

# Comprehensive Path-sensitive Data-flow Analysis

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## ABSTRACT

Data-flow analysis is an integral part of any aggressive optimizing compiler. We propose a framework for improving the precision of data-flow analysis in the presence of complex control-flow. We initially perform data-flow analysis to determine those control-flow merges which cause the loss in data-flow analysis precision. The control-flow graph of the program is then restructured such that performing data-flow analysis on the resulting restructured graph gives more precise results. The proposed framework is both simple, involving the familiar notion of product automata, and also general, since it is applicable to any forward data-flow analysis. Apart from proving that our restructuring process is correct, we also show that restructuring is effective in that it necessarily leads to more optimization opportunities. Furthermore, the framework handles the trade-off between the increase in data-flow precision and the code size increase inherent in the restructuring. We show that determining an optimal restructuring is NP-hard, and propose and evaluate a greedy strategy. The framework has been implemented in the Scale research compiler, and instantiated for the specific problem of Constant Propagation. On the SPECINT 2000 benchmark suite we observe an average speedup of 4% in the running times over Wegman-Zadeck conditional constant propagation algorithm and 2% over a purely path profile guided approach.

## Categories and Subject Descriptors

D.3.4 [PROGRAMMING LANGUAGES]: Processors—Code generation; Compilers; Optimization

## General Terms

Algorithms, Languages, Performance, Theory

## Keywords

Code Duplication, Data-flow analysis, Destructive Merge, Path-sensitive, Precision, Restructuring, Split Graph

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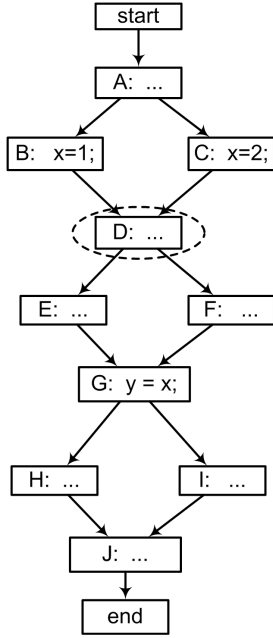
## 1. INTRODUCTION

It is becoming increasingly difficult to get performance improvement using compiler optimizations, as epitomized by Proebsting's Law [16]. But developers still want that extra 5-15% improvement in the running times of their applications, and the compiler optimizations are a safer alternative to manual optimizations carried out by developers which might introduce errors [17].

Information gathered using data-flow analysis is used by an aggressive optimizing compiler to perform code optimizations such as constant propagation, dead-code elimination, common sub-expression elimination. Imprecision in data-flow analysis leads to a reduction in optimization opportunities. The loss of data-flow precision occurs due to the approximation or merging of differing data-flow facts along incoming edges of a control-flow merge. In this paper, we present a new framework to overcome this imprecision. We initially perform data-flow analysis to determine those control-flow merges that cause the loss in data-flow analysis precision, which we call *Destructive Merges*. The control-flow graph (CFG) of the program is then restructured so that destructive merges are eliminated. Thus, performing data-flow analysis on the resulting restructured graph gives more precise results. This leads to more optimization opportunities in the restructured CFG.

The framework presented in this paper is simple and clean, and uses the familiar notion of product automata [11] to carry out the restructuring transformation. Further, the framework is general since it can be instantiated to any forward data-flow analysis. This clean formulation allows us to show that the restructuring is correct, in the sense that the original and restructured CFGs are equivalent. Also, we prove that the restructuring is effective in that it necessarily leads to more optimization opportunities in the restructured CFG. The restructuring inherently entails an increase in code size due to code duplication i.e., the increase in precision comes at the cost of an increase in code size. Furthermore, we show that determining an optimal restructuring is NP-Hard and, hence, propose a greedy heuristic for restructuring. Our framework explicitly handles the trade-off between precision and code size increase and also makes use of low-cost basic-block profile information.

We have implemented the proposed framework in the Scale research compiler [19] and instantiate it with the specific problem of Constant Propagation. We compare our technique with the Wegman-Zadeck conditional constant propagation algorithm [23](*Base*) and with the Hot Path Graph approach (*HPG*) [1] which is a purely path profile-guided



**Figure 1:** In program  $P$ , node  $D$  is a destructive merge. Use of variable  $x$  cannot be replaced by a constant at node  $G$ .

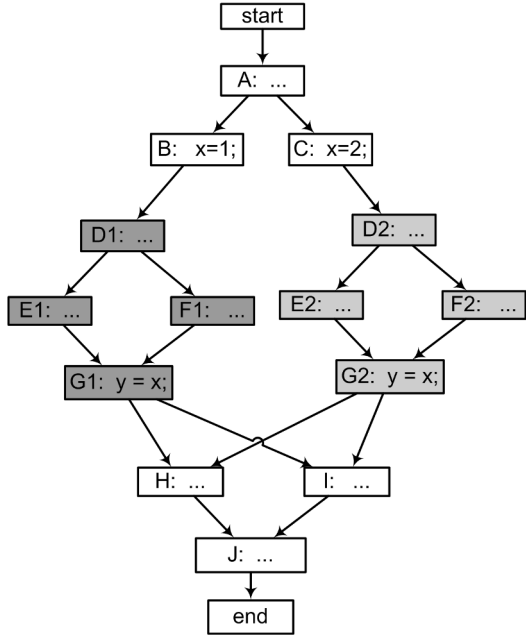
restructuring technique. On the SPECINT 2000 benchmark suite [20], our technique exposes, on an average, 3.5 times more dynamic constants as compared to the HPG technique. Further, we observe an average speedup of 4% in the running times over Base and 2% over the HPG technique.

