## Optimising Compilers

Computer Science Tripos Part II
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## Lecture I Introduction

An optimising compiler


## Optimisation

=
Analysis
$+$

## Transformation

## Analysis + Transformation

- An analysis shows that your program has some property...
- ...and the transformation is designed to be safe for all programs with that property...
- ...so it's safe to do the transformation.


## Analysis + Transformation

```
int main(void)
{
    return 42;
}
int f(int x)
{
    return x * 2;
}
```

Analysis + Transformation

```
int main(void)
{
    return f(21);
}
int f(int x)
{
    return x * 2;
}
```

Analysis + Transformation

```
while (i <= k*2) {
    j = j * i;
    i = i + 1;
}
```

```
Analysis + Transformation
    while (i <= k*2) {
    k = k - i;
    i = i + 1;
}
```


## Analysis + Transformation

```
int main(void)
{
    return f(21);
}
```



## Analysis + Transformation

```
int t = k * 2;
while (i <= t) {
    j = j * i;
    i = i + 1;
}
```


int t = k * 2;
int t = k * 2;
while (i <= t) {
while (i <= t) {
k = k - i;
k = k - i;
i = i + 1;
i = i + 1;
}
}

## Stack-oriented code



## 3-address code

MOV t32, arg1
MOV t33,arg2
ADD t34,t32,t33
MOV t35, arg3
MOV t36, arg4
ADD t37,t35,t36
MUL res1,t34,t37
EXIT

## C into 3-address code

ENTRY fact
MOV t32, arg1
CMPEQ t32, \#0, lab1
SUB arg1,t32, \#1
CALL fact
MUL res1,t32,res1
EXIT
lab1: MOV res1,\#1
EXIT

Flowgraphs


## Basic blocks



Basic blocks


Basic blocks

A basic block doesn't contain any interesting control flow.

Basic blocks
ENTRY fact
 CMPEQ t $32, \# 0$


Basic blocks

Reduce time and space requirements for analysis algorithms by calculating and storing data flow information once per block
(and recomputing within a block if required) instead of once per instruction.

Basic blocks


Types of analysis
(and hence optimisation)

Scope:

- Within basic blocks ("local"/ "peephole")
- Between basic blocks ("global" / "intra-procedural")
- e.g. live variable analysis, available expressions
- Whole program ("inter-procedural")
- e.g. unreachable-procedure elimination


Peephole optimisation

$$
\begin{aligned}
& \text { ADD t32, arg1,\#1 replace } \\
& \text { MOV rōri } \\
& \text { MOV r1, ro: matches } \\
& \text { MUL } t 33, r 0, t 32 \\
& \text { MOV x,y } \\
& \text { MOV y x } \\
& \text { with } \\
& \text { MOV } x, y \\
& \text { ADD t32, arg1, \#1 } \\
& \text { MOV-rorí } \\
& \text { MUL t } 33, r 0, t 32
\end{aligned}
$$

## Types of analysis

(and hence optimisation)
Type of information:

- Control flow
- Discovering control structure (basic blocks, loops, calls between procedures)
- Data flow
- Discovering data flow structure (variable uses, expression evaluation)


## Finding basic blocks

I. Find all the instructions which are leaders:

- the first instruction is a leader;
- the target of any branch is a leader; and
- any instruction immediately following a branch is a leader.

2. For each leader, its basic block consists of itself and all instructions up to the next leader.

## Finding basic blocks

## ENTRY fact

MOV t32, arg1
CMPEQ t32, \#0, lab1
SUB arg1,t32, \#1
CALL fact
:MUL res1,t3/2,res1 EXIT
lab1: MOV res1,\#1
EXIT

## Summary

- Structure of an optimising compiler
- Why optimise?
- Optimisation $=$ Analysis + Transformation
- 3-address code
- Flowgraphs
- Basic blocks
- Types of analysis
- Locating basic blocks


## Lecture 2 <br> Unreachable-code \& -procedure elimination

## Intra-procedural analysis

An intra-procedural analysis collects information about the code inside a single procedure.

We may repeat it many times (i.e. once per procedure), but information is only propagated within the boundaries of each procedure, not between procedures.

One example of an intra-procedural control-flow optimisation (an analysis and an accompanying transformation) is unreachable-code elimination.

## Dead vs. unreachable code

int $f(i n t \mathrm{x}$, int y$)$ \{
return $x+y$;
int $z=x$ * $y ;$ UNREACHABLE
\}

Unreachable code cannot possibly be executed.
(Waste of space.)

## Dead vs. unreachable code

Unreachability is a control-flow property:
"May control ever arrive here?"


Control-flow analysis


Discovering information about how control (e.g. the program counter) may move through a program.

## Dead vs. unreachable code

```
int f(int x, int y) {
    int z = x * y; DEAD
```

    return \(x+y\);
    \}

Dead code computes unused values. (Waste of time.)

## Dead vs. unreachable code

Deadness is a data-flow property:
"May this data ever arrive anywhere?"
int $f(i n t \quad x, i n t y) ~\{$
int $z=x$ * $y$;
$\vdots$

Safety of analysis

```
int f(int x, int y) {
    if (g(x))
        int z = x * y; UNREACHABLE?
    }
    return x + y;
}
bool g(int x) {
    return false;
}
```


## Safety of analysis

```
int f(int x, int y) {
    if (g(x)) {
        int z = x * y; UNREACHABLE?
    }
    return x + y;
}
bool g(int x) {
    return ...x...;
}
```


## Safety of analysis

- Many interesting properties of programs are undecidable and cannot be computed precisely...
- ...so they must be approximated.
- A broken program is much worse than an inefficient one...
- ...so we must err on the side of safety.


## Safety of analysis

Naïvely,

```
if (false) {
    int z = x * y;
}
```

this instruction is reachable,

```
while (true) {
    // Code without 'break'
}
int z = x * y;
```

and so is this one.


## Safety of analysis

```
int f(int x, int y) {
    if (g(x)) {
        int z = x * y; UNREACHABLE?
    }
    return x + y;
}
```

In general, this is undecidable.
(Arithmetic is undecidable; cf. halting problem.)

## Safety of analysis

- If we decide that code is unreachable then we may do something dangerous (e.g. remove it!)...
- ...so the safe strategy is to overestimate reachability.
- If we can't easily tell whether code is reachable, we just assume that it is. (This is conservative.)
- For example, we assume
- both branches of a conditional are reachable
- and that loops always terminate.


## Safety of analysis

Another source of uncertainty is encountered when constructing the original flowgraph: the presence of indirect branches (also known as "computed jumps").

## Safety of analysis




## Safety of analysis



## Unreachable code

This naïve reachability analysis is simplistic, but has the advantage of corresponding to a very straightforward operation on the flowgraph of a procedure:

I .mark the procedure's entry node as reachable;
2.mark every successor of a marked node as reachable and repeat until no further marking is required.

## Unreachable code



## Safety of analysis

Again, this is a conservative overestimation of reachability.
In the worst-case scenario in which branch-address computations are completely unrestricted (i.e. the target of a jump could be absolutely anywhere), the presence of an indirect branch forces us to assume that all instructions are potentially reachable in order to guarantee safety.

## Safety of analysis



## Unreachable code



## Unreachable code

Programmers rarely write code which is completely unreachable in this naïve sense. Why bother with this analysis?

- Naïvely unreachable code may be introduced as a result of other optimising transformations.
- With a little more effort, we can do a better job.


## Unreachable code

Obviously, if the conditional expression in an if statement is literally the constant "false", it's safe to assume that the statements within are unreachable.

```
if (false) {
    int z = x * y; UNREACHABLE
}
```

But programmers never write code like that either.

## Unreachable code

However, other optimisations might produce such code. For example, copy propagation:

```
\vdots
if (false) {
    int z = x * y; UNREACHABLE
}
```


## Unreachable code

Note, however, that the reachability analysis no longer consists simply of checking whether any paths to an instruction exist in the flowgraph, but whether any of the paths to an instruction are actually executable.

With more effort we may get arbitrarily clever at spotting non-executable paths in particular cases, but in general the undecidability of arithmetic means that we cannot always spot them all.

## Unreachable code

For example, straightening is an optimisation which can eliminate jumps between basic blocks by coalescing them:


## Unreachable code

However, other optimisations might produce such code.
For example, copy propagation:

```
bool debug = false;
\vdots
if (debug) {
    int z = x * y;
}
```


## Unreachable code

We can try to spot (slightly) more subtle things too.

- if (!true) \{... \}
- if (false \&\& ...) \{... \}
- if (x != x) \{... \}
- while (true) \{... \} ...
- ...


## Unreachable code

Although unreachable-code elimination can only make a program smaller, it may enable other optimisations which make the program faster.

## Unreachable code

For example, straightening is an optimisation which can eliminate jumps between basic blocks by coalescing them:


## Unreachable code

For example, straightening is an optimisation which can eliminate jumps between basic blocks by coalescing them:


Straightening has removed a branch instruction, so the new program will execute faster.

## Inter-procedural analysis

An inter-procedural analysis collects information about an entire program.

Information is collected from the instructions of each procedure and then propagated between procedures.

One example of an inter-procedural control-flow optimisation (an analysis and an accompanying transformation) is unreachable-procedure elimination.

## Unreachable procedures

Unreachable-procedure elimination is very similar in spirit to unreachable-code elimination, but relies on a different data structure known as a call graph.

## Call graphs

Again, the precision of the graph is compromised in the presence of indirect calls.


## Call graphs

In general, we assume that a procedure containing an indirect call has all address-taken procedures as successors in the call graph - i.e., it could call any of them.

This is obviously safe; it is also obviously imprecise.
As before, it might be possible to do better by application of more careful methods (e.g. tracking data-flow of procedure variables).

## Call graphs



## Call graphs

Again, the precision of the graph is compromised in the presence of indirect calls.


And as before, this is a safe overestimation of reachability.

## Unreachable procedures

The reachability analysis is virtually identical to that used in unreachable-code elimination, but this time operates on the call graph of the entire program (vs. the flowgraph of a single procedure):

I .mark procedure main as callable;
2.mark every successor of a marked node as callable and repeat until no further marking is required.

## Unreachable procedures



## Safety of transformations

- All instructions/procedures to which control may flow at execution time will definitely be marked by the reachability analyses...
- ...but not vice versa, since some marked nodes might never be executed.
- Both transformations will definitely not delete any instructions/procedures which are needed to execute the program...
- ...but they might leave others alone too.


## If simplication

Empty then in if-then
(Assuming that $f$ has no side effects.)

## If simplication

Empty then in if-then-else

```
if (!f(x)) {
} else {
    z = x * y;
}
```


## Unreachable procedures



## If simplification

- Let's look at another set of basic controlflow transformations that can be carried out with only small amounts of analysis
- In this case, if simplification, which alters the structure of if statements (or removes them altogether) when possible


## If simplication

Empty else in if-then-else

```
if (f(x)) {
        z = x * y;
} else
```

\}


## If simplication

Empty then and else in if-then-else

## If simplication

Constant condition
if (true)
$z=x$ * $y ;$
\}

## If simplication

Nested if with common subexpression

```
if (x > 3 && t) {
    if (x > 3)
            z = x * y;
        else
        z = y - x;
}
```


## Loop simplification

```
int x = 10;
int i = 4;
```


## Summary

- Control-flow analysis operates on the control structure of a program (flowgraphs and call graphs)
- Unreachable-code elimination is an intraprocedural optimisation which reduces code size
- Unreachable-procedure elimination is a similar, inter-procedural optimisation making use of the program's call graph
- Analyses for both optimisations must be imprecise in order to guarantee safety


## Lecture 3 <br> Live variable analysis

## Motivation

## Programs may contain

- code which gets executed but which has no useful effect on the program's overall result;
- occurrences of variables being used before they are defined; and
- many variables which need to be allocated registers and/or memory locations for compilation.

The concept of variable liveness is useful in dealing with all three of these situations.

## Liveness

At each instruction, each variable in the program is either live or dead.

We therefore usually consider liveness from an instruction's perspective: each instruction (or node of the
flowgraph) has an associated set of live variables.


```
    return S'+手;
                live(n)={s,t,x,y}
```


## Semantic vs. syntactic

A variable $x$ is semantically live at a node $n$ if there is some execution sequence starting at $n$ whose (externally observable) behaviour can be affected by changing the value of $x$.


## Data-flow analysis



Discovering information about how data (i.e. variables and their values) may move through a program.

## Liveness

Liveness is a data-flow property of variables:
"Is the value of this variable needed?" (cf. dead code)

```
int f(int x, int y) {
    int z = x * y;
    \vdots尔
        #&!
```


## Semantic vs. syntactic

There are two kinds of variable liveness:

- Semantic liveness
- Syntactic liveness


## Semantic vs. syntactic

A variable $x$ is semantically live at a node $n$ if there is some execution sequence starting at $n$ whose (externally observable) behaviour can be affected by changing the value of $x$.


## Semantic vs. syntactic

Semantic liveness is concerned with the execution behaviour of the program.

This is undecidable in general.
(e.g. Control flow may depend upon arithmetic.)

## Semantic vs. syntactic

A variable is syntactically live at a node if there is a path to the exit of the flowgraph along which its value may be used before it is redefined.

Syntactic liveness is concerned with properties of the syntactic structure of the program.

Of course, this is decidable.

So what's the difference?

```
Semantic vs. syntactic
int t = x * y; t DEAD
if ((x+1)* (x+1) == y) {
    t = 1;
}
if (x*x + 2*x + 1 != y) {
    t = 2;
}
return t;
```

Semantically: one of the conditions will be true, so on every execution path $t$ is redefined before it is returned. The value assigned by the first instruction is never used.

## Semantic vs. syntactic



On this path through the flowgraph, $t$ is not redefined before it's used, so $t$ is syntactically live at the first instruction.

Note that this path never actually occurs during execution.

Semantic vs. syntactic

lab2:
 IADD t32,x,\#1
'MUL t33,t32,t32

lab1:
t34, x, x
MUL t35,x,\#2
,ADD t36,t34,t35 IADD t37,t36,\#1
CMPEQ t37, Y 1 Iab2
MOV res1,t_-.....

## Semantic vs. syntactic

So, as we've seen before, syntactic liveness is a computable approximation of semantic liveness.

## Semantic vs. syntactic



Semantic vs. syntactic


## Semantic vs. syntactic

$$
\operatorname{sem-live}(n) \subseteq \operatorname{syn-live}(n)
$$

Using syntactic methods, we safely overestimate liveness.

## Live variable analysis

Variable liveness flows (backwards) through the program in a continuous stream.

Each instruction has an effect on the liveness information as it flows past.

## Live variable analysis



## Live variable analysis



## Live variable analysis

LVA is a backwards data-flow analysis: usage information from future instructions must be propagated backwards through the program to discover which variables are live.

## Live variable analysis

An instruction makes a variable live when it references (uses) it.

## Live variable analysis

An instruction makes a variable dead when it defines (assigns to) it.

## Live variable analysis

We can devise functions ref(n) and def(n) which give the sets of variables referenced and defined by the instruction at node $n$.

$$
\left.\begin{array}{rlrl}
\operatorname{ref}(x=3) & =\{ \} & \operatorname{ref}(\operatorname{print} x) & =\{x\} \\
\operatorname{def}(x=3) & =\{x\} & \operatorname{def}(\operatorname{print} x) & =\{ \}
\end{array}\right\} \begin{aligned}
\operatorname{ref}(x=x+y) & =\{x, y\} \\
& \operatorname{def}(x=x+y)=\{x\}
\end{aligned}
$$

## Live variable analysis

As liveness flows backwards past an instruction, we want to modify the liveness information by adding any variables which it references (they become live) and removing any which it defines (they become dead).

$\operatorname{ref}(\operatorname{print} x)=\{x\}$



$$
x\}
$$

def( x
$x=3)=\{x\}$
$\{\{x, y\}$

## Live variable analysis

So, if we consider in-live(n) and out-live( $n$ ), the sets of variables which are live immediately before and immediately after a node, the following equation must hold:
$\operatorname{in-live}(n)=(\operatorname{out-live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n)$

## Live variable analysis

If an instruction both references and defines variables, we must remove the defined variables before adding the referenced ones.

$\operatorname{def}(x=x+y)=\{x\}$
$\operatorname{ref}(x=x+y)=\{x, y\}$

## Live variable analysis

$\operatorname{in-live}(n)=(\operatorname{out}-\operatorname{live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n)$


$$
\operatorname{def}(n)=\{x\} \quad \operatorname{ref}(n)=\{x, y\}
$$

## Live variable analysis

In straight-line code each node has a unique successor, and the variables live at the exit of a node are exactly those variables live at the entry of its successor.

## Live variable analysis

In general, however, each node has an arbitrary number of successors, and the variables live at the exit of a node are exactly those variables live at the entry of any of its successors.

## Live variable analysis



## Data-flow equations

These are the data-flow equations for live variable analysis, and together they tell us everything we need to know about how to propagate liveness information through a program.
$\operatorname{in-live}(n)=(\operatorname{out}-\operatorname{live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n)$
$\operatorname{out-live}(n)=\bigcup_{s \in \operatorname{succ}(n)} \operatorname{in-live}(s)$

## Algorithm

We now have a formal description of liveness, but we need an actual algorithm in order to do the analysis.

## Live variable analysis

So the following equation must also hold:

$$
\text { out-live }(n)=\bigcup_{s \in \operatorname{succ}(n)} \operatorname{in-live}(s)
$$

## Data-flow equations

Each is expressed in terms of the other, so we can combine them to create one overall liveness equation.
$\operatorname{live}(n)=\left(\left(\bigcup_{s \in \operatorname{succ}(n)} \operatorname{live}(s)\right) \backslash \operatorname{def}(n)\right) \cup \operatorname{ref}(n)$

## Algorithm

"Doing the analysis" consists of computing a value live( $n$ ) for each node $n$ in a flowgraph such that the liveness data-flow equations are satisfied.

A simple way to solve the data-flow equations is to adopt an iterative strategy.

## Algorithm



Algorithm


## Algorithm

for $\mathrm{i}=1$ to n do live[i] := \{\} while (live[] changes) do
for $i=1$ to $n$ do

$$
\text { live }[i]:=\left(\left(\bigcup_{s \in s u c c(i)} \operatorname{live}[s]\right) \backslash \operatorname{def}(\mathrm{i})\right) \cup \operatorname{ref}(\mathrm{i})
$$

## Algorithm

Implementation notes:

- If the program has $n$ variables, we can implement each element of live [ ] as an n-bit value, with each bit representing the liveness of one variable.
- We can store liveness once per basic block and recompute inside a block when necessary. In this case, given a basic block $n$ of instructions $i_{i}, \ldots, i_{k}$ : $\operatorname{live}(n)=\left(\bigcup_{s \in \operatorname{succ}(n)} \operatorname{live}(s)\right) \backslash \operatorname{def}\left(i_{k}\right) \cup \operatorname{ref}\left(i_{k}\right) \cdots \backslash \operatorname{def}\left(i_{1}\right) \cup \operatorname{ref}\left(i_{1}\right)$


## Safety of analysis

```
    MOV x,#1
    MOV y,#2
    MOV z,#3
    MOV t32,#&x
    MOV t33,#&y
    MOV t34,#&z
\vdots
m.'\overline{STT}}\overline{+}\overline{3}\overline{5
    ref(m)={t35}
```



```
    ref(n)={t37,x,y,z}
```


## Algorithm

This algorithm is guaranteed to terminate since there are a finite number of variables in each program and the effect of one iteration is monotonic.

Furthermore, although any solution to the data-flow equations is safe, this algorithm is guaranteed to give the smallest (and therefore most precise) solution.
(See the Knaster-Tarski theorem if you're interested.)

## Safety of analysis

- Syntactic liveness safely overapproximates semantic liveness.
- The usual problem occurs in the presence of address-taken variables (cf. labels, procedures): ambiguous definitions and references. For safety we must
- overestimate ambiguous references (assume all address-taken variables are referenced) and
- underestimate ambiguous definitions (assume no variables are defined); this increases the size of the smallest solution.


## Summary

- Data-flow analysis collects information about how data moves through a program
- Variable liveness is a data-flow property
- Live variable analysis (LVA) is a backwards dataflow analysis for determining variable liveness
- LVA may be expressed as a pair of complementary data-flow equations, which can be combined
- A simple iterative algorithm can be used to find the smallest solution to the LVA data-flow equations


# Lecture 4 Available expression analysis 

## Expressions

Any given program contains a finite number of expressions (i.e. computations which potentially produce values), so we may talk about the set of all expressions of a program.

```
int z = x * y;
print s + t;
int w = u / v;
!
```

program contains expressions $\left\{\mathrm{x}^{*} \mathrm{y}, \mathrm{s}+\mathrm{t}, \mathrm{u} / \mathrm{v}, \ldots\right\}$

## Availability

At each instruction, each expression in the program is either available or unavailable.

