

# Course Script 

 Static analysis and all that INF5906 / autum 2017Martin Steffen

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## Chapter Introduction

## Learning Targets of this Chapter

Apart from a motivational introduction, the chapter gives a high-level overview over larger topics covered in the lecture. They are treated hear just as a teaser and in less depth compared to later but there is already technical content.

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What
is it
about?

### 1.1 Motivation

### 1.1.1 General remarks

Static analysis: why and what?

- what
- static: at "compile time"
- analysis: deduction of program properties
* automatic/decidable
* formally, based on semantics
- why
- error catching
- catching common "stupid" errors without bothering the user much
- spotting errors early
- certain similarities to model checking
- examples: type checking, uninitialized variables, potential nil-pointer deref's, unused code
- optimization: based on analysis, transform the "code" , such the the result is "better"
- examples: precalculation of results, optimized register allocation ...


## The nature of static analysis

- compiler with differerent phases
- corresponding to Chomsky's hierarchy
- static $=$ in principle: before run-time, but in praxis, "context-free" ${ }^{2}$
- since: run-time most often: undecidable
$\Rightarrow$ static analysis as approximation
- see also [8, Figure 1.1]


[^0]
## Phases



Static analysis as approximation


The figure is, of course, only an informal illustration. In general, program behavior is (for any non-trivial programming language) undecidable. It's a general fact ("Rice's theorem") that all non-trivial, semantical properties of programs are undecidable. Actually, there are exactly two properties which qualify as being trivial. That's the property "true" which holds for all progams,
and "false", which holds for none. That means basically every single semantical property about programs is undecidable up front. Semantical properties are those that refer to the behavior of a programs, its semantics. Of course, syntactical properties are decidable, even if they are non-trivial in the above technical sense.

## Optimal compiler?

Full employment theorem for compiler writers It's a (mathematically proven!) fact that for any compiler, there exists another one which beats it.

- slightly more than non-existance of optimal compiler or undecidability of such a compiler
- theorem
- just states that there room for improvement is always guaranteed
- does not say how!. Finding a better one: undecidable


### 1.2 Data flow analysis

### 1.2.1 A simplistic while-language

## While-language

- simple, prototypical imperative language
- "untyped"
- simple control structure: while, conditional, sequencing
- simple data (numerals, booleans)
- abstract syntax $\neq$ concrete syntax
- disambiguation when needed: (...), or $\{\ldots\}$ or begin ...end


#### Abstract

The given while-language here (and other languages later) is rather simplististic. This fact of being simple is, however, not the reason why we call the syntax here abstract (not like "it's a very abstract language, it has only 5 constructs"). Besides that: the language may have some restricted form of syntax. Nonetheless, if we don't assume an upper bound on the memory, it is Turing complete.

In this lecture, we generally work with abstract syntax. That's different from concrete syntax. Remember from the phases of a compiler, that abstract syntax trees are typically the result of the parsing phase and the input of the static analysis phase (aka. semantical analysis). Since we are focusing here on the semantic phase of a compiler, we assume that the lexer and parser have


done their thing, and we start the considerations on abstract syntax. Abstract syntax is specified by context-free grammars, which is a formalism to specify structured trees. Thus we are completely not interested in parse trees (sometimes known as contrete syntax trees), or issues of precendece, associativity, etc.

This program "written" in abstract syntax are thought of as trees. Since trees are notationally not easy to write down, we don't depict them as actual trees (even if that would be accurate). Instead, we use textual notations and to appeal to the understanding in which way that represents trees, and allow ourselves groupong constructs like parentheses (..) to disambiguate the tree structure, even if of course parentheses and similar constructs are general not part of abstract syntax trees; they are just needed to help the human reader here to understand the underlying tree structure.

Furthermore, and also as a general remark: the while language here (even in its restricted syntactic capabilities) resembles a "high-level" language, at least insofar that it supports "structured programming" in the form of while-loops, as opposed to conditional jumps. Later we might extend it also with further capabilites and programming abstractions, for instance, procedure calls. Anyway, techniques similar to the ones we cover, can also be applied to lower-level intermediate languages. not just abstract syntax of the source language. Also in that case, in a more formal account, one might start fixing the abstract syntax of the intermediate language for the programs one intends to analyze.

Types As mentioned, working directly with abstract syntax, we very much ignore syntactical questions. Besides that, and in particular for the whilelanguage, we more or less ignore another issue, that of typing. Typing and type checking is one very important form of static analysis and will be covered, just not really in connection with the while-language and its derivatives. The "type system" here is so impoverished, supporting basically only integers and booleans, that it's not much worth introducing a type system for doing that. Instead, sidestepping the question, we deal with it "syntactically". Arithmetic expressions are represented by the nonterminal $a$, boolean expressions by $b$, and, as it happens, we have only variables of arithmetic type. In that sense the language is typed, only not very interestingly, and we assume the proper types have somehow been figured out already, so we concetrate for the while language on other things. For functional languages, resp. calculi (variations of the $\lambda$-calculus), we will take care of type checking and extensions of traditional type checking, as it becomes more challenging.

## Labelling

- associate flow information
$\Rightarrow$ labels
- elementary block $=$ labelled item
- identify basic building blocks
- consistent/unique labelling


## Abstract syntax

$$
\left.\begin{array}{rll}
a: & :=x|n| a \text { op }_{a} a & \text { arithm. expressions } \\
b: & := & \text { true } \mid \text { false } \mid \text { not } b \mid b \text { op }_{b} b \mid a \text { op }_{r} a
\end{array}\right) \text { boolean expr. }
$$

Table 1.1: Abstract syntax

$$
\begin{aligned}
a & ::= & x|n| a \text { op }_{a} a & \\
b: & \text { true } \mid \text { false } \mid \text { not } b \mid b \text { op }_{b} b \mid a \text { op }_{r} a & & \text { boolean expr. } \\
S: & ::= & {[x:=a]^{l} \mid[\text { skip }]^{l} \mid S_{1} ; S_{2} } & \\
& \text { if }[b]^{l} \text { then } S \text { else } S \mid \text { while }[b]^{l} \text { do } S & &
\end{aligned}
$$

Table 1.2: Labelled abstract syntax

## Example factorial

$$
y:=x ; z:=1 ; \text { while } y>1 \operatorname{do}(z:=z * y ; y:=y-1) ; y:=0
$$

- input variable: $x$
- output variable: $z$

$$
\begin{align*}
& {[y:=x]^{0} ;}  \tag{1.1}\\
& {[z:=1]^{1} ;} \\
& \text { while }[y>1]^{2} \\
& \operatorname{do}\left([z:=z * y]^{3} ;[y:=y-1]^{4}\right) ; \\
& {[y:=0]^{5}}
\end{align*}
$$

## CFG factorial



Control flow graph The factorial was used to illustrate the important concept of control flow graphs. Later in the lecture we will have another look in more detail how to actually calculate a control-flow graph, given a program in abstract syntax (i.e., in the form of an abstract syntax tree). Actually, it's not very complex; basically, one has to traverse the tree, "label" the nodes appropriately, which typically means, creating one new node of the graph for each encountered basic construct. In the illustrations here, the nodes are identified by unique labels $l_{0}, l_{1}, l_{2} \ldots$ (in some way, the numbers serve themselves as identification). The above "labelled" abstract syntax from above is a different notation to illustrate the nodes of the control flow graph + the basic expression "contained in" or "associated with" the nodes (and the nodes of the graph are the same as the labels in the syntax notation). Missing in the labelled programm from equation (1.1) obviously are the edges of the graph, but, as said, it's not too hard to calculate them as well (while labelling the statements and traversing the syntax tree).

For participants of the course compiler construction (INF5110): the definition of the control flow graph was done there at a lower level, i.e., at some intermediate code (like three-address-code) without looping constructs, but conditional jumps instead. That intermediate code can be seen as a kind of "machineindependent machine language", i.e., an intermediate language already rather close to actual machine language, but not yet quite there. In any case, that intermediate language language had officially labels that could be used for conditional or unconditional jumps (which are like goto's). In other words, that intermediate language was "officially labelled" via a specific label-command (which was counted among the so-called pseudo-instructions). Because of that, that form of language almost immediately contains its control-flow graph, since
the jumps corresponds obviously to the edges. More precisely, for conditional jumps, one of the edges is goes from the conditional jump-statement to the target label, the other edge is the "fall-through".

A final remark on the nodes and the code contained therein. In the lecture here, we make the assumption for the imperative language: "one node, one basic statement". In practice, it's often the case that one groups together sequences of statements like assignments together into larger block which contains no branching or jumps into it. Those larger blocks are sometimes called basic blocks and the grouping is done for efficiency. In the factorial example, $l_{0}$ and $l_{1}$ could be lumped together (as well as $l_{3}$ and $l_{4}$ ). While a good idea in practice, we don't care much here, as the principles of data flow would not don't change with this refinement.

## Factorial: reaching definitions analysis

- "definition" of $x$ : assignment to $x: x:=a$
- better name: reaching assignment analysis
- first, simple example of data flow analysis

Reaching def's An assignment (="definition") $[x:=a]^{l}$ may reach a program point, if there exists an execution where $x$ was last assigned at $l$, when the mentioned program point is reached.

Reaching definitions The reaching definitions is just one of a family of similar analyses the lecture covers (and which themselves are only a small selection of many much more). It's a typical data flow problem and the techniques to solve this problem can be used analogously to a wide range of problem. This class of problems is known as monotone framework and goes back to Kildall [5].

The data which is imagined to "flow" through the program, resp. rather flow through the control flow graphs as the current intermediate representation of the program, is not so much the actual program data (i.e., here, the integer values stored in the variables). It's rather information about the data that flows through the graph, which may be seen as an abstraction. Of course, there may be different kinds of information one could be interested in, and the choice determines the analysis. In case of the reaching definition analysis here, we are interested in the places, where variables are "defined", i.e., the nodes or labels where variables are being assigned to.

## Factorial: reaching definitions



- data of interest: tuples of variable $\times$ label (or node)
- note: distinguish between the entry and the exit of a node.

Factorial Note in particular, that the exit of the node $l_{4}$ cannot be reached by the assignment to $y$ in the node $l_{1}$, whereas the entry of that node $l_{4}$ may well be reached. That's indicated by the dotted, resp. the solid arrows. Not all points which are reachable via the choices $(y, 1)$ are given in the figure, to keep it readable.

It should be also noted: accepting that the analysis works on tuples $(x, l)$, we immediately see that the problem intuitively is decidable: there are only finitely many variables and finitely many nodes (= labels) in the control flow graph. One could analyze thereby the problem, by making some graph seach for all combinations of variables with start nodes and end nodes. That would be a very wasteful approach, and data flow analysis is about more efficient techniques.

## Factorial: reaching assignments

- " points" in the program: entry and exit to elementary blocks/labels
- ?: special label (not occurring otherwise), representing entry to the program, i.e., $(x, ?)$ represents initial (uninitialized) value of $x$
- full information: pair of "functions"

$$
\begin{equation*}
\mathrm{RD}=\left(\mathrm{RD}_{\text {entry }}, \mathrm{RD}_{\text {exit }}\right) \tag{1.2}
\end{equation*}
$$

- tabular form (array): see next slide


## Factorial: reaching assignments table

| $l$ | $\mathrm{RD}_{\text {entry }}$ | $\mathrm{RD}_{\text {exit }}$ |
| :--- | :--- | :--- |
| 0 | $(x, ?),(y, ?),(z, ?)$ | $(x, ?),(y, 0),(z, ?)$ |
| 1 | $(x, ?),(y, 0),(z, ?)$ | $(x, ?),(y, 0),(z, 1)$ |
| 2 | $(x, ?),(y, 0),(y, 4),(z, 1),(z, 3)$ | $(x, ?),(y, 0),(y, 4),(z, 1),(z, 3)$ |
| 3 | $(x, ?),(y, 0),(y, 4),(z, 1),(z, 3)$ | $(x, ?),(y, 0),(y, 4), \quad(z, 3)$ |
| 4 | $(x, ?),(y, 0),(y, 4),(z, 3)$ | $(x, ?), \quad(y, 4), \quad(z, 3)$ |
| 5 | $(x, ?),(y, 0),(y, 4),(z, 1),(z, 3)$ | $(x, ?),(y, 5), \quad(z, 1),(z, 3)$ |

Remarks The highlighted information in the table is not $100 \%$ consistently done. The intention is to highlight the information which is generated new in the correspond block. That can be seen in the blocks (the lines in the table) corresponding to assignments (which are all except $l=2$ ). At the exit of those blocks, a pair containing the corresponding label and the assigned variable is injected. That explains the highlights in the exit-column. Note that at label 3 , whose block is side-effect free, the exit set corresponds to the entry set.

Now for the entry points: one needs to look at the graph, they are determined by the inter-block relations, i.e., the edges. In the easiest case (for instance $l=$ 1), the set just coincides with the post-set of the predecessor block. Interesting is, of course, label $l=2$, where the graph (interpreted forwardly) joins two arrows, i.e., the block 2 has two predecessors, 1 and 4 . The question then is: is it the union or the intersection of the information. Intuitively: it must be union. That's due to the nature of the analysis here: the question is, may an assignment reach a point in question.

## Reaching assignments: remarks

- elementary blocks of the form
- $[b]^{l}$ : entry/exit information coincides
$-[x:=a]^{l}:$ entry/exit information (in general) different
- at program exit: $(x, ?), x$ is input variable
- table:"best" information $=$ smallest sets:
- additional pairs in the table: still safe
- removing labels: unsafe
- note: still an approximation
- no real (= run time) data, no real execution, only data flow
- approximate since
* in concrete runs: at each point in that run, there is exactly one last assignment, not a set
* label represents (potentially infinitely many) runs
- e.g.: at program exit in concrete run: either $(z, 1)$ or else $(z, 3)$

Input variable As a result of the analysis, variable $x$ is marked as ?. That can be interpreted as a sign that it's not assigned to. More precisely, the fact that there is the tuple ( $x, ?$ ) at the program exist together with the fact that there is no other tuple containing $x$ indicates that $x$ is never assigned to (since we are dealing with "may" information"). That's the reason why we can interpret it as input variable: It is often a convention that input variables are not assigned to inside the program, they are assumed to be given a value up front (indicated by ?).

If one analyses a function body - each function conventionally is represented by its own control flow graph capturing the control-flow of its body- the formal parameters are given their initial value from "outside" via parameter passing. It's often considered as bad style, at least when passing parameters by value, to assign to the formal parameters.

Of course, non-input variables which are not assigned to make no sense. Actually, if such variables occur, it may indicate a problem (unitialized variables) which may lead to random results or errors like nil-pointer exceptions.

For participants of the course "compiler construction": the situation about unitialized variables may be compared to the analgous situation for liveness analysis. There, the core analysis concentrated completely on analysing one single node or basic block (there called local analysis). Some variables where marked as liveness status unknown as they where locally not used, but concentrating on one local node only, one cannot determined whether from the global perspective they are live or dead. A special information (like ? here) was used to indicate that "locally unclear" situation. As a side remark: one difference of the liveness analysis works compared to reaching definition analysis is that it works backward. Liveness analysis will be covered also here later as one instance of the mentioned monotone framework.

Optimal solution Currently it may not yet been completely clear in what sense is the solution is minimal/optimal. Much of the underlying theory (covered later) is about assuring that such a minimal solution demonstratibly and uniquely exists (it will be based on the notion of lattices). Intuitively, the solution from the table is minimal (and thus best) in that we cannot remove one single entry of the table, without making it wrong (unsound, unsafe). Very intuitively, it can be compared also to the picture static analysis as approximation. An over-approximation is an area fully "covering" the actually, irregularly shaped behavior. Restricting to a certain class of analysis (for instance simple data flow, specifically say simple reaching definition) can be visualize by allowed overapproximation only in the form of a circle, let's say. The optimal solution is then the smallest such covering circle (which still is an opproximation). If one invests in a more detailed analysis, that might correspond to having, lets say, ellipses or six-edged polygons as (illustration of)
allowed results of the analysis. In that case, potentially better and more precise approximations may be doable (typically at higher computational costs). It may also be the case, that no longer a single best solution exists in general. For instance, there may be more two covering polygons, both of which cannot be made smaller without becoming unsound, but on the other hand uncomparable (neither is contained in the other). In this lecture, we typically deal with settings were optimal solution do exists, though.