The rest of the paper is structured as follows. We first illustrate our technique using a simple example. We then go on to present the general framework. In Section 3, we describe the method to determine which nodes of the CFG are to be duplicated for restructuring. Section 4 discusses the actual restructuring algorithm. In this section, we demonstrate the correctness and efficacy of our approach. We then discuss the trade-off between the increase in precision and increase in code-size in Section 5. Section 6 discusses the related work and compares our technique to the HPG technique [1]. This is followed by the experimental results in Section 7, after which we conclude.

## 2. OVERVIEW

We illustrate our technique using the problem of Constant Propagation. Consider the program  $P$  in Figure 1. There are two definitions of variable  $x$  at nodes  $B$  and  $C$ , while node  $G$  contains a use of variable  $x$ . The remaining nodes do not define or use variables  $x$  or  $y$ . We see that node  $G$  cannot be optimized since the use of variable  $x$  cannot be replaced by a constant. Our technique would restructure the program  $P$  by duplicating node  $D$ ,  $E$ ,  $F$  and  $G$  to obtain the Split Graph  $P'$ , shown in Figure 2. After this restructuring, the uses of variable  $x$  at nodes  $G1$  and  $G2$  can now be replaced with constants 1 and 2 respectively.

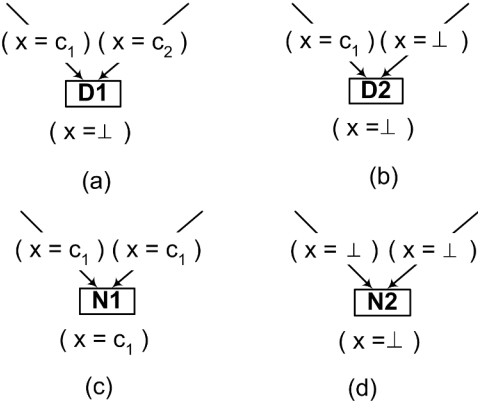
In order to understand how to obtain the program  $P'$ , we first look at control-flow merge  $D$  in program  $P$ . At node  $D$ , the incoming data-flow facts  $\{x = 1\}$  and  $\{x = 2\}$  are merged to get the data-flow fact  $\{x = nc\}$ , where  $nc$



**Figure 2:** In Split Graph  $P'$ , uses of variable  $x$  at  $G1$  and  $G2$  can now be replaced by constants 1 and 2 respectively.

stands for not-constant. We call such a control-flow merge where data-flow approximation leads to loss of precision a **Destructive Merge**. Due to this loss of precision at node  $D$ , only the data-flow fact  $\{x = nc\}$  reaches node  $G$ , and hence, node  $G$  cannot be optimized. Further, we also notice that if (somehow) the data-flow fact  $\{x = 1\}$  were to hold at node  $D$ , then the use of variable  $x$  at node  $G$  can be replaced by the constant 1. Thus, we see that in order to optimize node  $G$ , it is useful for data-flow fact  $\{x = 1\}$  to hold at node  $D$ . This also holds for the data-flow fact  $\{x = 2\}$ . On the other hand, it is not useful for the data-flow fact  $\{x = nc\}$  to hold at node  $D$ . Hence, in program  $P'$  in Figure 2 node  $D$  is duplicated and at the copies  $D1$  and  $D2$  of  $D$ , the data-flow facts  $\{x = 1\}$  and  $\{x = 2\}$  hold respectively. It can be seen that nodes  $E$ ,  $F$  and  $G$  also need to be duplicated to preserve the data-flow facts  $\{x = 1\}$  or  $\{x = 2\}$  at nodes  $G1$  and  $G2$  respectively. Further, nodes  $H$ ,  $I$  and  $J$  cannot be optimized even if the data-flow facts  $\{x = 1\}$  or  $\{x = 2\}$  hold at node  $D$ . Thus, these nodes are not duplicated in Program  $P'$  in Figure 2. We say that these nodes are not influenced by the destructive merge at node  $D$ . The nodes  $D$ ,  $E$ ,  $F$ , and  $G$  are called the *Region of Influence* for the destructive merge  $D$  and are exactly those nodes which have multiple copies in program  $P'$ .

In program  $P$ , there are four paths which reach node  $G$ . Along the two paths going through edge  $(B, D)$ , data-flow fact  $\{x = 1\}$  holds. Similarly, along the two paths going through edge  $(C, D)$ , data-flow fact  $\{x = 2\}$  holds. Hence, program  $P$  is restructured in such way these two sets of paths are separated and the differing data-flow facts along them do not merge in program  $P'$ . We use the concept of product automaton [11] to carry out this restructuring. The increase in the number of uses which can be replaced by constants comes at the cost of increase in code size.



**Figure 3:** For the problem of Constant Propagation, (a)-(b) show two destructive merges and (c)-(d) illustrate two control-flow merges which are not destructive.  $c_1$  and  $c_2$  are two differing constants, while  $\perp$  represents not-constant.

Though not illustrated in this pedantic example, our restructuring technique is also applicable when there are multiple destructive merges which we wish to eliminate and control-flow structures such as loops.

### 3. COMPUTING THE REGION OF INFLUENCE

Central to our approach, is the notion of a destructive merge. Intuitively, a destructive merge is a control-flow merge where data-flow analysis loses precision. The notion of precision is governed by the partial-order of the lattice of the data-flow problem.

Let  $\text{pred}(n)$  denotes the set of control-flow predecessors of a node  $n$  in the CFG,  $\text{in\_dff}(n)$  and  $\text{out\_dff}(n)$  denotes the data-flow facts holding true at the beginning and end of a node  $n$  in the fixed-point solution respectively, and  $\prec$  is the partial order of the lattice in the data-flow problem.

**DEFINITION 1. (DESTRUCTIVE MERGE)** For a given forward data-flow problem and its corresponding solution, a control-flow merge  $m$  is said to be a **Destructive Merge** if  $\exists p \in \text{pred}(m)$  s.t.  $\text{in\_dff}(m) \prec \text{out\_dff}(p)$ .