We therefore usually consider availability from an instruction's perspective: each instruction (or node of the flowgraph) has an associated set of available expressions.
int $z=x$ * $y$;
print_s_t_tí-
n: int w- $=$ ung $-\underline{i} \quad \operatorname{arail}(n)=\left\{x^{*} y, s+t\right\}$

## Motivation

Programs may contain code whose result is needed, but in which some computation is simply a redundant repetition of earlier computation within the same program.

The concept of expression availability is useful in dealing with this situation.

## Availability

Availability is a data-flow property of expressions:
"Has the value of this expression already been computed?"


So far, this is all familiar from live variable analysis.
Note that, while expression availability and variable liveness share many similarities (both are simple data-flow properties), they do differ in important ways.

By working through the low-level details of the availability property and its associated analysis we can see where the differences lie and get a feel for the capabilities of the general data-flow analysis framework.

## Semantic vs. syntactic

An expression is semantically available at a node $n$ if its value gets computed (and not subsequently invalidated) along every execution sequence ending at $n$.

## Semantic vs. syntactic

An expression is semantically available at a node $n$ if its value gets computed (and not subsequently invalidated) along every execution sequence ending at $n$.

## Semantic vs. syntactic

An expression is syntactically available at a node $n$ if its value gets computed (and not subsequently invalidated) along every path from the entry of the flowgraph to $n$.

As before, semantic availability is concerned with the execution behaviour of the program, whereas syntactic availability is concerned with the program's syntactic structure.

And, as expected, only the latter is decidable.

## Semantic vs. syntactic

```
    ADD t32,x,#1
    MUL t33,t32,t32
    CMPNE t33,y,lab1
    ADD s,x,y
lab1: MUL t34,x,x
    MUL t35,x,#2
    ADD t36,t34,t35
    ADD t37,t36,#1
    CMPEQ t37,y,lab2
    ADD t,x,y
lab2: ADD res1,x,y
```


## Semantic vs. syntactic

If an expression is deemed to be available, we may do something dangerous (e.g. remove an instruction which recomputes its value).

Whereas with live variable analysis we found safety in assuming that more variables were live, here we find safety in assuming that fewer expressions are available.

Semantic vs. syntactic


## Semantic vs. syntactic

sem-avail( $(n) \supseteq \operatorname{syn}-\operatorname{avail}(n)$

This time, we safely underestimate availability.

$$
(\text { cf. } \operatorname{sem-live}(n) \subseteq \operatorname{syn-live}(n))
$$

## Available expression analysis

Available expressions is a forwards data-flow analysis: information from past instructions must be propagated forwards through the program to discover which expressions are available.


## Available expression analysis

An instruction makes an expression available when it generates (computes) its current value.

## Available expression analysis

An instruction makes an expression unavailable when it kills (invalidates) its current value.

## Warning

Danger: there is a standard presentation of available expression analysis (textbooks, notes for this course) which is formally satisfying but contains an easily-overlooked subtlety.

We'll first look at an equivalent, more intuitive bottom-up presentation, then amend it slightly to match the version given in the literature.

## Available expression analysis

Unlike variable liveness, expression availability flows forwards through the program.

As in liveness, though, each instruction has an effect on the availability information as it flows past.


## Available expression analysis

As in LVA, we can devise functions gen( $n$ ) and kill( $n$ ) which give the sets of expressions generated and killed by the instruction at node $n$.

The situation is slightly more complicated this time: an assignment to a variable $x$ kills all expressions in the program which contain occurrences of $x$.

## Available expression analysis

So, in the following, $E_{x}$ is the set of expressions in the program which contain occurrences of $x$.

$$
\begin{aligned}
& \operatorname{gen}(x=3)=\{ \} \quad \operatorname{gen}(\operatorname{print} x+1)=\{x+1\} \\
& \operatorname{kill}(x=3)=E_{x} \quad \operatorname{kill}(\operatorname{print} x+1)=\{ \}
\end{aligned}
$$

$$
\begin{aligned}
\operatorname{gen}(x & =x+y) \\
\operatorname{kill}(x & =x+y+y\} \\
& =E_{x}
\end{aligned}
$$

## Available expression analysis

If an instruction both generates and kills expressions, we must remove the killed expressions after adding the generated ones (cf. removing def(n) before adding ref(n)).

$$
\begin{array}{r}
\{\{x+1, y+1\} \\
x \\
\left\{\begin{array}{l}
\{x+y \\
\{y+1\}
\end{array}\right.
\end{array}
$$

As availability flows forwards past an instruction, we expressions which it generates (they become available)

$$
\operatorname{gen}(x=x+y)=\{x+y\}
$$

$\operatorname{kill}(x=x+y)=E_{x}$ want to modify the availability information by adding any and removing any which it kills (they become unavailable).


## Available expression analysis

So, if we consider in-avail(n) and out-avail(n), the sets of expressions which are available immediately before and immediately after a node, the following equation must hold:
$\operatorname{out-avail}(n)=(\operatorname{in-avail}(n) \cup \operatorname{gen}(n)) \backslash \operatorname{kill}(n)$

## Available expression analysis

As in LVA, we have devised one equation for calculating out-avail(n) from the values of gen(n), kill(n) and in-avail(n), and now need another for calculating in-avail(n).


## Available expression analysis

$$
\begin{aligned}
& \text { out-avail }(n)=(\text { in-avail }(n) \cup \operatorname{gen}(n)) \backslash \operatorname{kill}(n) \\
& \left\{\begin{array}{l}
\text { in-avail }(n)=\{x+1, y+1\}
\end{array}\right. \\
& \qquad \begin{array}{l}
\text { ( } n=(\{x+1, y+1\} \cup\{x+y\}) \backslash\{x+1, x+y\} \\
=\{x+1, x+y, y+1\} \backslash\{x+1, x+y\}=\{y+1\}
\end{array} \\
& \operatorname{gen}(n)=\{x+y\} \quad \operatorname{kill}(n)=\{x+1, x+y\}
\end{aligned}
$$

## Available expression analysis

When a node $n$ has a single predecessor $m$, the information propagates along the control-flow edge as you would expect: in-avail(n) = out-avail( $m$ ).

When a node has multiple predecessors, the expressions available at the entry of that node are exactly those expressions available at the exit of all of its predecessors (cf."any of its successors" in LVA).

## Available expression analysis



## Data-flow equations

These are the data-flow equations for available expression analysis, and together they tell us everything we need to know about how to propagate availability information through a program.
$\operatorname{in-avail}(n)=\bigcap_{p \in \operatorname{pred}(n)} \operatorname{out-avail}(p)$
$\operatorname{out-avail}(n)=(\operatorname{in-avail}(n) \cup \operatorname{gen}(n)) \backslash \operatorname{kill}(n)$

## Data-flow equations

Danger: we have overlooked one important detail.


Clearly there should be no expressions available here, so we must stipulate explicitly that $\operatorname{avail}(n)=\{ \}$ if $\operatorname{pred}(n)=\{ \}$.

## Data-flow equations

The functions and equations presented so far are correct, and their definitions are fairly intuitive.

However, we may wish to have our data-flow equations in a form which more closely matches that of the LVA equations, since this emphasises the similarity between the two analyses and hence is how they are most often presented.

A few modifications are necessary to achieve this.

## Available expression analysis

So the following equation must also hold:

$$
\operatorname{in-avail}(n)=\bigcap_{p \in \operatorname{pred}(n)} \operatorname{out-avail}(p)
$$

## Data-flow equations

Each is expressed in terms of the other, so we can combine them to create one overall availability equation.
$\operatorname{avail}(n)=\bigcap_{p \in \operatorname{pred}(n)}((\operatorname{avail}(p) \cup \operatorname{gen}(p)) \backslash \operatorname{kill}(p))$

## Data-flow equations

With this correction, our data-flow equation for expression availability is
$\operatorname{avail}(n)= \begin{cases}\bigcap_{p \in \operatorname{pred}(n)}((\operatorname{avail}(p) \cup \operatorname{gen}(p)) \backslash \operatorname{kill}(p)) & \text { if } \operatorname{pred}(n) \neq\{ \} \\ \{ \} & \text { if } \operatorname{pred}(n)=\{ \}\end{cases}$

## Data-flow equations

$\operatorname{in-live}(n)=(\operatorname{out-live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n)$
out-live $(n)=\bigcup_{s \in \operatorname{succ}(n)} i n-\operatorname{live}(s)$
These differences are inherent in the analyses.
$\operatorname{in-avail}(n)=\bigcap_{p \in \overline{\operatorname{pred}(n)}} \operatorname{out-avail}(p)$
$\operatorname{out-avail}(n)=(\operatorname{in-avail}(n) \cup \operatorname{gen}(n)) \backslash \operatorname{kill}(n)$

## Data-flow equations

$$
\begin{aligned}
\operatorname{in-live}(n) & =(\operatorname{out-live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n) \\
\operatorname{out-live}(n) & =\bigcup_{s \in \operatorname{succ}(n)} \operatorname{in-live}(s)
\end{aligned}
$$

These differences are an arbitrary result of our definitions.

$$
\begin{aligned}
\operatorname{in-avail}(n) & =\bigcap_{p \in \operatorname{pred}(n)} \operatorname{out-avail}(p) \\
\operatorname{out-avail}(n) & =(\operatorname{in-avail}(n) \cup \operatorname{gen}(n)) \backslash \operatorname{kill}(n)
\end{aligned}
$$

## Data-flow equations

We might instead have decided to define gen( $n$ ) and kill(n) to coincide with the following (standard) definitions:

- A node generates an expression e if it must compute the value of $e$ and does not subsequently redefine any of the variables occuring in e.
- A node kills an expression e if it may redefine some of the variables occurring in $e$ and does not subseauently recompute the value of $e$.


## Data-flow equations

Since these new definitions take account of which expressions are generated overall by a node (and exclude those which are generated only to be immediately killed), we may propagate availability information through a node by removing the killed expressions before adding the generated ones, exactly as in LVA.
$\operatorname{out-avail}(n)=(\operatorname{in-avail(}(n) \backslash \operatorname{kill}(n)) \cup \operatorname{gen}(n)$ $\operatorname{in-live}(n)=(\operatorname{out-live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n)$

## Algorithm

- We again use an array, avail [ ], to store the available expressions for each node.
- We initialise avail [ ] such that each node has all expressions available (cf. LVA: no variables live).
- We again iterate application of the data-flow equation at each node until avail [ ] no longer changes.


## Algorithm

We can do better if we assume that the flowgraph has a single entry node (the first node in avail [ ]).

Then avail [ 1] may instead be initialised to the empty set, and we need not bother recalculating availability at the first node during each iteration.

## Algorithm

```
avail[1] := {}
for i = 2 to n do avail[i] := U
while (avail[] changes) do
    for i = 2 to n do
        avail[i] := \bigcap ((avail[p]\kill(p))\cupgen(p))
                    p\inpred(i)
```


## Algorithm

As with LVA, this algorithm is guaranteed to terminate since the effect of one iteration is monotonic (it only removes expressions from availability sets) and an empty availability set cannot get any smaller.

Any solution to the data-flow equations is safe, but this algorithm is guaranteed to give the largest (and therefore most precise) solution.

## Algorithm

Implementation notes:

- If we arrange our programs such that each assignment assigns to a distinct temporary variable, we may number these temporaries and hence number the expressions whose values are assigned to them.
- If the program has $n$ such expressions, we can implement each element of avail [ ] as an n-bit value, with the $m^{\text {th }}$ bit representing the availability of expression number $m$.


## Safety of analysis

- Syntactic availability safely underapproximates semantic availability.
- Address-taken variables are again a problem. For safety we must
- underestimate ambiguous generation (assume no expressions are generated) and
- overestimate ambiguous killing (assume all expressions containing address-taken variables are killed); this decreases the size of the largest solution.


## Analysis framework

$$
\begin{aligned}
\operatorname{in}-\operatorname{live}(n) & =(\operatorname{out-live}(n) \backslash \operatorname{def}(n)) \cup \operatorname{ref}(n) \\
\operatorname{out}-\operatorname{live}(n) & =\bigcup_{s \in \operatorname{succ}(n)} \operatorname{in-live}(s)
\end{aligned}
$$

$$
\operatorname{in-avail}(n)=\bigcap_{p \in \operatorname{pred}(n)} \text { out-avail }(p)
$$

$$
\operatorname{out-avail}(n)=(\operatorname{in-avail}(n) \backslash \operatorname{kill}(n)) \cup \operatorname{gen}(n)
$$

## Algorithm

Implementation notes:

- Again, we can store availability once per basic block and recompute inside a block when necessary. Given each basic block $n$ has $k_{n}$ instructions $n[I], \ldots, n\left[k_{n}\right]$ :

```
avail(n)=\bigcap(avail(p)\\operatorname{kill}(p[1])\cup\operatorname{gen}(p[1])\cdots\\operatorname{kill}(p[\mp@subsup{k}{p}{}])\cup\operatorname{gen}(p[\mp@subsup{k}{p}{}]))
``` \(p \in \operatorname{pred}(n)\)

\section*{Analysis framework}

The two data-flow analyses we've seen, LVA and AVAIL, clearly share many similarities.

In fact, they are both instances of the same simple dataflow analysis framework: some program property is computed by iteratively finding the most precise solution to data-flow equations, which express the relationships between values of that property immediately before and immediately after each node of a flowgraph.

\section*{Analysis framework}

LVA's data-flow equations have the form
\[
\operatorname{in}(n)=(\operatorname{out}(n) \backslash \ldots) \cup \ldots \quad \operatorname{out}(n)=\bigcup_{s \in \operatorname{suc}(n)} \operatorname{in}(s)
\]
union over successors

AVAIL's data-flow equations have the form
\[
\operatorname{out}(n)=(i n(n) \backslash \ldots) \cup \ldots \quad i n(n)=\bigcap_{p \in \operatorname{pred}(n)} \operatorname{out}(p)
\]
intersection over predecessors

\section*{Analysis framework}

...and others

\section*{Analysis framework}

So, given a single algorithm for iterative solution of data-flow equations of this form, we may compute all these analyses and any others which fit into the framework.

\section*{Summary}
- Expression availability is a data-flow property
- Available expression analysis (AVAIL) is a forwards data-flow analysis for determining expression availability
- AVAIL may be expressed as two complementary data-flow equations, which may be combined
- A simple iterative algorithm can be used to find the largest solution to the data-flow equations
- AVAIL and LVA are both instances (among others) of the same data-flow analysis framework

\section*{Lecture 5 Data-flow anomalies and clash graphs}

\section*{Optimisation vs. debugging}

Data-flow anomalies may manifest themselves in different ways: some may actually "break" the program (make it crash or exhibit undefined behaviour), others may just make the program "worse" (make it larger or slower than necessary).

Any compiler needs to be able to report when a program is broken (i.e. "compiler warnings"), so the identification of data-flow anomalies has applications in both optimisation and bug elimination.


\section*{Dead code}


Successive iterations may yield further improvements.

\section*{Motivation}

Both human- and computer-generated programs sometimes contain data-flow anomalies.

These anomalies result in the program being worse, in some sense, than it was intended to be.

Data-flow analysis is useful in locating, and sometimes correcting, these code anomalies.

\section*{Dead code}

Dead code is a simple example of a data-flow anomaly, and LVA allows us to identify it.

Recall that code is dead when its result goes unused; if the variable x is not live on exit
from an instruction which assigns some value to \(x\), then the whole instruction is dead.

\section*{Dead code}

For this kind of anomaly, an automatic remedy is not only feasible but also straightforward: dead code with no live side effects is useless and may be removed.

\section*{Dead code}

The program resulting from this transformation will remain correct and will be both smaller and faster than before (cf. just smaller in unreachable code elimination), and no programmer intervention is required.

\section*{Uninitialised variables}

In some languages, for example C and our 3-address intermediate code, it is syntactically legitimate for a program to read from a variable before it has definitely been initialised with a value.

If this situation occurs during execution, the effect of the read is usually undefined and depends upon unpredictable details of implementation and environment.

\section*{Uninitialised variables}

In a "healthy" program, variable liveness produced by later instructions is consumed by earlier ones; if an instruction demands the value of a variable (hence making it live), it is expected that an earlier instruction will define that variable (hence making it dead again).

\section*{Uninitialised variables}

If any variables are still live at the beginning of a program, they represent uses which are potentially unmatched by corresponding definitions, and hence indicate a program with potentially undefined (and therefore incorrect) behaviour.

\section*{Uninitialised variables}

This kind of behaviour is often undesirable, so we would like a compiler to be able to detect and warn of the situation.

Happily, the liveness information collected by LVA allows a compiler to see easily when a read from an undefined variable is possible.

Uninitialised variables


Uninitialised variables


\section*{Uninitialised variables}


\section*{Uninitialised variables}

Here the analysis is being too safe, and the warning is unnecessary, but this imprecision is the nature of our computable approximation to semantic liveness.

So the compiler must either risk giving unnecessary warnings about correct code ("false positives") or failing to give warnings about incorrect code ("false negatives").Which is worse?

Opinions differ.

\section*{Uninitialised variables}

Although dead code may easily be remedied by the compiler, it's not generally possible to automatically fix the problem of uninitialised variables.

As just demonstrated, even the decision as to whether a warning indicates a genuine problem must often be made by the programmer, who must also fix any such problems by hand.

\section*{Uninitialised variables}

Note that higher-level languages have the concept of (possibly nested) scope, and our expectations for variable initialisation in"healthy" programs can be extended to these.

In general we expect the set of live variables at the beginning of any scope to not contain any of the variables local to that scope.

\section*{Write-write anomalies}

While LVA is useful in these cases, some similar data-flow anomalies can only be spotted with a different analysis.

Write-write anomalies are an example of this. They occur when a variable may be written twice with no intervening read; the first write may then be considered unnecessary in some sense.
\[
\begin{aligned}
& \mathrm{x}=11 ; \\
& \mathrm{x}=13 ; \\
& \text { print } \mathrm{x} ;
\end{aligned}
\]

\section*{Uninitialised variables}
```

int x = 5;
int y = 7;

```
```

if (p) {

```
if (p) {
    \vdots
    \vdots
    print z;
    print z;
}
}
print x+y;
```

print x+y;

```


\section*{Write-write anomalies}

A simple data-flow analysis can be used to track which variables may have been written but not yet read at each node.

In a sense, this involves doing LVA in reverse (i.e. forwards!): at each node we should remove all variables which are referenced, then add all variables which are defined.

\section*{Write-write anomalies \\ \section*{Write-write anomalies}}

\section*{Write-write anomalies}
\[
\begin{aligned}
\operatorname{in-wnr}(n) & =\bigcup_{p \in \operatorname{pred}(n)} \text { out-wnr }(p) \\
\text { out-wnr }(n) & =(\operatorname{in-wnr}(n) \backslash \operatorname{ref}(n)) \cup \operatorname{def}(n)
\end{aligned}
\]
\[
w n r(n)=\bigcup_{p \in \operatorname{pred}(n)}((w n r(p) \backslash \operatorname{ref}(p)) \cup \operatorname{def}(p))
\]


\section*{Write-write anomalies}

But, although the second write to a variable may turn an earlier write into dead code, the presence of a write-write anomaly doesn't necessarily mean that a variable is dead hence the need for a different analysis.

\section*{Write-write anomalies}
```

if (p) {
x = 13;
} else {
x = 11;
}
print x;

```

This code does the same job, but avoids writing to x twice in succession on any control-flow path.

\section*{Write-write anomalies}
```

x = 11;
if (p) {
x = 13;
}
print x;

```
\(x\) is live throughout this code, but if \(p\) is true during execution, x will be written twice before it is read. In most cases, the programmer can remedy this.

\section*{Write-write anomalies}
```

if (p) {
x = 13;
}
if (!p) {
x = 11;
}
print x;

```

Again, the analysis may be too approximate to notice that a particular write-write anomaly may never occur during any execution, so warnings may be inaccurate.

\section*{Clash graphs}

The ability to detect data-flow anomalies is a nice compiler feature, but LVA's main utility is in deriving a data structure known as a clash graph (aka interference graph).

\section*{Clash graphs}

When generating intermediate code it is convenient to simply invent as many variables as necessary to hold the results of computations; the extreme of this is "normal form", in which a new temporary variable is used on each occasion that one is required, with none being reused.

\section*{Clash graphs}


\section*{Clash graphs}

This makes generating 3-address code as straightforward as possible, and assumes an imaginary target machine with an unlimited supply of "virtual registers", one to hold each variable (and temporary) in the program.

Such a naïve strategy is obviously wasteful, however, and won't generate good code for a real target machine.

\section*{Clash graphs}

Before we can work on improving the situation, we must collect information about which variables actually need to be allocated to different registers on the target machine, as opposed to having been incidentally placed in different registers by our translation to normal form.