## Data flow analysis

- standard: representation of program as control flow graph (aka flow graph)
- nodes: elementary blocks with labels (or basic block)
- edges: flow of control
- two approaches, both (especially here) quite similar
- equational approach
- constraint-based approach


### 1.2.2 Equational approach

## From flow graphs to equations

- associate an equation system with the flow graph:
- describing the "flow of information"
- here:
* the information related to reaching assignments
* information imagined to flow forwards
- solutions of the equations
- describe safe approximations
- not unique, interest in the least (or largest) solution
- here: give back RD of equation (1.2) on slide 9


## Equations for RD and factorial: intra-block

first type: local, intra-block":

- flow through each individual block
- relating for each elementary block its exit with its entry

$$
\begin{aligned}
& \text { elementary block: }[y:=x]^{0} \\
& \text { elementary block: }[y>1]^{2}
\end{aligned}
$$

all equations with $\mathrm{RD}_{\text {exit }}$ as "left-hand side"

$$
\begin{align*}
& \operatorname{RD}_{\text {exit }}(0)=\operatorname{RD}_{\text {entry }}(0) \backslash\{(y, l) \mid l \in \mathbf{L a b}\} \cup\{(y, 0)\}  \tag{1.3}\\
& \operatorname{RD}_{\text {exit }}(1)=\operatorname{RD}_{\text {entry }}(1) \backslash\{(z, l) \mid l \in \mathbf{L a b}\} \cup\{(z, 1)\} \\
& \operatorname{RD}_{\text {exit }}(2)=\operatorname{RD}_{\text {entry }}(2) \\
& \operatorname{RD}_{\text {exit }}(3)=\operatorname{RD}_{\text {entry }}(3) \backslash\{(z, l) \mid l \in \mathbf{L a b}\} \cup\{(z, 3)\} \\
& \operatorname{RD}_{\text {exit }}(4)=\operatorname{RD}_{\text {etrry }}(4) \backslash\{(y, l) \mid l \in \mathbf{L a b}\} \cup\{(y, 4)\} \\
& \operatorname{RD}_{\text {exit }}(5)=\operatorname{RD}_{\text {entry }}(5) \backslash\{(y, l) \mid l \in \mathbf{L a b}\} \cup\{(y, 5)\}
\end{align*}
$$

## Inter-block flow

second type: global, inter-block

- flow between the elementary blocks, following the control-flow edges
- relating the entry of each block with the exits of other blocks, that are connected via an edge (exception: the initial block has no uncoming edge)
- initial block: mark variables as uninitialized

$$
\begin{align*}
& \mathrm{RD}_{\text {entry }}(1)=\mathrm{RD}_{\text {exit }}(0)  \tag{1.4}\\
& \mathrm{RD}_{\text {entry }}(2)=\mathrm{RD}_{\text {exit }}(1) \cup \mathrm{RD}_{\text {exit }}(4) \\
& \mathrm{RD}_{\text {entry }}(3)=\mathrm{RD}_{\text {exit }}(2) \\
& \mathrm{RD}_{\text {entry }}(4)=\mathrm{RD}_{\text {exit }}(3) \\
& \mathrm{RD}_{\text {entry }}(5)=\mathrm{RD}_{\text {exit }}(2) \\
& \mathrm{RD}_{\text {entry }}(0)=\{(x, ?),(y, ?),(z, ?)\}
\end{align*}
$$

There are 6 equations as there are 6 nodes. As far as the right-hand sides are concened: there are 6 mentionings of $\mathrm{RD}_{\text {exit }}(l)$. That indicates that the graph has 6 edges (not counting the incoming edge into node $l_{0}$ and the outgoing edge at the exit).

The entry node has no internally incoming edge in this example (only one that is shown in the picture to come from "outside" which is therefore not part of the graph nor an edge, it's just a conventional, graphical indication of the initial node). This is not a coincidence, insofar that one generally assumes that the control flow graph does not have such an initial loop. In the unlikely event that the program would start immediately with a loop or similar, the compiler would arrange it so that there is an extra "skip" entry node (a sentinel). Alternatively, the generation of the control flow graph might simply automatically add some extra sentinel, just in case.

Technically, nothing would go wrong allowing such initial loops, it's just that the technical representation later (and the algorithms) are slighly simpler. Sure, not fundamentally simpler, only avoiding some extra corner cases, it just gets a tiny bit more smooth.

Later, we will encounter analyses, which work backwards, i.e., the flow will follow the edges of the flow graph in reverse direction. Live variable analysis is one prime example. In those cases, one tries to avoid "final loops" at the exit of the program.

For participants of "compiler constructions" (INF5110): that's reminiscent of the treatment of context-free grammars for some constructions (like the LR(0)-DFA construction). For some constructions, it was assumed that the grammar's start symbol, say $S$, does not show up on the right-hand side of any production. That would correspond to an loop back to the "intitial state" here. Since also there, one wanted to slightly simplify the treatment avoiding that case, the standard construction routinely simply added "another" start symbol $S^{\prime}$ and a production $S^{\prime}::=S$.

## General scheme (for RD)

Intra - for assignments $[x:=a]^{l}$

$$
\begin{equation*}
\mathrm{RD}_{\text {exit }}(l)=\mathrm{RD}_{\text {entry }}(l) \backslash\left\{\left(x, l^{\prime}\right) \mid l^{\prime} \in \mathbf{L a b}\right\} \cup\{(x, l)\} \tag{1.5}
\end{equation*}
$$

- for other blocks $[b]^{l}$ (side-effect free)

$$
\begin{equation*}
\mathrm{RD}_{\text {exit }}(l)=\mathrm{RD}_{\text {entry }}(l) \tag{1.6}
\end{equation*}
$$

## Inter

$$
\begin{equation*}
\mathrm{RD}_{\text {entry }}(l)=\bigcup_{l^{\prime} \rightarrow l} \mathrm{RD}_{\text {exit }}\left(l^{\prime}\right) \tag{1.7}
\end{equation*}
$$

Initial $l$ : label of the initial block (isolated entry)

$$
\begin{equation*}
\mathrm{RD}_{\text {entry }}(l)=\{(x, ?) \mid x \text { is a program variable }\} \tag{1.8}
\end{equation*}
$$

## The equation system as fix point

- RD example: solution to the equation system $=12$ sets

$$
\mathrm{RD}_{\text {entry }}(0), \ldots, \mathrm{RD}_{\text {exit }}(5)
$$

- i.e., the $\mathrm{RD}_{\text {entry }}(l), \mathrm{RD}_{\text {exit }}(l)$ are the variables of the equation system, of type: sets of pairs of the form $(x, l)$
- domain of the equation system:
- RD : the mentioned twelve-tuple of variables
$\Rightarrow$ equation system understood as function $F$


## Equations

$$
\overrightarrow{\mathrm{RD}}=F(\overrightarrow{\mathrm{RD}})
$$

Fix point equation The above fixpoint equation on vectors of $R D$ variables may be broken down more explicitly 12 parts (the individual "equations") like for instance

$$
F(\overrightarrow{\mathrm{RD}})=\left(F_{\text {entry }}(1)(\overrightarrow{\mathrm{RD}}), F_{\text {exit }}(0)(\overrightarrow{\mathrm{RD}}), \ldots, F_{\text {exit }}(5)(\overrightarrow{\mathrm{RD}})\right)
$$

After solving the equation system, we for instance could get as part of a solution:

$$
F_{\text {entry }}(2)=\left(\ldots, \mathrm{RD}_{\text {exit }}(1), \ldots, \mathrm{RD}_{\text {exit }}(4), \ldots\right)=\mathrm{RD}_{\text {exit }}(1) \cup \mathrm{RD}_{\text {exit }}(4)
$$

## The least solution

- Var $_{*}=$ variables "of interest" (i.e., occurring), Lab ${ }_{*}$ : labels of interest
- here $\mathbf{V a r}_{*}=\{x, y, z\}, \mathbf{L a b}_{*}=\{?, 1, \ldots, 6\}$

$$
\begin{equation*}
F:\left(2^{\mathbf{V a r}_{*} \times \mathbf{L a b}_{*}}\right)^{12} \rightarrow\left(2^{\mathbf{V a r}_{*} \times \mathbf{L a b}_{*}}\right)^{12} \tag{1.9}
\end{equation*}
$$

- domain $\left(2^{\mathbf{V a r}_{*} \times \mathbf{L a b}_{*}}\right)^{12}$ : partially ordered pointwise:

$$
\begin{equation*}
\overrightarrow{\mathrm{RD}} \sqsubseteq \overrightarrow{\mathrm{RD}}^{\prime} \text { iff } \forall i . \mathrm{RD}_{i} \subseteq \mathrm{RD}_{i}^{\prime} \tag{1.10}
\end{equation*}
$$

$\Rightarrow$ complete lattice

### 1.2.3 Constraint-based approach

This section is basically a rerun of the previous one: the constraint-based approach here is nothing much more than a variation of the equational approach. As for terminology, the distinction between "equational" vs. "constaint-based" here is slightly misleading. It's misleading insofar, as also the equational approach is based on constraints, namely equational constraints. In contrast, next we will use "inequations". In the RD analysis we are currently focussing on, the solutions are sets of pairs $(x, l)$. Consequently, the inequations will be subset constraints $\subseteq$ on sets, instead of equality $=$ on sets.

In the current setting, a basic data flow problem for an simplistic imperative language, there's not much difference between the two approaches. Not only is it straightforward on the surface, to replace the $=$ by $\subseteq$, and there you are. Also on a deeper level here, there are technical reasons resulting in the following fact: the best (here smallest) solution of the equational approach coincides with the smallest solution of the constraint-based approach. That means, for
the purpose of program analysis, which is after the best, safe approximation, both approaches are the same!

The setting here with simple control flow graphs, however, is indeed simple. In more complex setting (for instance functional languages with much more flexible control flow) it's not always the case that equational approaches and constraint based approaches give the same (best) analysis result. In general, the constraint based approach offers more flexibility (thus sometimes more precise analysis) in more complex settings, but not here. In the context of type and effect systems, we will encounter such more complex settings.

When stating that (here) the step from an equational to a constraint-based approach amounts to a trivial reformulation of the equations leaves, however, one question still to be answered:

$$
\text { should }=\text { be replaced by } \subseteq \text { or } \supseteq \text { ? }
$$

We will see that the answer to that question depends on the analysis we are doing (basically whether we are interested in safe over-approximations or underapproximation). In the case of the reaching definitions ("... may reach ..."), a solution larger than a safe one is safe, as well. As a consequence, the "lefthand sides" of the previous equations need to be $\supseteq$-larger than the "right-hand sides".

A final remark: the fact that we move from equations to $\supseteq$ allow allows for a simple rerrangement of the constraints. Basically, instead of saying equationally $s_{1}=s_{2} \cup s_{3}$, reformulated to $s_{1} \supseteq s_{2} \cup s_{3}$, the inequation is split into

$$
\begin{aligned}
& s_{1} \supseteq s_{2} \\
& s_{1} \supseteq s_{3}
\end{aligned}
$$

## Factorial program: intra-block constraints

> elementary block: $[y:=x]^{0}$
> elementary block: $[y>1]^{2}$
all equations with $\mathrm{RD}_{\text {exit }}$ as left-hand side

$$
\begin{aligned}
& \operatorname{RD}_{\text {exit }}(0) \supseteq \operatorname{RD}_{\text {entry }}(0) \backslash\{(y, l) \mid l \in \mathbf{L a b}\} \\
& \operatorname{RD}_{\text {exit }}(0) \supseteq\{(y, 0)\} \\
& \operatorname{RD}_{\text {exit }}(1) \supseteq \operatorname{RD}_{\text {entry }}(1) \backslash\{(z, l) \mid l \in \mathbf{L a b}\} \\
& \operatorname{RD}_{\text {exit }}(1) \supseteq\{(z, 1)\} \\
& \operatorname{RD}_{\text {exit }}(2) \supseteq \operatorname{RD}_{\text {entry }}(2) \\
& \operatorname{RD}_{\text {exit }}(3) \supseteq \operatorname{RD}_{\text {entry }}(3) \backslash\{(z, l) \mid l \in \mathbf{L a b}\} \\
& \operatorname{RD}_{\text {exit }}(3) \supseteq\{(z, 3)\} \\
& \operatorname{RD}_{\text {exit }}(4) \supseteq \operatorname{RD}_{\text {entry }}(4) \backslash\{(y, l) \mid l \in \mathbf{L a b}\} \\
& \operatorname{RD}_{\text {exit }}(4) \supseteq\{(y, 4)\} \\
& \operatorname{RD}_{\text {exit }}(5) \supseteq \operatorname{RD}_{\text {entry }}(5) \backslash\{(y, l) \mid l \in \mathbf{L a b}\} \\
& \operatorname{RD}_{\text {exit }}(5) \supseteq\{(y, 5)\}
\end{aligned}
$$

## Factorial program: inter-block constraints

cf. slide 13 ff.: inter-block equations:

$$
\begin{aligned}
& \mathrm{RD}_{\text {entry }}(1)=\mathrm{RD}_{\text {exit }}(0) \\
& \mathrm{RD}_{\text {entry }}(2)=\mathrm{RD}_{\text {exit }}(1) \cup \mathrm{RD}_{\text {exit }}(4) \\
& \mathrm{RD}_{\text {entry }}(3)=\mathrm{RD}_{\text {exit }}(2) \\
& \mathrm{RD}_{\text {entry }}(4)=\mathrm{RD}_{\text {exit }}(3) \\
& \mathrm{RD}_{\text {entry }}(5)=\mathrm{RD}_{\text {exit }}(2) \\
& \mathrm{RD}_{\text {entry }}(0)=\{(x, ?),(y, ?),(z, ?)\}
\end{aligned}
$$

splitting of composed right-hand sides + using $\supseteq$ instead of $=$ :

$$
\begin{array}{lll}
\mathrm{RD}_{\text {entry }}(1) & \supseteq \mathrm{RD}_{\text {exit }}(0) \\
\mathrm{RD}_{\text {entry }}(2) & \supseteq \mathrm{RD}_{\text {exit }}(1) \\
\mathrm{RD}_{\text {entry }}(2) & \supseteq \mathrm{RD}_{\text {exit }}(4) \\
\mathrm{RD}_{\text {entry }}(3) & \supseteq \mathrm{RD}_{\text {exit }}(2) \\
\mathrm{RD}_{\text {entry }}(4) & \supseteq \mathrm{RD}_{\text {exit }}(3) \\
\mathrm{RD}_{\text {entry }}(5) & \supseteq \mathrm{RD}_{\text {exit }}(2) \\
& \\
\mathrm{RD}_{\text {entry }}(1) & \supseteq\{(x, ?),(y, ?),(z, ?)\}
\end{array}
$$

## Least solution revisited

instead of $F(\overrightarrow{R D})=\overrightarrow{R D}$

- clear: solution to the equation system $\Rightarrow$ solution to the constraint system
- important: least solutions coincides!

$$
\begin{equation*}
F(\overrightarrow{\mathrm{RD}}) \sqsubseteq \mathrm{RD} \tag{1.11}
\end{equation*}
$$

### 1.3 Constraint-based analysis

### 1.3.1 Control-flow analysis

## Control-flow analysis

Goal CFA which elem. blocks lead to which other elem. blocks

- for while-language: immediate (labelled elem. blocks, resp., graph)
- complex for: more advanced features, higher-order languages, oo languages ...
- here: prototypical higher-order functional language $\lambda$-calculus
- formulated as constraint based analysis


## Introduction

This section is called constraint based analysis, but it's also mostly about functional, higher-order languages (and complications that entails for static analysis).

In the section about data-flow analysis, the analysis was building upon controlflow graphs. Given the control-flow graph of a program, the problem was turned into a set of data flow equations or data flow inequations (there called constraints), which in turn then have to be solved. While it was stressed there that the equatioan approach and the approach based in subset constraints give the same result (in that the best solution coincides in both representations). The analysis therefore proceeds in two clearly separated stages: first the control-flow graph is determined, and, based on that, the data flow is analized. We did not cover really the calculation of the control flow graph in first stage as it's rather simple.

For languages with higher-order functions, the dividing line between "data" and "'control" is blurred: a function on the one hand is "code", but on the other hand is also "data" in that funtctions can be passed as arguments to other functions and returned as values from function. As a consequence, one would expect static program analysis gets involved for higher-order functions. That's even true taking into account that the setting for data flow analysis in the introduction so far was really simple in that we did no even consider (non-higher-order) functions there. Adding those would complicate things later in the lecture we will cover that to some extent - but still the 2 stages
of determining the control-flow graph first, and afterwards doing data flow constraints are clearly discernible.

Here, determining the control flow is a problem in itself, unlike in the simpler setting before, where determining the CFG was so simple that would not even bother calling it control-flow analysis (even it it is). Here, we are facing a nontrivial control flow analysis problem and we use a constraint-based approach (not an equational) to tackle it.
Another way of seeing it is as follows: the control-flow problem here (which is inseparable from data-flow aspects as functions are higher-order), will not even result in a control-flow graph. For a non-higher order language, each function body has its own CFG, all clearly separated, but such a depiction makes no sense any longer. One the other hand: in the data flow section, we slightly touched upon the close connection between control-flow graph and the constraints (equational or otherwise). We did not explore it to the end, but one can view each edge of a control-flow graph as constraints connecting (in a forward analysis) the solutions of a target node dependent (via a constraint) on the solution on the source nodes and the nodes (= labels) play the role of variables in the constraint system. The connection is so close that a constraint systems in the form we covered in the data-flow section is nothing else than some representation of the control-flow graph (together with constraints connecting the nodes). As illustration, an inequation

$$
x_{1} \supseteq f\left(x_{2}, x_{3}\right)
$$

represents the dependence of (the solution for) $x_{1}$ on the solutions of $x_{2}$ and $x_{3}$ via some $f$, thereby representing two incoming edges in $x_{1}$, starting from $x_{2}$ resp. $x_{3}$. In Section 1.2, this representation was not made explicit, and we did not called the variables $x_{i}$ but $\mathrm{RD}_{i}$ (and split between entries and exits), but in general the connection is between variables of the constraint system and the nodes of the control flow graph.

Now, in the more complex situation of higher-order functions, we operate directly with constraints; since they are now more complicated, they can no longer be visualized as expressing relations on values in some graph.

## Simple example

```
let f = fn x => x 1;
    g = fn y }=>\textrm{y}+2
    h = fn z m z + 3;
in (f g) + (f h)
```

- higher-order function $f$
- for simplicity: untyped
- local definitions via let-in
- interesting above: $x 1$

Goal (more specifically) For each function application, which function may be applied.

Local definitions with let The above example uses let for local definition. That slighty different than assignments. For once, it's "single assignment", secondly it has a local scope associate with it. In later chapters, the functional language may support that construct, for the introduction we ignore it.