**DEFINITION 2. (DESTROYED DATA-FLOW FACTS)** Given a destructive merge  $m$ , a data-flow fact  $d$  is said to be a **Destroyed Data-flow Fact**, i.e.  $d \in \text{destroyed\_dff}(m)$ , iff  $d \in \text{out\_dff}(p)$ , where node  $p \in \text{pred}(m)$ , and  $\text{in\_dff}(m) \prec d$ .

*Example:* For the problem of Constant Propagation, Figure 3 illustrates the different scenarios possible at a control-flow merge. Nodes  $D_1$  and  $D_2$  are destructive, while nodes  $N_1$  and  $N_2$  are not.

More specifically, in Figure 1 node  $D$  is a destructive merge since  $\text{in\_dff}(D) = \{\perp\}$ , while  $\text{out\_dff}(B) = \{x = 1\}$ . Further,

$$\text{destroyed\_dff}(m) = \{x = 1, x = 2\}.$$

□

Our restructuring targets such destructive merges. In the rest of this section, we determine which nodes are influenced by the loss of precision lost at a destructive merge and which

nodes need to be duplicated in order to eliminate the effects of a destructive merge.

#### 3.1 Single Destructive Merge

The following definitions assume that we are given a CFG and a data-flow problem and a corresponding solution.

**DEFINITION 3. (USEFUL DATA-FLOW FACTS)** For a given node  $n$ , a data-flow fact is said to be a **Useful Data-flow Fact**, i.e.  $d \in \text{useful\_dff}(n)$ , iff data-flow fact  $d$  holding true at node  $n$  implies that the node  $n$  can be optimized.

The above definition implicitly captures the interaction between a data-flow analysis and the client optimization. It allows us to abstract out the details of this relation by providing an oracle which knows which data-flow facts enable the optimization for a particular program statement. We can now generalise this notion of useful data-flow facts.

**DEFINITION 4. (USEFUL DATA-FLOW FACTS)** Given nodes  $m$  and  $n$ , a data-flow fact is said to be **Useful Data-flow Fact for node  $n$  at node  $m$** , i.e.  $d \in \text{useful\_dff}(m, n)$ , iff data-flow fact  $d$  holding true at node  $m$  implies that the node  $n$  can be optimized.

It is easy to see that Definition 3 is a special case of Definition 4 with node  $m$  being the same node  $n$ .

*Example:* Consider the program in Figure 1,

$$\text{useful\_dff}(G) = \{\dots, x = -1, x = 0, x = 1, x = 2, \dots\},$$

$$\text{useful\_dff}(D, G) = \{\dots, x = -1, x = 0, x = 1, x = 2, \dots\}.$$

This is due to the fact that if the value of variable  $x$  were to be a constant at node  $D$ , then it would remain a constant at node  $G$ . This would enable us to optimize node  $G$  by replacing the use of variable  $x$  with a constant. □

**DEFINITION 5. (INFLUENCED NODES)** Given a destructive merge  $m$ , we say a node  $n$  is influenced by the destructive merge  $m$ , i.e.  $n \in \text{influenced\_nodes}(m)$ , iff

$$\text{destroyed\_dff}(m) \cap \text{useful\_dff}(m, n) \neq \emptyset.$$

In other words, a node  $n$  is *influenced* by a destructive merge  $m$  when, if one of the data-flow facts  $d$  destroyed by node  $m$  were to hold true at node  $n$ , then node  $n$  could be optimized. Intuitively, a node is *influenced* by a destructive merge if eliminating the destructive merge can enable the optimization of the node. Further, these are the only nodes which can be optimized if the destructive merge is eliminated.

**DEFINITION 6. (REVIVAL DATA-FLOW FACTS)** Given a destructive merge  $m$ , a data-flow fact is said to be a **Revival Data-flow Fact**, i.e.  $d \in \text{revival\_dff}(m)$ , iff there exists a node  $n \in \text{influenced\_nodes}(m)$  such that

$$d \in \text{destroyed\_dff}(m) \cap \text{useful\_dff}(m, n).$$

The Revival data-flow facts denote those data-flow facts which if they would hold true at the destructive merge  $m$  would enable the optimization of some influenced node.

Example: Consider Figure 1,

$$\text{influenced\_nodes}(m) = \{G\},$$

$$\text{destroyed\_dff}(m) = \{x = 1, x = 2\},$$

$$\text{useful\_dff}(D, G) = \{\dots, x = -1, x = 0, x = 1, x = 2, \dots\},$$

$$\text{revival\_dff}(m) = \{x = 1, x = 2\}.$$

□

Since the influenced nodes are the only nodes which can be optimized by eliminating the destructive merge, we would like to determine the largest such set of nodes. Unfortunately, for the specific problem of Constant Propagation determining whether a node  $n$  belongs to  $\text{influenced\_nodes}(m)$  for a destructive merge  $m$  is undecidable in general.

**THEOREM 1.** For the problem of Constant Propagation, given a node  $n$  and destructive merge  $m$  determining if  $n \in \text{influenced\_nodes}(m)$  is undecidable.

**PROOF.** (Sketch) The proof follows from the undecidability of constant propagation for programs with loops even if all branches are considered to be non-deterministic [14, 9, 18]. The reduction is based on the Post correspondence problem [11]. We show that the node  $n \in \text{influenced\_nodes}(m)$  iff the Post correspondence problem is not solvable. □

As we see in the Section 4, our restructuring is guaranteed to improve data-flow precision and optimize the nodes which are influenced by the destructive merge. The above theorem implies that we cannot determine all nodes which are influenced by a destructive merge. If we include a node which is not actually influenced by the destructive merge in  $\text{influenced\_nodes}(m)$ , then the restructuring will not enable any optimizations for that node. Thus, the corresponding increase in code size will be unnecessary. Hence, in practice, we under-approximate the set of influenced nodes. In particular, for Constant Propagation, if the data-flow fact for variable  $x$  is destroyed at merge node  $m$ , then the uses which are reachable only along paths from node  $m$  which do not contain any definitions of variable  $x$  are influenced by the destructive merge  $m$ . This set is an under-approximation and there might be nodes which are influenced by the destructive merge  $m$  and not be included in the set  $\text{influenced\_nodes}(m)$  which is actually determined. Note, this under-approximation is orthogonal to our overall approach.