LVA is useful here because it can tell us which variables are simultaneously live, and hence must be kept in separate virtual registers for later retrieval.

\section*{Clash graphs}
```

x = 11;
y = 13;
z = (x+y) * 2;
a = 17;
b = 19;
z = z + (a*b);

```

\section*{Clash graphs}
\begin{tabular}{cl} 
& \(\}\) \\
In a program's clash graph there is & \(\{x\}\) \\
one vertex for each virtual register & \(\{x, y\}\) \\
and an edge between vertices when & \(\{\mathrm{z}\}\) \\
their two registers are ever & \(\{a, z\}\) \\
simultaneously live. & \(\{a, b, z\}\) \\
& \(\{t 2, z\}\)
\end{tabular}

\section*{Clash graphs}

MOV \(a, \# 11\)
MOV \(b, \# 13\)
ADD \(a, a, b\)
MUL \(z, a, \# 2\)
MOV \(a, \# 17\)
MOV b, \#19
MUL \(a, a, b\)
ADD \(z, z, a\)

\section*{Clash graphs}

MOV x, \#11
MOV y, \#13
ADD t1,x,y
MUL \(z, t 1, \# 2\)
MOV a, \#17
MOV b, \#19
MUL t2, a,b
ADD \(z, z, t 2\)

\section*{Clash graphs}


\section*{Summary}
- Data-flow analysis is helpful in locating (and sometimes correcting) data-flow anomalies
- LVA allows us to identify dead code and possible uses of uninitialised variables
- Write-write anomalies can be identified with a similar analysis
- Imprecision may lead to overzealous warnings
- LVA allows us to construct a clash graph

\section*{Lecture 6 Register allocation}

\section*{Graph colouring}

Register allocation depends upon the solution of a closely related problem known as graph colouring.

\section*{Graph colouring}


\section*{Graph colouring}


\section*{Motivation}

Normal form is convenient for intermediate code.

However, it's extremely wasteful.
Real machines only have a small finite number of registers, so at some stage we need to analyse and transform the intermediate representation of a program so that it only requires as many (architectural) registers as are really available.

This task is called register allocation.


\section*{Graph colouring}

For general (non-planar) graphs, however, four colours are not sufficient; there is no bound on how many may be required.


\section*{Allocation by colouring}

This is essentially the same problem that we wish to solve for clash graphs.
- How many colours (i.e. architectural registers) are necessary to colour a clash graph such that no two connected vertices have the same colour (i.e. such that no two simultaneously live virtual registers are stored in the same arch. register)?
- What colour should each vertex be?

\section*{Allocation by colouring}

MOV r0, \#11
MOV r1, \#13
ADD r0,r0,r1
MUL r2,r0, \#2
MOV r0, \#17
MOV r1,\#19
MUL r0,r0,r1
ADD r2,r2,r0


\section*{Algorithm}
- Choose a vertex (i.e. virtual register) which has the least number of incident edges (i.e. clashes).
- Remove the vertex and its edges from the graph, and push the vertex onto a LIFO stack.
- Repeat until the graph is empty.
- Pop each vertex from the stack and colour it in the most conservative way which avoids the colours of its (already-coloured) neighbours.


\section*{Algorithm}

Bear in mind that this is only a heuristic.


A better (more minimal) colouring may exist.

\section*{Spilling}

This algorithm tries to find an approximately minimal colouring of the clash graph, but it assumes new colours are always available when required.

In reality we will usually have a finite number of colours (i.e. architectural registers) available; how should the algorithm cope when it runs out of colours?

\section*{Spilling}

The quantity of architectural registers is strictly limited, but it is usually reasonable to assume that fresh memory locations will always be available.

So, when the number of simultaneously live values exceeds the number of architectural registers, we may spill the excess values into memory.

Operating on values in memory is of course much slower, but it gets the job done.

\section*{Spilling}

ADD a,b,c
vs.
LDR t1, \#0xFFA4
LDR t2, \#0xFFA8
ADD t3,t1,t2
STR t3, \#0xFFA0

\section*{Algorithm}

a: 3, b: 5, c: 7, d: II, w: I3, x: I7, y: I9, z: 23

\section*{Algorithm}

r0
r1
\(a\) and \(b\)
spilled to memory
a: 3, b: 5, c: 7, d: II, w: I3, x: I7, y: I9, z: 23

\section*{Algorithm}
- Choose a vertex with the least number of edges.
- If it has fewer edges than there are colours,
- remove the vertex and push it onto a stack,
- otherwise choose a register to spill - e.g. the least-accessed one - and remove its vertex.
- Repeat until the graph is empty.
- Pop each vertex from the stack and colour it.
- Any uncoloured vertices must be spilled.

\section*{Algorithm}


\section*{Algorithm}

Choosing the right virtual register to spill will result in a faster, smaller program.

The static count of "how many accesses?" is a good start, but doesn't take account of more complex issues like loops and simultaneous liveness with other spilled values.

One easy heuristic is to treat one static access inside a loop as (say) 4 accesses; this generalises to \(4^{n}\) accesses inside a loop nested to level \(n\).

\section*{Algorithm}
"Slight lie": when spilling to memory, we (normally) need one free register to use as temporary storage for values loaded from and stored back into memory.

If any instructions operate on two spilled values simultaneously, we may need two such temporary registers to store both values.

So, in practise, when a spill is detected we may need to restart register allocation with one (or two) fewer architectural registers available so that these can be kept free for temporary storage of spilled values.

\section*{Algorithm}

When we are popping vertices from the stack and assigning colours to them, we sometimes have more than one colour to choose from.

If the program contains an instruction "MOV \(a, b\) " then storing a and b in the same arch. register (as long as they don't clash) will allow us to delete that instruction.

We can construct a preference graph to show which pairs of registers appear together in MOV instructions, and use it to guide colouring decisions.

\section*{Non-orthogonal instructions}

We can handle the situation tidily by pre-allocating a virtual register to each of the target machine's arch. registers, e.g. keep v0 in r0, v1 in r1, ..., v31 in r31.

When generating intermediate code in normal form, we avoid this set of registers, and use new ones (e.g. v32, v33, ...) for temporaries and user variables.

In this way, each architectural register is explicitly represented by a unique virtual register.

\section*{Non-orthogonal instructions}

If (hypothetically) \(A D D\) on the target architecture can only perform r0 \(=r 1+r 2\) :
\(\left.\left.\begin{array}{l}\mathrm{x}=19 ; \\
\mathrm{y}=23 ; \\
\mathrm{z}=\mathrm{x}+\mathrm{y} ;\end{array} \longrightarrow \begin{array}{l}\text { MOV v32,\#19 } \\
\text { MOV v33, \#23 }\end{array}\right\} \begin{array}{l}\text { MOV v1,v32 } \\
\text { MOV v2, v33 }\end{array}\right\}\)\begin{tabular}{l} 
ADD v0,v1,v2 \\
MOV v34, v0
\end{tabular} arch. register (e.g. x86 MUL), we generate a trailing MOV to transfer the result into a new virtual register.

\section*{Non-orthogonal instructions}

This may seem particularly wasteful, but many of the MOV instructions will be eliminated during register allocation if a preference graph is used.

```

MOV v32,\#19
MOV v33,\#23
MOV v1,v32
MOV v2,v33
ADD v0,v1,v2
MOV v34,v0

```

\section*{Non-orthogonal instructions}

This may seem particularly wasteful, but many of the MOV instructions will be eliminated during register allocation if a preference graph is used.

preference graph


\section*{Non-orthogonal instructions}

This may seem particularly wasteful, but many of the MOV instructions will be eliminated during register allocation if a preference graph is used.


\section*{Non-orthogonal instructions}

If (hypothetically) MUL on the target architecture corrupts the contents of r0:
```

MOV r1,\#6
MOV r2,\#7
MUL r0,r1,r2
\vdots

```


\section*{Procedure calling standards}

\section*{On the ARM, for example:}
- Arguments should be placed in r0-r3 before a procedure is called.
- Results should be returned in \(r 0\) and \(r 1\).
- r4-r8, r10 and r11 should be preserved over procedure calls, and r9 might be depending on the platform.
- r12-r15 are special registers, including the stack pointer and program counter.

\section*{Non-orthogonal instructions}

\section*{And finally,}
- When we know an instruction is going to corrupt the contents of an architectural register, we insert an edge on the clash graph between the corresponding virtual register and all other virtual registers live at that instruction - this prevents the register allocator from trying to store any live values in the corrupted register.

\section*{Procedure calling standards}

This final technique of synthesising edges on the clash graph in order to avoid corrupted registers is helpful for dealing with the procedure calling standard of the target architecture.

Such a standard will usually dictate that procedure calls (e.g. CALL and CALLI instructions in our 3-address code) should use certain registers for arguments and results, should preserve certain registers over a call, and may corrupt any other registers if necessary.

\section*{Procedure calling standards}

Since a procedure call instruction may corrupt some of the registers (r0-r3 and possibly r9 on the ARM), we can synthesise edges on the clash graph between the corrupted registers and all other virtual registers live at the call instruction.

As before, we may also synthesise MOV instructions to ensure that arguments and results end up in the correct registers, and use the preference graph to guide colouring such that most of these MOVs can be deleted again.

\section*{Procedure calling standards}
\(x=7\);
MOV v32,\#7
MOV v33,\#11
MOV v34,\#13
y = 11;
MOV v0,v32
MOV v1,v33
\(a=f(x, y)+z ; \quad\) CALL \(f\)
MOV v35,v0
ADD v36,v34,v35

\section*{Procedure calling standards}

```

MOV v32,\#7
MOV v33,\#11
MOV v34,\#13
MOV v0,v32
MOV v1,v33
CALL f
MOV v35,v0
ADD v36,v34,v35

```

\section*{Procedure calling standards}


MOV r0, \#7
MOV r1, \#11
MOV r4, \#13
MOV r0, r0
MOV rurr
CALL f
Mov ro, ro

\section*{Summary}
- A register allocation phase is required to assign each virtual register to an architectural one during compilation
- Registers may be allocated by colouring the vertices of a clash graph
- When the number of arch. registers is limited, some virtual registers may be spilled to memory
- Non-orthogonal instructions may be handled with additional MOVs and new edges on the clash graph
- Procedure calling standards also handled this way

\title{
Lecture 7 \\ Redundancy elimination
}

\section*{Motivation}

Some expressions in a program may cause redundant recomputation of values.

If such recomputation is safely eliminated, the program will usually become faster.

There exist several redundancy elimination optimisations which attempt to perform this task in different ways (and for different specific meanings of "redundancy").

\section*{Common subexpressions}

Common-subexpression elimination is a transformation which is enabled by available-expression analysis (AVAIL), in the same way as LVA enables dead-code elimination.

Since AVAIL discovers which expressions will have been computed by the time control arrives at an instruction in the program, we can use this information to spot and remove redundant computations.

\section*{Common subexpressions}

We consider this redundantly-computed expression to be a common subexpression: it is common to more than one instruction in the program, and in each of its occurrences it may appear as a subcomponent of some larger expression.
```

x = (a*b) +c;
\vdots
print a * b; a*b AVAILABLE

```

\section*{Algorithm}
- Find a node \(n\) which computes an alreadyavailable expression e
- Replace the occurrence of e with a new temporary variable \(t\)
- On each control path backwards from n, find the first instruction calculating \(e\) and add a new instruction to store its value into \(t\)
- Repeat until no more redundancy is found

\section*{Common subexpressions}

Recall that an expression is available at an instruction if its value has definitely already been computed and not been subsequently invalidated by assignments to any of the variables occurring in the expression.

If the expression \(e\) is available on entry to an instruction which computes e , the instruction is performing a redundant computation and can be modified or removed.

\section*{Common subexpressions}

We can eliminate a common subexpression by storing its value into a new temporary variable when it is first computed, and reusing that variable later when the same value is required.



\section*{Common subexpressions}

The program might have "got worse" as a result of performing common-subexpression elimination.

In particular, introducing a new variable increases register pressure, and might cause spilling.

Memory loads and stores are much more expensive than multiplication of registers!

\section*{Common subexpressions}

Our transformed program performs (statically) fewer arithmetic operations: \(y * z\) is now computed in three places rather than four.

However, three register copy instructions have also been generated; the program is now larger, and whether it is faster depends upon characteristics of the target architecture.

\section*{Copy propagation}

This simple formulation of CSE is fairly careless, and assumes that other compiler phases are going to tidy up afterwards.

In addition to register allocation, a transformation called copy propagation is often helpful here.

In copy propagation, we scan forwards from an \(x=y\) instruction and replace \(x\) with \(y\) wherever it appears (as long as neither x nor y have been modified).

\section*{Copy propagation}


\section*{Code motion}

Transformations such as CSE are known collectively as code motion transformations: they operate by moving instructions and computations around programs to take advantage of opportunities identified by control- and data-flow analysis.

Code motion is particularly useful in eliminating different kinds of redundancy.

It's worth looking at other kinds of code motion.

\section*{Code hoisting}

Code hoisting reduces the size of a program by moving duplicated expression computations to the same place, where they can be combined into a single instruction.

Hoisting relies on a data-flow analysis called very busy expressions (a backwards version of AVAIL) which finds expressions that are definitely going to be evaluated later in the program; these can be moved earlier and possibly combined with each other.

\section*{Code hoisting}


\section*{Loop-invariant code motion}

Some expressions inside loops are redundant in the sense that they get recomputed on every iteration even though their value never changes within the loop.

Loop-invariant code motion recognises these redundant computations and moves such expressions outside of loop bodies so that they are only evaluated once.

\section*{Loop-invariant code motion}
```

a = ...;
b = ....;
x = a + b;
while (...) {
...
}
print x;

```

Note: the loop must iterate at least once.

\section*{Code hoisting}


\section*{Code hoisting}

Hoisting may have a different effect on execution time depending on the exact nature of the code. The resulting program may be slower, faster, or just the same speed as before.

\section*{Loop-invariant code motion}
```

a = ...;
b = ....;
while (...) {
x = a + b;
}
print x;

```

\section*{Loop-invariant code motion}

This transformation depends upon a data-flow analysis to discover which assignments may affect the value of a variable ("reaching definitions").

If none of the variables in the expression are redefined inside the loop body (or are only redefined by computations involving other invariant values), the expression is invariant between loop iterations and may safely be relocated before the beginning of the loop.

\section*{Partial redundancy}

Partial redundancy elimination combines commonsubexpression elimination and loop-invariant code motion into one optimisation which improves the performance of code.

An expression is partially redundant when it is computed more than once on some (vs. all) paths through a flowgraph; this is often the case for code inside loops, for example.

\section*{Partial redundancy}
```

a = ...;
b = ....;
... = a + b;
while (...) {
a = ...;
... = a + b;
}

```

\section*{Partial redundancy}

This example gives a faster program of the same size.
Partial redundancy elimination can be achieved in its own right using a complex combination of several forwards and backwards data-flow analyses in order to locate partially redundant computations and discover the best places to add and delete instructions.

\section*{Partial redundancy}
```

a = ...;
b = ....;
while (...) {
... = a + b;
a = ...;
... = a + b;
}

```

\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Putting it all together}


\section*{Summary}
- Some optimisations exist to reduce or remove redundancy in programs
- One such optimisation, common-subexpression elimination, is enabled by AVAIL
- Copy propagation makes CSE practical
- Other code motion optimisations can also help to reduce redundancy
- These optimisations work together to improve code

\section*{Lecture 8 \\ Static single assignment and strength reduction}

\section*{Live ranges}

User variables are often reassigned and reused many times over the course of a program, so that they become live in many different places.

Our intermediate code generation scheme assumes that each user variable is kept in a single virtual register throughout the entire program.

This results in each virtual register having a large live range, which is likely to cause clashes.

\section*{Live ranges}

```

a = f(1);
b = f(2);
a, b CLASH

```
h (a,b);
- \(\begin{aligned} & b=f(3) ; \\ & c=f(4) ;\end{aligned}\)
b, c CLASH
h (b, c) ;
c \(=\mathrm{f}(5)\);
\(a=f(6) ;\)
c, a CLASH
h (c, a) ;

\section*{Live ranges}

We may remedy this situation by performing a transformation called live range splitting, in which live ranges are made smaller by using a different virtual register to store a variable's value at different times, thus reducing the potential for clashes.

\section*{Motivation}

Intermediate code in normal form permits maximum flexibility in allocating temporary variables to architectural registers.

This flexibility is not extended to user variables, and sometimes more registers than necessary will be used.

Register allocation can do a better job with user variables if we first translate code into SSA form.

\section*{Live ranges}
```

extern int f(int);

```
extern int f(int);
extern void h(int,int);
extern void h(int,int);
void g()
void g()
{
{
            int a,b,c;
            int a,b,c;
            a = f(1); b = f(2); h(a,b);
            a = f(1); b = f(2); h(a,b);
            b = f(3); c = f(4); h(b,c);
            b = f(3); c = f(4); h(b,c);
            c =f(5); a = f(6); h(c,a);
            c =f(5); a = f(6); h(c,a);
}
```

}

```


\section*{Live ranges}
```

extern int f(int);
extern void h(int,int);
void g()
{
int a1,a2, b1,b2, c1,c2;
a1 = f(1); b2 = f(2); h(a1,b2);
b1 = f(3); c2 = f(4); h(b1,c2);
c1 = f(5); a2 = f(6); h(c1,a2);
}

```

\section*{Live ranges}

\(a 1=\mathrm{f}(1)\); \(\mathrm{b} 2=\mathrm{f}(2) ; \quad \mathrm{a}, \mathrm{b} 2\) CLASH h (a1, b2) ;
b1 \(=\mathrm{f}(3)\);
\(\mathrm{c} 2=\mathrm{f}(4) ; \mathrm{b} 1, \mathrm{c} 2 \mathrm{CLASH}\)
h (b1, c2) ;

c1 \(=\mathrm{f}(5)\); a2 \(=\mathrm{f}(6) ; \mathrm{c} 1, \mathrm{a} 2\) CLASH h (c1, a2) ;

\section*{Live ranges}

\(\mathrm{a} 1=\mathrm{f}(1)\); \(\mathrm{b} 2=\mathrm{f}(2)\); h (a1, b2) ;

\(\mathrm{b} 1=\mathrm{f}(3)\);
\(\mathrm{c} 2=\mathrm{f}(4)\);
h (b1, c2) ;


2 registers needed

\section*{Static single-assignment}

Live range splitting is a useful transformation: it gives the same benefits for user variables as normal form gives for temporary variables.

However, if each virtual register is only ever assigned to once (statically), we needn't perform live range splitting, since the live ranges are already as small as possible.

Code in static single-assignment (SSA) form has this important property.

\section*{Static single-assignment}

When the program's control flow is more complex, extra effort is required to retain the original data-flow behaviour.

Where control-flow edges meet, two (or more) differently-named variables must now be merged together.

\section*{Static single-assignment}

It is straightforward to transform straight-line code into SSA form: each variable is renamed by being given a subscript, which is incremented every time that variable is assigned to.
\[
\begin{aligned}
\mathrm{V}_{1} & =3 ; \\
\mathrm{V}_{2} & =\mathrm{V}_{1}+1 ; \\
\mathrm{V}_{3} & =\mathrm{V}_{2}+\mathrm{W}_{1} ; \\
\mathrm{W}_{2} & =\mathrm{V}_{3}+2 ;
\end{aligned}
\]

\section*{Static single-assignment}


\section*{Static single-assignment}

The \(\phi\)-functions in SSA keep track of which variables are merged at control-flow join points.

They are not executable since they do not record which variable to choose (cf. gated SSA form).

\section*{Static single-assignment}
"Slight lie": SSA is useful for much more than register allocation!

In fact, the main advantage of SSA form is that, by representing data dependencies as precisely as possible, it makes many optimising transformations simpler and more effective, e.g. constant propagation, loop-invariant code motion, partial-redundancy elimination, and strength reduction.

\section*{Phase ordering}

We now have many optimisations which we can perform on intermediate code.

It is generally a difficult problem to decide in which order to perform these optimisations; different orders may be more appropriate for different programs.

Certain optimisations are antagonistic: for example, CSE may superficially improve a program at the expense of making the register allocation phase more difficult (resulting in spills to memory).

\section*{Higher-level optimisations}


\section*{Algebraic identities}

The idea behind peephole optimisation of intermediate code can also be applied to abstract syntax trees.

There are many trivial examples where one piece of syntax is always (algebraically) equivalent to another piece of syntax which may be smaller or otherwise "better"; simple rewriting of syntax trees with these rules may yield a smaller or faster program.

\section*{Algebraic identities}


\section*{Algebraic identities}

These optimisations are boring, however, since they are always applicable to any syntax tree.

We're interested in more powerful transformations which may only be applied when some analysis has confirmed that they are safe.