## Labelling

- more complex language $\Rightarrow$ more complex labelling
- "elem. blocks" can be nested
- all syntactic constructs (expressions) are labelled
- consider:


## Unlabelled abstract syntax

$$
(\mathrm{fn} x \Rightarrow x)(\mathrm{fn} y \Rightarrow y)
$$

## Full labelling

$$
\left[\left[\mathrm{fn} x \Rightarrow[x]^{1}\right]^{2}\left[\mathrm{fn} y \Rightarrow[y]^{3}\right]^{4}\right]^{5}
$$

- functional language: side effect free
$\Rightarrow$ no need to distinguish entry and exit of labelled blocks.


## Data of the analysis

Pairs $(\hat{C}, \hat{\rho})$ of mappings:
abstract cache: $\hat{C}(l)$ : set of values/function abstractions, the subexpression labelled $l$ may evaluate to
abstract env.: $\hat{\rho}$ : values, $x$ may be bound to

## The constraint system

- ignoring "let" here: three syntactic constructs $\Rightarrow$ three kinds of constraints
- relating $\hat{C}, \hat{\rho}$, and the program in form of subset constraints (subsets, order-relation)


## 3 syntactic classes

- function abstraction: $[\mathrm{fn} x \Rightarrow x]^{l}$
- variables: $[x]^{l}$
- application: $[f g]^{l}$


## Constraint system for the small example

## Labelled example

$$
\left[\left[\mathrm{fn} x \Rightarrow[x]^{1}\right]^{2}\left[\mathrm{fn} y \Rightarrow[y]^{3}\right]^{4}\right]^{5}
$$

- application: connecting function entry and (body) exit with the argument but:
- also $\left[\mathrm{fn} y \Rightarrow[y]^{3}\right]^{4}$ is a candidate at 2 ! (according to $\hat{C}(2)$ )
- function abstractions
- variables (occurrences of use)

$$
\begin{aligned}
\left\{\mathrm{fn} x \Rightarrow[x]^{1}\right\} & \subseteq \hat{C}(2) \\
\left\{\mathrm{fn} y \Rightarrow[y]^{3}\right\} & \subseteq \hat{C}(4) \\
\hat{\rho}(x) & \subseteq \hat{C}(1) \\
\hat{\rho}(y) & \subseteq \hat{C}(3) \\
\left\{\mathrm{fn} x \Rightarrow[x]^{1}\right\} \subseteq \hat{C}(2) \Rightarrow \hat{C}(4) & \subseteq \hat{\rho}(x) \\
\left\{\operatorname{fn} x \Rightarrow[x]^{1}\right\} \subseteq \hat{C}(2) \Rightarrow \hat{C}(1) & \subseteq \hat{C}(5) \\
\left\{\operatorname{fn} y \Rightarrow[y]^{3}\right\} \subseteq \hat{C}(2) \Rightarrow \hat{\mathbf{C}}(4) & \subseteq \hat{\rho}(y) \\
\left\{\mathrm{fn} y \Rightarrow[y]^{3}\right\} \subseteq \hat{C}(2) \Rightarrow \hat{C}(3) & \subseteq \hat{\mathbf{C}}(5)
\end{aligned}
$$

Explanation of the constraint system The example is rather small, but containts all constructs of the language (as we ignore let), therefore it illustrates how a label program gives rise to a constraint system, in this case for control-flow analysis. It's still a bit of an illustration only, as we don't give the general construction, we just show how it works in the given example.

The slides present the construction in 2 stages (for didactic reasons). The first stage gives "standard", unconditional constraints. The resulting system is find in the sense that, all solutions to it are safe approximations of the problem.

The second stage is an improvement in that it gives more precise solutions. It is formulated with conditional constraints.

In general, there are 3 different classes of constraints, since there are three classes of syntactial constructs. The classification is done according to the top-level construct at the label being considerd. Remember that abstract syntax represents trees, and now the labels indentify subtrees. The three cases are the following:

1. One for abstractions (the easiest one), one for
2. variables, and one for
3. applications (the most complex).

They are shown in this order via overlays.

1. Abstractions

We have two abstractions, at label 2 and 4 . The constraints in this category are relevant for the $\hat{C}$, only, as there are no variables "involved" (only the bound variables inside the function definition). Therefore, the abstract environment is not mentioned here. The inequations are easy. They just state that the expression at the given label (which labels, as said, an abstraction), at least evaluates to that abstraction. That's pretty obvious.
2. Variables

Variables are a bit more tricky, especially because the corresponding constraints relates the $\hat{C}$ and the abstract environment. Basically it relates directly the $\hat{C}$ for the label with the $\hat{\rho}$ variable carrying that label. The only perhaps tricky question is: how do they relate, in which order is the $\subseteq$. In the example, in the smallest solution, it's $=$, anyway, in the end. It's the way that one "visualizes" the flow. Here we have a variable, and the information flows from the variable to the label. We do not write to the variable, but read from it and get the value from the variable into the program (at the given label). Since we have a forward-may-analysis and go for the smallest value, the "post-state" (at l) must be equal or larger than the "pre-state" (at x).
3. Applications

Basically, we have one application (labelled 5). So we have to think of the flow. One is that the argument "flows into" the variable which constitues the formal argument of the function. The same reasoning (intuitive direction of the flow of information) determines the direction of the inequation. The second flow is the output: what comes out of the body of the function function comes \{out of the application\}. That relates the $\hat{C}$ of the corresponding labels. This explains the first two inequations (which are not yet complete, there will be 2 steps to capture the flow (more precisely)). The first step deals with the fact that we don't really know which function
a variable represents (higher-order). The second one is an optimization, introducing conditional constraints.
However, that is not all for application. So far we have just looked "intuitively" at the code, but we need to relate to the analysis/an algorithm. This means, we should not just look at what function there "/is/" according to our intuition at label 2 (the place for the function in the application) but formally we need to think, what functions location 2 evaluates according to the analysis. In other words, we need to consult $\hat{C}(2)$. We have only a lower bound for $\hat{C}(2)$ (some slides earlier), therefore, in principle all possible abstractions (there are 2) are candidates to be at the applicator position in the application. I.e., the one missing is $\left[\mathrm{fn} y \Rightarrow[\mathbf{y}]^{3}\right]^{4}$. This adds two more inequations, again one for input and the other for output, this time for the second abstraction.
What we have so far (especially for category 3) would work (in category 3 ), in the sense of leading to a sound analysis. As mentioned before, we can do better, however, in the category for applications. Namely that we formulate the inequations conditionally making the information about which function the $l=2$ evaluates to more precise (in the end: smaller). In the example, we will see that the analysis finds out that $\hat{C}(2)$ indeed only contains one function, the obvious one. This additional precision gives indeed a smaller solution.
The refinement can be explained as follows: it concerns the category for applications (location 5). Each "instance" of an application leads to two constraints "in" and "out" (for parameter passing and returning the data). The question only is (as discussed): which function definition is it actually where the data flows "in" and "out". What we know is that the function, whichever it is, is placed at location 2. Therefore the condition in the conditional refinement of the constraints expresses the following
"if such-and-such function occurs at 2 , then consider the "in" constraint (and "out" constraint) for that function"

## The least (= best) solution

| $\hat{C}(1)$ | $=\left\{\mathrm{fn} y \Rightarrow[y]^{3}\right\}$ |
| ---: | :--- |
| $\hat{C}(2)$ | $=\left\{\mathrm{fn} x \Rightarrow[x]^{1}\right\}$ |
| $\hat{C}(3)$ | $=\emptyset$ |
| $\hat{C}(4)$ | $=\left\{\mathrm{fn} y \Rightarrow[y]^{3}\right\}$ |
| $\hat{C}(5)$ | $=\left\{\mathrm{fn} y \Rightarrow[y]^{3}\right\}$ |
| $\hat{\rho}(x)$ | $=\left\{\mathrm{fn} y \Rightarrow[y]^{3}\right\}$ |
| $\hat{\rho}(y)$ | $=\emptyset$ |

One interesting bit here in the solution is: $\hat{\rho}(y)=\emptyset$ : that means, the variable $y$ never evaluated, i.e., the function is not applied at all.

### 1.4 Type and effect systems

### 1.4.1 Introduction

## Standard type systems and others

In this section, we are mainly dealing with "non-standard" type systems, but let's have a look at standard type systems first.

Types and type systems are a well-established and central part of static analysis. Before continuing further: It should go without saying that, as far as the lecture is concerned, we are dealing with static type systems. The typing part of the semantic analysis phase is also kind of familiar as all programmers are accustomed to in one form or the other, as they are often visible in the programming language and the programmer has to learn to deal with them. For languages being typed at run-time (for instance some scripting languages). types may be less visible, even if they are still there (but dynamically typed languages are out of scope for that lecture anyway). In any case, even novice programmers are aware that somewhere under the hood, the compiler checks whether the given program adheres to the language's typing discipline; that's the task of the type checker.

The "standard" role of types is to describe "data values" and type checking assures that meaningful use is made of the date. At the lowest level, it served the compiler, for instance, also to know the "size" of data so as to allocate adequate amount of memory for storing and accessing it. The lecture here covers types mostly on a higher abstraction level, and types are seen as specifications of allowed uses of data. For instance, the type system may allow to use the logical operations and and or on boolean typed values, but not addition. On that level, type and type systems are very rudimentary and potentially very rigid (but still important). Modern (standard) type system are far more complex, basically due to the wish to add flexibility (different forms of so-called polymorphism) but still maintaining efficient, scalable, decidable static type checking.

Type theory - the study of type systems, their expressiveness, efficiency etc.often deals with functional languages, one reason being that functional languages feature one particular expressive form of data, namely functions. There are other reasons why type systems for functional languages have been widely studied, but that's perhaps outside the scope of the lecture and we leave it at that.

At any rate, whether it's the more down to earth "size-of-memory" basic types for code generation, or advanced polymorphic type systems for some $\lambda$-calculus or other, standard type systems are always concerned with specifying sets of data values (with the purpose of regulating their usage).

So far for standard type systems, what about non-standard ones? The latter term here is used for all aspects different from describing values in a programming language. A value of a program is "what comes out at the end", that's why executing a program is sometimes called evaluation... Another name we will encounter is reduction, the intuition being that the program is "reduced" to its final value. Anyhow, non-standard type systems, in contrast, specify aspects that are not related to the final value. For instance, data flow or control flow information is certainly something that typically is not covered by type system. Effects, in particular, cover aspects that happen "during" the execution. It's not a coincidance that this is a very broad definition, as basically, everything that needs semantic analysis at compile time, which refers to "what happens when the program runs" (as opposed to "what's the final value") might be described by a effect system. Often, the desciption is combined with a type system, in which case it's called a type and effect system. The lecture will probably cover different effects (excepctions, communication, ...).

For Haskell progammers, the separation between the "pure", functional part and all "the other aspects" should be familiar. Haskell goes to great lengths to separate both aspects, encapsulating all impure effects in so-called monads. So one might call standard types all that captures the pure (effect-free) core of the language, and effects correspond to those covered by monads. The connection between effects and monads can even be made formal (giving a monadic interpretation to effects), but that's beyond this lecture.

## Effects: Intro

- type system: "classical" static analysis:

$$
t: T
$$

- judgment: "term or program phrase has type $T$ "
- in general: context-sensitive judgments (remember Chomsky ...)


## Judgement :

$$
\Gamma \vdash t: \tau
$$

- $\Gamma$ : assumption or context
- here: "non-standard" type systems: effects and annotations
- natural setting: typed languages, here: trivial! setting (while-language)

Type system is a classical context sensitive analysis, following parsing, which is the classical context-free analysis. It's thus not a coincidence, that the component $\Gamma$ in the judgements is called "context". In implementations, it directly
corresponds to the so-called symbol table. That's a data structure, often realized as hash table or similar, used to keep relevant information about syntactic entities during the semantical phase (and also for code generation), in particular about variables or other "symbols". One prominent piece of information is about the types, and that's what $\Gamma$ contains. When coming to effect systems and some non-standard types, also other kind of information may be stored in the types, and therefore in $\Gamma$, as well.

For the participants for the compiler construction course (INF5110): information attached to syntactic elements were there also called attributes. In that parlance, a type is an attribute of a variable (or an expression etc.). Note in passing, that also the labelling of expressions can well be seen as attribuation.

Context-sensitive information assosciated with syntactic entities was called attributes basically because in the static analysis part, the lecture was working with so-called attribute grammars. In this lecture we don't bother with that general formalism. The type and effect systems will be represented in the form of derivation systems, but in priciple that can be viewed as some special notation for specific attribute grammars, as well.

## "Trival" type system

- setting: while-language
- each statement maps: state to states
- $\Sigma$ : type of states
judgement

$$
\begin{equation*}
\vdash S: \Sigma \rightarrow \Sigma \tag{1.12}
\end{equation*}
$$

- specified as a derivation system
- note: partial correctness assertion

The "type system" here is trivial in a technical sense. Type systems are used to distinguish well-typed programs from ill-typed ones, and reject the latter. This is generally not a context-free task, and therefore cannot be done by the parser alone, even though, for simple languages, type checking can be done while parsing. Here, the "type system" (show below) accepts all programs, all syntactically correct programs are already well-typed (remember that we are dealing with abstract syntax, i.e., trees, which represent syntactically correct programs). Since all programs are well-typed, there is actually no need for a type system at all. It's only used here to illustrate how to extend a (in this case trivial) type system with extra information, leading to annotated type system and effect type system, which then yield useful information. Note in passing that the above judgment from equation (1.12) has no context $\Gamma$, reflecting the
fact that it's technically context-free and not context-sensitive, and thereby trivial.

The typing judgment, as is the case in general for standard type system, is intended to be a partial correctness assertion. The trivial type system here is a bad illustation for that fact (being so trivial), but in general, the meaning of a judgment $e: \tau$ is:
if the statement, expression or progam $e$ terminates, then the resulting value conforms to the type $\tau$.

Type systems typically don't attempt have an opinion about termination, they only guarantee that the data at the end is ok, provided the program terminates thereby yielding said value. This restricted form of assertion is known as partial correctness (as opposed to total correctness). Another word used for such specifications is, that typing is conventionally concerned with safety properties (as opposed to liveness properties).

This is not intended to say, that it's impossible to devise type systems that try to capture "total correctness" in that they would guarantee termination or would warn against possible non-termination. In general, that would require definitely "non-standard" augmented information.

Finally, as a side remark: there is a connection between termination and standard type system in the following way: for typed $\lambda$-calculi (without recursion), well-typed programs guarantee termination, which requires non-trivial techniques for proving that, and furthermore makes pure, typed $\lambda$-calculi (without recursion) no real programming languages, as they are not Turing-complete. The $\lambda$-calculus we will encounter later does have a recursion operator for that reason.

## "Trival" type system: rules

$$
\begin{aligned}
& \vdash[x:=a]^{l}: \Sigma \rightarrow \Sigma \quad \text { Ass } \\
& {[\text { skip }]^{l}: \Sigma \rightarrow \Sigma \quad \text { SKIP }} \\
& \frac{\vdash S_{1}: \Sigma \rightarrow \Sigma \quad S_{2}: \Sigma \rightarrow \Sigma}{\vdash S_{1} ; S_{2}: \Sigma \rightarrow \Sigma} \text { SEQ } \\
& \frac{\vdash S: \Sigma \rightarrow \Sigma}{\vdash \text { while }[b]^{l} \operatorname{do} S: \Sigma \rightarrow \Sigma} \text { WHILE } \\
& \frac{\vdash S_{1}: \Sigma \rightarrow \Sigma \quad \vdash S_{2}: \Sigma \rightarrow \Sigma}{\vdash \text { if }[b]^{l} \text { then } S_{1} \text { else } S_{2}: \Sigma \rightarrow \Sigma} \text { COND }
\end{aligned}
$$

As mentioned, the "type system" does not do anything useful. It shows, however, the general style of writing down type systems, namely in the form of derivation rules. In the current version, there are five rules, one for each construct. Each rule has a set of premises and one conclusion. In case, there are no premises, a rule is also called axiom. That's the case for Ass and SEQ, which are dealing with the basic constructs of the language. The compound statement are treated by the rules with a non-empty set of premises, where the premises deal with the sub-constructs.

The rules can be viewed a logical "implications", reading them from top to bottom: If $S_{1}$ is well-typed and $S_{2}$ is well-typed, then so is the sequential $S_{1} ; S_{2}$ composition (in rule SEQ). A program $S$ is well-typed, if there is a derivation tree using the givem rules such that $\vdash S: \Sigma \rightarrow \Sigma$ is derivable, i.e., is the root of the tree (derivation trees have their roots at the bottom and their leaves, corresponding to axioms, at the top... )

One can view the rules also as the specification of a recursive procedure: in order to establish that $S_{1} ; S_{2}$ is well-typed, one has to recursively check $S_{1}$ and $S_{2}$ for well-typedness. It's of course just a different angle on the same thing. This "reading" of the rules is sometimes called "goal-directed": in order to establish the conclusion, establish the premises first.

Seen in this goal-directed manner, the rules directly can be seen as a recursive procedure (corresponding to a tree-traversal of the abstract syntax tree). It's a very straight-forward divide and conquer strategy. Note in this context: the language has 5 syntactic constructs ( 2 basic ones and 3 compound ones) and
the derivation system has 5 rules (including 2 axioms), exactly one for each contruct.

That entails that the top-level constuct of a subprogram determines which rule to apply recursively (in the goal-directed reading of the rules). Furthermore, the premises always deals with proper subterms compared to the term or statement in the conclusion, which in particular guarantees termination. Type systems, or derivation systems in general with these "one-rule-per-construct" and "in-the-premises-subterms-only" properties are called syntax directed. In the terminology of attribute grammars: the types here correspond to inherited attributes (in their simplest form). But keep in mind that the "type system" here is trival to the point of being meaningless. More realistic type systems do not correspond to inherited attributes, they are more complex.

