rule for π -function assignments when constructing and evaluating the pointer assignment graph.

If a variable b1 is the result of an assignment from a π -function on a variable b,

$$x = \{1, 2, 3\}$$
 if (b != true)
b1 = Pi(b)
if (x > 2) use(b1)
x1 = Pi(x)
print(x1)

Figure 4.4: Example fragments where a comparison or inequality may reveal useful information for constant propagation. In the first case, x is a non-constant which may take 3 possible values, but x1 can be determined to be the constant 3. In the second case, b1 can be determined to be the constant false.

```
a = new A()
....
// b may be aliased to a
// b may point to object B
....
b.doSomething()
if(a == b)
```

print(b.type())

the original algorithm.

b

Figure 4.5: Example situation where points-to analysis might be improved. The information gained at the comparison statement is not taken into consideration in

and the π -function is associated with the true-branch of an equality statement which references **b** (or the false-branch of an inequality statement which references **b**), then a new *intersection* node is added to the pointer assignment graph with edges from the two nodes referenced in the equality statement (**b** and **a** in the figure). Finally, a node for **b1** is created with an edge from the new intersection node to b1.

If the given conditions are not met, the π -function assignment can be treated as a normal copy statement when building the pointer assignment graph.

An intersection node differs from a regular node in that the set of values associated with it is an intersection of the sets of values of the in-edges rather than a union. In our example, b1 can never point to objects that are not in the intersection of b and a given the equality constraint. The pointer assignment graph for the example is shown in Figure 4.6.

```
a = new A()
...
// b may be aliased to a
// b may point to object B
...
b.doSomething()
```

```
if (a == b)
b1 = Pi(b)
print(b1.type())
```

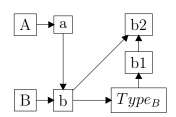
b2 = Phi(b, b1)

Figure 4.6: Example from Figure 4.5 shown in eSSA form. We take advantage of the introduction of π -functions and can therefore deduce that b1 can only point to object A.

Similarly, we might derive further information from comparison expressions of the form (a instanceof TypeX), a type of expression which tends to occur frequently

in Java code. Figure 4.7 illustrates an example which introduces a type filter node to the pointer assignment graph. The new node is simply responsible for filtering out all objects which do not match the type being tested against.

```
a = new A()
...
// b may be aliased to a
// b may point to object B
...
b.doSomething()
if (b instanceof B)
    b1 = Pi(b)
    print(b1.type())
```



b2 = Phi(b, b1)

Figure 4.7: As with direct comparison statements, we can also make use of information gained from **instanceof** tests. In this example, **b1** is guaranteed to be of type **B** in the context of the if-block. Accordingly, we have extended the pointer assignment graph to include a type filter node which outputs the out-set containing all objects from the in-set which match the given type.

The simple rules we have proposed to improve points-to analysis on eSSA form, while improving over our previous results on SSA form, can be extended or used in conjunction with other analyses such as copy-propagation and global value numbering to provide even better results e.g. in the presense of variable aliasing (Figure 4.8). Our purpose in this section has been to simply illustrate the possible use of the context information exposed by eSSA form.

```
a = new A()
...
// b may be aliased to a
// b may point to object B
...
c = b
b.doSomething()
if(c == a)
```

print(b.type())

Figure 4.8: Although c is aliased to b, our new rule for points-to analysis on eSSA form will not detect that b can only point to object A in the if-test context. We could perhaps use copy-propagation or global value numbering in conjunction with points-to analysis in order to improve the results.

4.1.4 Value Range Analysis

Value range analysis [Har77], also known as generalised constant propagation [VCH96], seeks to determine the set of values that might be associated with a variable rather than a simple determination of whether a variable is of constant or non-constant value. Value range analysis naturally subsumes constant propagation and an implementation can potentially detect more constants than a typical constant propagation algorithm.

Patterson [Pat95] details a value range analysis algorithm suitably adapted to eSSA form. While the original algorithm only made use of basic SSA form, the program code was augmented with assertion statements judiciously placed after comparison statements. The assertion statements served to identify context information on a variable, causing a variable split in an identical manner to π -functions.

The value range analysis algorithm is very similar to the SSA constant propagation algorithm we described in Section 3.3.2, with a few additional difficulties that need to be addressed.

Efficiency Concerns

In constant propagation we are only interested in whether a variable is of a constant or non-constant value and hence the semantic domain for the data flow analysis is very simple. Data flow information associated with a variable simply indicates whether the variable is assumed to be an unknown constant (\top) , a known constant, or a nonconstant (\bot) and the information is gradually refined in a monotonic order, from \top to \bot . Once a variable has been determined to be a non-constant, no further change in that assumption is possible, and hence constant propagation analysis converges very quickly even in the worst case.

In value range analysis, we must track sets of values instead of the simple constant information. A representative semantic domain might hence range from the empty set of values (\top) to the set of all possible values of a datatype (\bot) , with a partial ordering defined by set inclusion. This domain is very large, and considering that information associated with a variable may grow gradually from the empty set to the full set during an analysis, practical compromises are necessary in order to achieve acceptable space and time performance.

Value range analyses typically do not track sets of values, but simply one or more value ranges per variable that allow the most common cases (e.g. an arithmetic progression of values) to be represented. A range can be compactly represented without need of enumerating all the possible values in it and can hence conservatively represent the potential values a variable may hold.

Patterson [Pat95] represents value ranges as a three-tuple of the form [Lower:-Upper:Step]³ tracking the lower and upper bounds of a range as well as an arithmetic step value indicating the stride. The author chooses to associate a set, of maximum size 4, of these ranges per variable, widening existing ranges as necessary and decreasing the overall precision in order to guarantee conservativeness of results.

 $^{^{3}\}mathrm{The}$ author also tracks the probability associated with a value range at runtime, but we shall not consider this here.

Verbrugge *et al.* [VCH96] represent a value range for a variable by a single twotuple of the form [Lower:Upper]. We shall use the latter representation for the purposes of demonstrating the use of eSSA form in the context of value range analysis, although evidently the precision of results can be improved by using larger representations at the expense of efficiency.

Even with a compact value range representation, the semantic domain is still large enough to cause concerns; it can take a long time for ranges to converge during flow analysis.

Patterson [Pat95] addresses the issue by identifying loop carried expressions and attempting to *derive* their corresponding value ranges without data flow iteration. In cases where the derivation heuristic fails, the algorithm falls back to brute force iteration.

Loop-carried variables can easily be detected in SSA form by identifying ϕ -functions which have at least one in-edge determined to be a back edge by a depth-first search rooted at the start node of the control-flow graph. Variables that are the target definition of such a ϕ -function are evidently loop-carried.