\section*{Algebraic identities}

In a lazy functional language,
```

let x = e in if e' then ...x... else e"'
|
if e' then let x = e in ...x... else e"
provided e' and e"' do not contain x.

```

This is still quite boring.

\section*{Strength reduction}

For example, it may be advantageous to replace multiplication \((2 * e)\) with addition
(let \(\mathrm{x}=\mathrm{e}\) in \(\mathrm{x}+\mathrm{x}\) ) as before.
Multiplication may happen a lot inside loops (e.g. using the loop variable as an index into an array), so if we can spot a recurring multiplication and replace it with an addition we should get a faster program.

\section*{Strength reduction}
```

int i; char *p;
for (i = 0; i < 100; i++)
{
p = (char *)v + 4*i;
p[0] = 0; p[1] = 0;
p[2] = 0; p[3] = 0;
}

```

\section*{Strength reduction}
```

int i; char *p;

```
int i; char *p;
p = (char *)v;
p = (char *)v;
for (i = 0; i < 100; i++)
for (i = 0; i < 100; i++)
{
{
    p[0] = 0; p[1] = 0;
    p[0] = 0; p[1] = 0;
    p[2] = 0; p[3] = 0;
    p[2] = 0; p[3] = 0;
    p += 4;
    p += 4;
}
```

}

```

\section*{Strength reduction}
```

int i; char *p;
p = (char *)v;
for (i = 0; p < (char *)v + 400; i++)
{
p[0] = 0; p[1] = 0;
p[2] = 0; p[3] = 0;
p += 4;
}

```

\section*{Strength reduction}
```

int i; int *p;
p = v;
for (i = 0; p < v + 100; i++)
{
*p = 0;
p++;
}

```

\section*{Strength reduction}
```

int *p;
for (p = v; p < v + 100; p++)
{
*p = 0;
}

```

Multiplication has been replaced with addition.

\section*{Strength reduction}

We are not restricted to replacing multiplication with addition, as long as we have
- induction variable: \(\mathrm{i}=\mathrm{i} \oplus \mathrm{c}\)
- another variable: \(j=C_{2} \oplus\left(C_{1} \otimes i\right)\)
for some operations \(\oplus\) and \(\otimes\) such that \(x \otimes(y \oplus z)=(x \otimes y) \oplus(x \otimes z)\)

\section*{Strength reduction}
```

int i; int *p;
p = v;?
for (i= 0; p < v + 100; it+)
{
*p = 0
p++;
}

```

\section*{Strength reduction}

Note that, while this code is now almost optimal, it has obfuscated the intent of the original program.

Don't be tempted to write code like this!
For example, when targeting a 64-bit architecture, the compiler may be able to transform the original loop into fifty 64 -bit stores, but will have trouble with our more efficient version.

\section*{Strength reduction}

It might be easier to perform strength reduction on the intermediate code, but only if annotations have been placed on the flowchart to indicate loop structure.

At the syntax tree level, all loop structure is apparent.

\section*{Summary}
- Live range splitting reduces register pressure
- In SSA form, each variable is assigned to only once
- SSA uses \(\phi\)-functions to handle control-flow merges
- SSA aids register allocation and many optimisations
- Optimal ordering of compiler phases is difficult
- Algebraic identities enable code improvements
- Strength reduction uses them to improve loops

\title{
Lecture 9 Abstract interpretation
}

\section*{Motivation}

We reason about programs statically, but we are really trying to make predictions about their dynamic behaviour.

Why not examine this behaviour directly?
It isn't generally feasible (e.g. termination, inputs) to run an entire computation at compile-time, but we can find things out about it by running a simplified version.

This is the basic idea of abstract interpretation.

\section*{Abstract interpretation}

Warning: this will be a heavily simplified view of abstract interpretation; there is only time to give a brief introduction to the ideas, not explore them with depth or rigour.

\section*{Abstract interpretation}

For example, to plan a trip, you might use a map.
- A road map sacrifices a lot of detail -
- trees, road conditions, individual buildings;
- an entire dimension -
- but it retains most of the information which is important for planning a journey:
- place names;
- roads and how they are interconnected.

\section*{Abstract interpretation}

Trying to plan a journey by exploring the concrete world instead of the abstraction (i.e. driving around aimlessly) is either very expensive or virtually impossible.

A trustworthy map makes it possible - even easy.
This is a real application of abstract interpretation, but in this course we're more interested in computer programs.

\section*{Abstract interpretation}

The key idea is to use an abstraction: a model of (otherwise unmanageable) reality, which
- discards enough detail that the model becomes manageable (e.g. small enough, computable enough), but
- retains enough detail to provide useful insight into the real world.

\section*{Abstract interpretation}

Crucially, a road map is a useful abstraction because the route you plan is probably still valid back in reality.
- A cartographer creates an abstraction of reality (a map),
- you perform some computation on that abstraction (plan a route),
- and then you transfer the result of that computation back into the real world (drive to your destination).

\section*{Multiplying integers}

A canonical example is the multiplication of integers.
If we want to know whether \(-1515 \times 37\) is positive or negative, there are two ways to find out:
- Compute in the concrete world (arithmetic), using the standard interpretation of multiplication. \(-15 \mathrm{I} 5 \times 37=-56055\), which is negative.
- Compute in an abstract world, using an abstract interpretation of multiplication: call it \(\otimes\).

\section*{Multiplying integers}

In this example the magnitudes of the numbers are insignificant; we care only about their sign, so we can use this information to design our abstraction.
\((-)=\{z \in \mathbb{Z} \mid z<0\}\)
(0) \(=\{0\}\)
\((+)=\{z \in \mathbb{Z} \mid z>0\}\)
In the concrete world we have all the integers; in the abstract world we have only the values ( - ), ( 0 ) and ( + ).

\section*{Multiplying integers}

Armed with our abstraction, we can now tackle the original problem.
\[
\begin{aligned}
a b s(-1515) & =(-) \\
a b s(37) & =(+) \\
(-) \otimes(+) & =(-)
\end{aligned}
\]

So, without doing any concrete computation, we have discovered that \(-1515 \times 37\) has a negative result.

\section*{Safety}

As always, there are important safety issues.
Because an abstraction discards detail, a computation in the abstract world will necessarily produce less precise results than its concrete counterpart.

It is important to ensure that this imprecision is safe.

\section*{Adding integers}

A good example is the addition of integers.
How do we define the abstract operator \(\oplus\) ?
\begin{tabular}{c|ccc}
\(\oplus\) & \((-)\) & \((0)\) & \((+)\) \\
\hline\((-)\) & \((-)\) & \((-)\) & \((?)\) \\
\((0)\) & \((-)\) & \((0)\) & \((+)\) \\
\((+)\) & \((?)\) & \((+)\) & \((+)\)
\end{tabular}

\section*{Multiplying integers}

We need to define the abstract operator \(\otimes\). Luckily, we have been to primary school.
\begin{tabular}{c|ccc}
\(\otimes\) & \((-)\) & \((0)\) & \((+)\) \\
\hline\((-)\) & \((+)\) & \((0)\) & \((-)\) \\
\((0)\) & \((0)\) & \((0)\) & \((0)\) \\
\((+)\) & \((-)\) & \((0)\) & \((+)\)
\end{tabular}

\section*{Multiplying integers}

This is just a toy example, but it demonstrates the methodology: state a problem, devise an abstraction that retains the characteristics of that problem, solve the problem in the abstract world, and then interpret the solution back in the concrete world.

This abstraction has avoided doing arithmetic; in compilers, we will mostly be interested in avoiding expensive computation, nontermination or undecidability.

\section*{Safety}

We consider a particular abstraction to be safe if, whenever a property is true in the abstract world, it must also be true in the concrete world.

Our multiplication example is actually quite precise, and therefore trivially safe: the magnitudes of the original integers are irrelevant, so when the abstraction says that the result of a multiplication will be negative, it definitely will be.

In general, however, abstractions will be more approximate than this.

\section*{Adding integers}

When adding integers, their (relative) magnitudes are important in determining the sign of the result, but our abstraction has discarded this information.

As a result, we need a new abstract value: (?).
\((-)=\{z \in \mathbb{Z} \mid z<0\}\)
\((0)=\{0\}\)
\((+)=\{z \in \mathbb{Z} \mid z>0\}\)
(?) \(=\mathbb{Z}\)

\section*{Adding integers}
(?) is less precise than \((-),(0)\) and \((+)\); it means "I don't know", or "it could be anything".

Because we want the abstraction to be safe, we must put up with this weakness.

\section*{Adding integers}
\(a b s(-1515+37)=a b s(-1478)=(-)\)
\(a b s(-15 \mathrm{I} 5) \oplus a b s(37)=(-) \oplus(+)=(?)\)


\section*{Abstraction}

Formally, an abstraction of some concrete domain \(D\) (e.g. \(\wp(\mathbb{Z}))\) consists of
- an abstract domain \(D^{\#}(e . g .\{(-),(0),(+),(?)\})\),
- an abstraction function \(\alpha: D \rightarrow D^{\#}\) (e.g. \(a b s\) ), and
- a concretisation function \(\gamma: D^{\#} \rightarrow D\), e.g.:
- \((-) \mapsto\{z \in \mathbb{Z} \mid z<0\}\),
- (0) \(\mapsto\{0\}\), etc.

\section*{Adding integers}



\section*{Safety}

Here, safety is represented by the fact that \((-) \subseteq\) (?):
\[
\{z \in \mathbb{Z} \mid z<0\} \subseteq \mathbb{Z}
\]

The result of doing the abstract computation is less precise, but crucially includes the result of doing the concrete computation (and then abstracting), so the abstraction is safe and hasn't missed anything.


\section*{Abstraction}

So, for \(D=\wp(\mathbb{Z})\) and \(D^{\#}=\{(-),(0),(+),(?)\}\), we have:

where \(\alpha_{1,2}\) and \(\gamma_{1,2}\) are the appropriate abstraction and concretisation functions.

\section*{Abstraction}

These mathematical details are formally important, but are not examinable on this course.

Abstract interpretation can get very theoretical, but what's significant is the idea of using an abstraction to safely model reality.

Recognise that this is what we were doing in dataflow analysis: interpreting 3 -address instructions as operations on abstract values - e.g. live variable sets - and then "executing" this abstract program.

\section*{Summary}
- Abstractions are manageably simple models of unmanageably complex reality
- Abstract interpretation is a general technique for executing simplified versions of computations
- For example, the sign of an arithmetic result can be sometimes determined without doing any arithmetic
- Abstractions are approximate, but must be safe
- Data-flow analysis is a form of abstract interpretation

\section*{Lecture 10 \\ Strictness analysis}

\section*{Call-by-value evaluation}

Strict ("eager") functional languages (e.g. ML) use a call-by-value evaluation strategy:
\(\frac{e_{2} \Downarrow v_{2} e_{1}\left[v_{2} / x\right] \Downarrow v_{1}}{\left(\lambda x . e_{1}\right) e_{2} \Downarrow v_{1}}\)
- Efficient in space and time, but
- might evaluate more arguments than necessary.

\section*{Call-by-need evaluation}

One simple optimisation is to use call-by-need evaluation instead of call-by-name.

If the language has no side-effects, duplicated instances of an argument can be shared, evaluated once if required, and the resulting value reused.

This avoids recomputation and is better than call-byname, but is still more expensive than call-by-value.

\section*{Call-by-need evaluation}
```

plus(x,y) = if x=0 then y else plus(x-1,y+1)
Using call-by-need:
plus $(3,4) \mapsto$ if $3=0$ then 4 else plus (3-1,4+1)
$\mapsto$ plus (3-1,4+1)
$\mapsto$ plus (2-1, 4+1+1)
$\mapsto$ plus $(1-1,4+1+1+1)$
$\rightarrow 4+1+1+1$
$\mapsto 5+1+1$
$\rightarrow 6+1$
$\mapsto 7$

```

\section*{Motivation}

The operations and control structures of imperative languages are strongly influenced by the way most real computer hardware works.

This makes imperative languages relatively easy to compile, but (arguably) less expressive; many people use functional languages, but these are harder to compile into efficient imperative machine code.

Strictness optimisation can help to improve the efficiency of compiled functional code.

\section*{Call-by-name evaluation}

Non-strict ("lazy") functional languages (e.g. Haskell) use a call-by-name evaluation strategy:
\(\frac{e_{1}\left[e_{2} / x\right] \Downarrow v}{\left(\lambda x \cdot e_{1}\right) e_{2} \Downarrow v}\)
- Only evaluates arguments when necessary, but
- copies (and redundantly re-evaluates) arguments.

\section*{Call-by-need evaluation}
```

plus (x,y) = if x=0 then y else plus (x-1,y+1)

```

\section*{Using call-by-value:}
```

plus(3,4) }\mapsto\mathrm{ if 3=0 then 4 else plus(3-1,4+1)

```
\(\mapsto\) plus \((2,5)\)
\(\mapsto\) plus \((1,6)\)
\(\mapsto\) plus \((0,7)\)
\(\mapsto 7\)

\section*{Replacing CBN with CBV}

So why not just replace call-by-name with call-by-value?
Because, while replacing call-by-name with call-by-need never changes the semantics of the original program (in the absence of side-effects), replacing CBN with CBV does.

In particular, the program's termination behaviour changes.

\section*{Replacing CBN with CBV}

Assume we have some nonterminating expression, \(\Omega\).
- Using CBN, the expression ( \(\lambda_{x}\). 42) \(\Omega\) will evaluate to 42 .
- But using CBV, evaluation of ( \(\lambda x\). 42) \(\Omega\) will not terminate: \(\Omega\) gets evaluated first, even though its value is not needed here.

We should therefore try to use call-by-value wherever possible, but not when it will affect the termination behaviour of a program.

\section*{Neededness}

These needed arguments can safely be passed by value: if their evaluation causes nontermination, this will just happen sooner rather than later.

\section*{Strictness}

What we really want is a more refined notion:
It is safe to pass an argument by value when the function fails to terminate whenever the argument fails to terminate.

When this more general statement holds, we say the function is strict in that argument.

\section*{\(\lambda x, y, z\). if \(x\) then \(y\) else \(\Omega\)} is strict in \(x\) and strict in \(y\).

\section*{Neededness}

Intuitively, it will be safe to use CBV in place of CBN whenever an argument is definitely going to be evaluated.

We say that an argument is needed by a function if the function will always evaluate it.
- \(\lambda x, y, x+y\) needs both its arguments.
- \(\lambda x, y \cdot x+\mid\) needs only its first argument.
- \(\lambda x, y .42\) needs neither of its arguments.

\section*{Neededness}

In fact, neededness is too conservative:

\section*{\(\lambda x, y, z\). if \(x\) then \(y\) else \(\Omega\)}

This function might not evaluate \(y\), so only \(x\) is needed.
But actually it's still safe to pass y by value: if \(y\) doesn't get evaluated then the function doesn't terminate anyway, so it doesn't matter if eagerly evaluating \(y\) causes nontermination.

\section*{Strictness}

If we can develop an analysis that discovers which functions are strict in which arguments, we can use that information to selectively replace CBN with CBV and obtain a more efficient program.

\section*{Recursion equations}
\[
\begin{aligned}
& F_{1}\left(x_{1}, \ldots, x_{k_{1}}\right)=e_{1} \\
& \cdots=\cdots \\
& F_{n}\left(x_{1}, \ldots, x_{k_{n}}\right)=e_{n} \\
& e::=x_{i}\left|A_{i}\left(e_{1}, \ldots, e_{r_{i}}\right)\right| F_{i}\left(e_{1}, \ldots e_{k_{i}}\right)
\end{aligned}
\]
where each \(A_{i}\) is a symbol representing a built-in (predefined) function of arity \(r_{i}\).

\section*{Recursion equations}

For our earlier example,
```

plus(x,y) = if x=0 then y else plus(x-1,y+1)

```
we can write the recursion equation
\(\operatorname{plus}(x, y)=\operatorname{cond}(e q(x, 0), y, \operatorname{plus}(\operatorname{sub1}(x), \operatorname{add1}(y)))\)
where cond, eq, 0 , sub/ and add/ are built-in functions.

\section*{Standard interpretation}

Each built-in function needs a standard interpretation.
We will interpret each \(A_{i}\) as a function \(a_{i}\) on values in \(D\) :
```

cond}(\perp,x,y)=
cond}(0,x,y)=
cond}(p,x,y)=x otherwis
eq(\perp,y) = \perp
eq(x,\perp) = \perp

```


\section*{Abstract interpretation}

Our abstraction must capture the properties we're interested in, while discarding enough detail to make analysis computationally feasible.

Strictness is all about termination behaviour, and in fact this is all we care about: does evaluation of an expression definitely not terminate (as with \(\Omega\) ), or may it eventually terminate and return a result?

Our abstract domain \(D^{\#}\) is therefore \(\{0, I\}\).

\section*{Abstract interpretation}

A formal relationship exists between the standard and abstract interpretations of each built-in function; the mathematical details are in the lecture notes.

Informally, we use the same technique as for multiplication and addition of integers in the last lecture: we define the abstract operations using what we know about the behaviour of the concrete operations.

\section*{Standard interpretation}

We must have some representation of nontermination in our concrete domain.

As values we will consider the integer results of terminating computations, \(\mathbb{Z}\), and a single extra value to represent nonterminating computations: \(\perp\).

Our concrete domain \(D\) is therefore \(\mathbb{Z}_{\perp}=\mathbb{Z} \cup\{\perp\}\).

\section*{Standard interpretation}

The standard interpretation \(f_{i}\) of a user-defined function \(F_{i}\) is constructed from the built-in functions by composition and recursion according to its defining equation.
\(\operatorname{plus}(x, y)=\operatorname{cond}(e q(x, 0), y, \operatorname{plus}(\operatorname{sub1} 1(x), \operatorname{add1}(y)))\)

\section*{Abstract interpretation}

For each built-in function \(A_{i}\) we need a corresponding strictness function \(a_{i}^{\#}\) - this provides the strictness interpretation for \(A_{i}\).

Whereas the standard interpretation of each built-in is a function on concrete values from \(D\), the strictness interpretation of each will be a function on abstract values from \(D^{\#}\) (i.e. 0 and I).

\section*{Abstract interpretation}
\begin{tabular}{c|c|c}
\(x\) & \(y\) & eq\(^{\#}(x, y)\) \\
\hline 0 & 0 & 0 \\
\hline 0 & 1 & 0 \\
\hline 1 & 0 & 0 \\
\hline 1 & 1 & 1
\end{tabular}

\section*{Abstract interpretation}
\begin{tabular}{c|c|c|c}
\(p\) & \(x\) & \(y\) & cond \(^{\#}(p, x, y)\) \\
\hline 0 & 0 & 0 & 0 \\
\hline 0 & 0 & 1 & 0 \\
\hline 0 & I & 0 & 0 \\
\hline 0 & I & 1 & 0 \\
\hline I & 0 & 0 & 0 \\
\hline I & 0 & 1 & 1 \\
\hline I & I & 0 & 1 \\
\hline I & I & 1 & 1
\end{tabular}

\section*{Abstract interpretation}

These functions may be expressed more compactly as boolean expressions, treating 0 and I from \(D^{\#}\) as false and true respectively.

We can use Karnaugh maps (from IA DigElec) to turn each truth table into a simple boolean expression.

\section*{Abstract interpretation}

\[
e q^{\#}(x, y)=x \wedge y
\]

\section*{Strictness analysis}

The point of this analysis is to discover the missing piece: what is the strictness function \(f_{i}{ }^{\#}\) corresponding to each user-defined \(F_{i}\) ?

These strictness functions will show us exactly how each \(F_{i}\) is strict with respect to each of its arguments - and that's the information that tells us where we can replace lazy, CBN-style parameter passing with eager CBV.

\section*{Strictness analysis}
plus \({ }^{\sharp}(x, y)=\operatorname{cond}^{\sharp}\left(e q^{\sharp}\left(x, 0^{\sharp}\right), y, p l u s^{\sharp}\left(s u b 1^{\sharp}(x), a d d 1^{\sharp}(y)\right)\right)\)
We already know all the other strictness functions:
\[
\begin{aligned}
\operatorname{cond}^{\sharp}(p, x, y) & =p \wedge(x \vee y) \\
e q^{\sharp}(x, y) & =x \wedge y \\
0^{\sharp} & =1 \\
\operatorname{sub} 1^{\sharp}(x) & =x \\
\operatorname{add} 1^{\sharp}(x) & =x
\end{aligned}
\]

So we can use these to simplify the expression for plus\#.