Unfortunately, not all type systems are given in a syntax-directed manner. That means, not all type system specifications can be immediately understood as an algorithm. Sometimes, that is for fundamental reasons: the rules describe a type system so complex that the question whether $\Gamma \vdash e: \tau$ is derivable or not is undecidable. It won't be much the case in this lecture. A more common reason is: the rules of the type system are not a priori intended as an algorithm, they are rather intended as specification of the typing discipline and showing an algorithm would obscure that specification. A non-syntax directed specification for instance will be used when dealing with subtyping or other forms of polymorphism, resp. subeffecting. We will also touch upon the question: given a non-syntax-directed type system as specification, how can one turn it into a syntax-directed, thus algorithmic version. The algorithm, of course, corresponds to the type checker.

## Types, effects, and annotations

$$
\begin{equation*}
\vdash S: \Sigma_{1} \rightarrow \Sigma_{2} \tag{1.13}
\end{equation*}
$$

$$
\begin{equation*}
\vdash S: \Sigma \xrightarrow{\varphi} \Sigma \tag{1.14}
\end{equation*}
$$

type and effect system (TES)

- effect system + annotated type system
- borderline fuzzy
- annotated type system
- $\Sigma_{i}$ : property of state (" $\Sigma_{i} \subseteq \Sigma$ ")
- "abstract" properties: invariants, a variable is positive, etc.
- effect system
- "statement $S$ maps state to state, with (potential ...) effect $\varphi$ "
- effect $\varphi$ : e.g.: errors, exceptions, file/resource access, ...


### 1.4.2 Annotated type systems

## Annotated type systems

- example again: reaching definitions for while-language
- 2 flavors

1. annotated base types: $S: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2}$
2. annotated type constructors: $S: \Sigma \underset{\mathrm{RD}}{\underset{X}{X}} \Sigma$

Here we see that the border line between annotated type systems and effect system is fuzzy. The first sub-flavor corresponds to the intuition we have used so far: the states are restricted. The second one, if we think of it as functional type, can be seen as if the "functional" type is annotated. However, the annotation is like an effect (as we will see later).

## RD with annotated base types

judgement

$$
\begin{equation*}
\vdash S: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2} \tag{1.15}
\end{equation*}
$$

- $\mathrm{RD} \subseteq 2^{\mathrm{Var} \times \text { Lab }}$
- auxiliary functions
- note: every $S$ has one "initial" elementary block, potentially more than one "at the end"
- $\operatorname{init}(S)$ : the (unique) label at the entry of $S$
- final $(S)$ : the set of labels at the exits of $S$
"meaning" of judgment $\vdash S: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2}$ " $\mathrm{RD}_{1}$ is the set of var/label reaching the entry of $S$ and $\mathrm{RD}_{2}$ the corresponding set at the exit(s) of $S "$ :

$$
\begin{aligned}
& \mathrm{RD}_{1}=\mathrm{RD}_{\text {entry }}(\text { init }(S)) \\
& \mathrm{RD}_{2}=\bigcup\left\{\operatorname{RD}_{\text {exit }}(l) \mid l \in \operatorname{final}(S)\right\}
\end{aligned}
$$

Concerning the "meaning" of the judgment: the formulation is not $100 \%$ correct, it's too strict as we will see. The problem is the claim that RD is the set of ...... As in the data flow section, there is not just one single safe solution, but many. There is (as before) exactly one minimal, i.e., best one. However, the effect system is "lax" in that it specifies all safe ones. That is completely in analogy to the previous constraint system approaches.

One may compare the sets RD to the analysis data used in the "original" reaching definitions analysis (in the equational or constraint based approach, it does not matter which). The "functional" type here expresses the preand post-states. For elementary blocks, that corresponds to the entry and
the exit point of the node or label. As one rule (coming next) treats one block/statement (elementary or not) at a time, it's not the 12-tuple, of course, (taking the concrete factorial example) but just 2 generic slots of it: the preand the post-state.

Concerning the auxiliary functions (initial and final): They calculate implicitly the control flow graph.

## Rules

$$
\begin{aligned}
& \vdash[x:=a]^{l^{\prime}}: \mathrm{RD} \rightarrow \mathrm{RD} \backslash\{(x, l) \mid l \in \mathrm{Lab}\} \cup\left\{\left(x, l^{\prime}\right)\right\} \quad \text { ASS } \\
& \vdash[\text { skip }]^{l}: \mathrm{RD} \rightarrow \mathrm{RD} \quad \text { SKIP } \\
& \frac{\vdash S_{1}: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2} \quad \vdash S_{2}: \mathrm{RD}_{2} \rightarrow \mathrm{RD}_{3}}{\vdash S_{1} ; S_{2}: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{3}} \mathrm{SEQ} \\
& \frac{\vdash S_{1}: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2} \quad \vdash S_{2}: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2}}{\vdash \mathrm{if}[b]^{l} \text { then } S_{1} \text { else } S_{2}: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2}} \mathrm{IF} \\
& \frac{\vdash S: \mathrm{RD} \rightarrow \mathrm{RD}}{\vdash \text { while }[b]^{l} \text { do } S: \mathrm{RD} \rightarrow \mathrm{RD}} \mathrm{WHILE} \\
& \stackrel{\vdash S: \mathrm{RD}_{1}^{\prime} \rightarrow \mathrm{RD}_{2}^{\prime} \quad \mathrm{RD}_{1} \subseteq \mathrm{RD}_{1}^{\prime} \quad \mathrm{RD}_{2}^{\prime} \subseteq \mathrm{RD}_{2}}{\vdash S: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2}} \mathrm{SUB}
\end{aligned}
$$

The rules may be compared with the constraint-based formulation of the reaching definitions early, which was based on the control-flow graph. The intra-block equations are covered by the axioms here, and the inter-block equations are the compositional part, the rules.

Worth mentioning is also the fact that in the annotated type system now, we have 6 rules (for 5 syntactic constructs). So, there goes the syntaxdirectedness.... As a consequence, the type system does not (directly) describe an algorithm.

The culprit is rule SUB (for subsumption). We will encounter subsumption many times (for subeffecting, and also subtyping), and it's one classical reason why type systems are not syntax directed. It breaks it in two ways: first, for each construct, there are now two rules to choose from (if one thinks in a goal-directed manner), as subsumption is always possible. as well. Secondly, the core judgment in the premise does not assert well-typedness for a proper
sub-term of $S$ in the conclusion. So, going from conclusion to the premise, $S$ does not get "smaller". Suddenly, termination of type checking may become a non-trivial issue.

## Meaning of annotated judgments

"Meaning" of judgment $S: \mathrm{RD}_{1} \rightarrow \mathrm{RD}_{2}$ : " $\mathrm{RD}_{1}$ is the set of var/label reaching the entry of $S$ and $\mathrm{RD}_{2}$ the corresponding set at the exit(s) of $S "$ :

$$
\begin{aligned}
& \mathrm{RD}_{1}=\mathrm{RD}_{\text {entry }}(\text { init }(S)) \\
& \mathrm{RD}_{2}=\bigcup\left\{\mathrm{RD}_{\text {exit }} l \mid l \in \operatorname{final}(S)\right\}
\end{aligned}
$$

- Be careful:

$$
\text { if }[b]^{l} \text { then } S_{1} \text { else } S_{2}
$$

- more concretely

$$
\text { if }[b]^{l} \text { then }[x:=y]^{l_{1}} \text { else }[y:=x]^{l_{2}}
$$

## Derivation



- abbreviate $f_{3}=$ while $\ldots ;[y:=0]^{6}$
- loop invariant

$$
\mathrm{RD}_{\text {bod } y}=\left\{?_{x}, 1,5,2,4\right\}
$$

### 1.4.3 Annotated type constructors

## Annotated type constructors

- alternative approach of annotated type systems
- arrow constructor itself annotated
- annotion of $\rightarrow$ : flavor of effect system
- judgment

$$
S: \Sigma \underset{\mathrm{RD}}{X} \Sigma
$$

- annotation with RD (corresponding to the post-condition from above) alone is not enough
- also needed: the variables "being" changed

Intended meaning " $S$ maps states to states, where RD is the set of reaching definitions, $S$ may produce and $X$ the set of var's $S$ must (= unavoidably) assign.

In the previous formulation (with annotated base types), each judgment mentioned two versions of the RD information, the one before and the one after. Now, there is only one RD information, which is interpreted as the reaching definitions the statement of the rule may produce. That means, the generated RD is considered very much like an effect of the statement.

RD is indeed the information we are interested in for this analysis. However, to make the system technically work and fit together: It's not enough to keep track of "reaching definitions" which are generated per construct. If that were the only information, we would never "remove" any tuples, just add them. That can in particularly seen in the treatment of sequential composition $S_{1} ; S_{2}$, see rule SEQ below.

In order to capture that, the sets $X$ of assigned variables is kept track of, as well. In the monotone frameworks later, the removal of flow information is also called "killing" and, in many cases, the data flow equations and transfer functions can be decscribed as a combination of "generating" and "killing" data flow. It could also be noted: the RD here is "may" information, wheras the $X$ is "must" information.

## Rules

$$
\begin{aligned}
& {[x:=a]^{l}: \Sigma \underset{\{(x, l)\}}{\{x\}} \Sigma \quad \text { ASS } \quad[\text { skip }]^{l}: \Sigma \xrightarrow[\emptyset]{\emptyset} \Sigma \quad \text { SkIP }} \\
& \frac{S_{1}: \Sigma \underset{\mathrm{XD}_{1}}{\mathrm{XD}_{1}} \Sigma \quad S_{2}: \Sigma \frac{X_{2}}{\mathrm{RD}_{2}} \Sigma}{S_{1} ; S_{2}: \Sigma \underset{\mathrm{RD}_{1} \backslash X_{2} \backslash \mathrm{X}_{2} \mathrm{URD}_{2}}{X_{1} \cup X_{2}} \Sigma} \mathrm{SEQ}
\end{aligned}
$$

$$
\begin{aligned}
& \xrightarrow[{\text { while }[b]]^{l} \text { do } S: \Sigma \xrightarrow[\mathrm{RD}]{\xrightarrow{D}}} \Sigma]{S} \text { WHILE } \\
& \frac{S: \Sigma \underset{\mathrm{RD}^{\prime}}{X^{\prime}} \Sigma \quad X \subseteq X^{\prime} \quad \mathrm{RD}^{\prime} \subseteq \mathrm{RD}}{S: \Sigma \xrightarrow[\mathrm{RD}]{X} \Sigma} \mathrm{SUB}
\end{aligned}
$$

In [8], the IF rule is formulated more complex:

$$
\frac{S_{1}: \Sigma \underset{\mathrm{RD}_{1}}{X_{1}} \Sigma \quad S_{2}: \Sigma \underset{\mathrm{RD}}{\mathrm{RD}_{2}} \Sigma}{\underset{\mathrm{R}}{2}[b]} \text { then } S_{1} \text { else } S_{2}: \Sigma \underset{\mathrm{RD}_{1} \cup \mathrm{RD}_{2}}{\mathrm{X}_{2} \cap X_{2}} \Sigma \mathrm{IF}
$$

The formulations are, of course, equivalent. The one on the slides is a special case of the more complex one. For the reverse direction, one can use subsumption. In the presence of the more complex rule, one can remove subsumption!

Interesting is the While-rule, especially the fact that the set above the arrow is $\emptyset$. That's because the while-loop may not be taken at all, so that corresponds to the intersection of $X$ in the premise and $\emptyset$.

### 1.4.4 Effect systems

## Effect systems

- this time: back to the functional language
- starting point: simple type system
- judgment:

$$
\Gamma \vdash e: \tau
$$

- $\Gamma$ : type environment (or context), "mapping" from variable to types
- types: bool, int, and $\tau \rightarrow \tau$

The "language" here basically is known as the simply typed $\lambda$-calculus. It is a variant which is known as Curry-style formulation of the type system. That refers to the fact that abstractions are written as $\lambda x$.e instead of $\lambda x: \tau . e$. In the latter case, it's know as "Church style". In the less explicit Curry-style, one is facing a problem of so-called type inference, i.e., figuring out what the (omitted) type of $x$ should be. That in itself is an important and interesting problem, which we might cover later, but in the introduction, we focus on the effect part, not the typing.

As far as the syntax is concerned: The $\pi$-subscript of the abstraction is nonstandard as far as the typed $\lambda$-calculus and its typing is concerned. It's added for the purpose of the effect system, only.

In comparison to the earlier "type-system" we used a starting point for the while language, now the system is non-trivial (even if completely standard).

## Rules

$$
\begin{aligned}
& \frac{\Gamma(x)=\tau}{\Gamma \vdash x: \tau} \mathrm{VAR} \\
& \frac{\Gamma, x: \tau_{1} \vdash e: \tau_{2}}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e: \tau_{1} \rightarrow \tau_{2}} \mathrm{ABS} \\
& \frac{\Gamma \vdash e_{1}: \tau_{1} \rightarrow \tau_{2} \quad \Gamma \vdash e_{2}: \tau_{1}}{\Gamma \vdash e_{1} e_{2}: \tau_{2}} \mathrm{APP}
\end{aligned}
$$

As mentioned, the flavor of the (simply typed) $\lambda$-calculus here is Curry-style. As before, an easy way to interpret the rules is as derivation rules, where the premises imply the conclusion.

It may be also intructive, to look at them in a goal directed manner, as if it where a recursive procedure: what is needed in order to derive a conclusion.

But even more instructive is to think what such a recursive interpretation is supposed to achieve. To be useful as conventional type checker, it's not meant as: check if the following is true

$$
\begin{equation*}
\Gamma \vdash e: \tau ? \tag{1.16}
\end{equation*}
$$

like confirming the user's guess whether or not it's true that $e$ has type $\tau$ (given the assumptions $\Gamma$ ). Instead, more useful is the interpretation

$$
\begin{equation*}
\Gamma \vdash e: ? \tag{1.17}
\end{equation*}
$$

i.e., an answer to the question: is $e$ well-typed under the given assumption, and if so, what's its type? In other words, when interpreting the rules, one useful reading is to see $\Gamma$ and $e$ as input of a recursive procedure, and $\tau$ as the output.

With this interpretation in mind, especially rule ABS is interesting. Here, in the Curry-style formulation, the formal parameter $x$ is mentioned in the conclusion without type. That means, that the "recusive call" corresponding to the premise guesses a type $\tau_{1}$ for it. This guess may be right in that it leads to a successful type check of the premise and thus a sucessful type check and returning $\tau_{1} \rightarrow \tau_{2}$ for the abstraction in the conclusion. Or it may also fail. The form of abstraction may determined that a recursive call according rule APP is in place, however, not information is given in the input $\Gamma$ and $\lambda x$.e to determine the subesequent recusive call (as $\tau_{1}$ has to be guess). Note that there are infinitely many types to choose from.... In other words, the rules (in the interpretation given) do not really qualify as algorithm, and they would not qualify as syntax-directed (if we interpret the rules as describing a problem as in equation (1.17) and not as confirming the type as in equation (1.16).

Solving the "guessing problem" in a proper manner is known as type inference or type reconstruction and we might return to it later. Also note: if the abstraction would be of the form $\lambda x: \tau . e$, the guessing problem would go away as well. This form is also known a Church-style typing (as opposed to Currystyle).

## Effects: Call tracking analysis

Call tracking analysis: Determine: for each subexpression: which function abstractions may be applied, i.e., called, during the subexpression's evaluation.
$\Rightarrow$ set of function names
annotate: function type with latent effect
$\Rightarrow$ annotated types: $\hat{\tau}$ : base types as before, arrow types:

$$
\begin{equation*}
\hat{\tau}_{1} \xrightarrow{\varphi} \hat{\tau}_{2} \tag{1.18}
\end{equation*}
$$

- functions from $\tau_{1}$ to $\tau_{2}$, where in the execution, functions from set $\varphi$ are called.


## Judgment

$$
\begin{equation*}
\hat{\Gamma} \vdash e: \hat{\tau}::: \varphi \tag{1.19}
\end{equation*}
$$

It may be worthwhile to reflect what the connection is between the call-tracking analysis here and the constraint-based analysis we had before. Both analyses are not the same. In the constraint-based analysis from earlier, we wanted to know at each point to which functions it evaluates to answer the question for applications $f a$, which functions are actually called. Here, the perspective is different: we take an expression and think of it as being evaluated and ask, which functions are being called during evaluation. Both questions, however, are related. Remember that the analysis earlier was not formulated as a type system.

Later in the lecture, we may revisit the control-flow analysis in a type-based formulation. This will be easier to compare to the call-tracking analysis from here, basically because it's formulated similarly, namely as an effect system quite similar to the one here. This allows to see the differences and similarities more clearly. The earlier control-flow analysis was not formalized as type system, but with constraints. It's not, however, meant to mean that constraint based systems are fundamentally different from type and effect systems. After all they may tackle the same problems. Again it's more a stylistic question and type systems as the one here can also be seen as a particular way (using logical derivation rules as used for type system) so specify the contraint system.