Having determined which nodes are influenced, we now find those nodes which need to be duplicated in order to eliminate the effects of a destructive merge. Let  $\text{reachable}_l$  denotes the set of nodes reachable from node  $l$  in the CFG.

**DEFINITION 7. (REGION OF INFLUENCE)** Given a destructive merge  $m$ , a node  $n$  is said be in the Region of Influence, i.e.  $n \in \text{RoI}(m)$ , iff  $n \in \text{reachable}_n$  and there exists a node  $u \in \text{influenced}_n$  and  $u \in \text{reachable}_n$ .

Figure 4 shows a schematic diagram showing  $\text{RoI}(m)$  for the destructive merge  $m$ . Node  $n$  belongs to the  $\text{RoI}(m)$  since node  $n$  is reachable from node  $m$  and node  $u_4 \in \text{influenced}_n$  is reachable from node  $n$ .

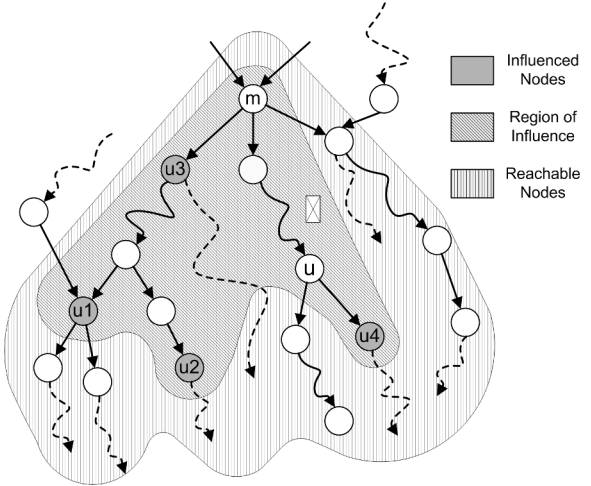


Figure 4: Schematic diagram showing the Region of Influence for a destructive merge  $m$ .

Having determined the set of influenced nodes, the Region of Influence consists of those nodes which are sufficient and necessary to be duplicated in order to improve precision and optimize the influenced nodes.

**Example:** In Figure 1, node  $D$  is a destructive merge.

$$\text{influenced\_nodes}(D) = \{G\},$$

$$\text{reachable}_n(D) = \{D, E, F, G, H, I, J\},$$

$$G \in \text{reachable}_n(D), G \in \text{reachable}_n(E),$$

$$G \in \text{reachable}_n(F), G \in \text{reachable}_n(G),$$

$$\text{RoI}(m) = \{D, E, F, G\}.$$

□

## 3.2 Multiple Destructive Merges

In the previous section, we dealt with the situation where we were given a single destructive merge to eliminate. In practice, we will have a set of destructive merges  $\mathcal{M} = \{m_1, m_2, \dots, m_k\}$  which are to be eliminated. In this section, we extend the concepts of the previous section to handle multiple destructive merges.

**DEFINITION 8. (INFLUENCED NODES)** Given a set of destructive merges  $\mathcal{M} = \{m_1, m_2, \dots, m_k\}$ , the set of Influenced Nodes is defined as

$$\text{influenced}_n(\mathcal{M}) = \bigcup_{m \in \mathcal{M}} \text{influenced}_n(m)$$

**DEFINITION 9. (REGION OF INFLUENCE)** Given a set of destructive merges  $\mathcal{M} = \{m_1, m_2, \dots, m_k\}$ , the Region of Influence corresponding to  $\mathcal{M}$  is defined as

$$\text{RoI}(\mathcal{M}) = \bigcup_{m \in \mathcal{M}} \text{RoI}(m)$$

Extending the definitions to handle multiple destructive merges is straight forward. A node  $n$  belongs to  $\text{influenced}_n(\mathcal{M})$  iff it is influenced by at least one destructive merge  $m \in \mathcal{M}$ . Similarly, a node  $n$  belongs to  $\text{RoI}(\mathcal{M})$  iff it belongs to the Region of Influence of at least one destructive merge  $m \in \mathcal{M}$ .

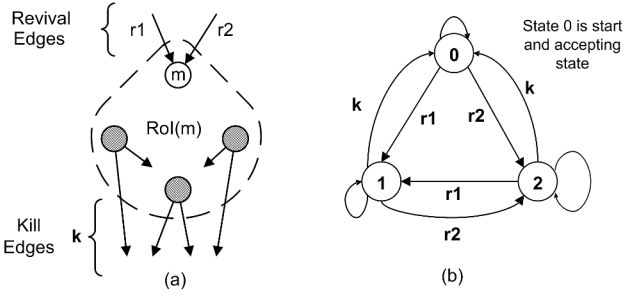


Figure 5: (a) Schematic diagram of the Region of Influence for a destructive merge  $m$  showing the Revival and Kill edges. (b) The corresponding Split Automaton  $A_m$ .

## 4. THE CFG RESTRUCTURING

In this section, we discuss the actual restructuring algorithm. There are two main constraints which such a restructuring transformation should satisfy.

- **Equivalence.** The original and transformed programs should be equivalent.
- **Efficacy.** The transformation should guarantee that the improvement in data-flow precision leads to optimization opportunities.

We explain a clean and simple technique which makes use of familiar notions of product automaton [11], and show that indeed our restructuring satisfies both the constraints. As before, we first restrict ourselves to a single destructive merge, and then generalise this to multiple destructive merges.