If the Patterson [Pat95] algorithm comes across a loop-carried variable for the first time, it examines all statements operating on the variable and employs a heuristicbased template-matching approach in order to derive a value range for the variable. If a variable has had its value range successfully derived it is not subsequently reevaluated, otherwise, the variable is marked as impossible to derive and is treated normally during data flow analysis.

Verbrugge *et al.* [VCH96] are concerned with optimising the data flow analysis convergence even in the worst case scenario, by artificially limiting the number of iterations. The main approach in the paper is to allow iteration to proceed for a fixed number of times, and if convergence has not yet been achieved, to artificially 'step up' key ranges monotonically in the semantic domain. The analysis then proceeds for a fixed number of times, and if convergence has still not been reached, another 'step up' is executed. Since ranges will eventually be stepped up to the maximum range $([-\infty, \infty]$ where ∞ is typically limited by the size of the datatype), the analysis will proceed for a reasonable number of times before forcibly converging.

Maintaining Value Range Assumptions

As in constant propagation, we track and update value range assumptions for each variable as the analysis proceeds; assumptions are updated each time an assignment statement is processed.

We identify four main types of assignment statements.

• Simple assignments of the form a = b involving either a constant, a variable or a function on the right-hand side.

If **b** is a constant, then the value range for **a** is [b:b]. If **b** is a variable, **a** inherits the value range for **b**. Otherwise, if we have no further information, **b** is assumed to be $[-\infty, \infty]$ (i.e. \perp).

• General binary arithmetic statements of the form a = b OP c, or similarly, unary operations.

The value range for a can be derived from the value ranges for b and c.

For example, if OP is the additive operator + and the value ranges for b and c are [b1:b2] and [c1:c2] respectively, then the value range for a would be [(b1+c1):(b2+c2)] representing the range with the minimum and maximum possible results when b is added to c.⁴

All arithmetic operations can be handled in this manner, including unary and binary operations.

• Assignments of the form v = Phi(a, ..., z).

The value range of \mathbf{v} is the range with the lowest and highest boundaries as extracted from the ranges associated with the reachable arguments to the ϕ function.

• Assignments of the form a = Pi(b).

In general, a will inherit the value range for b using any facts associated with the π -function to improve the range when possible.

⁴Assuming that no underflow or overflow occurs.

 π -functions are a potentially interesting source of information for value range analysis. In particular, information might be extracted from any numerical comparison associated with a π -function, as was the case in limited situations with constant propagation analysis (e.g. Figures 4.3 and 4.4).

In Figure 4.9, the π -function is associated with the comparison statement i < 7. Hence we have information about the possible values of i1 which should be taken into consideration during flow analysis or when ancilliary procedures such as the range 'stepping up' process we previously described are performed (e.g. we will never need to step up to ∞ , since 7 is known to be the range upper bound for i1).

if(i < 7)
i1 = Pi(i)
print(i1)
...</pre>

Figure 4.9: In value range analysis, the π -function is associated with the knowledge that i is always less than 7 in the context of the if-block. In a similar manner, many other numerical comparisons can provide useful value range constraints.

Algorithm Outline

We model the algorithm for value range analysis after the algorithm we outlined in Section 3.3.2 for constant propagation [CFR⁺91].

Hence, we keep track of two separate sets of assumptions, our value range assumptions for each variable as well as a set of assumptions determining whether an edge is executable or not. At initialisation time, all edges are deemed unexecutable and the value range assumption for each variable is the null or undefined range.

A worklist of nodes is maintained; the list contains the start node at initialisation time and nodes are removed and processed one at a time from the list. The algorithm is outlined in Figures 4.10 and 4.11.

while nodes list not empty:

remove first node from list:

examine each statement in node:

if assignment statement:

process(stmt) and update value assumptions if necessary if change, add reachable affected uses to node list

- if branching statement:
 - if value assumptions have changed or first time visit: process and update edge assumptions if necessary add nodes reachable from statement to node list

Figure 4.10: Outline of algorithm for value range analysis on eSSA form – the process function is outlined in Figure 4.11.

4.2 SSI Form

4.2.1 Overview

SSA form can be seen as enabling forward sparse data flow analysis since information can be flowed from variable definitions directly to their uses, in contrast to data flow analysis on non-SSA forms where information must generally be flowed to and from every node in the CFG.

The main reason forwards sparse data flow analysis is possible in SSA form is the presence of ϕ -functions which identify the key positions where flow information must be merged. For instance, in Figure 2.2, a (naive) sparse constant propagation algorithm might flow information from the definitions of i1 and i2 directly to i3 at the merge point identified by the ϕ -function.

process (statement):

if value is loop-derived: increment iteration count for value if count is larger than predefined limit: step up range assumption associated with value return new assumption immediately

compute and return new value range assumption per normal rules

Figure 4.11: Function for processing an assignment statement. If the value is a loopderived one and the iteration count associated with the statement has exceeded a given limit, the current value range assumption is stepped up in the semantic domain as necessary, for efficiency reasons. Otherwise, the new value range assumption is computed according to the type of assignment statement as we previously described.

Static Single Information (SSI) form [Ana99] introduces σ -functions which can be seen as serving to identify control-flow *split* points, hence potentially enabling sparse *backwards* data flow analysis in addition to forwards analysis. We note that eSSA form, as originally formulated, may not prove as suitable for backwards flow analysis given that eSSA π -functions are only inserted if directly referred to in the conditional at the split point.

SSI is also useful for the purpose of predicated data flow analysis, similarly to eSSA form; the two representations can in fact be considered related to each other, with SSI being the more expressive of the two.

4.2.2 σ -functions

Figure 4.12 shows the SSI version of the code from Figure 4.1. We have placed it side by side with the eSSA version for the purposes of comparison.

As can be seen from the figure, where eSSA form inserts separate nodes for each branch split of a variable, SSI form introduces a single σ -function with multiple

if $(x > 0)$	if $(x > 0)$
x1 = Pi(x)	x1, $x2 = Sigma(x)$
print(x1)	y1, y2 = Sigma(y)
print(y)	print(x1)
else	print(y1)
x2 = Pi(x)	else
print(x2)	print(x2)
\mathbf{D}	\mathbf{D} \mathbf{D} $(1, 0)$
x3 = Phi(x1, x2)	x3 = Phi(x1, x2)
print(x3)	y3 = Phi(y1, y2)
	print(x3)

Figure 4.12: The code from Figure 4.1 shown here in both eSSA and SSI forms. The σ -functions are placed at the end of the control-flow block containing the if statement i.e. they are executed before the control-flow split.

targets. Also notable is the fact that SSI form will split variable y, although eSSA form will not.

Where ϕ -functions serve to select an argument or use depending on which branch control flow comes from at runtime, σ -functions select which target variable is assigned depending on which control flow branch is to be taken at runtime. Table 4.1 [Sin] further contrasts the differences between ϕ -functions and σ -functions.