\section*{Strictness analysis}
```

plus }\mp@subsup{}{}{\sharp}(x,y)=\mp@subsup{\operatorname{cond}}{}{\sharp}(e\mp@subsup{q}{}{\sharp}(x,\mp@subsup{0}{}{\sharp}),y,plu\mp@subsup{s}{}{\sharp}(sub1\mp@subsup{1}{}{\sharp}(x),add\mp@subsup{1}{}{\sharp}(y))
= eq}\mp@subsup{}{}{\sharp}(x,\mp@subsup{0}{}{\sharp})\wedge(y\vee\mp@subsup{plus}{\#}{\sharp}(\mathrm{ sub1 }\mp@subsup{1}{}{\sharp}(x),add1 1 (y))
=eq}\mp@subsup{q}{}{\sharp}(x,1)\wedge(y\veeplus\mp@subsup{s}{}{\sharp}(x,y)
= x^1\wedge(y\vee plus }\mp@subsup{}{}{\sharp}(x,y)
= x^(y\vee plus

```

\section*{Algorithm}
for \(i=1\) to \(n\) do \(\mathrm{f} \#[\mathrm{i}]:=\lambda \vec{x} .0\)
while (f\#[] changes) do for \(i=1\) to \(n\) do
\(\mathrm{f} \#[\mathrm{i}]:=\lambda \vec{x} . \mathrm{e}_{\mathrm{i}}{ }^{\#}\)
\(e_{i}^{\#}\) means " \(e_{i}\) (from the recursion equations) with each \(A_{j}\) replaced with \(a_{j}^{\#}\) and each \(F_{j}\) replaced with \(\mathrm{f} \#[j\) ]".

\section*{Algorithm}

On the first iteration, we calculate e। \({ }^{\#}\) :
- The recursion equations say \(e_{I}=\operatorname{cond}(e q(x, 0), y\), plus(subl(x), addI(y)))
- The current contents of \(\mathrm{f} \#\left[\right.\) ] say \(\mathrm{fi}^{\#}\) is \(\lambda x, y .0\)
- So:
\(\mathrm{e}^{\#}=\operatorname{cond}^{\#}\left(\mathrm{eq}^{\#}\left(x, 0^{\#}\right), y,(\lambda x, y, 0)\left(\operatorname{sub}^{\#}(x)\right.\right.\), add \(\left.\left.I^{\#}(y)\right)\right)\)

\section*{Algorithm}

So, at the end of the first iteration,
\[
\mathrm{f} \#[1]:=\lambda x, y \cdot x \wedge y
\]

\section*{Strictness analysis}
\[
p l u s^{\sharp}(x, y)=x \wedge\left(y \vee p l u s^{\sharp}(x, y)\right)
\]

This is a recursive definition, and so unfortunately doesn't provide us with the strictness function directly.

We want a definition of plus\# which satisfies this equation - actually we want the least fixed point of this equation, which (as ever!) we can compute iteratively.

We have only one user-defined function, plus, and so only one recursion equation:
\(\operatorname{plus}(x, y)=\operatorname{cond}(e q(x, 0), y, \operatorname{plus}(\operatorname{sub1} 1(x), \operatorname{add} 1(y)))\)

We initialise the corresponding element of our f \# [ ] array to contain the always-0 strictness function of
array to contain the always- 0 strictn
the appropriate arity:
\[
\mathrm{f} \#[1]:=\lambda x, y .0
\]

\section*{Algorithm}

\section*{Algorithm}
\(\mathrm{e}^{\#}=\operatorname{cond}^{\#}\left(\mathrm{eq}^{\#}\left(x, 0^{\#}\right), y,(\lambda x, y \cdot 0)\left(s_{1} I^{\#}(x), a d d I^{\#}(y)\right)\right)\)
Simplifying:
\[
e_{1}^{\#}=\operatorname{cond}^{\#}\left(e q^{\#}\left(x, 0^{\#}\right), y, 0\right)
\]

Using definitions of cond \({ }^{\#}, q^{\#}\) and \(0^{\#}\) :
\[
e I^{\#}=(x \wedge I) \wedge(y \vee 0)
\]

Simplifying again:
\[
\mathrm{e}_{1}{ }^{\#}=x \wedge y
\]

\section*{Algorithm}

On the second iteration, we recalculate el \({ }^{\#}\) :
- The recursion equations still say \(e_{I}=\operatorname{cond}(e q(x, 0), y\), plus(subl(x), add \(\left.I(y))\right)\)
- The current contents of \(\mathrm{f} \#\left[\mathrm{]}\right.\) say \(\mathrm{fi}^{\#}\) is \(\lambda x, y . x \wedge y\)
- So:
\(\mathrm{e}^{\#}{ }^{\#}=\operatorname{cond}^{\#}\left(\mathrm{eq}^{\#}\left(x, 0^{\#}\right), y,(\lambda x, y \cdot x \wedge y)\left(\operatorname{sub} I^{\#}(x), a d d I^{\#}(y)\right)\right)\)

\section*{Algorithm}
\[
\mathrm{e} \mathrm{I}^{\#}=\operatorname{cond} \mathrm{d}^{\#}\left(\mathrm{eq} q^{\#}\left(x, 0^{\#}\right), y,(\lambda x, y \cdot x \wedge y)\left(\operatorname{sub} I^{\#}(x), a d d I^{\#}(y)\right)\right)
\]

Simplifying:
\[
e I^{\#}=\operatorname{cond} d^{\#}\left(q^{\#}\left(x, 0^{\#}\right), y, \operatorname{sub} I^{\#}(x) \wedge a d d I \#(y)\right)
\]

Using definitions of cond\#, eq\#, \(0^{\#,}\) sub।\# and addI\#:
\[
\mathrm{e} \mathrm{l}^{\#}=(x \wedge \mathrm{I}) \wedge(y \vee(x \wedge y))
\]

Simplifying again:
\[
e 1^{\#}=x \wedge y
\]

\section*{Algorithm}

So, at the end of the second iteration,
\[
\mathrm{f} \#[1]:=\lambda x, y \cdot x \wedge y
\]

This is the same result as last time, so we stop.

\section*{Optimisation}

So now, finally, we can see that
\[
\begin{gathered}
\operatorname{plus}^{\#}(\mathrm{I}, 0)=\mathrm{I} \wedge 0=0 \\
\text { and } \\
\operatorname{plus}^{\#}(0, \mathrm{I})=0 \wedge \mathrm{I}=0
\end{gathered}
\]
which means our concrete plus function is strict in its first argument and strict in its second argument: we may always safely use \(C B V\) when passing either.

\section*{Summary}
- Functional languages can use CBV or CBN evaluation
- CBV is more efficient but can only be used in place of CBN if termination behaviour is unaffected
- Strictness shows dependencies of termination
- Abstract interpretation may be used to perform strictness analysis of user-defined functions
- The resulting strictness functions tell us when it is safe to use CBV in place of CBN

\title{
Lecture II Constraint-based analysis
}

\section*{Imprecise control flow}


\section*{OCFA}

0CFA - "zeroth-order control-flow analysis" - is a constraint-based analysis for discovering which values may reach different places in a program.

When functions (or pointers to functions) are present, this provides information about which functions may potentially be called at each call site.

We can then build a more precise call graph.

\section*{Specimen program}

\section*{let id \(=\lambda x . x\) in id id 7}

\section*{Motivation}

Intra-procedural analysis depends upon accurate control-flow information.

In the presence of certain language features (e.g. indirect calls) it is nontrivial to predict accurately how control may flow at execution time - the naïve strategy is very imprecise.

A constraint-based analysis called 0CFA can compute a more precise estimate of this information.

\section*{Constraint-based analysis}

Many of the analyses in this course can be thought of in terms of solving systems of constraints.

For example, in LVA, we generate equality constraints from each instruction in the program:
```

in-live}(m)=(\mathrm{ out-live }(m)<br>operatorname{def}(m))\cup\operatorname{ref}(m
out-live(m) = in-live(n) \cup in-live(o)
in-live(n)=(out-live(n)}<br>operatorname{def}(n))\cupref(n

```
and then iteratively compute their minimal solution.

\section*{Specimen language}

Functional languages are a good candidate for this kind of analysis; they have functions as first-class values, so control flow may be complex.

We will use a minimal syntax for expressions:
\[
\mathrm{e}::=x|c| \lambda x . \mathrm{e}\left|\mathrm{e}_{1} \mathrm{e}_{2}\right| \text { let } x=\mathrm{e}_{1} \text { in } \mathrm{e}_{2}
\]

A program in this language is a closed expression.

\section*{Program points}
let id \(=\lambda x . x\) in id id 7


\section*{Program points}
\(\left(\text { let id }{ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\mid}\)


\section*{Program points}
\(\left(\text { let } \mathrm{id}^{2}=\left(\lambda x^{4} . \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{1}\)
The precise value of each \(\alpha_{i}\) is undecidable in general, so our analysis will compute a safe overapproximation.

From the structure of the program we can generate a set of constraints on the flow variables, which we can then treat as data-flow inequations and iteratively compute their least solution.

\section*{Generating constraints}
\(\left(\text { let id }{ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}\)
\[
\mathfrak{c}^{3} \longrightarrow \alpha_{a} \supseteq\left\{c^{a}\right\}
\]

\section*{Generating constraints}
\(\left(\text { let } \mathrm{id}^{2}=\left(\lambda x^{4} \cdot \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {। }}\)
\(\left(\lambda x^{a} \cdot e^{b}\right) \quad \alpha_{c} \supseteq\left\{\left(\lambda x^{a} \cdot e^{b}\right)^{c}\right\}\)
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)

\section*{Program points}
\(\left(\text { let } \mathrm{id}^{2}=\left(\lambda \mathrm{x}^{4} \cdot \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}\)
Each program point \(i\) has an associated flow variable \(\alpha_{i}\).
Each \(\alpha_{i}\) represents the set of flow values which may be yielded at program point \(i\) during execution.

For this language the flow values are integers and function closures; in this particular program, the only values available are \(7^{10}\) and \(\left(\lambda x^{4} . x^{5}\right)^{3}\).

\section*{Generating constraints} \(\left(\text { let } \mathrm{id}^{2}=\left(\lambda x^{4} . \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}\)

\section*{Generating constraints}
\(\left(\text { let } \mathrm{id}^{2}=\left(\lambda \mathrm{x}^{4} \cdot \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}\)
\[
7^{10} \longrightarrow \alpha_{10} \supseteq\left\{7^{10}\right\}
\]
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)

\section*{Generating constraints} \(\left(\text { let id }{ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}\)
\(\left(\lambda x^{4} \cdot x^{5}\right)^{3} \longrightarrow \alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)

\section*{Generating constraints} \(\left(\right.\) let id \({ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3}\) in \(\left.\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)\)

\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)

\section*{Generating constraints} \(\left(\text { let } \mathrm{id}{ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{1}\)

\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)
\(\alpha_{8} \supseteq \alpha_{2}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{9} \supseteq \alpha_{2}\)
\(\alpha_{5} \supseteq \alpha_{4}\)

\section*{Generating constraints}
\(\left(\text { let } \mathrm{id}^{2}=\left(\lambda \mathrm{x}^{4} \cdot \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{1}\)
\[
\lambda x^{4} \ldots x^{5} \ldots \longrightarrow \alpha_{5} \supseteq \alpha_{4}
\]
\[
\text { let } \mathrm{id}^{2}=\ldots \mathrm{id}^{8} \ldots \longrightarrow \boldsymbol{\alpha}_{8} \supseteq \boldsymbol{\alpha}_{2}
\]
\[
\text { let } \mathrm{id}^{2}=\ldots \mathrm{id}^{9} \ldots \longrightarrow \alpha_{9} \supseteq \boldsymbol{\alpha}_{2}
\]
\begin{tabular}{ll}
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\) & \(\alpha_{8} \supseteq \alpha_{2}\) \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) &
\end{tabular}

\section*{Generating constraints} \(\left(\text { let } \mathrm{id}^{2}=\left(\lambda \mathrm{x}^{4} \cdot \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}\)

\(\alpha_{10} \supseteq\{710\}\)
\(\alpha_{8} \supseteq \alpha_{2}\)
\(\alpha_{2} \supseteq \alpha_{3}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{5} \supseteq \alpha_{4}\)
\(\alpha_{9} \supseteq \alpha_{2}\)
\(\alpha_{1} \supseteq \alpha_{6}\)

\section*{Generating constraints} \(\left(\right.\) let id \({ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3}\) in \(\left.\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)\)

\(\alpha_{8} \supseteq \alpha_{2}\)
\(\alpha_{2} \supseteq \alpha_{3}\)
\(\alpha_{9} \supseteq \alpha_{2}\)
\(\alpha_{5} \supseteq \alpha_{4}\)
\(\alpha_{\mid} \supseteq \alpha_{6}\)

\section*{Generating constraints} \(\left(\right.\) let id \({ }^{2}=\left(\lambda x^{4} . x^{5}\right)^{3}\) in \(\left.\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)\)
\begin{tabular}{rlr}
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\) & \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{1} \supseteq \alpha_{6}\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{2} \supseteq \alpha_{3}\) \\
\(\alpha_{8} \supseteq \alpha_{2}\) & \(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \\
\(\alpha_{9} \supseteq \alpha_{2}\) & \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)
\end{tabular}

\section*{Generating constraints} \(\left(\text { let id }{ }^{2}=\left(\lambda x^{4} \cdot x^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{1}\)
\(\left(\_^{8}-{ }^{9}\right)^{7} \longrightarrow\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\)
\(\left(\sim^{7}-^{10}\right)^{6} \longrightarrow\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{8} \supseteq \alpha_{2}\)
\(\alpha_{9} \supseteq \alpha_{2}\)
\(\alpha_{2} \supseteq \alpha_{3}\)
\(\alpha_{5} \supseteq \alpha_{4}\)
\(\alpha_{1} \supseteq \alpha_{6}\)
\(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\)
\(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)

\section*{Solving constraints}
\begin{tabular}{llr}
\(\alpha_{10} \supseteq\{710\}\) & \(\alpha_{8} \supseteq \alpha_{2}\) & \(\alpha_{2} \supseteq \alpha_{3}\) \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) & \(\left(\alpha_{9 \mapsto} \mapsto \alpha_{7} \supseteq \alpha_{8}\right.\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{1 \supseteq \alpha_{6}}\) & \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\) \\
\(\alpha_{1}=\{ \}\) & \(\alpha_{6}=\{ \}\) \\
\(\alpha_{2}=\{ \}\) & \(\alpha_{7}=\{ \}\) \\
\(\alpha_{3}=\{ \}\) & \(\alpha_{8}=\{ \}\) \\
\(\alpha_{4}=\{ \}\) & \(\alpha_{9}=\{ \}\) \\
\(\alpha_{5}=\{ \}\) & \(\alpha_{10}=\{ \}\)
\end{tabular}

\section*{Solving constraints}
\(\alpha_{10} \supseteq\{710\}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{5} \supseteq \alpha_{4}\)
\(\alpha_{8} \supseteq \alpha_{2}\) \(\alpha_{9} \supseteq \alpha_{2}\) \(\alpha_{1} \supseteq \alpha_{6}\)
\(\alpha_{2} \supseteq \alpha_{3}\)
\(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)
\(\boldsymbol{\alpha}_{1}=\{ \}\)
\(\alpha_{6}=\{ \}\)
\(\boldsymbol{\alpha}_{2}=\{ \}\)
\(\alpha_{7}=\{ \}\)
\(\alpha_{3}=\{ \}\)
\(\alpha_{8}=\{ \}\)
\(\alpha_{9}=\{ \}\)
\(\alpha_{10}=\left\{7^{10}\right\}\)

\section*{Solving constraints}


\section*{Solving constraints}
\begin{tabular}{llr}
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\) & \(\alpha_{8} \supseteq \alpha_{2}\) & \(\alpha_{2} \supseteq \alpha_{3}\) \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) & \(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{1} \supseteq \alpha_{6}\) & \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)
\end{tabular}
\begin{tabular}{ll}
\(\alpha_{1}=\{ \}\) & \(\alpha_{6}=\{ \}\) \\
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{7}=\{ \}\) \\
\(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{8}=\{ \}\) \\
\(\alpha_{4}=\{ \}\) & \(\alpha_{9}=\{ \}\) \\
\(\alpha_{5}=\{ \}\) & \(\alpha_{10}=\left\{7^{10}\right\}\)
\end{tabular}

\section*{Solving constraints}
\begin{tabular}{llr}
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\) & \(\alpha_{8} \supseteq \alpha_{2}\) & \(\alpha_{2} \supseteq \alpha_{3}\) \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) & \begin{tabular}{rl}
\(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{1} \supseteq \alpha_{6}\)
\end{tabular} \\
\(\alpha_{1}=\{ \}\) & \(\alpha_{6}=\{ \}\) \\
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{7}=\{ \}\) \\
\(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\(\alpha_{4}=\{ \}\) & \(\alpha_{9}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\(\alpha_{5}=\{ \}\) & \(\alpha_{10}=\{710\}\)
\end{tabular}

\section*{Solving constraints}
\begin{tabular}{crr}
\(\alpha_{10} \supseteq\left\{7_{10}\right\}\) & \(\alpha_{8} \supseteq \alpha_{2}\) & \(\alpha_{2} \supseteq \alpha_{3}\) \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) & \(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{1} \supseteq \alpha_{6}\) & \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\) \\
\hdashline\(\alpha_{4} \supseteq \alpha_{9}\) & \(\alpha_{7} \supseteq \alpha_{5}\) &
\end{tabular}
\(\alpha_{1}=\{ \}\)
\[
\alpha_{6}=\{ \}
\]
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{7}=\{ \}\)
\(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{4}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{9}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{5}=\{ \}\)
\(\alpha_{10}=\left\{7^{10}\right\}\)

\section*{Solving constraints}


\section*{Solving constraints}
\(\begin{aligned} & \alpha_{10} \supseteq\left\{7^{10}\right\} \\ & \alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\} \\ & \alpha_{5} \supseteq \alpha_{4}\end{aligned}\)
\(\alpha_{8} \supseteq \alpha_{2}\) \(\alpha_{9} \supseteq \alpha_{2}\) \(\alpha_{1} \supseteq \alpha_{6}\)
\(\alpha_{2} \supseteq \alpha_{3}\)
\(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\)
\(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)
\(\alpha_{1}=\{ \}\)
\(\alpha_{6}=\{ \}\)
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{7}=\{ \}\)
\(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{4}=\{ \}\)
\(\alpha_{5}=\{ \}\)
\(\alpha_{9}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{10}=\left\{7^{10}\right\}\)

\section*{Solving constraints}
\begin{tabular}{llr}
\(\alpha_{10} \supseteq\{710\}\) & \(\alpha_{8} \supseteq \alpha_{2}\) & \(\alpha_{2} \supseteq \alpha_{3}\) \\
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) & \(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \\
\(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{1} \supseteq \alpha_{6}\) & \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\) \\
\(\alpha_{4} \supseteq \alpha_{9}\) & \(\alpha_{7} \supseteq \alpha_{5}\) & \(\alpha_{6}=\{ \}\) \\
\(\alpha_{1}=\{ \}\) & \(\alpha_{7}=\{ \}\) \\
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & \(\alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\(\alpha_{3}=\left\{\left(\lambda \mathbf{x}^{4} \cdot \mathbf{x}^{5}\right)^{3}\right\}\) & \(\alpha_{9}=\left\{\left(\lambda \mathbf{x}^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\(\alpha_{4}=\left\{\left(\lambda \mathbf{x}^{4} \cdot \mathbf{x}^{5}\right)^{3}\right\}\) & \(\alpha_{10}=\{710\}\)
\end{tabular}

\section*{Solving constraints}
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\) \(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{5} \supseteq \alpha_{4}\)
\(\alpha_{7} \supseteq \alpha_{5}\)
\(\alpha_{\mid}=\{ \}\)
\(\alpha_{6}=\{ \}\)
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{7}=\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\}\)
\(\alpha_{3}=\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\}\)
\(\alpha_{8}=\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\}\)
\(\alpha_{4}=\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\}\)
\(\alpha_{9}=\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\}\)
\(\alpha_{5}=\left\{\left(\lambda x^{4} \cdot \mathbf{x}^{5}\right)^{3}\right\}\)
\(\alpha_{8} \supseteq \alpha_{2}\) \(\alpha_{9} \supseteq \alpha_{2}\) \(\alpha_{1} \supseteq \alpha_{6}\)
\(\alpha_{2} \supseteq \alpha_{3}\) \(\left(\alpha_{9} \mapsto \alpha_{7}\right) \supseteq \alpha_{8}\) \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\)
\(\alpha_{4} \supseteq \alpha_{9}\)
\(\alpha_{10}=\left\{7^{10}\right\}\)