## Call tracking rules

$$
\begin{aligned}
& \frac{\hat{\Gamma}(x)=\hat{\tau}}{\hat{\Gamma} \vdash x: \hat{\tau}:: \emptyset} \mathrm{VAR} \\
& \frac{\Gamma, x: \hat{\tau}_{1} \vdash e: \hat{\tau}_{2}:: ~ \varphi}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e: \hat{\tau}_{1} \xrightarrow{\varphi \cup \pi\}} \hat{\tau}_{2}:: \emptyset} \mathrm{ABS} \\
& \frac{\hat{\Gamma} \vdash e_{1}: \hat{\tau}_{1} \xrightarrow{\varphi} \hat{\tau}_{2}:: \varphi_{1} \quad \hat{\Gamma} \vdash e_{2}: \hat{\tau}_{1}:: \varphi_{2}}{\hat{\Gamma} \vdash e_{1} e_{2}: \hat{\tau}_{2}:: \varphi \cup \varphi_{1} \cup \varphi_{2}} \mathrm{APP}
\end{aligned}
$$

## Call tracking: example

$$
\frac{x: \text { int } \xrightarrow{\{Y\}} \text { int } \vdash x: \text { int } \xrightarrow{\{Y\}} \text { int }:: \emptyset}{\vdash\left(\mathrm{fn}_{X} x \Rightarrow x\right):(\text { int } \xrightarrow{\{Y\}} \text { int }) \xrightarrow{\{X\}}(\text { int } \xrightarrow{\{Y\}} \text { int }):: \emptyset \quad \vdash\left(\mathrm{fn}_{Y} y \Rightarrow y\right): \text { int } \xrightarrow{\{Y\}} \text { int }:: \emptyset}
$$

### 1.5 Algorithms

## Introduction

This part is rather short. So far, we touched upon here and there on issues how to algorithmically treat the problems we encountered. In particular in the context of the type systems, some remarks tried to raise awareness under which circumstances the derivation rules could straightforwardly be interpreted as a recursive procedure, and when not (basically in all interesting cases). But we never really tackled the issue, especially not for the "flow problems" (data flow, effects etc.) in which we are specifically interested in.

In particular: we saw how data flow problems such as reaching definitions can be described or specified as constraint systems (equational or otherwise). What we did not do so far is giving hints of how to actually solve such constraint system, i.e., how do do constraint solving.

The notion of "constraint system" is extremely broad and can capture all sorts of problems. The lecture is mostly concerned with particular forms of constraint systems which capture flow problems (or program analysis problem in general). In addition, we discuss under which circumstances those kind of constraint systems have solutions, in particular have unique best solutions. This is a very welcome sitation and deserve a thorough treatment.

Here, we just hint at a very simple strategy (also only sketched on a very highlevel), which is non-deterministic. Being non-deterministic makes it not really directly an implementation unless one wishes to make use of some randomgenerator which helps to implement the non-determinism. There would in practice no point in doing so, instead one would aim for deterministic solution, perhaps realizing specific stratetgies or heuristic.

Why then bother with a non-deterministic description at all? Basically it's to separate the question of correctness of the algorithm from the question of efficiency. As it turns out, the kind of constraints we are considering and the solution domains have the very, very desirable property that
when facing a non-deterministic, choice, no matter how one resolve it, the one never make a "wrong choice".

So the choice does not really matter except for how fast the algorithm terminates, i.e., how efficient the algo runs. Even there are perhaps smarter and less smart choices at each point, there are not real wrong ones. As a consequence, there is no backtracking. And that's crucial for being usable. As a side remark: it's well-known that many forms of constraints, even simple ones as boolean constraints ("SAT-solving") don't have this favorable property and there is basically no way around solving those by brute force combinatorial exploration. Indeed, SAT (boolean satisfiability) one of the most famous NP complete problems there is (and the first one for which that was proven). Fortunately, for simple data flow equations, things look much brighter.

## Chaotic iteration

- back to data flow/reaching def's
- goal: solve

$$
\overrightarrow{R D}=F(\mathrm{RD}) \quad \text { or } \quad \overrightarrow{R D} \sqsubseteq F(\mathrm{RD})
$$

- $F$ : monotone, finite domain


## straightforward approach

init $\overrightarrow{R D}_{0}=F^{0}(\emptyset)$
iterate $\overrightarrow{\mathrm{RD}}_{n+1}=F\left(\overrightarrow{\mathrm{RD}}_{n}\right)=F^{n+1}(\emptyset)$ until stabilization

- approach to implement that: chaotic iteration
- non-deterministic stategy
- abbreviate:

$$
\overrightarrow{\mathrm{RD}}=\left(\mathrm{RD}_{1}, \ldots, \mathrm{RD}_{12}\right)
$$

## Chaotic iteration (for RD)

```
Input: equations for reaching defs
    for the given program
Output: least solution: R\vec{D}=(R\mp@subsup{D}{1}{},\ldots,R\mp@subsup{D}{12}{})
```

Initialization:

$$
\mathrm{RD}_{1}:=\emptyset ; \ldots ; \mathrm{RD}_{12}:=\emptyset
$$

Iteration:
while $\mathrm{RD}_{j} \neq F_{j}\left(\mathrm{RD}_{1}, \ldots, \mathrm{RD}_{12}\right)$ for some $j$
do

$$
\mathrm{RD}_{j}:=F_{j}\left(\mathrm{RD}_{1}, \ldots, \mathrm{RD}_{12}\right)
$$

### 1.6 Conclusion

The introductory part touched upon different topics. The approaches are also related, i.e., it's sometimes a bit of a matter of preference if one represents a problem directly as flow equations, type systems etc. Things not covered in the introduction (but probably later are complications in the while language, like procedure calls or pointers). Also, there will be other analyses besides reaching definitions (and a systematic common overview over similar analyses (knows as monotone framework). Also we go into the underlying theory (lattices) as well has considering in which way to establish that the various analysis are actually a sound overapproximation of the program behavior ("soundness", "correctness", "safe approximation", all mean the same).

## Learning Targets of this Chapter

various DFAs
monotone frameworks operational semantics
foundations
special topics (SSA, context-sensitive analysis ...)

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What
is it
about?

### 2.1 Introduction

In this part we cover classical data flow analysis, first in a few special, specific analyses, among other ones, more time reaching definitions. Besides that, also different other well-known ones. Those analyses are based on very similar common principles, which then lead to the notion of monotone framework. All of this is done for the simple while language from the general introduction. We also have a look at important extensions. One is the treatment of procedures. Those will be first-order procedures, not higher-order procedures. Nonetheless, they are already complicating the data flow problem (and its complexity), leading to what is known as context-sensitive analysis. Another extension deals with dynamically allocated memory on heaps. Analyses that deal with that particular language feature are known as alias analysis, pointer analysis, and shape analysis. Also we might cover SSA this time.

### 2.2 Intraprocedural analysis

### 2.2.1 Determining the control flow graph

## While language and control flow graph

- starting point: while language from the intro
- labelled syntax (unique labels)
- labels $=$ nodes of the cfg
- initial and final labels
- edges of a cfg: given by function flow

Determining the edges of the control-flow graph Given an program in labelled (and abstract) syntax, the control-flow graph is easily calculated. The nodes we have already (in the form of the labels), the edges are given by a function flow. This function needs, as auxiliary functions, the functions init and final

The latter 2 functions are of the following type:

$$
\begin{equation*}
\text { init }: \text { Stmt } \rightarrow \text { Lab } \quad \text { final }: \text { Stmt } \rightarrow 2^{\text {Lab }} \tag{2.1}
\end{equation*}
$$

Their definition is straightforward, by induction on the labelled syntax:

|  | init | final |
| :---: | :---: | :---: |
| $[x:=a]^{l}$ | $l$ | $\{l\}$ |
| $[\text { skip }]^{l}$ | $l$ | $\{l\}$ |
| $S_{1} ; S_{2}$ | init $\left(S_{1}\right)$ | final $\left(S_{2}\right)$ |
| if $[b]^{l}$ then $S_{1}$ else $S_{2}$ | $l$ | final $\left(S_{1}\right) \cup$ final $\left(S_{2}\right)$ |
| while $[b]^{l}$ do $S$ | $l$ | $\{l\}$ |

The label $\operatorname{init}(S)$ is the entry node to the graph of $S$. The language is simple and initial nodes are unique, but "exits" are not. Note that unique entry is not the same as the notion of "isolated" entry (mentioned already in the introduction). Isolated would mean: the entry is not the target of any edge. That's not the case, for instance for the while loop. In general, however, it may be preferable to have an isolated entry, as well, and one can arrange easily for that, adding one extra sentinel node.

Using those, determining the edges, by a function

$$
\text { flow }: \mathbf{S t m t} \rightarrow 2^{\text {Lab } \times \text { Lab }}
$$

works as follows:

$$
\begin{align*}
\operatorname{flow}\left([x:=a]^{l}\right)= & \emptyset  \tag{2.3}\\
\text { flow }\left([\text { skip }]^{l}\right)= & \emptyset \\
\operatorname{flow}\left(S_{1} ; S_{2}\right)= & \operatorname{flow}\left(S_{1}\right) \cup \text { flow }\left(S_{2}\right) \\
& \cup\left\{\left(l, \text { init }\left(S_{2}\right)\right) \mid l \in \operatorname{final}\left(S_{1}\right)\right\} \\
\text { flow }\left(\text { if }[b]^{l} \text { then } S_{1} \text { else } S_{2}\right)= & \operatorname{flow}\left(S_{1}\right) \cup \text { flow }\left(S_{2}\right) \\
& \cup\left\{\left(l, \text { init }\left(S_{1}\right)\right),\left(l, \text { init }\left(S_{2}\right)\right)\right\} \\
\text { flow }\left(\text { while }[b]^{l} \text { do } S\right)= & \operatorname{flow}\left(S_{1}\right) \cup\{l, \text { init }(S)\} \\
& \cup\left\{\left(l^{\prime}, l\right) \mid l^{\prime} \in \operatorname{final}(S)\right\}
\end{align*}
$$

Two further helpful functions In the following, we make use of two further (very easy) functions with the following types

$$
\text { labels: Stmt } \rightarrow 2^{\text {Lab }} \text { and } \quad \text { blocks }: \text { Stmt } \rightarrow 2^{\text {Stmt }}
$$

They are defined straightforwardly as follows:

$$
\begin{align*}
\text { blocks }\left([x:=a]^{l}\right) & =[x:=a]^{l}  \tag{2.4}\\
\text { blocks }\left([\text { skip }]^{l}\right) & =[\text { skip }]^{l} \\
\text { blocks }\left(S_{1} ; S_{2}\right) & =\text { blocks }\left(S_{1}\right) \cup \text { blocks }\left(S_{2}\right) \\
\text { blocks }\left(\text { if }[b]^{l} \text { then } S_{1} \text { else } S_{2}\right) & =\left\{[b]^{l}\right\} \cup \text { blocks }\left(S_{1}\right) \cup \text { blocks }\left(S_{2}\right) \\
\text { blocks }\left(\text { while }[b]^{l} \text { do } S\right) & =\left\{[b]^{l}\right\} \cup \text { blocks }(S)
\end{align*}
$$

$$
\begin{equation*}
\operatorname{labels}(S)=\left\{l \mid[B]^{l} \in \operatorname{blocks}(S)\right\} \tag{2.5}
\end{equation*}
$$

All the definitions and concepts are really straightforward and should be intuitively clear almost without giving a definition at all. One point with those definitions, though is the following: the given definitions are all "constructive". They are given by structural induction over the labelled syntax. That means, they directly describe recurvise procedures on the syntax trees. It's a leitmotif of the lecture: we are dealing with static analysis, which is a phase of a compiler, which means, all definitions and concepts need to be realized in the form of algorithms and data structures: there must be a concrete control-flow graph data structure and there must be a function that determines it.

## Flow and reverse flow

$$
\operatorname{labels}(S)=\operatorname{init}(S) \cup\left\{l \mid\left(l, l^{\prime}\right) \in \operatorname{flow}(S)\right\} \cup\left\{l^{\prime} \mid\left(l, l^{\prime}\right) \in \operatorname{flow}(S)\right\}
$$

- data flow analysis can be forward (like RD) or backward
- flow: for forward analyses
- for backward analyses: reverse flow $f l o w^{R}$, simply invert the edges


## Program of interest

- $S_{*}$ : program being analysed, top-level statement
- analogously Lab $_{*}$, Var $_{*}$, Blocks $_{*}$
- trivial expression: a single variable or constant
- $\operatorname{AExp}_{*}$ : non-trivial arithmetic sub-expr. of $S_{*}$, analogous for $\operatorname{AExp}(a)$ and $\operatorname{AExp}(b)$.
- useful restrictions
- isolated entries: $\quad\left(l, \operatorname{init}\left(S_{*}\right)\right) \notin \operatorname{flow}\left(S_{*}\right)$
- isolated exits $\quad \forall l_{1} \in \operatorname{final}\left(S_{*}\right) . \quad\left(l_{1}, l_{2}\right) \notin \operatorname{flow}\left(S_{*}\right)$
- label consistency

$$
\left[B_{1}\right]^{l},\left[B_{2}\right]^{l} \in \operatorname{blocks}(S) \quad \text { then } \quad B_{1}=B_{2}
$$

" labels the block $B$ "

- even better: unique labelling

Concerning label consistency: indeed, unique labelling is better . Otherwise nodes of the graph are "overlaid", i.e., there will be confusion wrt. predecessors and successors. Unique labelling is an very natural condition. When labelling the syntax (and building the control flow graph), one simply generate one label or node after the other, then naturally the blocks are unqiuely labelled. Label
consistency, but with non-unique labelling looks rather unnatural, at the first sight.

See also the operational semantics later, which preserves label consistency but not unique labelling (in the case of unrolling a while-construct). Actually, the semantics would preserve a better property, it seems to me. Not only is the labelling "consistent" in the sense defined here. But also the edges and neighbors of a node remains comparable. But the book does not point that out.

### 2.2.2 Availabe expressions

This is the first of a few classical data flow analyses we cover (like reaching definitions as well). The analysis can be used for a so-called common subexpression elimination. CSE is a program transformation or optimization which makes use of the available expression analysis. The idea is easy: if in a program, the analysis finds out that an expression is computed twice, it may pay off to store it the first time it's computed, and in the second occurence, look it up again.

Of course, it not just a syntactical problem, i.e., it's not enough to find syntactical occurrences of the same expression. In an imperative language and for expression containing variable, the content of variables mentioned in such expression may or may not have changed comparing different occurences of the same expression, and that has to be figured out via a specific data flow analysis, namely "available expressions" analysis.

## Avoid recomputation: Available expressions

$$
\begin{array}{lll}
{[x:=a+b]^{0} ;[y:=a * b]^{1} ;} & \text { while } & {[y>a+b]^{2}} \\
& \text { do } & \left([a:=a+1]^{3} ;[x:=a+b]^{4}\right)
\end{array}
$$

Goal For each program point: which expressions must have already been computed (and not later modified), on all paths to the program point.

- usage: avoid re-computation

One important aspect in the (informal) goal of the analysis is the use of the word "must". That's different form what was done for reaching definitions. There, it was about if a "definition" may reach a point in questions. It's also worthwhile to reflect about "approximation". As always, exact information is not possible, what's why we content ourselves with "must" information (or "may" in other case). In the case here (and related to it): if we have some safe set of available expressions, then a smaller set it safe, too. Again, for the
may-setting for reaching definition, enlarging sets was safe. The situation here is therefore dual,

What obviously also different is the nature or type of the information of interests. Here it's sets of expressions, in the reaching definitions it was sets containing pairs of locations and variables.

## Available expressions: general

- given as flow equations (not constraints, but not too crucial, as we know already)
- uniform representation of effect of basic blocks (= intra-block flow)


## 2 ingredients of intra-block flow

- kill: flow information "eliminated" passing through the basic blocks
- generate: flow information "generated new" passing through the basic blocks
- later analyses: presented similarly
- different analyses $\Rightarrow$ different kind of flow information + different killand generate-functions

In the introduction, the reaching definition analysis was done without explicitly mentioning kill and generate, but they where there implicitly anyway (for the intra-block equations).

## Available expressions: types

- interested in sets of expressions: $2^{\mathbf{A E x p}_{*}}$
- generation and killing:

$$
\text { kill }_{\mathrm{AE}}, \text { gen }_{\mathrm{AE}}: \mathbf{B l o c k s}_{*} \rightarrow 2^{\mathbf{A E x p}_{*}}
$$

- analysis: pair of functions

$$
\mathrm{AE}_{\text {entry }}, \mathrm{AE}_{\text {exit }}: \mathbf{L a b}_{*} \rightarrow 2^{\mathbf{A E x p}_{*}}
$$

Explanations $\mathbf{A E x p}_{*}$ can be taken as all arithmetic expressions occuring in the program, including all their subsexpression. To be hyper-precise, one may refine it in that trivial (sub-)expressions don't count. Trivial expressions are constants and single variables. Those trivial expressions are uninteresting from the perspective of available expressions and therefore are left out. They are likewise left out for the very busy expression analysis which will be discussed soon.

## Intra-block flow specification: Kill and generate

$$
\begin{aligned}
\operatorname{kill}_{\mathrm{AE}}\left([x:=a]^{l}\right) & =\left\{a^{\prime} \in \operatorname{AExp}_{*} \mid x \in f v\left(a^{\prime}\right)\right\} \\
\operatorname{kill}_{\mathrm{AE}}\left([\operatorname{skip}]^{l}\right) & =\emptyset \\
\operatorname{kill}_{\mathrm{AE}}\left([b]^{l}\right) & =\emptyset \\
\operatorname{gen}_{\mathrm{AE}}\left([x:=a]^{l}\right) & =\left\{a^{\prime} \in \operatorname{AExp}(a) \mid x \notin f v\left(a^{\prime}\right)\right\} \\
\operatorname{gen}_{\mathrm{AE}}\left([\operatorname{skip}]^{l}\right) & =\emptyset \\
g e n_{\mathrm{AE}}\left([b]^{l}\right) & =\boldsymbol{\operatorname { E E x p }}(b)
\end{aligned}
$$

Explanation The interesting case is of course the one for assignments (for generation, also the boolean equations are similar). An assignment kills all expressions, which contain the variable assigned to, and generates all (non-trivial) sub-expressions of the expression on the right-hand side of the assignment.

For generation, we have, however, to be careful: those sub-expressions of $a$ which contain the variable $x$ are of course not generated (because they are no longer "valid" after the assignment); note (on the next slide): the flow in a block is forward, and the flow at the exits depends on the in-flow in the following order:

1. first kill, and
2. then generate.