4.2.3 Computing SSI Form

In SSI form, a reverse symmetry has become evident with regards to σ -functions and ϕ -functions. Indeed, the symmetry is intentional given that σ -functions are intended to perform the same function for sparse backwards analysis as ϕ -functions perform for sparse forwards analysis.

The placement of nodes in SSI form is required for a variable V as follows [Ana99]:

• ϕ -functions are placed at a control-flow merge when disjoint paths from a con-

ϕ -function	σ -function
Inserted at control-flow merge points.	Inserted at control-flow split points.
Placed at start of basic block.	Placed at end of basic block.
Single destination operand.	n destination operands, where n is the
	number of successors to the basic block
	that contains the σ -function.
n source operands, where n is the number	Single source operand.
of predecessors to the basic block that con-	
tains the ϕ -function.	
Takes the value of one of its source	Takes the value of its source operand and
operands (dependent on control-flow) and	assigns the value to one of the destination
assigns the value to the destination	operands (dependent on control-flow).
operand.	

Table 4.1: Differences between ϕ -functions and σ -functions [Sin].

ditional branch come together and at least one of the paths contains a definition of V.

• σ -functions are placed at locations where control-flow splits and at least one of the disjoint paths from the split uses the value of V.

It should come as no surprise that the algorithm for computing SSI form can be formulated as a natural extension to the algorithm previously described for computing SSA form [Sin02].

We can retool the algorithm for inserting trivial ϕ -functions from Figure 2.8 to insert trivial σ -functions as shown in Figure 4.13. In particular, instead of tracking variable definitions we track variable uses, and instead of using the dominance frontier, we use the reverse dominance frontier which is simply computed by reversing the edges in a control flow graph and computing dominance frontiers on the nodes of the reversed graph.

The algorithm for computing SSI form might hence be formulated as first inserting

for each variable V do for each node X that uses V: add X to worklist W for each node X in worklist W: for each node Y in reverse dominance frontier of X: if node Y does not already have a σ -function for V: append ''V,..., $V = \sigma(V)$ '' to Y if Y has never been added to worklist W: add Y to worklist W

Figure 4.13: Algorithm for inserting σ -functions.

trivial ϕ -functions, inserting trivial σ -functions, followed by the variable renaming process we described in Section 2.3.2. A complication arises, however, since inserting ϕ -functions potentially introduces new variable uses to the code and inserting σ functions introduces new variable definitions; new uses may require new σ -functions while new definitions may require new ϕ -functions.

Hence, after one pass of ϕ -function and σ -function insertion another may be required and so on until a fixed point is reached. Since the number of possible node insertions is limited by the size of the control-flow graph, termination is guaranteed. If we assume the generic placement algorithm is linear complexity (in terms of the program size) in practice [CFR⁺91], we can say that the SSI node placement algorithm might be of quadractic complexity in the worst case [Sin02]. It has been shown that in practice, computation of SSI form need not be significantly more expensive than computation of SSA form [Sin].

An algorithm for computing SSI form [Sin02] is outlined in Figure 4.14. The algorithm results in the construction of *minimal* SSI form, paralleling our definition for minimal SSA form. By subsequently eliminating unused ϕ -functions or σ -functions we can obtain a *pruned* SSI form which might prove more convenient for a particular analysis [Ana99].

```
insertTrivialPhiNodes()
change = insertTrivialPiNodes()
```

```
while(change):
    change = insertTrivialPhiNodes()
    if(change):
        change = insertTrivialPiNodes()
```

performVariableRenaming()

Figure 4.14: Algorithm for computing SSI form.

4.2.4 SSI Analyses

All the analyses we previously detailed for eSSA form can be retooled for SSI form by essentially operating on σ -functions instead of π -functions. In this section we briefly outline how a simple sparse backwards analysis might work on SSI form.

Resource Unlocked Analysis

The objective of the analysis is to determine whether a resource is guaranteed to be unlocked by the exit of a program. We assume that once a variable is unlocked, it cannot be re-locked.

Figure 4.15 shows an example program in its original version and in minimal SSI form. A backwards analysis can be used to determine that the SSI variable \mathbf{x} is in fact not always properly unlocked by the end of the program as well as help pinpoint the context in which it is not unlocked by indicating that $\mathbf{x}2$, as split from \mathbf{x} , is not locked before exit.

The backwards analysis only needs to process statements which use or define \mathbf{x} (as well as any variables related to \mathbf{x}), as opposed to having to flow information for all nodes in the control flow graph, making it a sparse analysis.

The σ -functions are located such that backwards flow information can be ap-

```
x = resource
                                  x = resource
                                  if(b)
if(b) exit
                                  x1, x2 = Sigma(x)
if(c)
                                      exit
  use(x)
                                  else
  unlock(x)
                                     if(c)
                                     x3, x4 = Sigma(x1)
else
  use(x)
                                         use(x3)
  unlock(x)
                                         unlock(x3)
                                      else
                                         use(x4)
                                         unlock(x4)
                                     x5 = Phi(x3, x4)
```

Figure 4.15: Example target program for our resource unlocked analysis in its original version and SSI form. The objective of the analysis is to determine whether the SSI variable x is properly unlocked on all paths to the exit.

propriately merged. Any information known about variables on the left side of the statement can be propagated to the variable on the right side. In a backwards analysis, ϕ -functions must also be handled by propagating information from the variable on the left side to variables on the right side of the statement.

Our rules for the analysis are as follows.

- All variables are assumed to be locked by default (\top) .
- A use of the form unlock(v) generates a new fact that v has been unlocked.
- A statement of the form v = Phi(a, b) causes all known facts for v to be propagated to a and b i.e. if v is known to be unlocked than so are a and b, otherwise any known facts for a and b are unchanged.⁵
- A statement of the form a, b = Sigma(v) propagates a fact to v if it is known to hold for both a and b. Copy statements of the form a = v can be handled similarly by propagating facts from a to v.

Informally, we can see in our example that the analysis would learn that x3 and x4 are unlocked, allowing it to deduce that x1 is unlocked. When processing the σ -function on x however, the analysis will deduce that x may not be properly unlocked given that variable x2 is not unlocked before exit.

As a last point, it is worth noting again that we would not be able to formulate this backwards analysis in a similar manner on eSSA form; consider in our example that neither **b** nor **c** in the conditional statements refer to **x** directly and hence would not require π -functions to be inserted.

4.3 Implementation of Extended Shimple

In Extended Shimple, we have implemented what we consider to be the best (or more practical) features of the eSSA and SSI forms. Furthermore, we have successfully implemented value range analysis as previously described.

⁵We might simplify this rule to simply "all facts known about v are propagated to a and b" if we use pruned SSI form where we would not have dead ϕ -functions as we do in the example.