### 4.1 Single Destructive Merge

Corresponding to each destructive merge, we construct a **Split Automaton** which will then be used to restructure the CFG and eliminate the effects of the destructive merge.

**DEFINITION 10. (KILL EDGES)** Given a Region of Influence for a destructive merge  $m$ , an edge  $e = (u, v)$  is a Kill Edge, i.e.  $e \in \text{kill\_edges}(m)$ , iff  $u \in \text{RoI}(m)$  and  $v \notin \text{RoI}(m)$ .

In other words, Kill Edges are those edges whose source node is in the Region of Influence and target node is not in the Region of Influence for a destructive merge  $m$ .

**DEFINITION 11. (REVIVAL EDGES)** Given a Region of Influence for a destructive merge  $m$ , an edge  $e = (u, m)$  is said to be a Revival Edge, i.e.  $e \in \text{revival\_edges}(m)$ , iff

$$\text{out\_dff}(u) \in \text{revival\_dff}(m).$$

In other words, Revival Edges are those incoming edges of the destructive merge which correspond to Revival Data-flow facts. Further, let  $d_1, d_2, \dots, d_k$  be the  $k$  distinct Revival Data-flow facts for destructive merge  $m$ . We can partition the incoming edges of destructive merge  $m$  into  $k+1$  equivalence classes  $R_0, R_1, \dots, R_k$ . An edge  $(u, m) \in R_i$ ,  $1 \leq i \leq k$  if  $\text{out\_dff}(u) = d_i$ , and  $(u, m) \in R_0$  otherwise.

Figure 5(a) shows a schematic diagram illustrating the above concepts.

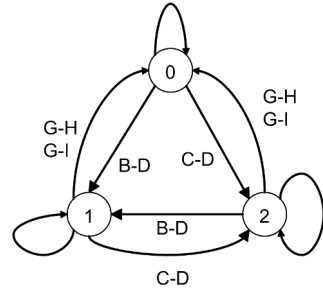


Figure 6: The Split Automaton  $A_D$  corresponding to destructive merge  $D$  in Figure 1.

*Example:* In Figure 1,

$$\text{out\_dff}(B) = \{x = 1\}, \text{out\_dff}(C) = \{x = 2\},$$

$$\text{revival\_dff}(m) = \{x = 1, x = 2\},$$

$$\text{revival\_edges}(m) = \{(B, D), (C, D)\},$$

$$\text{RoI}(m) = \{D, E, F, G\},$$

$$\text{kill\_edges}(m) = \{(G, H), (G, I)\}.$$

□

Armed with the above concepts we are now ready to define the Split Automaton.

**DEFINITION 12. (SPLIT AUTOMATON)** The Split Automaton  $A_m$  corresponding to a destructive merge  $m$  is a finite-state automaton defined as:

- the input alphabet  $\Sigma = E$ , the set of all edges in the CFG.
- a set of  $k+1$  states  $Q = \{s_0, s_1, s_2, \dots, s_k\}$ .
- $s_0 \in Q$  is the initial and accepting state.
- the transition function  $\delta : Q \times \Sigma \rightarrow Q$  defined as
  - $(s_i, e) \rightarrow s_j, e \in R_j$  (Revival Transitions)
  - $(s_i, e) \rightarrow s_0, e \in \text{kill\_edges}(m)$  (Kill Transitions)
  - $(s_i, e) \rightarrow s_i$ , otherwise (No Transition)

Intuitively, state  $s_i$ ,  $1 \leq i \leq k$ , corresponds to the Revival Data-flow fact  $d_i$  and whenever an edge  $e \in R_i$  is seen, the Split Automaton makes a transition to state  $s_i$ . We call this the Revival Transitions. Further, whenever a Kill edge is seen, the Split Automaton transitions to state  $s_0$ . We call these the Kill Transitions. In all other cases, the automaton remains in the same state, and makes no transitions.

*Example:* Figure 5(b) shows a schematic diagram with the Split Automaton corresponding to destructive merge  $m$  in Figure 5(a).

Figure 6 shows the Split Automaton corresponding to destructive merge  $D$  in Figure 1. Intuitively, in the Split Automaton the transitions from states 0 and 2 to state 1 revive the data-flow fact  $\{x = 1\}$  and stop it from being destroyed due to a merge. The transition from state 1 to state 0 finally kills this data-flow fact. Similarly, the transitions from states 0 and 1 to state 2 revive the data-flow fact  $\{x = 2\}$ . The transition from state 2 to state 0 finally kills this data-flow fact. The self-loops represent no transitions. □

A CFG can be viewed as a finite-state automaton with nodes of the CFG corresponding to the state of the automaton, and the edges defining the alphabet as well as the transition function. The entry and exit node of the CFG correspond to the start and accepting states in the automaton respectively.

**DEFINITION 13. (SPLIT GRAPH)** Given a CFG  $P$  and a Split Automaton  $A_m$ , we define the Split Graph  $S$  to be  $S = P \times A_m$ , where  $\times$  is the product automaton operator [11].

Each node  $n$  in the Region of Influence of node  $m$  in  $P$  will have multiple copies  $n_i$  in the Split Graph  $S$  corresponding to the states in the Split Automaton.

*Example:* The Split Graph  $P'$  in Figure 2 is obtained by performing the product of the CFG  $P$  in Figure 1 and the Split Automaton  $A_D$  in Figure 6.  $\square$

**THEOREM 2.** The Split Graph  $S$  is equivalent to the original CFG  $P$ , where  $S = P \times A_m$ .

**PROOF.** (Sketch) Viewing CFG as finite-state automaton, we say that the two CFGs  $G_1$  and  $G_2$  are *equivalent* if the languages accepted by  $G_1$  is equal to language accepted  $G_2$ , i.e.,  $L(G_1) = L(G_2)$ . Such a notion of equivalence suffices since the restructuring only duplicates nodes and does not change the semantics of the individual nodes. Since  $S$  is the product of  $P$  and  $A_m$ , we can say that  $L(S) = L(P) \cap L(A_m)$  [11]. Further, we can show that  $L(P) \subset L(A_m)$ . Intuitively,  $A_m$  is a control-flow abstraction of  $P$  and accepts more words. Thus, it follows that  $L(S) = L(P)$ .  $\square$

Thus, using simple concepts from automaton theory [11], we are able to prove the correctness of our restructuring transformation. Next, we prove the efficacy of the restructuring.