Because of this order, we cannot generate sub-expressions which contain $x$. The data flow analysis, at least those which are formulated with the help of kill and generate function, use them in that order. One might as well use killing and generating in the opposite order, but obviously, in that case, the exact definition of the kill and generate functions needs to take that into account and would have to be adapted to reflect that.

## Flow equations: $A E^{=}$

split into
nodes: intra-block equations, using kill and generate
edges: inter-block equations, using flow

## Flow equations for AE

$$
\begin{aligned}
\mathrm{AE}_{\text {entry }}(l) & = \begin{cases}\emptyset & l=\operatorname{init}\left(S_{*}\right) \\
\cap\left\{\mathrm{AE}_{\text {exit }}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in \operatorname{flow}\left(S_{*}\right)\right\} & \text { otherwise }\end{cases} \\
\mathrm{AE}_{\text {exit }}(l) & =\mathrm{AE}_{\text {entry }}(l) \backslash \operatorname{kill}_{\mathrm{AE}}\left(B^{l}\right) \cup \operatorname{gen}_{\mathrm{AE}}\left(B^{l}\right)
\end{aligned}
$$

where $B^{l} \in \operatorname{blocks}\left(S_{*}\right)$

- note the "order" of kill and generate

Explanation Apart from the fact that before we did not make use of some explicit kill and generate funnctions, the flow equations here are pretty similar to the ones for available expressions. One conceptual difference is the replacement of $\cap$ (must) by $\cup$ (may).

Note that the definition of the flow equations assume isolated entries, which can be seen at the equation for $\mathrm{AE}_{\text {entry }}(l)$, in the case where $l$ is the initial label (otherwise it would be a bit more complex). Note also: for $\mathrm{AE}_{\text {entry }}$, we must make the case distinction of initial nodes (no incoming edges) and others, otherwise: the empty intersection would be something like the "full set" of expressions.

As subtle and perhaps not too relevant remark in that condition: that the empty intersection corresponds to the "full set" is by definition (of ultimately dealing with lattices). That sounds strange, but it's ok due to the following observation: the initial node is the only node in the control flow graph which -being isolated- has no incoming edge. It's straightforward to see that all cfgs from the given syntax have that property. The one and only node without incoming edge is of course $\operatorname{init}\left(S_{*}\right)$ (if we assume isolated entries), Having an isolated entry is not guaranteed by the syntax, which means, we have to additionally assume it resp. ensure it.

As mentioned: be aware of the order of kill and generate in the equation for the exit: first, the killed ones are removed, then the generated ones are added. Because of that order, one must make sure, that no expressions are generated that contain the assigned variable.

## Available expressions

- forward analysis (as RD)
- interest in largest solution (unlike RD)
$\Rightarrow$ must analysis (as opposed to may)
- expression is available: if no path kills it
- remember: informal description of AE: expression available on all paths (i.e., not killed on any)
- illustration


## Example AE

$$
\begin{array}{lll}
{[x:=a+b]^{0} ;[y:=a * b]^{1} ;} & \text { while } & {[y>a+b]^{2}} \\
& \text { do } & \left([a:=a+1]^{3} ;[x:=a+b]^{4}\right)
\end{array}
$$



Worthwhile is (for instance) the entry of node / block $l_{2}$. At that point, expression $a+b$ is available. That's despite the fact that $a$ is changed inside the body of the loop!

As a side remark: before we mentioned that available expressions analysis is useful for common sub-expression elimination. The example shows that one has to be careful with that, nonetheless.

### 2.2.3 Reaching definitions

## Reaching definitions

- remember the intro
- here: the same analysis, but based on the new definitions: kill, generate, flow...

$$
[x:=5]^{0} ;[y:=1]^{1} ; \text { while }[x>1]^{2} \operatorname{do}\left([y:=x * y]^{3} ;[x:=x-1]^{4}\right)
$$



## Reaching definitions: types

- interest in sets of tuples of var's and program points i.e., labels:

$$
2^{\operatorname{Var}_{*} \times \mathbf{L a b}_{*}^{?}} \text { where } \mathbf{L a b}_{*}^{?}=\mathbf{L a b}_{*}+\{?\}
$$

- generation and killing:

$$
\text { kill }_{\mathrm{RD}}, \text { gen }_{\mathrm{RD}}: \text { Blocks }_{*} \rightarrow 2^{\mathbf{V a r}_{*} \times \mathbf{L a b}_{*}^{?}}
$$

- analysis: pair of mappings

$$
\mathrm{RD}_{\text {entry }}, \mathrm{RD}_{\text {exit }}: \mathbf{L a b}_{*} \rightarrow 2^{\mathbf{V a r}_{*} \times \mathbf{L a b}_{*}^{?}}
$$

The information is the same as in the introduction (except here, we are explict that it should be not just sets of variables, but that only the sets of variables of the program are of interests, which here is denoted as Var $_{*}$. Similarly for $\mathbf{L a b}_{*}$ ). But that's just a bit more precise (perhaps overly so).

Af ar as the mappings or functions $\mathrm{RD}_{\text {entry }}$ and $\mathrm{RD}_{\text {exit }}$ are concerned: In a practical implementation, one might use arrays for that. If the implementation identifies nodes by "numbers", one can have an integer-indexed standard array, which typically is fast way of prepresenting that information.

## Reaching defs: kill and generate

$$
\begin{aligned}
\operatorname{kill}_{\mathrm{RD}}\left([x:=a]^{l}\right) & =\{(x, ?)\} \cup \\
& \bigcup\left\{\left(x, l^{\prime}\right) \mid B^{l^{\prime}} \text { is assgm. to } x \text { in } S_{*}\right\} \\
\operatorname{kill}_{\mathrm{RD}}\left([\mathrm{skip}]^{l}\right) & =\emptyset \\
k i l l_{\mathrm{RD}}\left([b]^{l}\right) & =\emptyset \\
\operatorname{gen}_{\mathrm{RD}}\left([x:=a]^{l}\right) & =\{(x, l)\} \\
\operatorname{gen}_{\mathrm{RD}}\left([\mathrm{skip}]^{l}\right) & =\emptyset \\
\operatorname{gen}_{\mathrm{RD}}\left([b]^{l}\right) & =\emptyset
\end{aligned}
$$

Similar to the AE analysis: the interesting case is of course the one for assignments. The generation and killing is indeed also quite similar to before. It is the assignment to $x$ which affects the flow, of course. Now, it eliminates all pairs of similar assignments, in the AE-analysis, it invalidates all expressions, which mention $x$. For the generation, the AE has been a bit more complex than the analysis here: here, just the current pair of label and the variable is added (actually, for unique labelling, even the label alone would suffice). For AE , the relevant generated information is not drawn from $x$ in an assignment $x:=a$, but from $a$ (its non-trivial sub-expressions).

Flow equations: $\mathrm{RD}^{=}$
split into

- intra-block equations, using kill and generate
- inter-block equations, using flow


## Flow equations for RD

$$
\begin{aligned}
\mathrm{RD}_{\text {entry }}(l) & = \begin{cases}\left\{(x, ?) \mid x \in f v\left(S_{*}\right)\right\} & l=\operatorname{init}\left(S_{*}\right) \\
\bigcup\left\{\mathrm{RD}_{\text {exit }}\left(l^{\prime}\right) \mid\left(l, l^{\prime}\right) \in \operatorname{flow}\left(S_{*}\right)\right\} & \text { otherwise }\end{cases} \\
\mathrm{RD}_{\text {exit }}(l) & =\mathrm{RD}_{\text {entry }}(l) \backslash \operatorname{kill}_{\mathrm{RD}}\left(B^{l}\right) \cup \operatorname{gen}_{\mathrm{RD}}\left(B^{l}\right)
\end{aligned}
$$

where $B^{l} \in \operatorname{blocks}\left(S_{*}\right)$

- same order of kill/generate


### 2.2.4 Very busy expressions

## Introduction

This is a another example of a classical data flow analysis. As for AE, one is interested in expressions (not assignments). This time it's about if an expression is "needed" in the future. Compared to AE, the perspective has changed. It's not about if an expression that has been evaluated in the past is still available as some given point. It the opposite: will an expression be of use in the future.

This change of perspective also means, that VB is an example of a backward analysis. The natural way of analysing very busy expressions is: at the place where an expression is actually used, immediately in front of that place it's definitely very busy. And then from there, let the information flow backward: in the previous location, it's also very busy (unless relevant variables are change which "destroy" the the "busy-ness"), then the continue the argument.

Being very busy also means an expression is used on all future paths, which makes it a must analysis.

One can make use of very busy information as follow: if an expression is very busy, it may pay off to calculate it already now, i.e., it can be used for a program transformation, that moves the calculation of expression in an "eager" fashion as early as posssible. Transformations like this are known as expression "hoisting".

This is may lead to shorter code of an expression, which is being calculated in two branches of a conditional, for example, can be move earlier outside the branching construct. Note that while that may be reduce the code size but not really the run-time for executing the code.

Transformations like the one mentioned are often done (also) on low level code (like machine-code or low-level intermediate reprentations which are already close to machine code, but still machine-independent). Executing one command ("one line of machine code") costs clock-cycle(s) already, since the command itself needs to be loaded to the processor; on top of that comes costs for loading the operands. So, shortening straight-line code may well improve the execution time. However, hoisting an expression out from both branches of a conditional and position it in front of the branch shorten the size of the code without making it faster.

## Very busy expressions

$$
\begin{array}{ll}
\text { if } & {[a>b]^{1}} \\
\text { then } & {[x:=b-a]^{2} ;[y:=a-b]^{3}} \\
\text { else } & {[a:=b-a]^{4} ;[x:=a-b]^{5}}
\end{array}
$$

Definition 2.2.1 (Very busy expression). An expression is very busy at the exit of a label, if for all paths from that label, the expression is used before any of its variables is "redefined" (= overwritten).

- usage: expression "hoisting"

Goal For each program point, which expressions are very busy at the exit of that point.

Note that the definition and the goal are formulated in a subtle way. It's about information at the exits of the basic blocks, not the entry. In principle, and as far as the equations or constraints are concerned, the formulation will mention $\mathrm{VB}_{\text {entry }}$ and $\mathrm{VB}_{\text {entry }}$ (see later) in the same way as the equations for reaching definitions, for example, mentioned $\mathrm{RD}_{\text {entry }}$ and $\mathrm{RD}_{\text {exit }}$. So it seems to be, one calculates exits and entries.

Nothing wrong with that, but looking carefully to the pseudo-code formulation of the algorithms later, a refinement of the rather sketchy random iteration of the introduction, we will see that what is given back is indeed only the very busy information at the exits of the basic blocks. The reason why it's the exits (and not the entries) is because the very busy expression analysis works backwards, for forward analyses it's the corresponding information at the entries of the blocks.

Why is that? Basically (in the case of the backward analysis), having the solution at the exits allows to reconstruct immediately the solution values at the entries per block (via the kill and generate function attached to the block, something which will be called the transfer function of the block). The pseudocode will indeed work with "arrays" $\mathrm{VB}_{\text {exit }}$ and $\mathrm{VB}_{\text {entry }}$, it's only that what the algo will give back $\mathrm{VB}_{\text {exit }}$, only. One can, however, implement basically the same algorithm leaving out $\mathrm{VB}_{\text {entry }}$, storing only $\mathrm{VB}_{\text {exit }}$ throughout the run.

Anyway, the reason why the goal is formulated like that is as (for backward analysis) the exit information is the crucial one, if one has that, the entry information follows by applying the transfer function (a combination of kill and generate) to the exit communcation, so there is not need to store it seperately in $\mathrm{RD}_{\text {entry }}$ if one wants to do without a second array during the run.

## Very busy expressions: types

- interested in: sets of expressions: $2^{\mathbf{A E x p}_{*}}$
- generation and killing:

$$
\text { kill }_{\mathrm{VB}}, \text { gen }_{\mathrm{VB}}: \text { Blocks }_{*} \rightarrow 2^{\mathbf{A E x p}_{*}}
$$

- analysis: pair of mappings

$$
\mathrm{VB}_{\text {entry }}, \mathrm{VB}_{\text {exit }}: \mathbf{L a b}_{*} \rightarrow 2^{\mathbf{A E x p}_{*}}
$$

## Very busy expr.: kill and generate

core of the intra-block flow specification

$$
\begin{aligned}
\text { kill }_{\mathrm{VB}}\left([x:=a]^{l}\right) & =\left\{a^{\prime} \in \operatorname{AExp}_{*} \mid x \in f v\left(a^{\prime}\right)\right\} \\
\text { kill }_{\mathrm{VB}}\left([\text { skip }]^{l}\right) & =\emptyset \\
\text { kill }_{\mathrm{VB}}\left([b]^{l}\right) & =\emptyset \\
\text { gen }_{\mathrm{VB}}\left([x:=a]^{l}\right) & =\operatorname{AExp}(a) \\
\operatorname{gen}_{\mathrm{VB}}\left([\operatorname{skip}]^{l}\right) & =\emptyset \\
g e n_{\mathrm{VB}}\left([b]^{l}\right) & =\operatorname{AExp}(b)
\end{aligned}
$$

A comparison with the kill and generate functions for AE might be interesting. First of all, in both cases, the functions have the same types, i.e., operate on the same domains. Of course, one difference is, that now the flow is backwards. For the blocks without side effects, this does not matter, i.e., the generate function is identical in both cases (the kill-function as well, of course). For the assignment, there are obviously differences. Let's first look at the kill-case. Literally, the two definitions coincide, but they have a different intuition (backward vs. forward). Here for VB we ask, because we are thinking backwards, which expressions are very busy at the entry of that block. Of course, also the killing works backwards: whatever was very busy at the exits of the block, all expressions that contain $x$ are modified and thus are not very busy at the entry of the block (one could say, as there is no branching withing one block, they are not even busy at all), and thus the kill-function removes those. The reasoning for the AE case is similar, only working forward.

For the generation function, as we are working backwards, the assignment generates $a$ as very busy at the entry of the block. Unlike for AE, the free occurrence of $x$ does not play a role. That's because the order of the applications of first kill and then generate
: $\mathrm{BEAMER}_{\text {env }}$ : againframe : $\mathrm{BEAMER}_{\text {ref }}$ : frame.AE.killgenerate

Flow equations.: $\mathrm{VB}^{=}$
split into

- intra-block equations, using kill/generate
- inter-block equations, using flow
however: everything works backwards now


## Flow equations: VB

$$
\begin{aligned}
\mathrm{VB}_{\text {exit }}(l) & = \begin{cases}\emptyset & l \in \text { final }\left(S_{*}\right) \\
\cap\left\{\mathrm{VB}_{\text {entry }}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in \text { flow }^{R}\left(S_{*}\right)\right\} & \text { otherwise }\end{cases} \\
\mathrm{VB}_{\text {entry }}(l) & =\mathrm{VB}_{\text {exit }}(l) \backslash \operatorname{kill}_{\mathrm{VB}}\left(B^{l}\right) \cup \text { gen }_{\mathrm{VB}}\left(B^{l}\right)
\end{aligned}
$$

where $B^{l} \in \operatorname{blocks}\left(S_{*}\right)$
Note: Doing a backward analysis, the roles of entries and exits are now reversed. The kill and generate functions now calculate the entry as function of the exit point. Analogously, the inter-block flow equations (of the graph) calculate the exit of a block as function of the entries of others.

## Example




Since the very busy expression analysis works backwards, the illustration show the reversed control flow graph.

Besides that: The looping example is quite instructive. It illustrates a subtle point which might not immediately clear from the informal formulation of what "very busy" means. The example is a bit artificial, and the only expression occuring at all is $x+1$ in node $l_{2}$. Now the question is:

Is expression $x+1$ very busy at the beginning of the program or not?
Assuming that $x>0$, there is obviously an infinite loop and the assignment of $l_{2}$ will never be executed. Consequently, the expression will not be needed anyhow. That's of course naive in that standard data flow analysis does not try to figure out if a left-branch or a right branch is taken; in the case of the example, whether the loop body is ignored or not.

On the other hand: it seems that the analysis could make the assumption that there is actually a path on which the $x+1$ is never, ever evaluated. That seems to indicate that one should intuitively consider the expression $x+1$ not very busy. If we don't know how often the loop body is executed (if at all), and since we cannot exclude that the body is taken infinitely (as in this case), it seems plausible to say, there's a chance that $x+1$ may not executed and therefor count it as not very busy.

Plausible as that argument is: it's wrong and $x+1$ is indeed very busy! Informally, the reason being that in a way, "infinite paths don't count" (like the one cycling infinitely many times through the skip-body). Formally, the fact comes from fact that we are interest in the largest safe solution and the way the largest fixpoint it defined (and then they way that the fixpoint iteration, like the chaotic iteration calculates is).

Later, the same example will be used for live variable analysis. Like the one here, it's a backward analysis. Different from very busy expressions, it's a may
analysis (and consequently it's about the smallest possible safe solution). Being a may analysis will make use of $\cup$. Anyway, the variable $x$ will be counted as live and the beginning of the program, as there is a possibility that $x$ is used (in $l_{2}$ ) and that possiblity does not involve making an argument about infinite paths. Unlike the situation of the very busy expressions, this seems intuitively plausible.

### 2.2.5 Live variable analysis

## Introduction

This analysis focuses of variables again (not on expressions). If we use "dead" for being not live, a variable is dead intuitively if its value is definitely ("must") not used in the future. This is very important information, in that the memory bound to the variable can be "deallocated".