4.3.1 Disadvantages of σ -functions

While SSI form is more powerful than eSSA form in that it supports sparse backwards flow analyses as well as predicated data flow analyses, we consider σ -functions to be awkward to use in a practical implementation such as Shimple for the following reasons.

 σ-functions result in one or more target variables being defined in a statement. Although Soot can support multiple variable definitions per statement, in practice it would be awkward or unexpected to have to manipulate such statements in an analysis.

Furthermore, it would be hard to indicate the scoping of the newly defined variables in a σ -function i.e. there is little clue in the σ -function notation as to which block a particular variable definition pertains to.

• σ -functions must be placed at the end of a branching block.

As we have previously mentioned in Section 3.2.1, in Soot, Jimple and Shimple are represented internally by a chain of statements which can be either branching or non-branching. Any block structure is consequently implicit rather than explicit since blocks are not directly represented – although they can be inferred by a separate analysis.

Hence, Extended Shimple would require a significant deviation from the Jimple IR if we were to admit the possibility that formerly branching statements in a chain might defer branching to an assignment statement (σ -function) which would have been formerly expected to be a non-branching statement.

We believe Soot developers would not take kindly to such an abrupt change in representation.

4.3.2 Placement of π -functions

Given the disadvantages associated with σ -functions in the context of Soot, we have adopted the π -functions from eSSA form for Extended Shimple. A statement of the form $a_1, ..., a_n = \sigma(v)$ can be represented by n statements of the form $a_x = \pi(v)$ placed on the appropriate edge of the branching block.

A slight difficulty in Soot is that it is not possible to place instructions directly on an edge since internally Jimple, and hence Shimple, is represented by a simple chain of statements rather than a CFG (Section 3.2.1). Although this is generally not an issue, consider Figure 4.16 where a target block may have one or more sources.

goto target	goto target
if(x > 0) goto target	if(x > 0) goto newtarget
II(X > 0) goto target	
print(x)	x1 = Pi(x)
	print(x1)
target:	
doSomething(x)	newtarget:
	x2 = Pi(x)
	tangat
	target:
	x3 = Phi(x2, x)
	doSomething(x3)

Figure 4.16: Example of a target block with more than one source in original version and Extended Shimple form. We cannot simply prepend a π -function to the target block in Extended Shimple since another statement may reach the block with another context value for **x**.

The if-statement requires placement of two π -functions in Extended Shimple, however as we cannot place these π -functions on the control-flow edges from the statement, we generally place them directly in the destination blocks. In this particular situation however, we cannot simply prepend a π -function to the target since control flow may reach the block from another statement with potentially another context value for \mathbf{x} .

As shown, we resolve this situation in Extended Shimple by creating a new target block containing the π -function and updating the relevant branching statement to use the new target. A ϕ -function may consequently be required in the original target block.

4.3.3 Representation of π -functions

As we have seen, it is often useful in predicated data flow analyses to be able to access the conditional statement as well as the relevant value of the conditional which caused a split. Hence, as a convenience to the Soot developer, we store this information directly in the π -functions in Extended Shimple.

Figure 4.17 is a sample of Extended Shimple code showing π -functions in both an if-statement and switch-block context.

4.3.4 Computing Extended Shimple

In computing Extended Shimple, we had the choice between the eSSA algorithm and the SSI algorithm.

As we had noted in Section 4.1.2, the original approach to computing eSSA form was to first insert π -functions for all variables mentioned in the conditional branch at a split point, and subsequently compute SSA form in the usual manner. The drawbacks of this approach are that nodes may be inserted where they are otherwise not needed, causing excessive variable splits and merges, as well as the fact that potentially interesting nodes for both predicated and backwards flow analysis may not be placed.

Hence, we have chosen to implement the SSI algorithm which allows Extended Shimple to support both predicated analyses as well as sparse backwards flow analyses, and further resolving situations such as that seen in Figure 4.2.

Finally, Extended Shimple eliminates π -functions by treating them as copy statements. π -functions can simply be removed with the corresponding variable uses re(0)if x = 0 goto label0; x1 = Pi(x #0 [false]);print(x1); (1)goto label1; label0: x2 = Pi(x #0 [true]);(2)print(x2); label1: $x3 = Phi(x1 \ \#1, \ x2 \ \#2);$ (3)lookupswitch(y) case 5: goto label2; default: goto label3; label2: y1 = Pi(y #3 [5]);(4)goto label5; label3: y2 = Pi(y #3 [default]);print(y3); (5)label4: $y_3 = Phi(y_1 \# 4, y_2 \# 5);$ return;

Figure 4.17: Sample Extended Shimple code showing a simple if statement followed by an example switch statement. The π -functions include a label indicating the branching statement which caused the split and the value of the branch expression relevant to the branch context. named to the original name, followed by clean up operations such as dead assignment elimination.

4.4 Related Work

It is perhaps no surprise that eSSA form and SSI form are so similar, given that they both postdate and reference work by Johnson and Pingali [JP93] on dependencebased program analysis. In their paper, Johnson and Pingali introduce a structure known as the Dependence Flow Graph (DFG), which is in essence a generalisation of SSA form. The DFG supports forwards flow analysis, like SSA form, and also backwards flow analysis, like eSSA and SSI forms. The DFG introduces *merge* and *switch* operators; the merge operators perform the same function as ϕ -functions while the switch operators are similar to σ -functions and π -functions.

Bodik *et al.* first introduced eSSA form [BGS00] as a means to simplify the formulation of an algorithm for eliminating array-bounds checks on demand in JikesRVM [AAB⁺00]. The additional variable splitting introduced by the form allowed the variable scope constraints to be implicitly represented.

Ananian introduced SSI form [Ana99] as the primary IR for efficient analysis and optimisation in the FLEX compiler infrastructure for Java [Gro]. Singer later performed work to clarify the relationship between SSI and SSA form [Sin], particularly reconciling the algorithm for computing SSI with that for SSA [Sin02].

Chapter 5 Array Shimple

So far in our exposition of SSA form and its variants, we have treated arrays in the same manner that we have treated scalars i.e. we have only considered 'whole' array assignments or allocations while ignoring any 'partial' assignments within an array. In this chapter we will overview a possible approach [CFR⁺91] towards handling arrays such that data flow analysis can be facilitated at the array-element level.

In Section 5.1 of this chapter we present an overview of the issue of dealing with arrays in SSA form and a suitable alternative array notation. Section 5.2 covers the various difficulties of dealing with arrays in Java and introduces Array Shimple. Section 5.3 is a discussion of the possible applications of Array Shimple. Finally, Section 5.4 is an overview of some of the related work on arrays in SSA form.

5.1 Array Notation

As shown in Figure 5.1, our treatment of arrays in Shimple has so far been at a coarse level. While assignment statements of the form array = v affect the SSA renaming of an array variable, assignment statements of the form array[index] = v have not been considered.