**THEOREM 3.** If node  $n \in \text{influenced\_nodes}(m)$  in CFG  $P$ , then in the Split Graph  $S$ , node  $n_i, i \neq 0$  can be optimized, where  $S = P \times A_m$ .

**PROOF.** (Sketch) Let  $d_i$  be a Revival Data-flow fact and  $R_i$  be the corresponding equivalence class of edges. Since  $n \in \text{influenced\_nodes}(m)$ , by definition, if  $d_i$  (somehow) holds true at node  $m$ , then node  $n$  can be optimized in CFG  $P$ . The theorem proceeds in two steps. We first show that data-flow fact  $d_i$  holds true at the start of node  $m_i$  in the Split Graph  $S$ . Then, we show that this data-flow fact does not merge with other differing data-flow facts. This is due to the nature of Revival Transitions.  $\square$

## 4.2 Multiple Destructive Merges

We now consider eliminating multiple destructive merges. Consider the set of destructive merges  $\mathcal{M} = \{m_1, m_2, \dots, m_k\}$  which are to be eliminated. Let  $A_1, A_2, \dots, A_k$  be the corresponding Split Automata.

**DEFINITION 14. (SPLIT GRAPH)** Given a CFG  $P$  and a set of Split Automata  $A_1, A_2, \dots, A_k$ , we define the Split Graph  $S$  to be  $S = P \times A_1 \times A_2 \times \dots \times A_k$ , where  $\times$  is the product automaton operator [11].

The correctness and efficacy results also hold in this general setting. It is interesting to note that we placed no restriction of the nature of the CFG. Thus, our restructuring can handle programs with complex loop structures.

## 5. PRECISION-CODE SIZE TRADE-OFF

The discussion till now has not dealt with the trade-off between the data-flow precision achieved and the increase in the code size due to the restructuring inherent to this approach. We have seen that the benefit of eliminating a destructive merge  $m$  is related to the number of nodes influenced by node  $m$ . Also, the size of the *Region of Influence* of the destructive merge determines the resulting cost in terms of increase code size. Thus, this trade-off depends directly on the set of destructive merges we choose to eliminate to form the Split Graph. In this section, we will prove that the problem of picking the best set of destructive merges which maximize the benefit in terms of data-flow precision, for a given cost in terms of code size, is *NP-Hard*. We then proceed to describe certain heuristics for the same.

**DEFINITION 15.** The problem *SPLIT* is defined by the triple  $(P, \mathcal{A}, C)$  where:

- $P$  is a program,
- $\mathcal{A}$  is a set of split automata corresponding to the various destructive merges, and
- $C$  is maximum increase in code size that is permitted.

A solution to *SPLIT* is a subset  $B$  of  $\mathcal{A}$  such that applying  $B$  to the program  $P$  does not increase the code-size by more than  $C$  and which maximizes the number of influenced nodes which can be optimized in the resulting program  $P'$ .

**THEOREM 4.** *SPLIT* is *NP-Hard*.

**PROOF.** We shall reduce *KNAPSACK* [15] to it. We are given a set  $\mathcal{I}$  of  $n$  items, each item  $i$  having a specific weight  $w_i$  and a profit  $p_i$ . The goal of *KNAPSACK* is to pick a subset  $J \subseteq \mathcal{I}$  of the items so as to maximize the total profit subject to the condition that the total weight is less than a specified weight  $W$ .

We will only give a brief outline of the proof due to lack of space. Intuitively, in our reduction, picking an item  $i$  in *KNAPSACK* will correspond to selecting an split automaton in the solution of *SPLIT*. Thus, we construct a program  $P$ , in which for each item  $i$  in *KNAPSACK* there exists a destructive merge  $D_i$  and a split automaton  $a_i$  so that  $|\text{influenced\_nodes}(D_i)| = p_i$  and  $|\text{RoI}(D_i)| = w_i$ . Further, the profits and costs of items in *KNAPSACK* are independent of each other i.e. the cost of picking an item  $i$  does not depend on whether item  $j$  has been placed in the knapsack. To ensure this the constructed regions are such that  $\text{RoI}(D_i) \cap \text{RoI}(D_j) = \emptyset, i \neq j$ . The constraint of the total weight  $W$  of the knapsack is mapped to the increase in code size  $C$  which we are allowed in *SPLIT*. It can be shown that split automaton  $a_i$  is in the optimal solution of *SPLIT* if and only if item  $i$  is selected in the optimal solution of *KNAPSACK*.  $\square$

It is interesting to note that this hardness result does not rely on the complexity of the underlying data-flow analysis used, since we are already given the set of influenced nodes and the Region of Influence for each destructive merge. Further, the program  $P$  does not even contain any loops, and is acyclic. Thus, restricting the problem *SPLIT* any further does not result in a computationally tractable problem.

This lead us to device an aggressive greedy heuristic to solve this problem. Our approach is based on estimating

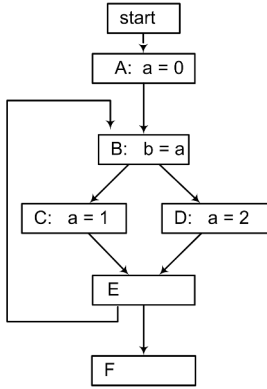


Figure 7: Node  $E$  is a destructive merge. Use of variable  $a$  at node  $B$  cannot be replaced with a constant.