\section*{Solving constraints}
\begin{tabular}{|c|c|c|}
\hline \multicolumn{3}{|r|}{\(\alpha_{2} \supseteq \alpha_{3}\)} \\
\hline \(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\}\) & \(\alpha_{9} \supseteq \alpha_{2}\) & \(\left(\alpha_{9} \leftrightarrow \alpha_{7}\right) \supseteq \alpha_{8}\) \\
\hline \(\alpha_{5} \supseteq \alpha_{4}\) & \(\alpha_{1} \supseteq \alpha_{6}\) & \(\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha_{7}\) \\
\hline \multicolumn{3}{|l|}{\(\alpha_{4} \supseteq \alpha_{9} \quad \alpha_{7} \supseteq \alpha_{5}\)} \\
\hline \(\alpha_{1}=\{ \}\) & & \(\alpha_{6}=\{ \}\) \\
\hline \(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & & \(\alpha_{7}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\hline \(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & & \(\alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\hline \(\alpha_{4}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & & \(\alpha_{9}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) \\
\hline \(\alpha_{5}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\) & & \(\alpha_{10}=\left\{7^{10}\right\}\) \\
\hline
\end{tabular}

\section*{Solving constraints}


\section*{Solving constraints}


\section*{Solving constraints}


\section*{Solving constraints}
\begin{tabular}{|c|c|c|c|}
\hline \(\alpha_{10} \supseteq\left\{7^{1}\right.\) & \multicolumn{2}{|c|}{\(\alpha_{8} \supseteq \alpha_{2}\)} & \\
\hline \(\alpha_{3} \supseteq\{(\lambda\) & \multicolumn{2}{|c|}{\(\alpha_{9} \supseteq \alpha_{2}\)} & ( \(\alpha_{9}\) \\
\hline \(\alpha_{5} \supseteq \alpha_{4}\) & \multicolumn{2}{|c|}{\(\alpha_{1} \supseteq \alpha_{6}\)} & ( \(\alpha_{10}\) \\
\hline \(\alpha_{4} \supseteq \alpha_{9}\) & \(\alpha_{7} \supseteq \alpha_{5}\) & \(\alpha_{4} \supseteq \alpha_{10}\) & \\
\hline \multicolumn{2}{|l|}{\(\alpha_{1}=\{ \}\)} & \multicolumn{2}{|l|}{\(\alpha_{6}=\left\{\left(\lambda x^{4} \cdot \mathbf{x}^{5}\right)^{3}\right\}\)} \\
\hline \multicolumn{2}{|l|}{\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)} & \multicolumn{2}{|l|}{\(\alpha_{7}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)} \\
\hline \multicolumn{2}{|l|}{\(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)} & \multicolumn{2}{|l|}{\(\alpha_{8}=\left\{\left(\lambda \mathbf{x}^{4} \cdot \mathbf{x}^{5}\right)^{3}\right\}\)} \\
\hline \(\alpha_{4}=\{\) & \(\left.\mathrm{x}^{5}\right)^{3}\), & \(\alpha_{9}=\{\) & \(\lambda\) \\
\hline \(\alpha_{5}=\{\) & \(\left.\left.x^{5}\right)^{3}\right\}\) & \(\alpha_{10}=\) & 710 \\
\hline
\end{tabular}

\section*{Solving constraints}


\section*{Solving constraints}


\section*{Solving constraints}
\(\alpha_{10} \supseteq\left\{7^{10}\right\}\)
\(\alpha_{8} \supseteq \alpha_{2}\)
\(\alpha_{2} \supseteq \alpha_{3}\)
\(\alpha_{3} \supseteq\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\} \quad \alpha_{9} \supseteq \alpha_{2} \quad\left(\alpha_{9} \mapsto \alpha_{7} \supseteq \alpha_{8}\right.\)
\(\alpha_{5} \supseteq \alpha_{4} \quad \alpha_{1} \supseteq \alpha_{6} \quad\left(\alpha_{10} \mapsto \alpha_{6}\right) \supseteq \alpha^{2}\)
\(\alpha_{।}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}, 710\right\} \quad \alpha_{6}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}, 710\right\}\)
\(\alpha_{2}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\} \quad \alpha_{7}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}, 7^{10}\right\}\)
\(\alpha_{3}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\} \quad \alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{4}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}, 7^{10}\right\} \quad \alpha_{9}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\}\)
\(\alpha_{5}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}, 7^{10}\right\} \quad \alpha_{10}=\left\{7^{10}\right\}\)

\section*{Limitations}

OCFA is still imprecise because it is monovariant: each expression has only one flow variable associated with it, so multiple calls to the same function allow multiple values into the single flow variable for the function body, and these values "leak out" at all potential call sites.

\section*{ICFA}

OCFA is still imprecise because it is monovariant: each expression has only one flow variable associated with it, so multiple calls to the same function allow multiple values into the single flow variable for the function body, and these values "leak out" at all potential call sites.

A better approximation is given by ICFA ("first-order..."), in which a function has a separate flow variable for each call site in the program; this isolates separate calls to the same function, and so produces a more precise result.
\[
\begin{aligned}
& \alpha_{10}=\left\{7^{10}\right\} \\
& \alpha_{2}, \alpha_{3}, \alpha_{8}, \alpha_{9}=\left\{\left(\lambda x^{4} . x^{5}\right)^{3}\right\} \\
& \alpha_{1}, \alpha_{4}, \alpha_{5}, \alpha_{6}, \alpha_{7}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}, 7^{10}\right\} \\
& \left(\text { let id }{ }^{2}=\left(\lambda x^{4} . x^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {| }}
\end{aligned}
\]


\section*{ICFA}

ICFA is a polyvariant approach.
Another alternative is to use a polymorphic approach, in which the values themselves are enriched to support specialisation at different call sites (cf. ML polymorphic types).

It's unclear which approach is "best".
\[
\begin{aligned}
& \alpha_{10} \supseteq\left\{7^{10}\right\} \\
& \text { and } \\
& \alpha_{5} \supseteq \alpha_{4} \\
& \alpha_{7} \supseteq \alpha_{5} \\
& \alpha_{8}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\} \\
& \alpha_{9}=\left\{\left(\lambda x^{4} \cdot x^{5}\right)^{3}\right\} \\
& \alpha_{10}=\left\{7^{10}\right\}
\end{aligned}
\]
\(\left(\text { let } \mathrm{id}^{2}=\left(\lambda \mathrm{x}^{4} \cdot \mathrm{x}^{5}\right)^{3} \text { in }\left(\left(\mathrm{id}^{8} \mathrm{id}^{9}\right)^{7} 7^{10}\right)^{6}\right)^{\text {l }}\)

\section*{Summary}
- Many analyses can be formulated using constraints
- OCFA is a constraint-based analysis
- Inequality constraints are generated from the syntax of a program
- A minimal solution to the constraints provides a safe approximation to dynamic control-flow behaviour
- Polyvariant (as in ICFA) and polymorphic approaches may improve precision

\title{
Lecture 12 \\ Inference-based analysis
}

\section*{Motivation}

In this part of the course were examining several methods of higher-level program analysis.

We have so far seen abstract interpretation and constraintbased analysis, two general frameworks for formally specifying (and performing) analyses of programs.

Another alternative framework is inference-based analysis.

\section*{Inference-based analysis}

Inference systems consist of sets of rules for determining program properties.

Typically such a property of an entire program depends recursively upon the properties of the program's subexpressions; inference systems can directly express this relationship, and show how to recursively compute the property.

\section*{Type systems}

Consider the ML type system, for example.
This particular inference system specifies judgements about a well-typedness property:

means "under the assumptions in Г, the expression e has type \(t\) ".

\section*{Type systems}
\(\Gamma\) is a set of type assumptions of the form
\[
\left\{x_{1}: t_{1}, \ldots, x_{\mathrm{n}}: t_{\mathrm{n}}\right\}
\]
where each identifier \(x_{i}\) is assumed to have type \(t_{i}\).

\section*{We write}
\[
\Gamma[x: t]
\]
to mean \(\Gamma\) with the additional assumption that \(x\) has type \(t\) (overriding any other assumption about \(x\) ).

\section*{Inference-based analysis}

An inference system specifies judgements:
\(\Gamma\)


\(\%\)
- e is an expression (e.g. a complete program)
- \(\Gamma\) is a set of assumptions about free variables of \(e\)
- \(\phi\) is a program property

\section*{Type systems}

We will avoid the more complicated ML typing issues (see Types course for details) and just consider the expressions in the lambda calculus:
\[
\mathrm{e}::=x|\lambda x . \mathrm{e}| \mathrm{e}_{1} \mathrm{e}_{2}
\]

Our program properties are types \(t\) :
\[
t::=\alpha \mid \text { int } \mid t_{1} \rightarrow t_{2}
\]

\section*{Type systems}

In all inference systems, we use a set of rules to inductively define which judgements are valid.

In a type system, these are the typing rules.

\section*{Type systems}
\[
\begin{gather*}
\overline{\Gamma[x: t] \vdash x: t} \quad(\mathrm{VAR}) \\
\frac{\Gamma[x: t] \vdash e: t^{\prime}}{\Gamma \vdash \lambda x \cdot e: t \rightarrow t^{\prime}} \quad(\mathrm{LAM}  \tag{LAM}\\
\frac{\Gamma \vdash e_{1}: t \rightarrow t^{\prime} \quad \Gamma \vdash e_{2}: t}{\Gamma \vdash e_{1} e_{2}: t^{\prime}} \tag{APP}
\end{gather*}
\]

\section*{Optimisation}

In the absence of a compile-time type checker, all values must be tagged with their types and run-time checks must be performed to ensure types match appropriately.

If a type system has shown that the program is well-typed, execution can proceed safely without these tags and checks; if necessary, the final result of evaluation can be tagged with its inferred type.

Hence the final result of evaluation is identical, but less run-time computation is required to produce it.

\section*{Type systems}
\(\Gamma=\{2\) : int, add : int \(\rightarrow\) int \(\rightarrow\) int, multiply : int \(\rightarrow\) int \(\rightarrow\) int \(\}\)
\(\mathrm{e}=\lambda x \cdot \lambda y\). add (multiply \(2 x) y\)
\(t=\) int \(\rightarrow\) int \(\rightarrow\) int
\(\Gamma[x:\) int \(][y:\) int \(] \vdash\) add \(:\) int \(\rightarrow\) int \(\rightarrow\) int \(\quad \Gamma[x:\) int \(][y:\) int \(] \vdash\) multiply \(2 x:\) int
\(\underline{\Gamma[x: \text { int }][y: \text { int }] \vdash \text { add (multiply } 2 x): \text { int } \rightarrow \text { int }} \overline{\Gamma[x: \text { int }][y: \text { int }] \vdash y: \text { int }}\)
\(\Gamma[x:\) int \(][y:\) int \(] \vdash\) add (multiply \(2 x) y:\) int
\(\overline{\Gamma[x: \text { int }] \vdash \lambda y \text {.add (multiply } 2 x) y: \text { int } \rightarrow \text { int }}\)
\(\overline{\Gamma \vdash \lambda x . \lambda y \text {.add (multiply } 2 x) y: \text { int } \rightarrow \text { int } \rightarrow \text { int }}\)

\section*{Safety}

The safety condition for this inference system is
\[
(\} \vdash e: t) \Rightarrow(\llbracket e \rrbracket \in \llbracket t \rrbracket)
\]
where 【e】 and \(\llbracket t \rrbracket\) are the denotations of e and \(t\) respectively: \(\llbracket \mathrm{e} \rrbracket\) is the value obtained by evaluating \(e\), and \(\llbracket t \rrbracket\) is the set of all values of type \(t\).

This condition asserts that the run-time behaviour of the program will agree with the type system's prediction.

\section*{Odds and evens}

This time, the program property \(\phi\) has the form
\[
\phi::=\text { odd } \mid \text { even } \mid \phi_{1} \rightarrow \phi_{2}
\]

\section*{Odds and evens}
\[
\begin{gather*}
\overline{\Gamma[x: \phi] \vdash x: \phi} \\
\frac{\Gamma[x: \phi] \vdash e: \phi^{\prime}}{\Gamma \vdash \lambda x . e: \phi \rightarrow \phi^{\prime}}  \tag{LAM}\\
\frac{\Gamma \vdash e_{1}: \phi \rightarrow \phi^{\prime} \quad \Gamma \vdash e_{2}: \phi}{\Gamma \vdash e_{1} e_{2}: \phi^{\prime}}
\end{gather*}
\]
(APP)

\section*{Odds and evens}
\[
\begin{aligned}
& \Gamma=\{2: \text { even, add : even } \rightarrow \text { even } \rightarrow \text { even, } \\
&\text { multiply }: \text { even } \rightarrow \text { odd } \rightarrow \text { even }\} \\
& \mathrm{e}=\lambda x \cdot \lambda y \cdot \text { add (multiply } 2 x) y \\
& \phi=\text { odd } \rightarrow \text { even } \rightarrow \text { even }
\end{aligned}
\]
\(\overline{\Gamma[x: \text { odd }][y: \text { even }] \vdash \text { add }: \text { even } \rightarrow \text { even } \rightarrow \text { even }} \overline{\Gamma[x: \text { odd }][y: \text { even }] \vdash \text { multiply } 2 x: \text { even }}\)
\(\uparrow \mid x:\) oda \(][y:\) even \(] \vdash\) add : even \(\rightarrow\) even \(\rightarrow\) even \(\bar{\Gamma}[x:\) odd \(][y:\) even \(\mid \vdash\) multiply \(2 x:\) even
\(\frac{\Gamma[x: \text { odd }][y: \text { even }] \vdash \text { add ( }}{\Gamma[x: \text { odd }][y: \text { even }] \vdash \text { add (multiply } 2 x) y: \text { even }}\)
\(\lceil x\) :oda \([y:\) even \(] \vdash\) add (multiply \(2 x\) ) \(y\) : even
\(\frac{\overline{\lceil[x: \text { odd }] \vdash \lambda y \text {. add }(\text { multipl } 2 x) y: \text { even } \rightarrow \text { even }}}{\Gamma \vdash \lambda x . \lambda y \text { add (multiply } 2 x) y: \text { odd } \rightarrow \text { even } \rightarrow \text { even }}\)

\section*{Safety}

The safety condition for this inference system is
\[
(\} \vdash e: \phi) \Rightarrow(\llbracket \in\rfloor \in \llbracket \phi \rrbracket)
\]
where \(\llbracket \phi \rrbracket\) is the denotation of \(\phi\) :
\(\llbracket o d d \rrbracket=\{z \in \mathbb{Z} \mid z\) is odd \(\}\),
\(\llbracket e v e n \rrbracket=\{\mathbf{z} \in \mathbb{Z} \mid \mathbf{z}\) is even \(\}\),
\(\llbracket \phi_{1} \rightarrow \phi_{2} \rrbracket=\llbracket \phi_{1} \rrbracket \rightarrow \llbracket \phi_{2} \rrbracket\)

\section*{Richer properties}

Note that if we want to show a judgement like
\(\Gamma \vdash \lambda x\). \(\lambda y\).add (multiply \(2 x\) ) (multiply \(3 y\) ) : even \(\rightarrow\) even \(\rightarrow\) even
we need more than one assumption about multiply:
\[
\begin{aligned}
\Gamma=\{ & \ldots, \text { multiply : even } \rightarrow \text { even } \rightarrow \text { even }, \\
& \text { multiply : odd } \rightarrow \text { even } \rightarrow \text { even, } . . .\}
\end{aligned}
\]

\section*{Summary}
- Inference-based analysis is another useful framework
- Inference rules are used to produce judgements about programs and their properties
- Type systems are the best-known example
- Richer properties give more detailed information
- An inference system used for analysis has an associated safety condition

\section*{Lecture 13 Effect systems}

\section*{Side-effects}

A side-effect is some event - typically a change of state
- which occurs as a result of evaluating an expression.
- " \(x++\) " changes the value of variable \(x\).
- "malloc(42)" allocates some memory.
- "print 42 " outputs a value to a stream.

\section*{Side-effects}

Some example expressions:

\section*{\}?x.x}
read an integer from channel \(\xi\) and return it
\(\xi!x . y\)
write the (integer) value of \(x\) to channel \(\xi\) and return the value of \(y\)
そ!x. ऽ!x.x
read an integer from channel \(\xi\), write it to channel \(\zeta\) and return it

\section*{Motivation}

We have so far seen many analyses which deal with control- and data-flow properties of pure languages.

However, many languages contain operations with sideeffects, so we must also be able to analyse and safely transform these impure programs.

Effect systems, a form of inference-based analysis, are often used for this purpose.

\section*{Side-effects}

As an example language, we will use the lambda calculus extended with read and write operations on "channels".
е \(::=x|\lambda x . е| e_{1} e_{2}\)
छ?x.e
\(\xi!e_{1 . e}\)
- \(\xi\) represents some channel name.
- \(\xi\) ?x.e reads an integer from the channel named \(\xi\), binds it to \(x\), and returns the result of evaluating \(e\).
- \(\xi!e_{ı} . e_{2}\) evaluates \(e_{1}\), writes the resulting integer to channel \(\xi\), and returns the result of evaluating \(\mathrm{e}_{2}\).

\section*{Side-effects}

Ignoring their side-effects, the typing rules for these new operations are straightforward.

\section*{Effect systems}

However, in order to perform any transformations on a program in this language it would be necessary to pay attention to its potential side-effects.

For example, we might need to devise an analysis to tell us which channels may be read or written during evaluation of an expression.

We can do this by modifying our existing type system to create an effect system (or "type and effect system").

\section*{Effect systems}

First we must formally define our effects:
An expression has effects \(F\).
\(F\) is a set containing elements of the form
\(R_{\xi}\) read from channel \(\xi\)
\(W_{\xi}\) write to channel \(\xi\)

\section*{Effect systems}

But we also need to be able to handle expressions like
\[
\lambda x . \xi!x . x
\]
whose evaluation doesn't have any immediate effects.
In this case, the effect \(W_{\xi}\) may occur later, whenever this newly-created function is applied.

\section*{Effect systems}

So, although it has no immediate effects, the type of
\[
\lambda x . \xi!x . x
\]
is

\[
\begin{gather*}
\text { Effect systems } \\
\frac{\Gamma[x: i n t] \vdash e: t}{\Gamma \vdash \xi ? x \cdot e: t}  \tag{READ}\\
\frac{\Gamma \vdash e_{1}: i n t \quad \Gamma \vdash e_{2}: t}{\Gamma \vdash \xi!e_{1} \cdot e_{2}: t}
\end{gather*}
\]
(Write)

\section*{Effect systems}

For example:
\(\xi ? x . x\)
\(F=\left\{R_{\xi}\right\}\)
\(\xi!x . y\)
\(F=\left\{W_{\xi}\right\}\)
\(\zeta ? X . \zeta!X . X \quad F=\left\{R \xi, W_{\zeta}\right\}\)

\section*{Effect systems}

To handle these latent effects we extend the syntax of types so that function types are annotated with the effects that may occur when a function is applied:
\[
t::=\operatorname{int} \mid t_{1} \xrightarrow{F} t_{2}
\]

\section*{Effect systems}

We can now modify the existing type system to make an effect system - an inference system which produces judgements about the type and effects of an expression:


Effect systems
\[
\begin{equation*}
\frac{\Gamma[x: i n t] \vdash e: t, F}{\Gamma \vdash \xi ? x . e: t,\left\{R_{\xi}\right\} \cup F} \tag{READ}
\end{equation*}
\]
\(\frac{\Gamma \vdash e_{1}: \text { int }, F \quad \Gamma \vdash e_{2}: t, F^{\prime}}{\Gamma \vdash \xi!e_{1} \cdot e_{2}: t, F \cup\left\{W_{\xi}\right\} \cup F^{\prime}}\)

\section*{Effect systems}
\[
\begin{array}{r}
\overline{\Gamma[x: t] \vdash x: t,\{ \}} \\
\frac{\Gamma[\mathrm{VAR})}{\Gamma \vdash \lambda x] \vdash e: t^{\prime}, F} \\
\frac{\Gamma \vdash e_{1}: t \xrightarrow{F} t^{\prime},\{ \}}{\Gamma \vdash e_{1} e_{2}: t^{\prime \prime}, F \cup F^{\prime} \cup F^{\prime \prime}} \tag{App}
\end{array}
\]

\section*{Effect subtyping}

We would probably want more expressive control structure in a real programming language.

For example, we could add if-then-else: \(\mathrm{e}:==x|\lambda x . e| \mathrm{e}_{1} \mathrm{e}_{2}|\xi!x . \mathrm{e}| \xi!\mathrm{e}_{\mathrm{e} . \mathrm{e}_{2} \mid \text { if } \mathrm{e}_{1} \text { then } \mathrm{e}_{2} \text { else } e_{3} .}\)

\section*{Effect subtyping}

However, there are some valid uses of if-then-else which this rule cannot handle by itself.