Although the previous approach does expose useful information about array allocations, potentially improving analyses such as points-to analysis (Section 3.3.1), the

```
if(bool)
array1 = new int[6]
else
array2 = otherArray
array3 = Phi(array1, array2)
array3[4] = 5
array3[2] = 3
```

Figure 5.1: Shimple example with no special array support. As shown, only 'whole' array assignments are considered for SSA renaming, while assignments to elements within an array are not.

resulting IR might not be considered as representing true SSA form given that multiple element-level assignments may occur to the same array variable hence potentially changing the array's matrix of values without a corresponding name change. This is a familiar indication that potentially useful information is not yet being exposed in the IR.

Examining a statement of the form array[index] = v, it is not clear how one might perform a renaming of the array variable. Simply assigning the array a new name such as in array1[index] = v has no meaning given that array1 refers to a non-existent array. The problem can be solved by a closer examination of arrays.

An array can be considered an abstract data type with a special notation. The two abstract operations that can be performed on an array (assuming it has already been created or allocated) are array element accesses, denoted in Jimple by statements of the form tmp = array[index], as well as array element updates, denoted in Jimple by statements of the form array[index] = data. The array notation tends to obscure the fact that these are two distinct operations although it does have the advantage of being similarly uniform to the syntax for scalar variable accesses and updates.

We can explicitly represent these two distinct single-dimensional array operations

with expressions of the form Access(array, index), which returns the value of the element found in array at index, and Update(array, index, value), which takes array and inserts value at the location index but has no return value.

By a slight modification of the new notation, we can make it more convenient for SSA renaming. Update(array, index, value) can be expressed in functional form as array = Update(array, index, value) where the expression Update(array, index, value) no longer has the side-effect of directly updating array but instead *returns* a *copy* of array with the location at index updated with value.¹ In this new notation, an array update no longer implicitly modifies an existing array, making it more convenient for SSA notation since it allows the possibility of assigning a new name to the resulting array after an assignment.

Figure 5.2 shows the example from Figure 5.1 using the new array notation in SSA form. In the new notation, the array is now split at all the definition points including array element-level assignments.

```
if(bool)
array1 = new int[6]
else
array2 = otherArray
array3 = Phi(array1, array2)
array4 = Update(array3, 4, 5)
array5 = Update(array4, 2, 3)
```

Figure 5.2: SSA example from Figure 5.1 using the new array notation. In this example, any change to the array variable whether a whole or partial assignment is reflected in the IR with a new SSA variable.

¹This is the array syntax proposed by Cytron *et al.* $[CFR^+91]$ and others for data flow analysis on arrays and aggregate structures.

5.2 Implementation of Array Shimple

Once Update and Access statements have been inserted, Array Shimple can be computed by the insertion of ϕ -functions (and π -functions if desired) followed by the SSA renaming algorithm described in Section 2.3.2.

However, we run into several difficulties when attempting to adopt the array notation described, requiring additional analyses as well as certain compromises. We have therefore chosen to first compute Shimple, determine where and how to insert Update/Access statements, and then recompute SSA form by adding any additional ϕ -functions and π -functions required, followed again by the SSA renaming algorithm.

The advantage of our approach of computing Shimple prior to computing Array Shimple is that we can formulate any supporting analyses on the assumption that the IR is in SSA form, allowing us to simplify algorithms, gain precision of results and leverage our previous work. The disadvantage is that we will have to subsequently recompute SSA form, which may be an additional inefficiency since the SSA algorithm is essentially being applied twice. Although we have chosen ease of implementation over efficiency, it would be a straightforward matter to update Array Shimple such that pre-computation of SSA form is not required.

In the rest of the section we detail the major difficulties involved in computing Array Shimple.

5.2.1 Multi-Dimensional Arrays

While arrays in the Java language appear to be multi-dimensional at first glance (e.g. an array access might be of the form array[index1][index2]...[indexn]), Java in fact only supports arrays of arrays [GJS05] and consequently at the bytecode and Jimple levels array accesses and updates are always single dimensional. Hence, although no extension of the Access/Update array syntax is required, array updates may become more expensive for Array Shimple.

Figure 5.3 shows an example of how arrays of arrays are treated in Shimple and Array Shimple. We can see that one write to a 2-dimensional array in Java requires one access and one write in Shimple, whereas Array Shimple requires one access and *two* updates since updates in Array Shimple do not have the side-effect of updating the original array, and therefore any update to the internal array must be propagated to its parent or containing array.

// a[4][5] = 10;	// a[4][5] = 10;
tmp1 = a[4];	tmp1 = Access(a, 4);
tmp1[5] = 10;	$tmp1_1 = Update(tmp1, 5, 10);$
	$a_1 = Update(a, 4, tmp1_1);$
// x = a [0] [1];	// x = a [0] [1];
tmp2 = a [0];	$tmp2 = Access(a_1, 0);$
x = tmp2 [1];	x = Access(tmp2, 1);

Figure 5.3: Code snippet demonstrating how arrays of arrays are updated and accessed in Shimple and Array Shimple. The original Java statements are shown as comments in the code.

In general in Array Shimple, a write to an *n*-dimensional array in Java requires n-1 accesses in order to obtain the relevant innermost array followed by *n* updates in order to propagate the write from the copy of the innermost array to a copy of the outermost array.

An initial pass over the statements in a method allows us to determine the set of top-level or outermost arrays by identifying all local array variables that are either created in the method itself, obtained externally (e.g. as the result of a field or parameter read), or are the result of a simple array local to array local copy (e.g. a0 = a1). More simply stated, any array local that is not the result of an assignment from an array access is considered a top-level array.

In addition to a set of top-level arrays, we also construct a table mapping inner arrays to their parent or containing array (and corresponding index) e.g. a statement of the form a = o[i] results in a table entry from array local a to array reference o[i] which allows us to determine that o is the parent array of a.

Note that in this section we will assume that **a** only points to one internal array,

since the intention is to handle the particular situation that arises when Java's multiarray syntax is translated to the array of arrays form at the bytecode (and hence Jimple/Shimple) level. We will deal with the situation where a local array variable **a** might point to one or more structures in the next sections.

Given these structures, Figure 5.4 illustrates the algorithm we have conceived to appropriately insert array update statements for array writes. Array reads in Shimple are trivially converted to array accesses in Array Shimple by a simple change in syntax, given the direct mapping of the notations.

```
process statement of the form ``a[i] = v``:
replace statement with ``a = Update(a, i, v)``
runner = a
while runner is not an outermost array:
    parent, index = get parent array and index of runner
    append statement ``parent = Update(parent, index, runner)``
    runner = parent
```

Figure 5.4: Algorithm for inserting array update statements for multi-dimensional arrays.