Ball-Larus Acyclic Path	Frequency
$A \rightarrow B \rightarrow C \rightarrow E$	10
$B \rightarrow C \rightarrow E$	60
$B \rightarrow D \rightarrow E$	20
$B \rightarrow D \rightarrow E \rightarrow F$	10

Table 1: A path profile for the example in Figure 7. The frequency of the acyclic path denotes the number of times it was taken at run-time.

the benefit obtained and cost incurred by eliminating a destructive merge. In the absence of profile information, we define the *fitness* of a destructive merge  $m$  to be

$$\text{fitness}(m) = |\text{influenced\_nodes}(m)| / |\text{RoI}(m)|.$$

Otherwise, we can make use of a low-cost basic-block profile to estimate the potential run-time benefit of eliminating a destructive merge. Let  $\text{count}(m)$  be the number of times the destructive merge was executed in the profile run. We now define the fitness to be

$$\text{fitness}(m) = \text{count}(m) * |\text{influenced\_nodes}(m)| / |\text{RoI}(m)|.$$

In this way, frequently executed destructive merges are more likely to be eliminated, and our approach can concentrate on the hot regions of code. Finally, we choose the  $k$  fittest destructive merges to be eliminated. It should be noted that while this heuristic method does not guarantee that the code size increase is within some bound ( $C$ ), it ensures that the code growth is not unbounded and eliminating two fittest destructive merges per function works well in practice as seen in Section 7.

## 6. RELATED WORK

### 6.1 Hot Path Graph Approach

An earlier proposal by Ammons and Larus [1] uses an acyclic path profile to try and improve the precision of the data-flow solution along hot paths. The approach consists of first using a Ball-Larus path profile [2] to determine the hot acyclic paths in the program. The next step in [1] consists of constructing a new CFG, called the *Hot Path Graph*

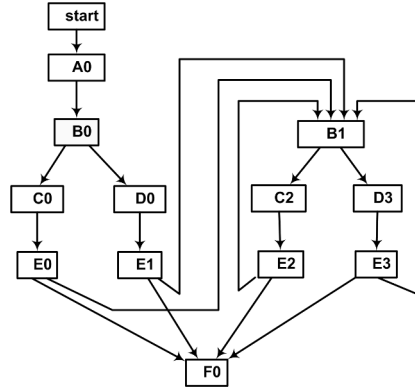


Figure 8: The *Hot Path Graph* corresponding to program  $P_1$ . Node  $B_1$  is a destructive merge, and use of variable  $a$  still cannot be replaced by a constant.

(HPG), in which each hot path is duplicated. The duplication eliminates control-flow merges along hot paths. The assumption being that this will alone will improve precision of data-flow analysis on the hot paths.

Consider the example code in Figure 7. Assume a path profile as shown in Table 1. Figure 8 shows the resulting HPG constructed assuming 100% coverage i.e. all taken paths are considered. Notice that in the HPG there are no control-flow merges along any of the acyclic paths listed in Table 1. For example, the two overlapping acyclic paths  $B \rightarrow C \rightarrow E$  and  $B \rightarrow D \rightarrow E$  in Figure 7 are separated into two separate paths  $B_1 \rightarrow C_2 \rightarrow E_2$  and  $B_1 \rightarrow D_3 \rightarrow E_3$  in the HPG. After performing conventional data-flow analysis on the HPG, the use of the variable  $a$  at node  $B_0$  can be replaced by the constant 0. However, the restructuring failed to optimize the two hot paths  $B \rightarrow C \rightarrow E$  and  $B \rightarrow D \rightarrow E$ , and could not replace the use at node  $B_1$  with a constant value. The destructive merge  $E$  in the original CFG is removed in the HPG by duplicating code and creating two copies,  $E_2$  and  $E_3$ . But the effect of the destructive merge has shifted to node  $B_1$ , which is now a destructive merge since the data-flow facts  $a = 1$  and  $a = 2$  flowing along the incoming edges are merged at node  $B_1$ . Thus, we see that simply duplicating acyclic paths does not always guarantee an increase in data-flow precision. Also, concentrating only on acyclic paths implies that all loop-back edges ( $E_2, B_1$  and  $E_3, B_1$  in the HPG) merge at a common loop-header (node  $B_1$  in the HPG). Thus, loop-headers which are destructive merges cannot be eliminated by the Ammons-Larus approach and data-flow precision is lost in these cases.

In comparison, the Split Graph constructed is shown in Figure 9. The destructive merge at node  $E$  is completely eliminated. In the Split Graph, uses of variable  $a$  at nodes  $B_0$ ,  $B_1$  and  $B_2$  can be replaced by constants 0, 1 and 2 respectively. Thus, we see that our approach effectively handles loop structures, guarantees additional optimization opportunities, and does not rely on expensive path profile information. Note that conventional control-flow graph restructuring approaches such as tail-duplication, superblock formation can be seen to be similar to the HPG approach since they use only profile information. We compare the HPG method with our approach quantitatively in Section 7.

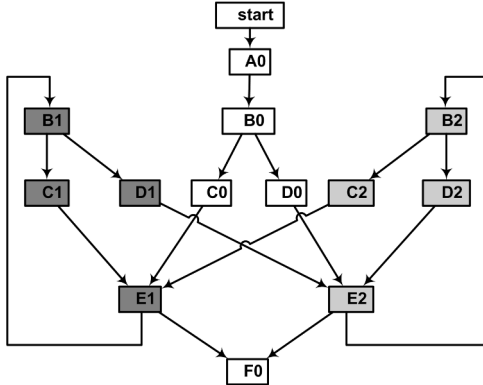


Figure 9: The *Split Graph* constructed from program  $P_1$  in which the *uses* of variable  $a$  at nodes  $B_0$ ,  $B_1$  and  $B_2$  can be replaced by constants 0, 1 and 2 respectively.