\section*{Effect subtyping}

if \(x\) then \(\lambda x . \xi!3 . x+1\) else \(\lambda x . x+2\)
\begin{tabular}{|c|}
\hline Effect subtyping \\
\hline  \\
\hline
\end{tabular}

\section*{Effect systems}
\[
\frac{\frac{\{x: \text { int, y:int }\} \vdash x: \text { int },\{ \}}{\{x: \text { int }, y: \text { int }\} \vdash \xi!x \cdot x: \text { int },\left\{W_{\xi}\right\}}}{\frac{\{y: \text { int }\} \vdash \lambda x . \xi!x . x: i n t \stackrel{\left\{W_{\xi}\right\}}{\rightarrow} \text { int, }\}}{} \quad \overline{\{y: i n t\} \vdash y: i n t,\{ \}}}
\]

\section*{Effect subtyping}
\[
\begin{equation*}
\frac{\Gamma \vdash e_{1}: \text { int }, F \quad \Gamma \vdash e_{2}: t, F^{\prime} \quad \Gamma \vdash e_{3}: t, F^{\prime \prime}}{\Gamma \vdash \text { if } e_{1} \text { then } e_{2} \text { else } e_{3}: t, F \cup F^{\prime} \cup F^{\prime \prime}} \tag{COND}
\end{equation*}
\]

\section*{Effect subtyping}
\[
\left\{W_{\xi}\right\}
\]
\[
\frac{\Gamma \vdash e_{1}: \text { int }, F \quad \Gamma \vdash e_{2} \in(t) F^{\prime} \quad \Gamma \vdash e_{3} \in F^{\prime \prime}}{\Gamma \vdash \text { if } e_{1} \text { then } e_{2} \text { else } e_{3}: t, F \cup F^{\prime} \cup F^{\prime \prime}}
\]
(Cond)

\section*{Effect subtyping}

We can solve this problem by adding a new rule to handle subtyping.
\begin{tabular}{|r|}
\hline Effect subtyping \\
\(\frac{\Gamma \vdash e: t \xrightarrow{F^{\prime}} t^{\prime}, F}{\Gamma \vdash e: t \xrightarrow{F^{\prime \prime}} t^{\prime}, F} \quad F^{\prime} \subseteq F^{\prime \prime}\) \\
\end{tabular}

\section*{Effect subtyping}

if \(x\) then \(\lambda x \cdot \xi!3 \cdot x+1\) else \(\lambda x \cdot x+2\)

\section*{Safety}
\[
\begin{gathered}
(\} \vdash e: t, F) \Rightarrow \\
(v \in \llbracket t \rrbracket \wedge f \subseteq F \text { where }(v, f)=\llbracket e \rrbracket)
\end{gathered}
\]

\section*{Extra structure}

If we use a different representation of effects, and use different operations on them, we can keep track of more information.

One option is to use sequences of effects and use an append operation when combining them.

\section*{Effect subtyping}

if \(x\) then \(\lambda x \cdot \xi!3 \cdot x+1\) else \(\lambda x \cdot x+2\)

\section*{Optimisation}

The information discovered by the effect system is useful when deciding whether particular transformations are safe.

An expression with no immediate side-effects is referentially transparent: it can safely be replaced with another expression (with the same value and type) with no change to the semantics of the program.

For example, referentially transparent expressions may safely be removed if LVA says they are dead.

\section*{Extra structure}

In this analysis we are using sets of effects.
As a result, we aren't collecting any information about how many times each effect may occur, or the order in which they may happen.
\[
\begin{array}{rlrl}
\xi \text { ?x. } \zeta!x . x & F & =\left\{R_{\xi}, W_{\zeta}\right\} \\
\zeta!y \cdot \xi ? x . x & F & =\left\{R_{\xi}, W_{\zeta}\right\} \\
\zeta!y . \xi ? x . \zeta!x . x & F & =\left\{R_{\xi}, W_{\zeta}\right\}
\end{array}
\]

\section*{Extra structure}
\[
\begin{equation*}
\frac{\Gamma[x: i n t] \vdash e: t, F}{\Gamma \vdash \xi ? x . e: t,\left\langle R_{\xi}\right\rangle @ F} \tag{READ}
\end{equation*}
\]
\(\frac{\Gamma \vdash e_{1}: \text { int }, F \quad \Gamma \vdash e_{2}: t, F^{\prime}}{\Gamma \vdash \xi!e_{1} \cdot e_{2}: t, F @\left\langle W_{\xi}\right\rangle @ F^{\prime}}\)

\section*{Extra structure}

In the new system, these expressions all have different effects:
\[
\begin{array}{rl}
\xi!x . \zeta!x . x & F=\left\langle R_{\xi} ; W_{\zeta}\right\rangle \\
\zeta!y . \xi!x \cdot x & F=\left\langle W_{\xi} ; R_{\xi}\right\rangle \\
\zeta!y \cdot \xi!x . \zeta!x \cdot x & F=\left\langle W_{\zeta} ; R_{\xi} ; W_{\zeta}\right\rangle
\end{array}
\]

\section*{Extra structure}

Whether we use sequences instead of sets depends upon whether we care about the order and number of effects. In the channel example, we probably don't.

But if we were tracking file accesses, it would be important to ensure that no further read or write effects occurred after a file had been closed.

And if we were tracking memory allocation, we would want to ensure that no block of memory got deallocated twice.

\section*{Summary}
- Effect systems are a form of inference-based analysis
- Side-effects occur when expressions are evaluated
- Function types must be annotated to account for latent effects
- A type system can be modified to produce judgements about both types and effects
- Subtyping may be required to handle annotated types
- Different effect structures may give more information

\title{
Lecture 13a \\ Alias and points-to analysis
}

\section*{Example}

As a simple example, consider some MP3 player code:
```

for (channel = 0; channel < 2; channel++)

```
    process_audio(channel);

\section*{Or even}
process_audio_left(); process_audio_right();

Can we run these two calls in parallel? In other words, when is it safe to do so?

\section*{Memory accessed}

We can reduce this problem to finding locations accessed by each instruction, then combining for all instructions within a procedure.

So, given a pointer value, we are interested in finding a
finite description of the locations it might point to.
If two such descriptions have an empty intersection then we can parallelise / reorder the instructions / ...

\section*{Motivation}

We've seen a number of different analyses that are affected by ambiguity in variables accessed (e.g. in LVA we assume all address-taken variables are referenced).

Alongside this, in modern machines we would like to exploit parallelism where possible, either by running code in separate threads on a multi-core, or in separate lanes using short vector (SIMD) instructions.

Our ability to do this depends on us being able to tell whether memory-access instructions alias (i.e. access the same memory location).

\section*{Memory accessed}

In general we can parallelise if neither call writes to a memory location read or written by the other.

We therefore want to know, at compile time, what memory locations a procedure might read from and write to at run time.

Essentially, we're asking what locations the procedure's instructions access at run time.

\section*{Andersen’s analysis}

Andersen's points-to analysis is an \(O\left(n^{3}\right)\) analysis-the underlying problem is the same as 0-CFA.

We'll only look at the intra-procedural case.
We won't consider pointer arithmetic or functions returning pointers.

All object fields are conflated, although a 'field-sensitive' analysis is possible too.

\section*{Andersen's analysis}

Assume the program has been re-written so that all pointer-typed operations are of the form:
\begin{tabular}{ll}
\(x:=\) new \(\ell\) & \\
\(x\) is a program point \\
\(x:=\) null & \\
optional, a variant of new \\
\(x:=\& y\) & C-like languages, also like new \\
\(x:=y\) & \\
\(x:=* y\) & field access of an object \\
\(* x:=y\) & \\
field access of an object
\end{tabular}
\(x:=\) new \(_{\ell} \quad \ell\) is a program point
\(x:=\) null optional, a variant of new
\(x:=\& y \quad\) C-like languages, also like new
\(x:=y \quad\) copy
\(* x:=y \quad\) field access of an object

\section*{Andersen's analysis}

We first define a set of abstract values:
\[
V=\operatorname{Var} \cup\left\{\text { new }_{\ell} \mid \ell \in \operatorname{Prog}\right\} \cup\{\text { null }\}
\]

Note that all allocations at program point \(\ell\) are conflated, which makes things finite but loses precision.

We create the points-to relation as a function:
\[
p t: V \rightarrow \mathcal{P}(V)
\]

Some analyses have a different \(p t\) at each program point (like LVA); Andersen keeps one per function.

\section*{Andersen's analysis}

We could use type-like constraints (one per source-level assignment):
\(\overline{\vdash x:=\& y: y \in p t(x)}\)
\(\overline{\vdash x:=\text { new }_{\ell}: \text { new }_{\ell} \in p t(x)}\)
\(\frac{z \in p t(y)}{\vdash x:=* y: p t(z) \subseteq p t(x)}\)
\(\overline{\vdash x:=\text { null }: \text { null } \in p t(x)}\)
\(\overline{\vdash x:=y: p t(y) \subseteq p t(x)}\)
\(\frac{z \in p t(x)}{\vdash * x:=y: p t(y) \subseteq p t(z)}\)

\section*{Andersen's analysis}

Or use the style of 0-CFA:


\section*{Andersen's analysis}

Or use the style of 0 -CFA:


\section*{Andersen's analysis}

The algorithm is flow-insensitive-it only considers the statements and not the order in which they occur. This is faster but less precise.

Property inference rules are then essentially:
\[
\begin{gathered}
(\mathrm{ASS}) \frac{}{\vdash x:=e: \ldots} \quad(\mathrm{SEQ}) \frac{\vdash C: S}{\vdash C \cdot C^{\prime}: S^{\prime}} \\
(\mathrm{COND}) \frac{\vdash C: S \cup S^{\prime}}{\vdash \text { if } e \text { then } C \text { else } C^{\prime}: S \cup S^{\prime}} \\
(\text { WHILE }) \frac{\vdash C: S}{\vdash \text { while } e \text { do } C: S}
\end{gathered}
\]

\section*{Andersen example}

Consider the following code:
\[
\begin{aligned}
& \mathrm{a}=\& \mathrm{~b} ; \\
& \mathrm{c}=\& \mathrm{~d} ; \\
& \mathrm{d}=\& \mathrm{a} ; \\
& \mathrm{e}=\mathrm{c} ; \\
& \mathrm{c}=\star \mathrm{e} ; \\
& \star \mathrm{a}=\mathrm{d} ;
\end{aligned}
\]

\section*{Andersen example}
\(\mathrm{a}=\& \mathrm{~b} ;\)
\(\mathrm{c}=\& \mathrm{~d}\);
\(d=\& a ;\)
e \(=c\);
\(c=* e ;\)
*a \(=d\);
\[
\begin{array}{cr}
\operatorname{pt}(\mathrm{a})=\{ \} \quad \operatorname{pt}(\mathrm{c})=\{ \} \\
\operatorname{pt}(\mathrm{b})=\{ \} & \operatorname{pt}(\mathrm{d})=\{ \} \\
\operatorname{pt}(\mathrm{e})=\{ \}
\end{array}
\]

\section*{Andersen example}
```

a = \&b;
c = \&d;
d = \&a; }\quad->\quadpt(a)\supseteq{b
e = c;
c = *e;
*a = d;

$$
\begin{array}{ll}
\operatorname{pt}(\mathrm{a})=\{\mathrm{b}\} & \operatorname{pt}(\mathrm{c})=\{ \} \\
\operatorname{pt}(\mathrm{b})=\{ \} & \operatorname{Pt}(\mathrm{d})=\{ \} \\
\operatorname{pt}(\mathrm{e})=\{ \} &
\end{array}
$$

```

\section*{Andersen example}
```

a = \&b;
c = \&d;
d=\&a; }->\quadpt(d)\supseteq{a
e = c;
c = *e;
*a = d;
pt(a)={b}
pt(e)={}

```

\section*{Andersen example}
```

a = \&b;
c = \&d;
$d=\& a ; \quad \rightarrow \quad p t(e) \supseteq\{d\}$
$e=c ; \quad \Rightarrow p t(c) \supseteq p t(d)$
c $=$ *e;
*a $=$ d;

```
```

$\mathrm{pt}(\mathrm{a})=\{\mathrm{b}\}$

```
\(\mathrm{pt}(\mathrm{a})=\{\mathrm{b}\}\)
    \(\operatorname{pt}(\mathrm{c})=\{\mathrm{a}, \mathrm{d}\}\)
    \(\operatorname{pt}(\mathrm{c})=\{\mathrm{a}, \mathrm{d}\}\)
\(\mathrm{pt}(\mathrm{b})=\{ \}\)
\(\mathrm{pt}(\mathrm{b})=\{ \}\)
    \(\mathrm{pt}(\mathrm{d})=\{\mathrm{a}\}\)
    \(\mathrm{pt}(\mathrm{d})=\{\mathrm{a}\}\)
\(\mathrm{pt}(\mathrm{e})=\{\mathrm{d}\}\)
```

$\mathrm{pt}(\mathrm{e})=\{\mathrm{d}\}$

```

\section*{Andersen example}
```

a = \&b;
c = \&d;
d = \&a; }->\quad->\quadpt(e)\supseteqpt(c
e = c;
c = *e;
*a = d;
pt(a)={b}

```
    \(\mathrm{pt}(\mathrm{e})=\{\mathrm{a}, \mathrm{d}\}\)

\section*{Andersen example}
```

a = \&b;
c = \&d;
d=\&a; }\quad->\quadpt(c)\supseteq{d
e = c;
c = *e;
*a = d;

$$
\begin{array}{cl}
\operatorname{pt}(\mathrm{a})=\{\mathrm{b}\} & \operatorname{pt}(\mathrm{c})=\{\mathrm{d}\} \\
\operatorname{pt}(\mathrm{b})=\{ \} & \operatorname{pt}(\mathrm{e})=\{ \}
\end{array}
$$

```

\section*{Andersen example}
```

a = \&b;
c = \&d;
d = \&a; }\quad->\quadpt(e)\supseteqpt(c
c = *e;
*a = d;

| $\operatorname{pt}(\mathrm{a})=\{\mathrm{b}\}$ | $\operatorname{pt}(\mathrm{c})=\{\mathrm{d}\}$ |
| :--- | :--- |
| $\operatorname{pt}(\mathrm{b})=\{ \}$ | $\operatorname{pt}(\mathrm{d})=\{\mathrm{a}\}$ |

```

\section*{Andersen example}
a \(=\& b ;\)
\(c=\& d ;\)
\(\mathrm{d}=\& a ; \quad \rightarrow \quad p t(a) \supseteq\{b\}\)
\(e=c ; \quad \Rightarrow p t(b) \supseteq p t(d)\)
c \(=\) *e;
*a \(=d\);
\[
\begin{gathered}
\operatorname{pt}(\mathrm{a})=\{\mathrm{b}\} \\
\operatorname{pt}(\mathrm{b})=\{\mathrm{at}\} \\
\operatorname{pt}(\mathrm{e})=\{\mathrm{pt}(\mathrm{c})=\{\mathrm{d}, \mathrm{~d})=\{\mathrm{a}\}
\end{gathered}
\]

\section*{Andersen example}


\section*{Andersen example}
\[
\begin{aligned}
& \text { a = \&b; } \\
& \text { c = \&d; } \\
& \mathrm{d}=\mathrm{q} a ; \quad \rightarrow \quad p t(\mathrm{e}) \supseteq p t(c) \\
& \text { e = c; } \\
& \text { c }=\text { *e; } \\
& \text { *a }=d ; \\
& \begin{array}{c}
\operatorname{pt}(\mathrm{a})=\{\mathrm{b}\} \quad \mathrm{pt}(\mathrm{c})=\{\mathrm{a}, \mathrm{~b}, \mathrm{~d}\} \\
\operatorname{pt}(\mathrm{b})=\{\mathrm{a}\} \quad \mathrm{pt}(\mathrm{~d})=\{\mathrm{a}\} \\
\operatorname{pt}(\mathrm{e})=\{\mathrm{a}, \mathrm{~b}, \mathrm{~d}\}
\end{array}
\end{aligned}
\]

\section*{Other approaches}

Steensgaard's algorithm merges \(a\) and \(b\) if any pointer can reference both. This is less accurate but runs in almost linear time.

Shape analysis (Sagiv, Wilhelm, Reps) models abstract heap nodes and edges between them representing must or may point-to. More accurate but the abstract heaps can get very large.

In general, coarse techniques give poor results whereas more sophisticated techniques are expensive.

\section*{Andersen example}
\[
\begin{gathered}
\operatorname{pt}(a)=\{b\} \quad \operatorname{pt}(c)=\{a, b, d\} \\
\operatorname{pt}(b)=\{a\} \quad \operatorname{pt}(d)=\{a\} \\
\operatorname{pt}(e)=\{a, b, d\}
\end{gathered}
\]

Note that a flow-sensitive algorithm would give
\[
\operatorname{pt}(\mathrm{c})=\{\mathrm{a}, \mathrm{~d}\} \text { and } \mathrm{pt}(\mathrm{e})=\{\mathrm{d}\}
\]
assuming the statements appear in the given order in a single basic block that is not in a loop.

\section*{Summary}
- Points-to analysis identifies which memory locations variables (and other memory locations) point to
- We can use this information to improve datadependence analysis
- This allows us to reorder loads and stores, or parallelise / vectorise parts of the code
- Andersen's analysis is a flow-insensitive algorithm that works in \(O\left(n^{3}\right)\)


Instruction scheduling


\section*{Single-cycle implementation}

In single-cycle processor designs, an entire instruction is executed in a single clock cycle.

Each instruction will use some of the processor's processing stages:
\begin{tabular}{|c|c|c|c|c|}
\hline \begin{tabular}{c} 
Instruction \\
fetch \\
\((I F)\)
\end{tabular} & \begin{tabular}{c} 
Register \\
fetch \\
\((R F)\)
\end{tabular} & \begin{tabular}{c} 
Execute \\
\((E X)\)
\end{tabular} & \begin{tabular}{c} 
Memory \\
access \\
\((M E M)\)
\end{tabular} & \begin{tabular}{c} 
Register \\
write-back \\
\((W B)\)
\end{tabular} \\
\hline
\end{tabular}

For example, a load instruction uses all five.

\section*{Single-cycle implementation}

On these processors, the order of instructions doesn't make any difference to execution time: each instruction takes one clock cycle, so \(n\) instructions will take \(n\) cycles and can be executed in any (correct) order.

In this case we can naïvely translate our optimised 3address code by expanding each intermediate instruction into the appropriate sequence of target instructions; clever reordering is unlikely to yield any benefits.

\title{
Part C
}

\section*{Instruction scheduling}

\section*{Motivation}

We have seen optimisation techniques which involve removing and reordering code at both the source- and intermediate-language levels in an attempt to achieve the smallest and fastest correct program.

These techniques are platform-independent, and pay little attention to the details of the target architecture.

We can improve target code if we consider the architectural characteristics of the target processor.

\section*{Single-cycle implementation}
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \multicolumn{5}{|r|}{lw \$1,0(\$0)} & \multicolumn{5}{|l|}{lw \$2,4(\$0)} & \multicolumn{5}{|l|}{lw \$3,8(\$0)} \\
\hline IF & RF & EX & MEM & WB & IF & RF & EX & MEM & WB & IF & RF & EX & MEM & WB \\
\hline
\end{tabular}

\section*{Pipelined implementation}

In pipelined processor designs (e.g. MIPS R2000), each processing stage works independently and does its job in a single clock cycle, so different stages can be handling different instructions simultaneously.

These stages are arranged in a pipeline, and the result from each unit is passed to the next one via a pipeline register before the next clock cycle.

Pipelined implementation


\section*{Pipeline hazards}

However, it is not always possible to run the pipeline at full capacity.

Some situations prevent the next instruction from executing in the next clock cycle: this is a pipeline hazard.

On interlocked hardware (e.g. SPARC) a hazard will cause a pipeline stall; on non-interlocked hardware (e.g. MIPS) the compiler must generate explicit NOPs to avoid errors.


\section*{Pipelined implementation}

In this multicycle design the clock cycle is much shorter (one pipeline stage vs. one complete instruction) and ideally we can still execute one instruction per cycle when the pipeline is full.

Programs will therefore execute more quickly.

\section*{Pipeline hazards}

Consider data hazards: these occur when an instruction depends upon the result of an earlier one.
\[
\begin{aligned}
& \text { add } \$ 3, \$ 1, \$ 2 \\
& \text { add } \$ 5, \$ 3, \$ 4
\end{aligned}
\]

The pipeline must stall until the result of the first add has been written back into register \(\$ 3\).

\section*{Pipeline hazards}

The severity of this effect can be reduced by using bypassing: extra paths are added between
functional units, allowing data to be used before it has been written back into registers.

\section*{Pipeline hazards}

But even when bypassing is used, some combinations of instructions will always result in a stall.