That is in particular done at lower levels of the compiler. There, the compiler attempts to generate code which make "optimal" use of avaible registers (except that real optimality is out of reach, so it's more like the compiler typically makes a decent effort in making good use of registers (at least on average). A register currently containing a dead variable can be recycled (to be very precise: the register can be recycled if it contains only dead variables as in some cases, a register can hold the content of more than one variable ...). So a variable is live if there is a potential use of it in the future ("may").

Referring to the future use of variables entails that the question for liveness of variables leads to a backward analysis, similar to the situation of very busy expressions, which was also backwards.

For the participants of the compiler construction lecture (INF5110). That lecture covered live variable analysis, as well, namely in a local variant for elementary blocks of straight-line code. Additionally, a "global" live analysis was sketched, which correspond to the one here.

## When can var's be "recycled": Live variable analysis

$$
\begin{aligned}
& {[x:=2]^{0} ;[y:=4]^{1} ;[x:=1]^{2} ;} \\
& \left(\operatorname{if}[y>x]^{3} \text { then }[z:=y]^{4} \text { else }[z:=y * y]^{5}\right) ;[x:=z]^{6}
\end{aligned}
$$

Live variable A variable is live (at the exit of a label) if there exists a path from the mentioned exit to the use of that variable which does not assign to the variable (i.e., redefines its value)

Goal therefore for each program point: which variables may be live at the exit of that point.

- use: register allocation

Live variables are about: when is a variable still "needed". If not needed, one can free the space. In some sense and very generally, the question resembles reaching definition in a superficial sense, at least, in that it's again about "variables" not expressions. In both cases we like to connect the assignment (also called definition of a variabe to its use). The perspective here is different, though. For RD, the question is: given an assignment, what locations can it reach. For LV it's the opposite: given a location, which assignments can have reached me. This switch in perspective is the difference between forward and backward analysis.

Unlike the informal definition of very busy expressions, here the word is may. With the may-word, the intuition is, that making the solution larger is ok, therefore we are interested in the smallest solution. This is consistent with the fact of making use of live variable analysis for recycling variables. If we estimate too many variables as live, we cannot reuse their memory, which is safe, we only may loose efficiency. Making the opposite approximation, marking an actually live variable erronously as non-live, may lead to errors and is therefore unsafe.

Note again at the goal: "backward" corresponds to "we are interested at the exit".

## Live variables: types

- interested in sets of variables $2^{\mathrm{Var}_{*}}$
- generation and killing:

$$
\text { kill }_{\mathrm{LV}}, \text { gen }_{\mathrm{LV}}: \text { Blockss }_{*} \rightarrow 2^{\mathbf{V a r}_{*}}
$$

- analysis: pair of functions

$$
\mathrm{LV}_{\text {entry }}, \mathrm{LV}_{\text {exit }}: \mathbf{L a b}_{*} \rightarrow 2^{\mathbf{V a r}_{*}}
$$

## Live variables: kill and generate

$$
\begin{aligned}
\operatorname{kill}_{\mathrm{AE}}\left([x:=a]^{l}\right) & =\{x\} \\
\operatorname{kill}_{\mathrm{LV}}\left([\text { skip }]^{l}\right) & =\emptyset \\
{\operatorname{kill} l_{\mathrm{LV}}\left([b]^{l}\right)}^{l} & =\emptyset \\
\operatorname{gen}_{\mathrm{LV}}\left([x:=a]^{l}\right) & =f v(a) \\
\operatorname{gen}_{\mathrm{LV}}\left([\operatorname{skip}]^{l}\right) & =\emptyset \\
\operatorname{gen}_{\mathrm{LV}}\left([b]^{l}\right) & =f v(b)
\end{aligned}
$$

We need to remember that the calculation is backwards. As for kill: in the only interesting case of assignment, the question is: given the live variables at the end, which ones are live at the entry. Certainly, $x$ is no longer live, as it is not used (forward) before overwritten.

That also explains the generation: all free variables in $a$, resp. in $b$ are live at the beginning of a block that mentions the resp. expression. In particular, the $x$ does not play a role in the generation function for assignments, as we are working backwards.

## Flow equations LV $=$

split into

- important\{intra\}-block equations, using kill/generate
- inter-block equations, using flow
however: everything works backwards now


## Flow equations LV

$$
\begin{aligned}
\mathrm{LV}_{\text {exit }}(l) & = \begin{cases}\emptyset & l \in \text { final }\left(S_{*}\right) \\
\bigcup\left\{\mathrm{LV}_{\text {entry }}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in \operatorname{flow}^{R}\left(S_{*}\right)\right\} & \text { otherwise }\end{cases} \\
\mathrm{LV}_{\text {entry }}(l) & =\mathrm{LV}_{\text {exit }}(l) \backslash \operatorname{kill}_{\mathrm{LV}}\left(B^{l}\right) \cup \operatorname{gen}_{\mathrm{LV}}\left(B^{l}\right)
\end{aligned}
$$

where $B^{l} \in \operatorname{blocks}\left(S_{*}\right)$
The example, why one is this time interested in the smallest solution is the same program as for the VB: a simple recursive equation (induced by a trivial while-loop). This time the loop contains a $\cup$. We can make the solution as large as possible (but not as small as possible, the empty set is not a solution). However, the smallest set is the most informative one. That can be guessed from the words "may be live" already. Also the intended use of freeing/re-using "non-live" variables makes clear that it's "larger is less precise".

## Example

$$
\text { (while } \left.[x>1]^{l_{0}} \text { do }[\text { skip }]^{l_{1}}\right) ;[x:=x+1]^{l_{2}}
$$

As one can see in the flow equations, especially the case dealing with the final nodes, variables are considered dead at the end. One may also have the intuition, the variables (or some) are returned to somewhere, in which case they are still needed "after" the final node (for being returned) and hence they
are marked live. In the latter initution, it would probably be clearer to have an explicit return statement (after which the variables are really dead).

For the participants of the compiler construction course (INF5110). The course presented a local live variable analysis, which concentrated on straight line code. The code was so-called three-address code and had two types of variables: normal ones and so-called "temporaries" (temporary variables). The standard variables were assumed live at the end of the straight-line code. The reason being that the SLC is code contained inside one basic block. Since the end if the block is not necessarily the end of the program, the analysis had to assume conservatively that chances are, that variables may by used by any potentially following block, and thus they variables were assumed live. Temporaries, on the other hand, were treated as dead at the end, which was justfied by the fact that the code never used temporaries from a previous block. That, of course, depended on the knowledge how this code was actually generated. The general point is that of course the formulation of the live variable analysis (or others) must go hand in hand with what is actually going on, i.e., the semantics of the language and the assumptions about how the program is used ("will the content of the variables be returned to a caller after the program code or not", "might there by a block after the code being analyzed or not, and if so, will it make use of temporaries resp. variables or not").

## Looping example



### 2.3 Theoretical properties and semantics

### 2.3.1 Semantics

## Introduction

So far we have formulated a number of analyses (using flow equations or constraints). We also stressed the importance that the analyses are safe (or correct or sound), meaning that the information given back from the analysis says something "true" about the program, more precisely about the program's behavior. So far, that is an empty claim as we have not fixed what the behavior actually is. Doing so may look superfluous, in particular as the while language we are currently dealing with is so simple that its semantics seems pretty "obvious" for most. That, however, may no longer be the case when dealing with more advanced or novel features or non-standard syntax etc. Being clear about what the semantics is supposed to be also pays off when implementing a language, after all, the ultimately running program is expecteed to implement exactly the specified semantics, down to the actual machine code on a particular platform. Leaving the semantics up-to the implementation or the platform is not considered a very dignified engineering approach ("the semantics of a program is what happens if you run it, you'll see.").

In this section we will precisely define the semantics of the while-language. The semantics is defined on a rather high-level, on the level of the abstract syntax, and the task of the compiler would be to preserve exactly the syntax through all its phases. The task of the static analyses is to soundly approximate this semantics. If optimizations and transformations are done, for example based on some static analyses, it's the task of the optmization to preserve the semantics, as well. So the semantics is the yardstick which all further actions of the compiler are measured against.

The semantics of a programming language can be specified in different styles or flavors. We make use of operational semantics, a style of semantics described the steps a program does. In particular, we make use of strucutural operational semantics (SOS), which refers to the fact that the steps are described making inductive use of the structure of the program (i.e., it's abstract syntax).

That's arguably a straightforward way for fixing the semantics. It basically descibes the semantics as steps tranforming an abstract syntax tree step by step and can be seen as an formal description of an interpreter.

There is, however, also not one unique way how such an operational semantics is defined, even there different flavors and styles exists. Perhaps later, for the more complex functional languages, the lecture covers some variations.

## Relating programs with analyses

- analyses
- intended as (static) abstraction or overapprox. of real program behavior
- so far: without real connection to programs
- soundness of the analysis: safe analysis
- but: behavior or semantics of programs not yet defined
- here: "easiest" semantics: operational
- more precisely: small-step SOS (structural operational semantics)


## States, configs, and transitions

fixing some data types

- state $\sigma:$ State $=\mathbf{V a r} \rightarrow \mathbf{Z}$
- configuration: pair of statement $\times$ state or (terminal) just a state


## Transitions

$$
\langle S, \sigma\rangle \rightarrow \dot{\sigma} \quad \text { or } \quad\langle S, \sigma\rangle \rightarrow\langle\dot{S}, \dot{\sigma}\rangle
$$

## Semantics of expressions

$$
\begin{aligned}
& {[]^{\mathcal{A}}: \mathbf{A E x p} \rightarrow(\text { State } \rightarrow \mathbf{Z})} \\
& {[-]_{-}^{\mathcal{B}}: \mathbf{B E x p} \rightarrow(\text { State } \rightarrow \mathbf{B})}
\end{aligned}
$$

simplifying assumption: no errors

$$
\begin{aligned}
{[x]_{\sigma}^{\mathcal{A}} } & =\sigma(x) \\
{[n]_{\sigma}^{\mathcal{A}} } & =\mathcal{N}(n) \\
{\left[a_{1} \mathrm{op}_{a} a_{2}\right]_{\sigma}^{\mathcal{A}} } & =\left[a_{1}\right]_{\sigma}^{\mathcal{A}} \mathbf{o p}_{a}\left[a_{2}\right]_{\sigma}^{\mathcal{A}} \\
{[\operatorname{not} b]_{\sigma}^{\mathcal{B}} } & =\neg[b]_{\sigma}^{\mathcal{B}} \\
{\left[b_{1} \mathrm{op}_{b} b_{2}\right]_{\sigma}^{\mathcal{B}} } & =\left[b_{1}\right]_{\sigma}^{\mathcal{B}} \mathbf{o p}_{b}\left[b_{2}\right]_{\sigma}^{\mathcal{B}} \\
{\left[a_{1} \mathrm{op}_{r} a_{2}\right]_{\sigma}^{\mathcal{B}} } & =\left[a_{1}\right]_{\sigma}^{\mathcal{A}} \mathbf{o p}_{r}\left[a_{2}\right]_{\sigma}^{\mathcal{A}}
\end{aligned}
$$

clearly:

$$
\forall x \in f v(a) \cdot \sigma_{1}(x)=\sigma_{2}(x) \text { then }[a]_{\sigma_{1}}^{\mathcal{A}}=[a]_{\sigma_{2}}^{\mathcal{A}}
$$

In the intro, we mentioned that we will do some specific form of semantics, namely an operational semantics. That's not $100 \%$ true. For dealing with the control-flow structure of the while language, we will indeed formulate operational rules to describe the transitions. We must, however, also give meaning
to expressions $a$ and $b$. To do that operationally would be possible, but perhaps an overkill. More straightforward is a inductive definition in the way given. That style corresponds more to a denotational semantics. It should be noted that expressions in the while language are side-effect free. So things like $\mathrm{x}:=5$ * $\mathrm{y}^{++}$which can be found in for example C-like languages, where the right-hand side of the assignment is at the same time an expression as well as having a side effect, are not welcome here. Without such side effects, the denotation-style semantics for expressions is just the easiest way of specifying their meaning. That way we can focus on the part that is more interesting for us, the steps or transitions of the operational semantics.

It would be possible to also specify it meaning of expressions in an operational, step-wise manner. In this way there would me more transitions explicitly mentioned in the semantics (maybe distinguising the transitions between configurations and "micro-transitions" evaluating the expressions).

For participants of the compiler construction course (INF5110): an operational semantics for the expressions showed up in some way in the lecture, when translating expression into three address code. Since there is no recursion and deeply nested expressions in three-address code (which is used in the definition of $[a]_{\sigma}^{\mathcal{A}}$ ), the expression has to be "expanded" into sequences of non-nested expression together with temporary variables ("temporaries") to hold intermediate results of subexpressions. That in a way corresponds to an explicit, step-by-step execution of a compound expression. It's not the same as an operational semantics, as it does not specify "transitions", but it's "code generation", but each single three-address-code instruction would correspond to one transition.

## SOS

$$
\begin{aligned}
& \left\langle[x:=a]^{l}, \sigma\right\rangle \rightarrow \sigma\left[x \mapsto[a]_{\sigma}^{\mathcal{A}}\right] \quad \text { ASS } \quad\left\langle[\text { skip }]^{l}, \sigma\right\rangle \rightarrow \sigma \quad \text { SKIP } \\
& \frac{\left\langle S_{1}, \sigma\right\rangle \rightarrow\left\langle\dot{S}_{1}, \sigma\right.}{\left\langle S_{1} ; S_{2}, \sigma\right\rangle \rightarrow\left\langle\dot{S}_{1} ; S_{2}, \dot{\sigma}\right\rangle} \mathrm{SEQ}_{1} \quad \frac{\left\langle S_{1}, \sigma\right\rangle \rightarrow \dot{\sigma}}{\left\langle S_{1} ; S_{2}, \sigma\right\rangle \rightarrow\left\langle S_{2}, \dot{\sigma}\right\rangle} \mathrm{SEQ}_{2} \\
& \frac{[b]_{\sigma}^{\mathcal{B}}=\mathrm{T}}{\left\langle\mathrm{if}[b]^{l} \text { then } S_{1} \text { else } S_{2}, \sigma\right\rangle \rightarrow\left\langle S_{1}, \sigma\right\rangle} \mathrm{IF}_{1} \\
& \frac{[b]_{\sigma}^{\mathcal{B}}=\mathrm{T}}{\left\langle\text { while }[b]^{l} \text { do } S, \sigma\right\rangle \rightarrow\left\langle S ; \text { while }[b]^{l} \text { do } S, \sigma\right\rangle} \text { WHILE }_{1} \\
& \frac{[b]_{\sigma}^{\mathcal{B}}=\perp}{\left\langle\text { while }[b]^{l} \text { do } S, \sigma\right\rangle \rightarrow \sigma} \text { WHiLE }_{2}
\end{aligned}
$$

## Derivation sequences

- derivation sequence: "completed" execution:
- finite sequence: $\left\langle S_{1}, \sigma_{1}\right\rangle, \ldots,\left\langle S_{n}, \sigma_{n}\right\rangle, \sigma_{n+1}$
- infinite sequence: $\left\langle S_{1}, \sigma_{1}\right\rangle, \ldots,\left\langle S_{i}, \sigma_{i}\right\rangle, \ldots$
- note: labels do not influence the semantics
- CFG for the "rest" of the program only gets "smaller" when running:


## Lemma 2.3.1.

1. $\langle S, \sigma\rangle \rightarrow \sigma^{\prime}$, then final $(S)=\{\operatorname{init}(S)\}$
2. Assume $\langle S, \sigma\rangle \rightarrow\langle\dot{S}, \dot{\sigma}\rangle$, then
a) $\operatorname{final}(S) \supseteq\{\operatorname{final}(S)\}$
b) flow $(S) \supseteq\left\{\right.$ flow $\left.\left(\begin{array}{c}S\end{array}\right)\right\}$,
c) blocks $(S) \supseteq$ blocks $(S)$; if $S$ is label consistent, then so is $\dot{S}$

## Correctness of live analysis

- LV as example
- given as constraint system (not as equational system)


## LV constraint system

$$
\begin{aligned}
& \mathrm{LV}_{\text {exit }}(l) \supseteq \begin{cases}\emptyset & l \in \text { final }\left(S_{*}\right) \\
\bigcup\left\{\mathrm{LV}_{\text {entry }}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in \text { flow }^{R}\left(S_{*}\right)\right\} & \text { otherwise }\end{cases} \\
& \mathrm{LV}_{\text {entry }}(l) \supseteq \quad \mathrm{LV}_{\text {exit }}(l) \backslash \text { kill }_{\mathrm{LV}}\left(B^{l}\right) \cup \operatorname{gen}_{\mathrm{LV}}\left(B^{l}\right) \\
& \operatorname{live}_{\text {entry }^{l}, \text { live }_{\text {exit }}: \mathbf{L a b}_{*} \rightarrow 2^{\text {Var }_{*}}}
\end{aligned}
$$

"live solves constraint system $\mathrm{LV} \subseteq(S)$ "

$$
\text { live } \models \mathrm{LV} \subseteq(S)
$$

(analogously for equations $\mathbf{L V}=(S)$ )

## Equational vs. constraint analysis

Lemma 2.3.2. 1. If live $\models \mathrm{LV}^{=}$, then live $\models \mathrm{LV} \subseteq$
2. The least solutions of live $\models \mathrm{LV}=$ and live $\models \mathrm{LV} \subseteq$ coincide.

### 2.3.2 Intermezzo: Lattices

## Intermezzo: orders, lattices. etc.

as a reminder:

- partial order $(L, \sqsubseteq)$
- upper bound $l$ of $Y \subseteq L$ :
- least upper bound (lub): $\bigsqcup Y$ (or join)
- dually: lower bounds and greatest lower bounds: $\Pi Y$ (or meet
- complete lattice $L=(L, \sqsubseteq)=(L, \sqsubseteq, \sqcap, \sqcup, \perp, \top)$ : a partially ordered set where meets and joins exist for all subsets, furthermore $\top=\Pi \emptyset$ and $\perp=\bigsqcup \emptyset$.