## 6.2 Other Related Approaches

In [5], an approach for complete removal of partial redundancy is described. Data-flow analysis is used to identify those regions of code which obstruct code motion. Code duplication and code motion are then used to eliminate the partial redundancy. Another approach targeted for PRE is discussed in [21]. Unlike our approach, [21] will result in unbounded code growth for finite height infinite lattices such as those seen in Constant Propagation. Further, [21] does not handle the precision versus code size trade-off and it is not clear from [21] how this can be accomplished. Martel [12] present an algorithm which uses graph substitution to unroll loops so as to improve the precision of invariants detected via static analysis. Our control-graph restructuring is more general and can handle more cases. For example, the programs in Figure 1 and Figure 7 cannot be optimized using such loop unrolling.

Code restructuring need not necessarily be limited to within a procedure. An extension of Ammons-Larus approach to the inter-procedural case is described in [13]. A more recent framework [22] for whole-program optimization also considers code duplication to perform *area specialization*, which is purely profile-driven.

There have also been several approaches which do not restructure the CFG, which are complementary to our approach. Holley and Rosen presented a general approach to improve data-flow precision by adding a finite set of predicates [10]. In [4], the precision of def-use analysis is improved by determining infeasible paths by using a low overhead technique based on detection of static branch correlations. Interestingly, path-sensitivity can also be obtained by synthesizing the name space of the data-flow analysis [3]. *Property simulation* is introduced in ESP [6] and is used to verify temporal safety properties. This approach keeps track of the correlation between “property state” and certain execution states. In [7], data-flow analysis is performed over a *predicated lattice*. The predicates used are determined automatically using a counterexample refinement technique. In [8], the context-sensitivity of the pointer analysis is adjusted based on the requirements of the client application.

Benchmark	Split over Base	Split over HPG
175.vpr	5	1
186.crafty	-2	2
197.parser	2	3
256.bzip2	0	3
300.twolf	3	-2
181.mcf	12	4
164.gzip	3	2
average	4	2

Table 2: Percentage speedup in the running times using Split in comparison to Base and HPG.

Benchmark	Split / HPG
175.vpr	1.15
186.crafty	1.10
197.parser	1.27
256.bzip2	1.11
300.twolf	0.93
181.mcf	13.75
164.gzip	2.32
average	3.5

Table 3: Ratio of the number of dynamic instructions with constant *uses* in Split over HPG.

## 7 EXPERIMENTAL RESULTS

We have implemented our approach in the Scale research compiler framework [19]. The framework is parameterized with the definition of a destructive merge, which depends on the data-flow analysis used, and on the definition of influenced nodes, which captures the interaction between the specific optimization and analysis. We present experimental results for the specific problem of Constant Propagation. We compare our approach (*Split*) with the Wegman-Zadeck conditional constant propagation algorithm [23](*Base*) and the Hot Path Graph approach(*HPG*) [1] using the SPECINT 2000 benchmark suite [20].

We instantiate the constant propagation phase of the O1 pass of the Scale compiler with the default approach (*Base*), the *HPG* approach, and *Split*. For our approach we eliminate the two fittest destructive merges per function using the heuristic explained in Section 5. The HPG approach uses a path profile generated using the *train* inputs for the respective programs, while the our Split approach uses a basic block profile from the same *train* inputs. The benchmarks are compiled for DEC ALPHA and were run on the 500MHz 21264 Alpha workstation. Running times were measures as average over multiple runs using the larger *ref* inputs.

### 7.1 Benefits of Split Approach

Table 2 shows the speedup obtained by our Split approach over the *Base* approach and over the *HPG* approach. Split gives an average speedup of 4% over the *Base* case, and it gives an average speedup of 2% over the *HPG* approach.

To understand where the speedup comes from, we calculate the number of dynamic instructions which have constant *uses* identified by the restructuring transformation. This is computed by first performing constant propagation and replacing all constant *uses* in the original program. Restructuring (*HPG* or *Split*) is then carried out. The constant *uses* discovered can be attributed only to the restructuring.

Benchmark	Split / Base	Split / HPG
175.vpr	1.5	0.7
186.crafty	2.0	0.7
197.parser	1.8	1.1
256.bzip2	1.9	0.5
300.twolf	2.0	0.7
181.mcf	1.9	1.0
164.gzip	1.5	0.8
average	1.8	0.65

**Table 4: Ratio of code size increase of Split over Base, and of Split over HPG.**

Thus, each instruction is weighted by the product of its execution count (using the *ref* inputs) and the number of new constant *uses*. The sum over all instructions gives us the number of **dynamic instructions which have constant *uses* only because of restructuring**. This metric has also been used in [1]. Table 3 shows the ratio of these instructions for Split over than of HPG. **We observe an average of 3.5 times more dynamic instructions with constants *uses* in Split as compared to HPG.** In the 181.mcf Split results in as many as 13.75 times dynamic constant use instructions. This is because Split can handle cyclic structures effectively.

## 7.2 Cost of Split Approach

As mentioned earlier, the increase in precision comes at the cost of code duplication. We measured the code size in terms of the number of Scale intermediate instructions. Table 4 shows the ratio of the code size of Split over that of Base. We observe an average of  $1.8 \times$  (80%) *increase* due to Split. The Table also shows the ratio of code size of Split over that of HPG. We notice that Split incurs less code size increase in comparison to HPG. Split shows an average of  $0.65 \times$  (35%) *decrease in code size* as compared to HPG.

## 8. CONCLUSION

We proposed a general framework to improve data-flow analysis precision based on restructuring the CFG of the program. The framework can be instantiated to any data-flow analysis. The actual transformation uses a known concepts of product automaton. We have proved that the transformation guarantees increase in optimization opportunities. Further, we showed that getting the optimal restructuring is NP-hard and proposed and evaluated a greedy heuristic. Our results indicate that our approach performs better than existing path profile driven approach [1].

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