Instruction order
lw \$1,0(\$0)
lw \$3,4(\$0)
add \(\$ 2, \$ 2, \$ 1\)
add \(\$ 4, \$ 4, \$ 3\)


\section*{Instruction order}

Since particular combinations of instructions cause this problem on pipelined architectures, we can achieve better performance by reordering instructions where possible.
```

lw \$1,0(\$0)
add \$2,\$2,\$1
lw \$3,4(\$0)
add \$4,\$4,\$3

```

\section*{Instruction order}
lw \(\$ 1,0(\$ 0)\)
\(C\) lw \(\$ 3,4(\$ 0)\)
add \(\$ 2, \$ 2, \$ 1\)
add \(\$ 4, \$ 4, \$ 3\)

STALL FOR \$1

STALL FOR \$3

\section*{Instruction dependencies}

We can only reorder target-code instructions if the meaning of the program is preserved.

We must therefore identify and respect the data dependencies which exist between instructions.

In particular, whenever an instruction is dependent upon an earlier one, the order of these two instructions must not be reversed.

\section*{Instruction dependencies}

Read after write:
An instruction reads from a location after an earlier instruction has written to it.


Reads old value

\section*{Instruction dependencies}

Write after read:
An instruction writes to a location after an earlier instruction has read from it.


Reads new value \(X\)

\section*{Instruction scheduling}

We would like to reorder the instructions within each basic block in a way which
- preserves the dependencies between those instructions (and hence the correctness of the program), and
- achieves the minimum possible number of pipeline stalls.

We can address these two goals separately.

\section*{Preserving dependencies}
| lw \$1,0(\$0)
2 lw \(\$ 2,4(\$ 0)\)
3 add \$3,\$1,\$2
4 sw \(\$ 3,12(\$ 0)\)
5 lw \$4,8(\$0)
6 add \$3,\$1,\$4
7 sw \$3,16(\$0)


\section*{Preserving dependencies}


I, 2, 3, 4, 5, 6, 7
2, I, 3, 4, 5, 6, 7
I, 2, 3, 5, 4, 6, 7
I, 2, 5, 3, 4, 6, 7
I, 5, 2, 3, 4, 6, 7
5, I, 2, 3, 4, 6, 7
2, I, 3, 5, 4, 6, 7
2, I, 5, 3, 4, 6, 7
2, 5, I, 3, 4, 6, 7
5, 2, I, 3, 4, 6, 7

\section*{Instruction dependencies}

Write after write:
An instruction writes to a location after an earlier instruction has written to it.


\section*{Preserving dependencies}

Firstly, we can construct a directed acyclic graph (DAG) to represent the dependencies between instructions:
- For each instruction in the basic block, create a corresponding vertex in the graph.
- For each dependency between two instructions, create a corresponding edge in the graph.

This edge is directed: it goes from the earlier instruction to the later one.

\section*{Preserving dependencies}

Any topological sort of this DAG (i.e. any linear ordering of the vertices which keeps all the edges "pointing forwards") will maintain the dependencies and hence preserve the correctness of the program.

\section*{Minimising stalls}

Secondly, we want to choose an instruction order which causes the fewest possible pipeline stalls.

Unfortunately, this problem is (as usual) NP-complete and hence difficult to solve in a reasonable amount of time for realistic quantities of instructions.

However, we can devise some static scheduling heuristics to help guide us; we will hence choose a sensible and reasonably optimal instruction order, if not necessarily the absolute best one possible.

\section*{Minimising stalls}

Each time we're emitting the next instruction, we should try to choose one which:
- does not conflict with the previous emitted instruction
- is most likely to conflict if first of a pair (e.g. prefer lw to add)
- is as far away as possible (along paths in the DAG) from an instruction which can validly be scheduled last

\section*{Algorithm}
- Construct the scheduling DAG.
- We can do this in \(O\left(n^{2}\right)\) by scanning backwards through the basic block and adding edges as dependencies arise.
- Initialise the candidate list to contain the minimal elements of the DAG.

\section*{Algorithm}

Armed with the scheduling DAG and the static scheduling heuristics, we can now devise an algorithm to perform instruction scheduling.

\section*{Algorithm}
- While the candidate list is non-empty:
- If possible, emit a candidate instruction satisfying all three of the static scheduling heuristics;
- if no instruction satisfies all the heuristics, either emit NOP (on MIPS) or an instruction satisfying only the last two heuristics (on SPARC).
- Remove the instruction from the DAG and insert the newly minimal elements into the candidate list.


\section*{Algorithm}



Instruction scheduling is important for getting the best performance out of a processor; if the compiler does a bad job (or doesn't even try), performance will suffer.

As a result, modern processors have dedicated hardware for performing instruction scheduling dynamically as the code is executing.

This may appear to render compile-time scheduling rather redundant.

\section*{Dynamic scheduling}


\section*{Algorithm}


Candidates: \{6\}
\| 1 w \$1,0(\$0)
2 lw \$2,4(\$0)
5 lw \$4,8(\$0)
3 add \$3,\$1,\$2
4 sw \$3,12 (\$0)
6 add \(\$ 3, \$ 1, \$ 4\)

\section*{Algorithm}
\begin{tabular}{|c|c|}
\hline Original code: & Scheduled code: \\
\hline \| lw \$1,0(\$0) & \| lw \$1,0(\$0) \\
\hline \(2 \mathrm{lw} \mathrm{\$ 2,4( } \mathrm{\$ 0)}\) & \(2 \mathrm{lw} \mathrm{\$ 2,4( } \mathrm{\$ 0)}\) \\
\hline 3 add \$3,\$1,\$2 & 5 lw \$4,8(\$0) \\
\hline 4 sw \$3,12(\$0) & 3 add \$3, \$1, \$2 \\
\hline 5 lw \$4,8(\$0) & 4 sw \$3,12(\$0) \\
\hline 6 add \$3,\$1,\$4 & 6 add \$3, \$1, \$4 \\
\hline 7 sw \$3,16(\$0) & 7 sw \$3,16(\$0) \\
\hline 2 stalls & no stalls \\
\hline 13 cycles & I I cycles \\
\hline
\end{tabular}

\section*{Scheduled code:}
| lw \$1,0(\$0)
\$1,0(\$0)
2 lw \$2,4(\$0)
5 lw \$4,8(\$0)
3 add \(\$ 3, \$ 1, \$ 2\)
4 sw \$3,12(\$0)
6 add \$3,\$1,\$4
no stalls
I | cycles

\section*{Dynamic scheduling}

\author{
But:
}
- This is still compiler technology, just increasingly being implemented in hardware.
- Somebody - now hardware designers - must still understand the principles.
- Embedded processors may not do dynamic scheduling, or may have the option to turn the feature off completely to save power, so it's still worth doing at compile-time.

\section*{Summary}
- Instruction pipelines allow a processor to work on executing several instructions at once
- Pipeline hazards cause stalls and impede optimal throughput, even when bypassing is used
- Instructions may be reordered to avoid stalls
- Dependencies between instructions limit reordering
- Static scheduling heuristics may be used to achieve near-optimal scheduling with an \(O\left(n^{2}\right)\) algorithm
\begin{tabular}{|c|}
\hline \\
Lecture I5 \\
Register allocation vs \\
instruction scheduling, \\
reverse engineering \\
\hline
\end{tabular}

\section*{Allocation vs. scheduling}

We have seen why register allocation is a useful compilation phase: when done well, it can make the best use of available registers and hence reduce the number of spills to memory.

Unfortunately, by maximising the utilisation of architectural registers, register allocation makes instruction scheduling significantly more difficult.

\section*{Allocation vs. scheduling}


\section*{Allocation vs. scheduling}

We might have done better if register \(\$ 5\) wasn't so heavily used.

If only our register allocation had been less aggressive!



\section*{Allocation vs. scheduling}
\begin{tabular}{|c|c|c|}
\hline *x : = *a; & lexing, parsing, translation & \[
\begin{aligned}
& \text { LDR v36,v32 } \\
& \text { STR v36,v33 }
\end{aligned}
\] \\
\hline * y : = *b; & & LDR v37,v34 \\
\hline & & \[
\begin{gathered}
\text { STR v37, v35 } \\
\mid \text { register allocation }
\end{gathered}
\] \\
\hline 1w \$5,0(\$1) & & LDR v5, v1 \\
\hline sw \$5,0(\$2) & & STR v5,v2 \\
\hline lw \$6,0(\$3) & code & LDR v6,v3 \\
\hline sw \$6,0(\$4) & & STR v6,v4 \\
\hline
\end{tabular}

\section*{Allocation vs. scheduling}

(3)


। lw \$5,0(\$1)
I, 2, 3, 4
2 sw \$5,0(\$2)
I, 3, 2, 4
3, I, 2, 4
I, 3, 4, 2
3, I, 4, 2
3, 4, I, 2

\section*{Allocation vs. scheduling}

```

lw \$5,0(\$1)
2 sw \$5,0(\$2)
3 lw \$6,0(\$3)
4 sw \$6,0(\$4)

```

\section*{Allocation vs. scheduling}
lw \$5,0(\$1)
1w \$6,0(\$3)
sw \(\$ 5,0(\$ 2)\)
sw \$6,0(\$4)

\[
\begin{array}{cl}
\text { This schedule of the } & \text { lw } \$ 5,0(\$ 1) \\
\text { new instructions } & \text { lw } \$ 6,0(\$ 3) \\
\text { produces no stalls. } & \text { sw } \$ 5,0(\$ 2) \\
& \text { sw } \$ 6,0(\$ 4)
\end{array}
\]

\title{
Allocation vs. scheduling
}

One option is to try to allocate architectural registers cyclically rather than re-using them at the earliest opportunity.

It is this eager re-use of registers that causes stalls, so if we can avoid it - and still not spill any virtual registers to memory - we will have a better chance of producing an efficient program.

\section*{Allocation vs. scheduling}

So, if we are less zealous about reusing registers, this should hopefully result in a better instruction schedule while not incurring any extra spills.

In general, however, it is rather difficult to predict exactly how our allocation and scheduling phases will interact, and this particular solution is quite ad hoc.

Some (fairly old) research (e.g. CRAIG system in 1995,

\section*{Allocation vs. scheduling}

The same problem also shows up in dynamic scheduling done by hardware.

Executable \(\times 86\) code, for example, has lots of register reuse because of the small number of architectural registers available.

Modern machines cope by actually having more registers than advertised; it does dynamic
recolouring using this larger register set, which then enables more effective scheduling.

\section*{Allocation vs. scheduling}

There is clearly antagonism between register allocation and instruction scheduling: one reduces spills by using fewer registers, but the other can better reduce stalls when more registers are used.

This is related to the phase-order problem discussed earlier in the course, in which we would like to defer optimisation decisions until we know how they will affect later phases in the compiler.

It's not clear how best to resolve the problem.

\section*{Allocation vs. scheduling}

In practise this means that, when doing register allocation by colouring for a basic block, we should
- satisfy all of the important constraints as usual (i.e. clash graph, preference graph),
- see how many spare architectural registers we still have left over, and then
- for each unallocated virtual register, try to choose an architectural register distinct from all others allocated in the same basic block.

Touati's PhD thesis in 2002) has improved the situation.

\section*{Decompilation}


\section*{Reverse engineering}

In general terms, engineering is a process which decreases the level of abstraction of some system.

\section*{Reverse engineering}

In contrast, reverse engineering is the process of increasing the level of abstraction of some system, making it less suitable for implementation but more suitable for comprehension and modification.

\section*{Legality and ethics}

It is quite feasible to decompile and otherwise reverseengineer most software.

So if reverse-engineering software is technologically possible, is there any ethical barrier to doing it?

In particular, when is it legal to do so?

\section*{Motivation}

The job of an optimising compiler is to turn humanreadable source code into efficient, executable target code.

Although executable code is useful, software is most valuable in source code form, where it can be easily read and modified.

The source code corresponding to an executable is not always available - it may be lost, missing or secret - so we might want to use decompilation to recover it.


\section*{Reverse engineering}


\section*{Legality and ethics}

Companies and individuals responsible for creating software generally consider source code to be their confidential intellectual property; they will not make it available, and they do not want you to reconstruct it.
(There are some well-known exceptions.)
Usually this desire is expressed via an end-user license agreement, either as part of a shrink-wrapped software package or as an agreement to be made at installation time ("click-wrap").

\section*{Legality and ethics}

However, the European Union Software Directive of 1991 (9I/250/EC) says:

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\section*{Legality and ethics}
"The authorization of the rightholder shall not be required where [...] translation [of a program is] necessary to achieve the interoperability of [that program] with other programs, provided [...] these acts are performed by [a] person having a right to use a copy of the program"

\section*{Legality and ethics}

The European Union Copyright Directive of 2001
(200 I/29/EC, aka "EUCD") is the EU's implementation of the 1996 WIPO Copyright Treaty.

It is again concerned with the ownership rights of technological IP, but Recital 50 states that:
"[this] legal protection does not affect the specific provisions [of the EUSD]. In particular, it should not apply to [...] computer programs [and shouldn't] prevent [...] the use of any means of circumventing a technological measure [allowed by the EUSD]."

\section*{Legality and ethics}

Predictably enough, the interaction between the EUSD, EUCD and DMCA is complex and unclear, particularly at the increasingly-blurred interfaces between geographical jurisdictions (cf. Dmitry
Sklyarov), and between software and other forms of technology (cf. Jon Johansen).

Get a lawyer.

\section*{Summary}
- Register allocation makes scheduling harder by creating extra dependencies between instructions
- Less aggressive register allocation may be desirable
- Some processors allocate and schedule dynamically
- Reverse engineering is used to extract source code and specifications from executable code
- Existing copyright legislation may permit limited reverse engineering for interoperability purposes

This technique is effective in legally circumventing copyrights and trade secrets, although not patents.

\section*{Clean room design}

Despite the complexity of legislation, it is possible to do useful reverse-engineering without breaking the law.

In 1982, Compaq produced the first fully IBMcompatible personal computer by using clean room design (aka "Chinese wall technique") to reverseengineer the proprietary IBM BIOS.

\section*{Legality and ethics}

And the USA has its own implementation of the WIPO Copyright Treaty: the Digital Millennium Copyright Act of 1998 (DMCA), which contains a similar exception for reverse engineering:
"This exception permits circumvention [...] for the sole purpose of identifying and analyzing elements of the program necessary to achieve interoperability with other programs, to the extent that such acts are permitted under copyright law."
\(\qquad\)

\section*{Lecture 16 \\ Decompilation}

\section*{The decompilation problem}

Even simple compilation discards a lot of information:
- Comments
- Function and variable names
- Structured control flow
- Type information

\section*{The decompilation problem}

Some of this information is never going to be automatically recoverable (e.g. comments, variable names); some of it we may be able to partially recover if our techniques are sophisticated enough.

Compilation is not injective. Many different source programs may result in the same compiled code, so the best we can do is to pick a reasonable representative source program.

\section*{Intermediate code}

For many purposes (e.g. simplicity, retargetability) it might be beneficial to convert the target instructions back into 3 -address code when storing it into the flowgraph.

This presents its own problems: for example, many architectures include instructions which test or set condition flags in a status register, so it may be necessary to laboriously reconstruct this behaviour with extra virtual registers and then use dead-code elimination to remove all unnecessary instructions thus generated.

\section*{Why decompilation?}

This course is ostensibly about Optimising Compilers.
It is really about program analysis and transformation.

Decompilation is achieved through analysis and transformation of target code; the transformations just work in the opposite direction.

\section*{The decompilation problem}

Optimising compilation is even worse:
- Dead code and common subexpressions are eliminated
- Algebraic expressions are rewritten
- Code and data are inlined; loops are unrolled
- Unrelated local variables are allocated to the same architectural register
- Instructions are reordered by code motion optimisations and instruction scheduling

\section*{Intermediate code}

It is relatively straightforward to extract a flowgraph from an assembler program.

Basic blocks are located in the same way as during forward compilation; we must simply deal with the semantics of the target instructions rather than our intermediate 3 -address code.

\section*{Control reconstruction}

A compiler apparently destroys the high-level control structure which is evident in a program's source code.

After building a flowgraph during decompilation, we can recover some of this structure by attempting to match intervals of the flowgraph against some fixed set of familiar syntactic forms from our high-level language.

\section*{Finding loops}

Any structured loops from the original program will have been compiled into tests and branches; they will look like arbitrary ("spaghetti") control flow.

In order to recover the high-level structure of these loops, we must use dominance.

\section*{Dominance}

A node \(n\) is in the dominance frontier of a node \(m\) if \(m\) does not strictly dominate \(n\) but does dominate an immediate predecessor of \(n\).

Intuitively this is the set of nodes where m's dominance stops.

We can represent this dominance relation with a dominance tree in which each edge connects a node with its immediate dominator.

\section*{Dominance}


\section*{Back edges}


\section*{Dominance}

In a flowgraph, we say a node \(m\) dominates another node \(n\) if control must go through \(m\) before it can reach \(n\).

A node \(m\) strictly dominates another node \(n\) if \(m\) dominates \(n\) and \(m \neq n\).

The immediate dominator of a node \(n\) is the unique node that strictly dominates \(n\) but doesn't dominate any other strict dominator of \(n\).


\section*{Back edges}

We can now define the concept of a back edge.
In a flowgraph, a back edge is one whose head dominates its tail.

\section*{Finding loops}

Each back edge has an associated loop.

The head of a back edge points to the loop header, and the loop body consists of all the nodes from which the tail of the back edge can be reached without passing through the loop header.


\section*{Finding loops}

Here, the loop header contains a conditional which determines whether the loop body is executed, and the last node of the body unconditionally transfers control back to the header.


This structure corresponds to source-leve while (...) \{...\} syntax.

\section*{Finding loops}

Once each loop has been identified, we can examine its structure to determine what kind of loop it is, and hence how best to represent it in source code.

\section*{Finding loops}

Here, the loop header unconditionally allows the body to execute, and the last node of the body tests whether the loop should execute again.


This structure corresponds to source-level do \{...\} while (...) syntax.


The first node in this interval transfers control to one node if some condition is true, otherwise it transfers control to another node (which control also eventually reaches along the first branch)

This structure corresponds to source-level
if (...) then \{...\} syntax.

\section*{Control reconstruction}

We can keep doing this for whatever other controlflow constructs are available in our source language.

Once an interval of the flowgraph has been matched against a higher-level control structure in this way, its entire subgraph can be replaced with a single node which represents that structure and contains all of the information necessary to generate the appropriate source code.

\section*{Type reconstruction}

Many source languages also contain rich information about the types of variables: integers, booleans, arrays, pointers, and more elaborate data-structure types such as unions and structs.

At the target code level there are no variables, only registers and memory locations.

Types barely exist here: memory contains arbitrary bytes, and registers contain integers of various bitwidths (possibly floating-point values too).

\section*{Type reconstruction}

Reconstruction of the types of source-level variables is made more difficult by the combination of SSA and register allocation performed by an optimising compiler.

SSA splits one user variable into many variables - one for each static assignment - and any of these variables with disjoint live ranges may be allocated to the same architectural register.

\section*{Type reconstruction}

Happily, we can undo the damage by once again converting to SSA form: this will split a single register into many registers, each of which can be assigned a different type if necessary.


\section*{Type reconstruction}
int \(f\) (int r0) \{ r0 = *(int *) (r0 + 4);
C \(\quad r 0=r 0+2\);
return r0;
\}
compile

ARM
f: ldr r0,[r0,\#4]
add r0,r0,\#2 mov r15,r14

\section*{Type reconstruction}
int foo (int *x) \{ return \(x[1]+2\);
\}

\section*{Type reconstruction}

So each user variable may be spread between several registers - and each register may hold the value of different variables at different times.

It's therefore a bit hopeless to try to give a type to each architectural register; the notional type of the value held by any given register will change during execution.
```

int x = 42; MOV r3,\#42
char *y = "42"; MOV r3,\#0xFF34

```

\section*{Type reconstruction}
```

int f (int *rOa) {
int r0b = *(r0a + 1);
int rOc}=r\mp@subsup{O}{b}{}+2
return rOc;
}
int f (int *r0a) {
int r0b = r0a[1];
int r00}=r\mp@subsup{O}{b}{}+2
return r0c;
}

```

\section*{Type reconstruction}
```

int f (int *r0a) {
return rOa[1] + 2;
}

```

In fact, the return type could be anything, so more generally:
```

T f (T *r0a) {
return r0a[1] + 2;
}

```

\section*{Summary}
- Decompilation is another application of program analysis and transformation
- Compilation discards lots of information about programs, some of which can be recovered
- Loops can be identified by using dominator trees
- Other control structure can also be recovered
- Types can be partially reconstructed with constraint-based analysis

\section*{Type reconstruction}
```

int f (int *rOa) {
return rOa[1] + 2;
}
propagate copies
int f (int *rOa) {
int r0b = rOa[1];
int r0ccrerob + 2;
return r0c;

```
\}

\section*{Type reconstruction}

This is all achieved using constraint-based analysis: each target instruction generates constraints on the types of the registers, and we then solve these constraints in order to assign types at the source level.

Typing information is often incomplete intraprocedurally (as in the example); constraints generated at call sites help to fill in the gaps.

We can also infer unions, structs, etc.```