Finally, we note that although Array Shimple appears to make multi-indexed arrays harder to analyse due to the introduction of multiple update statements, this is mainly a consequence of the fact that Java does not support true multi-dimensional arrays. It has been widely noted in the literature $[MMG^+00]$ that this lack of support for multi-dimensional arrays makes multi-indexed arrays harder to analyse and optimise in general. Array Shimple is simply exposing the flow of information – it is up to an analysis to make good use of it.

5.2.2 Fields, Side-effects and Concurrency

The goal of Array Shimple is to provide, as far as possible, the same guarantees for array locals as those provided for scalar locals in Shimple. We will refer to the vector or matrix of values representing an array simply as the value of an array, and we will consider any new assignment to a memory location within an array or to the array itself as a change in the value of the array. For example, a statement of the form a[i] = v changes the value of an array, but a statement of the form a[i].field = v where a is an array of objects will not be considered a change in the value of the array since a still points to the same objects.

Any modification to the value of an array must result in a new local array variable in that static context i.e. once an array local has been created and assigned its value in a static context, that value must not be changed within the context. Given that arrays are objects in Java, there are cases where it is especially hard to guarantee that this "same name same value" aspect [LH96] of SSA form holds for arrays. Recall that Shimple is essentially designed for intra-procedural analysis and as such only tends to represent the information available within a method itself.

If an array was created externally to a method currently being analysed, the array might be modified asynchronously by different threads, or in the presence of method calls with side-effects, a method call might unexpectedly change the value of an array. Similarly, given that arrays are objects in Java and that they can be passed by reference, it is possible that arrays may 'escape' to external fields or methods. Once an array has escaped from a method it too might be modified by other threads or as side-effect of a method call.

Consider the example in Figure 5.5. Normally an analysis might deduce that the value of variable t is the constant 3. Unfortunately, this is an unsafe assumption given that the array escapes to a field, since in the presence of concurrency, the array may well be modified before it is accessed again in the method.

A further problem demonstrated by the example is that it is unclear which array should be assigned to the field in Array Shimple – in the Shimple version, the field is simply assigned \mathbf{a} hence any changes to the field will be reflected in all uses of \mathbf{a} in the method, whereas in Array Shimple \mathbf{a} might be split into different variables and typically only one of them can be assigned to the field [LH96].

In general, arrays that are originally created from outside of a method, or that escape either by being assigned to a field or by being passed as a parameter to

a[4] = 3	a1 = Update(a, 4, 3)
this.field = a	this.field = $a1$
t = a [4]	t = Access(a1, 4)

Figure 5.5: Example where an array object might 'escape' to a field, shown in both Shimple and Array Shimple forms. No safe assumptions can be made about the value of t without additional analysis.

)

a method call, can all be considered 'unsafe' in the sense that array modifications outside of the method being analysed may occur.

As a matter of pragmatism, we have decided that our approach in Array Shimple is to identify all potentially unsafe arrays and leave them unmodified i.e. using the original array syntax. Our reasoning is that the information available is either insufficient or too complex to be represented at the IR level, and even if it were to be represented, it would be unlikely that an analysis would be able to make use of such partial and conditional information in the IR. Hence, an analysis might operate solely on the safe arrays represented in Array Shimple, while making no assumptions about the unsafe ones.

The algorithm we conceived for *conservatively* determining unsafe array locals is fairly straightforward.

Recall that in Chapter 3 we mentioned the availability of Soot's Spark interprocedural points-to analysis [Lho02] as well as our own simplistic SSA-based intraprocedural analysis. Depending on whether Soot is running in whole-program mode or intraprocedural mode, by default Array Shimple makes use of the most accurate points-to information available in order to determine the relevant alias sets for each local variable.

An array local is determined to be unsafe if it is the result of a field read, the result of a method invocation, obtained as a parameter to the current method, potentially aliased to a local already determined to be unsafe, or if the local (or a potential alias of the local) escapes either through a field or method invocation.

Our algorithm for detecting unsafe locals, and hence unsafe array locals, is outlined

```
in Figure 5.6.
assume all locals are initially safe
examine each statement in current method:
    if local in statement is of unsafe origin:
        mark local as unsafe
        mark all locals potentially aliased to local as unsafe
        if local in statement escapes from method scope:
        mark local as unsafe
        mark all locals potentially aliased to local as unsafe
        mark local as unsafe
        mark all locals potentially aliased to local as unsafe
        mark all locals potentially aliased to local as unsafe
        mark all locals potentially aliased to local as unsafe
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        mark all locals potentially aliased to local as unsafe
        mark all locals potentially aliased to local as
```

Figure 5.6: Algorithm to determine all unsafe locals. Unsafe array locals are not translated to Array Shimple form.

5.2.3 Variable Aliasing

Having side-stepped the previous issues, we must now deal with any variable aliasing that may occur amongst the 'safe' array locals.

Consider that the statement a = Update(a, i, v) is *not* identical in meaning to a[i] = v when variable a happens to be aliased to a variable b. Whereas the latter statement will result in b automatically sharing the update, no such update occurs in the first statement since the Update function has no side-effect.

There are three main cases to consider when dealing with the possibility of variable aliasing between two array locals **a** and **b**:

- 1. a and b can be proven to be unaliased variables.
- 2. a and b are *definitely* aliased when both variables are defined.
- 3. a and b may be aliased variables.

The three sets we have enumerated above are not necessarily mutually exclusive. In particular, the may-alias set may by definition contain variables that can be shown to be equivalent (i.e. definitely aliased) when both variables are defined, and the mayalias set may also include variables that might be proven to be definitely non-aliased, since it is computed conservatively.

As we have mentioned, by default in Array Shimple, we obtain may-alias information from Soot's Spark interprocedural points-to analysis if Soot is running in wholeprogram mode, or we use a simplistic intraprocedural points-to analysis on Shimple.² A potential source of definitely-aliased information is from the Global Value Numbering analysis implementation we described in Section 3.3.3 [FKS00], and if necessary we can obtain definitely-different information from simple Soot analyses such as a type-based analysis which might determine that two variables cannot be aliased if they can only hold objects of incompatible types.

Definitely-different information is only used to refine our may-alias information. If two variables are definitely different, the appropriate information can be filtered from the may-alias sets if necessary.

Figure 5.7 illustrates the case where we have definitely-aliased information for variables a and b. We insert a copy statement of the form b = a after an update to a, and prior to the SSA-renaming step for Array Shimple, ensuring that any changes to a will be propagated to new uses of b. In simpler cases, we might simply replace all uses of b with uses of a (or vice-versa) if the definition of a dominates all uses of b (or vice-versa).

Having dealt with definitely-aliased variables, we do not need to reconsider any occurrences of the same in the may-alias sets. In Array Shimple, we simply subtract all sets of definitely-aliased variables from the may-alias information before-hand.

If variable a may-aliases b, then a or b may or may not be pointing to the same object (array) – one can only be certain at runtime. Hence, if a is updated, it is not possible to determine whether b requires a similar update or not before runtime. For this reason, we will introduce the IfAlias construct as shown in Figure $5.8.^3$

²The actual analysis we use is also configurable at the Shimple API level.

³The IfAlias construct is based on the IsAlias construct [CG93] proposed by Cytron and Gersh-

if
$$(a = b)$$

 $a[1] = 4$
 $b = a$
if $(a = b)$
 $af(a = b)$
 $bf(a =$

Figure 5.7: Variables **a** and **b** are known to be aliased in the context shown in this example. The code fragments are shown in Shimple form, intermediate and final Array Shimple forms respectively. We have introduced a new assignment statement in order to propagate any changes made to **a** to new uses of **b**.

// a may alias b// a may alias b
$$a[2] = 5$$
(1) $a1 = Update(a, 2, 5)$ $b1 = IfAlias(b, a, a1)$

Figure 5.8: If **b** is not aliased to **a** at runtime, then IfAlias(**b**, **a**, **a**1) simply returns **b** itself, otherwise it returns **a**1, which is the updated value for **a**.

IfAlias(b, a, a1) is simply a shorthand for a runtime check that b is aliased to a. If it is, then IfAlias returns the newly updated array for a i.e. a1, otherwise it returns b itself. An IfAlias expression is added for any local variable that may be aliased to a local array variable that is being updated.

For simplicity, an analysis on Array Shimple may choose to consider only the first argument to IfAlias while ignoring the other ones e.g. IfAlias(b) (as an abbreviation to IfAlias(b, a, a1)) could be considered simply another SSA node which returns a potentially modified copy of b. Although simpler, this is a conservative approach since with slightly more effort an analysis can obtain more information on the scope of any changes to b.

Internally in Array Shimple, the IfAlias expression stores a link to the variable that might be modified (**b** in this case) and the update statement that might cause the modification – the other variables are extracted from the update statement and made available to the user through the API. The advantage of this approach is that $\frac{1}{1 + \frac{1}{1 + \frac{$

bein.

we do not have to modify the SSA renaming algorithm in order to handle the IfAlias expression as a special case, since any name changes to variables in the update statement are automatically reflected in the IfAlias i.e. the algorithm does not need to know that the second argument in the IfAlias expression is referencing an 'old' variable (**a** in the example) which has been renamed in the update statement (**a1** in the example).

5.2.4 Deconstructing Array Shimple

It suffices to describe how we can translate out of Array Shimple back to Shimple or Extended Shimple, since we already know how to translate out of the latter to Jimple. In particular, we need to convert the Access, Update and IfAlias statements we have introduced to at least their Jimple equivalents.

Access Statements

An expression of the form v = Access(a, i) can be trivially converted to the Jimple statement v = a[i].

Update and IfAlias Statements

Before attempting to eliminate Update and IfAlias statements, we apply a variable packing algorithm that will optimise storage allocation e.g. by using a graph colouring algorithm [Muc97]. Typically we expect to see the results shown in Figure 5.9 after applying variable packing especially if any 'old' variables (a and b in the example) are not subsequently used in the program.

a1 = Update(a, i, v)	a = Update(a, i, v)
b1 = IfAlias(b, a, a1)	b = IfAlias(b, a, a)

Figure 5.9: Array Shimple code before and after applying a variable packing algorithm. If **a** and **b** are not subsequently reused in the program, **a1** and **b1** will be collapsed into the 'old' variable names.

If an Update statement can be transformed as shown in the figure, then it is trivial to convert the Update statement as well as any associated IfAlias statements to Jimple syntax. In this example the Update statement and any associated IfAlias statements can be replaced by the Jimple array statement a[i] = v which will update array a as well as any variables that happen to be aliased to a.

Note that it may happen that the target variable of the IfAlias statement and the first argument to the IfAlias expression are not the same variable, even though the second and third arguments to the expression are the same variable (since these latter arguments are linked to the Update statement). In such cases, a statement such as b1 = IfAlias(b, a, a) could be replaced by a copy statement of the form b1 = b. A subsequent optimisation could eliminate the copy altogether.

Any analysis that performs significant transformations on Array Shimple must take into account that such transformations may come at a cost. Particularly, if Update statements and IfAlias statements cannot be eliminated in the manner described above due to any transformations that would prevent the variable packing algorithm from collapsing relevant variables, Array Shimple will need to replace the statements with equivalent and potentially costly Jimple statements.

It is perhaps worth noting at this point that any dead IfAlias or Update statements can simply be removed to avoid incurring unnecessary performance penalties e.g. any IfAlias (or Update) statements at the end of a method body can be eliminated, since they only apply to 'safe' arrays and hence have no effect. Any of Soot's existing dead code elimination analyses may be applied here.

Figures 5.10 and 5.11 show the mapping from Update and IfAlias statements to the equivalent Jimple code used by Array Shimple.

tmp1 = a.clone()
a1 = (ArrayType) tmp1
a1[i] = v

Figure 5.10: A statement of the form a1 = Update(a, i, v) is replaced by three Jimple statements in the worst case scenario.

```
begin:

if (b == a) goto ifalias

b1 = b

goto exit

ifalias:

b1 = a1

exit:

...
```

Figure 5.11: A statement of the form b1 = IfAlias(b, a, a1) is replaced by an equivalent control structure in the worst case scenario.

As shown, in the worst case array clone operations as well as new control structures may be added to the code. An analysis designer must carefully balance the pros and cons of performing elaborate transformations on Array Shimple.

In general, analyses such as constant propagation or points-to analysis which do not perform significant transformations on the program structure will not have the mentioned undesireable effects.

Multi-Indexed Arrays

Currently we have implemented a simple heuristic approach for eliminating multiple updates that result when writing to a multi-indexed array.

After applying the variable-packing algorithm and eliminating Access, Update and IfAlias statements, we might typically expect to see the code shown in Figure 5.12 if multi-indexed arrays are involved. By identifying typical patterns of redundant stores on the arrays we have processed (i.e. the 'safe' arrays) and eliminating these, we get code that is reasonably similar to the original code, provided no significant transformations have occurred.

 $\begin{array}{ll} tmp1 \ = \ a \ [4]; \\ tmp1 \ [5] \ = \ 10; \\ a \ [4] \ = \ tmp1; \end{array} \\ \begin{array}{ll} tmp1 \ [5] \ = \ 10; \\ tmp1 \ [5] \ = \ 10; \end{array} \\ \end{array}$

Figure 5.12: Resulting code after variable packing and elimination of Access, Update and IfAlias syntax, shown before and after elimination of redundant array store.

5.3 Overview of the Applicability of Array Shimple

Array Shimple introduces additional array variable splitting to the basic SSA form and hence has the potential to expose context information relevant to array elementlevel analysis.

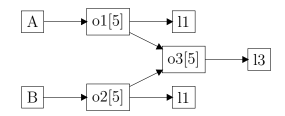
Figures 5.13 and 5.14 illustrate the previously described example on scalar locals (Figures 3.9 and Figure 3.10) but instead using array locals and Array Shimple. This example is nearly identical to the previous one on which it is based, with the exception that array reads and array writes are involved.

if (bool)

$$o[5] = A$$
 [1]
 $l1 = o[5]$
print(l1.type())
else
 $o[5] = B$ [2]
 $l2 = o[5]$
print(l2.type())
 $l3 = o[5]$

Figure 5.13: Points-to example, code and pointer assignment graph. o[5] may point to objects A and B due to the statements [1] and [2]. An analysis on the graph would also conclude that 11, 12, and 13 may also point to A and B.

print(l3.type())



```
o3 = \phi(o1, o2)

l3 = Access(o3, 5)

print(l3.type())
```

Figure 5.14: Points-to example from Figure 5.13 in Array Shimple form. An analysis on the pointer assignment graph can potentially obtain more precise points-to information, here 11 and 12 can only point to objects A and B respectively. Although Array Shimple helps expose more context information for array locals, it is up to the analysis to track the various array elements during data flow analysis. For example, an analysis building the points-to graph for Figure 5.14 would need to create a node for element o1[5] if it is interested in tracking information at that level. Furthermore, the analysis has to consider how to select or merge information in ϕ -functions on arrays if it is interested in computing this information at the arrayelement level.

Clearly, an analysis must be designed to balance the efficiency of an analysis with its precision. Sarkar and Knobe [SK98] consider some of these issues when implementing sparse conditional constant propation for array elements on their variant of Array SSA form [KS98].

Most of the analyses on scalar locals can similarly be formulated to operate at the array element level – however we note that array analysis can certainly get much more complex.

Consider an array access such as a[i] where the array is no longer being accessed by a constant but by a potentially dynamically assigned variable i. One might be able to make deductions about the value of a[i] even without specific information on variable i.

Although the Array Shimple IR does not aid directly in complex array analysis, the use of SSA can still be useful in this context. Consider if one wishes to determine whether array references a[i] and a[j] are equivalent. One test for this would be to test whether indices i and j themselves are equivalent. In Shimple and Array Shimple, one might make use of a Global Value Numbering algorithm as described in Section 3.3.3 in order to determine this information.

Other difficulties involved with analysing arrays are assignments that occur in a loop using variable indices – again, Array Shimple does not provide direct support for such analysis although it may be of use by exposing context information and by virtue of being in SSA form.

Finally, we note that just as Shimple can be extended to Extended Shimple through the introduction of π -functions, similarly Array Shimple can be computed as Extended Array Shimple if so desired.

5.4 Related Work

Cytron *et al.* point out [CFR⁺91] that the Update/Access syntax for accessing arrays is not new and is similar to notation used in previous work [FOW87, Den74] for the analysis of arrays and aggregate structures. The IfAlias syntax we have used to encode may-alias information in Array Shimple is also based on the IsAlias construct proposed by Cytron and Gershbein [CG93].

Knobe and Sarkar develop Array SSA form [KS98], redefining ϕ -functions on array variables such that an expression of the form Phi (a1, a2) is said to perform an arrayelement level merge where a new array is returned with elements taken from a1 and a2 such that the elements are the ones with the most recent 'timestamp' as defined by the 'time' at which they were written to an array. Knobe and Sarkar fully develop the semantics for Array SSA form including additional structures and computations for the IR in order to support analysis and execution at runtime. However, the additional structures introduced by Knobe and Sarkar do not appear to be entirely useful for static analyses, and once removed, Array SSA form appears to be extremely similar to scalar SSA form augmented with the Update/Access notation for arrays. Sarkar and Knobe also develop a sparse constant propagation algorithm of array elements for Array SSA form [SK98]. This algorithm may also be retooled for use on Array Shimple.

Finally, we note that Lapkowski and Hendren [LH96] have confronted some of the same issues we did in designing Array Shimple, including variable aliasing, side-effects and additional issues in languages with pointer support such as C. Lapkowski and Hendren devised extended SSA numbering where instead of computing and representing full SSA form (and any pointer and may alias information), only SSA numbers for each variable were stored as annotations in a secondary structure. The primary SSA number of a variable corresponded to the integer subscript that would normally be generated by the SSA algorithm for renaming the variable, and where necessary, a secondary SSA number was also computed and stored for a variable using points-to and aliasing information. The idea was that if a variable or pointer was changed by a direct assignment, it would have a new primary SSA number, but if the contents of a variable was or might have been changed through aliasing or otherwise, a new secondary SSA number would be generated. Thus extended SSA numbering provided the guarantee that a variable could be assumed to hold the same value in the context of different uses, provided those uses were associated with the same SSA numbers. The advantage of extended SSA numbering is that it sidesteps some of the issues we had to deal with (or avoid altogether) in Array Shimple, the disadvantages are that it only provides some of the advantages of SSA form since it does not represent ϕ -functions and useful data flow information is not represented in the IR.

Chapter 6 Summary and Conclusions

Our aim in this thesis has been to investigate and implement an SSA framework for the Soot compiler toolkit.

SSA form exposes context and flow information of variables in a program, facilitating the implementation of powerful and efficient compiler analyses and optimisations. The basic concept of exposing variable flow information in the IR was found to be extensible. We detailed the key concepts behind SSA form, following the evolution from simple variable splitting to basic SSA form, eSSA form, SSI form, and support for array elements.

In the pursuit of our investigation, we implemented the Shimple framework and presented the 3 IRs we implemented in bottom-up fashion – Simple Shimple, Extended Shimple and Array Shimple. Each IR implementation tended to reuse the work done in the previous implementation. As we progressed from Simple Shimple to Array Shimple, we encountered difficulties that could be solved by well-known algorithms or that needed new approaches specific to the situation. Furthermore, it became evident that many variations and sub-variations of the IRs were possible – consequently, we designed the Shimple framework to be extensible and to promote reusability. Shimple makes use of many existing Soot analyses, automatically benefitting from any improvements of the same, while several of the analyses that were implemented specifically for Shimple are available for general use in Soot.

The existing Soot and Jimple user will be at ease with Shimple. Many existing

analyses and optimisations can reap at least some of the benefits of SSA form with no further modification. By being retooled to take advantage of the new structures introduced by Shimple, Extended Shimple or Array Shimple, analyses can be simplified or made even more powerful. We implemented several analyses and proposed several interesting extensions of our analyses to take advantage of our new IRs, such as a possible extension of points-to analysis to use Extended Shimple form.

Shimple has been available in various incarnations in recent Soot releases and has been successfully used, tested and even improved by several Soot users. The final version described in this thesis will be integrated in an upcoming release of Soot.

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