Here we are working with a specific form of lattice, called complete lattice. It's a very nicely behaved lattice, which makes it useful for the monotone framework. The important condition, which makes the lattice actually complete is rather hidden in the above definition. It's that meets and joins exists for all subsets. If we drop the "completeness", then one would still requires joins or "least upper bounds" $a \sqcup b$ and or "greatest lower bounds" $a \sqcap b$ to exists. The difference here is we speak about binary such operations. To be complete, it's also required the meets and joins exists also for infinite sets. A priori a lattice is not required to be finite (and many interesting ones are not). For those situations, the existance of binary meets and joins implies the existance of finite meets and joins (those can always be expressed by a number of binary ones). What is not guaranteed it the existance of arbitrary meets and joins, including infinite ones. But that's required for a lattice to be complete.

There are also other forms of lattices, for instance, if one only needs joints, but not meets, one can get away with a semi-lattice, and there are many more variations. For the lecture, we generally simply assume complete lattices and the montone framework is happy. In particular, if we are dealing with finite lattices, which is an important case, we don't need to consider infinite sets, and "standard" lattices with binary meets and joins (and least and largest elements) are complete already.

## Fixpoints

given complete lattice $L$ and monotone $f: L \rightarrow L$.

- fixpoint: $f(l)=l$

$$
\operatorname{Fix}(f)=\{l \mid f(l)=l\}
$$

- $f$ reductive at $l, l$ is a pre-fixpoint of $f: f(l) \sqsubseteq l$ :

$$
\operatorname{Red}(f)=\{l \mid f(l) \sqsubseteq l\}
$$

- $f$ extensive at $l, l$ is a post-fixpoint of $f: f(l) \sqsupseteq l$ :

$$
\operatorname{Ext}(f)=\{l \mid f(l) \sqsupseteq l\}
$$

## Define "Ifp" / "gfp"

$$
l f p(f) \triangleq \bigcap \operatorname{Fix}(f) \quad \text { and } \quad g f p(f) \triangleq \bigsqcup \operatorname{Fix}(f)
$$

The last display just gives the names to the two elements of the lattice defined by the corresponding right-hand sides. We know tha those elements are existing thanks to the fact that $L$ is a complete lattice (and it's very easy to see that meets and joins are unique, that means the $l f p(f)$ and $g f p f$ are welldefined elements of the lattice. The chosen names somehow suggest that the two thusly defined elements are the least fixpoint, resp. the greatest fixpoint of the monotone function $f$.

But, so far $l f p$ and $g f p$ is just a suggestive choice of name. It requires an separate argument that the elements are actually fixpoints, and the least, resp. the largest fixpoint as that as well. Finally, if we take it really serious, an argument should be found that allows to speaking of the least fixpoint. If there is more than one least fixpoint, one should avoid talking about "the least fixpoint" (same for the largest fixpoint). The argument for uniquess of least fixpoints (or for greatest fixpoint) is very simple though, similar to arguing for the uniqueness of "the least upper bound" etc.

If one would carry out the argument, i.e., the proof, that all fits together in the sense that the $l f p(f)$ and $g f p(f)$ defined above are actually the least fixpoint and the largest fixpoint, and if one would carefully keep track of what is actually needed to make the proof go through step by step, then one would see that every single condition for being a complete lattice is needed (plus the fact that $f$ is monotone). If one removes one condition, the argument fails! Conversely that means the following: We are interested in uniquely "best approximations" (least or greatest fixpoints depending in whether it's a may or a must analysis),
and, having a monotone $f$, a complete lattices is exactly what guarantees that those fixpoints exists. Exactly that, nothing less and nothing more, If your framework has monotone functions and is based on a complete lattice, it works. If not, it does not work, very simple.

That explains the importance of lattices and monotone function. Also, I would guess that historically, the need to assure existance of fixpoints has led Tarski (the mathematician whose concepts we are currently covering) exactly to the definition of lattice, not the other way around ("oh, someone defined some lattice, let's see what I can find out about them, perhaps I could define some $l f p(f)$ like above and see if I could prove something iteresting about it, perhaps it's a fixpoint?". But as said, that is speculation.

Having stressed the importance of complete lattices, for fairness sake it should be said that there's also a place for analyses which fail to meet those conditions. In that case, one might not have a (unique) best solution. Perhaps even worse (and related to that), one might need combinatorial techniques (like backtracking), i.e., checking all possible solutions to find an acceptable one. If that happens, the cost of the analysis may explode. To avoid that one may give up to look for a "best solution" and settle for a "good enough" one and heuristics that hopefully find an acceptable one efficiently, or even throw the towel and give up "soundness". Anyway and fortunately, plenty of important analyses fit well into the monotone framework with its lattices, its unique best solution and -perhaps best of all- its efficient solving techniques. Therefore this lecture will cover only those here. Those are called classical data flow analyses.

## Tarski's theorem

Core Perhaps core insight of the whole lattice/fixpoint business: not only does the $\rceil$ of all pre-fixpoints uniquely exist (that's what the lattice is for), but -and that's the trick - it's a pre-fixpoint itself (ultimately due to montonicity of $f$ ).

Theorem 2.3.3. L: complete lattice, $f: L \rightarrow L$ monotone.

$$
\begin{align*}
\operatorname{lfp}(f) & \triangleq \sqcap \operatorname{Red}(f)  \tag{2.6}\\
g f p(f) & \triangleq \quad \operatorname{Fix}(f) \\
g \operatorname{Ext}(f) & \in \quad \operatorname{Fix}(f)
\end{align*}
$$

- Note: lfp (despite the name) is defined as glb of all pre-fixpoints
- The theorem (more or less directly) implies lfp is the least fixpoint


## Fixpoint iteration

- often: iterate, approximate least fixed point from below $\left(f^{n}(\perp)\right)_{n}$ :

$$
\perp \sqsubseteq f(\perp) \sqsubseteq f^{2}(\perp) \sqsubseteq \ldots
$$

- not assured that we "reach" the fixpoint ("within" $\omega$ )

$$
\perp \sqsubseteq f^{n}(\perp) \sqsubseteq \sqcup_{n} f^{n}(\perp) \sqsubseteq \underset{g f p(f)}{l f(f)} \sqsubseteq \prod_{n} f^{n}(\top) \sqsubseteq f^{n}(\top) \sqsubseteq(\top)
$$

- additional requirement: continuity on $f$ for all ascending chains $\left(l_{n}\right)_{n}$

$$
f\left(\bigsqcup_{n}\left(l_{n}\right)\right)=\bigsqcup\left(f\left(l_{n}\right)\right)
$$

- ascending chain condition ("stabilization"): $f^{n}(\perp)=f^{n+1}(\perp)$, i.e., $l f p(f)=$ $f^{n}(\perp)$
- descending chain condition: dually


## Basic preservation results

Lemma 2.3.4 ("Smaller" graph $\rightarrow$ less constraints). Assume live $\models \mathrm{LV} \subseteq\left(S_{1}\right)$. If flow $\left(S_{1}\right) \supseteq$ flow $\left(S_{2}\right)$ and blocks $\left(S_{1}\right) \supseteq \operatorname{blocks}\left(S_{2}\right)$, then live $\models \mathrm{LV} \subseteq\left(S_{2}\right)$.

Corollary 2.3.5 ("subject reduction"). If live $\models \mathrm{LV} \subseteq(S)$ and $\langle S, \sigma\rangle \rightarrow\langle\dot{S}, \sigma \bar{\sigma}\rangle$, then live $\models \mathrm{LV} \subseteq(S)$

Lemma 2.3.6 (Flow). Assume live $\models \mathrm{LV} \subseteq(S)$. If $l \rightarrow_{\text {flow }} l^{\prime}$, then live $_{\text {exit }}(l) \supseteq$ live $_{\text {entry }}\left(l^{\prime}\right)$.

The three mentioned results are actually pretty straightforward, resp. express properties of the live variable analysis which should be (after some reflection) pretty obvious. Analgous results would hold for other data flow analysis. Lemma 2.3.4 compares the analyses results for two programs $S_{1}$ and $S_{2}$, where $S_{2}$ has a "smaller" control-flow graph (less edges and/or less blocks). Since the control flow graph directly corresponds to sets of constraints, removing parts of the graph means removing constraints. That means, more solutions are possible, which is expressed by the lemma (live $\models \mathrm{LV} \subseteq(S)$ means that live (an assignment of liveness information to all variables of the constraint system) satisfies the constraint system of the program $S$.

It's probably obvious: the variables of the type system are (of course) not the program variables of the live variable analysis. The constraint variables are the (entry and exit points of the) nodes of the graph (which in turn correspond to the labels in the labelled abstract syntax).

The Corollary 2.3.5 is a direct consequence of that. In general, that's what the term "corollary" means: an immediate interesting follow-up of a preceding lemma or theorem etc.

However, the result is not without subtelty. It has to do with the step $\langle S, \sigma\rangle \rightarrow$ $\langle\dot{S}, \sigma\rangle$, resp, what this step does to the (labelled) program $S$. The interesting case for that is step covered by one of the rules dealing with the while-loop, namely While ${ }_{1}$. It's interesting insofar as that it duplicates the body of the
loop. That leads to a program what is no longer uniquely labelled (even if $S$ had been)! It's however still label consistent.

The last lemma is a direct consequence of the construction (backward may analysis).

These lemmas as such are not interesting in themselves.

## Correctness relation

- basic intuitition: only live variables influence the program
- proof by induction
$\Rightarrow$

Correctness relation on states: Given $V=$ set of variables:

$$
\begin{equation*}
\sigma_{1} \sim_{V} \sigma_{2} \text { iff } \forall x \in V . \sigma_{1}(x)=\sigma_{2}(x) \tag{2.7}
\end{equation*}
$$



Notation: $N(l)=$ live $_{\text {entry }}(l), X(l)=$ live $_{\text {exit }}(l)$
In the definition of $\sim_{V}$ above, $V$ is an arbitrary set of "variables". The intention (in the overall argument) will be, that the $V$ 's are those variable that are live (resp. variables that the analysis has marked as live). Of course, the set of variables being determined as live changes during execution.

In the figure above, the "control-part" of the component, i.e., the code $S, S^{\prime}$ etc., are identical step by step for both versions. Both program execute the very same steps.

As a side remark; while language is deterministic, meaning a program code $S$ and a state $\sigma$ determines the successor configuration (if we are not yet at the final configuration). Note also: the intra-block (and backward) definition of liveness directly gives that for an assignment $x:=a$, the free variables in $a$ are live right in front of the assignment. Likewise, variables in a boolean condition $b$ are live right in front of a conditinal or loop, to which $b$ belongs. Those variables therefore are contained in the $\$ \mathrm{~V} \$$-set directly before a step for the two variants of the system. Consequently, both system do exactly the same next step. And then the next step is the same again, and then the next
.... I.e., by induction both systems behave the same, which is exactly what we want to establish ("dead variables don't matter").

## Correctness (1)

Lemma 2.3.7 (Preservation inter-block flow). Assume live $\models \mathrm{LV} \subseteq$. If $\sigma_{1} \sim_{X(l)}$ $\sigma_{2}$ and $l \rightarrow_{\text {fow }} l^{\prime}$, then $\sigma_{1} \sim_{N\left(l^{\prime}\right)} \sigma_{2}$.

## Correctness

Theorem 2.3.8 (Correctness). Assume live $\models \mathrm{LV}^{\subseteq}(S)$.

- If $\left\langle S, \sigma_{1}\right\rangle \rightarrow\left\langle\dot{S}, \dot{\sigma}_{1}\right\rangle$ and $\sigma_{1} \sim_{N(i n i t(S))} \sigma_{2}$, then there exists $\sigma_{2}$ s.t. $\left\langle S, \sigma_{2}\right\rangle \rightarrow$ $\left\langle\dot{S}, \sigma_{2}\right\rangle$ and $\hat{\sigma}_{1} \sim_{N(i n i t(S))} \hat{\sigma}_{2}$.
- If $\left\langle S, \sigma_{1}\right\rangle \rightarrow \sigma_{1}$ and $\sigma_{1} \sim_{N(i n i t(S))} \sigma_{2}$, then there exists $\sigma_{2}$ s.t. $\left\langle S, \sigma_{2}\right\rangle \rightarrow \sigma_{2}$ and $\sigma_{1} \sim_{X(i n i t(S))} \sigma_{2}$.

$$
\begin{aligned}
& \left\langle S, \sigma_{1}\right\rangle \longrightarrow\left\langle\dot{S}, \dot{\sigma}_{1}\right\rangle \quad\left\langle S, \sigma_{1}\right\rangle \longrightarrow \dot{\sigma}_{1} \\
& \begin{array}{|l|l|l}
\sim_{N(i n i t(S))} & \sim_{N(i n i t(S))} & \mid \sim_{N(i n i t(S))} \quad \vdots \\
\sim_{X(i n i t(S))}
\end{array} \\
& \left.\left\langle S, \sigma_{2}\right\rangle \cdots \cdots \cdots\right\rangle\left\langle\dot{S}, \sigma_{2}\right\rangle \quad\left\langle S, \sigma_{2}\right\rangle \cdots \cdots \cdots \dot{\sigma}_{2}
\end{aligned}
$$

The picture are drawn in a "specific" manner to capture the formulation of the theorem. In particular see the use of "solid" arrows and lines vs. "dotted" ones. That a diagrammatic way to indicate the "for all such ..." (solid) and "... there exists some ..." (dotted). This notation is rather standard, and allows to express such properties in a short diagrammatic but still precise manner.

## Correctness (many steps)

Assume live $\models \mathrm{LV}^{\subseteq}(S)$

- If $\left\langle S, \sigma_{1}\right\rangle \rightarrow^{*}\left\langle\dot{S}, \hat{\sigma}_{1}\right\rangle$ and $\sigma_{1} \sim_{N(\text { init }(S))} \sigma_{2}$, then there exists $\sigma_{2}$ s.t. $\left\langle S, \sigma_{2}\right\rangle \rightarrow^{*}$ $\left\langle\dot{S}, \sigma_{2}\right\rangle$ and $\dot{\sigma}_{1} \sim_{N(i n i t(S))} \dot{\sigma}_{2}$.
- If $\left\langle S, \sigma_{1}\right\rangle \rightarrow^{*} \sigma_{1}$ and $\sigma_{1} \sim_{N(i n i t(S))} \sigma_{2}$, then there exists $\dot{\sigma}_{2}$ s.t. $\left\langle S, \sigma_{2}\right\rangle \rightarrow^{*}$ $\sigma_{2}$ and $\sigma_{1} \sim_{X(l)} \sigma_{2}$ for some $l \in \operatorname{final}(S)$.


### 2.4 Monotone frameworks

We have seen 4 different classical analyses, which all shared some similarities. In this section, those analyses will be systematically put into a larger context, which is known as monotone framework. As unifying principle, this was first formulated by Kildall [5] and constitutes in a way the common orthodox and completely standardized understanding of what classical data flow analysis is.

Besides that it capture many known analyses, it's also a "recipe" for designing other data flow analyses, starting from the program given in the form of a control flow graph. Indeed, the 4 analyses we have seen are only (important) representatives of the 4 classes of analyses that can be formulated as monotone framework. The analysis can be forward or backward, and it can be "may" or "must". That's about it, and that gives 4 different classes.

Besides that, the monotone framework concept lays down exactly what needs to be assumed about the structure of the information that is given back from the analysis. All four analyses somehow dealt with sets, like "sets of variables such that this and that" or "sets of expressions such that this or that". Dealing with sets reflected the fact that, being static, the analysis does not exactly knows what the program does and has to approximate. Dealing with sets of pieces of flow information also allows to enlarge or shrink the information via taking the subset or the superset. Which direction is safe in a given analysis depends on whether on whether it's a "may" or a "must" analysis. At any rate, sets and the subset relations is a special case of the notion of the more general notion of lattice, which is exactly the notion needed to make the monotone framework work.

Even if the monotone framework is based on the general notion of lattice for good reason, the special case of sets und subsets is an important one. Not only is it conceptually simple, it also allows efficient implementations. Finite sets over a given domain may be implemented as bit vectors and union and intersection, two crucial operations for "may" resp. "must" analyses can efficiently be implement via logical bitwise "or" resp. "and" on bitvectors.

## Monotone framework: general pattern

$$
\begin{aligned}
& \text { Analysis }_{\circ}(l)= \begin{cases}\iota & \text { if } l \in E \\
\left.\bigsqcup_{\text {Analysis }}^{\bullet}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in F\right\} & \text { otherwise }\end{cases} \\
& {\text { Analysis } \bullet_{\bullet}(l)}^{=} f_{l}\left(\text { Analysis }_{\circ}(l)\right)
\end{aligned}
$$

- $\bigsqcup$ : either $\cup$ or $\cap$
- F: either flow $\left(S_{*}\right)$ or $\operatorname{flow}^{R}\left(S_{*}\right)$.
- $E$ : either $\left\{\operatorname{init}\left(S_{*}\right)\right\}$ or $\operatorname{final}\left(S_{*}\right)$
- $\iota$ : either the initial or final information
- $f_{l}$ : transfer function for $[B]^{l} \in \operatorname{blocks}\left(S_{*}\right)$.

The definition is "generic" as it leaves open the alternatives "may" vs. "must" as well as "forward" vs. "backard". Also the domain flow information of interest is not fixed, neither is the special case of what the information at initial node resp. the final node is supposed to be. As we have discussed especially in connection with live variable analysis, this has to be decided an a case-by-case consideration, depending one specific conditions of the intended analysis and the language. One final ingredient is the transfer function. We have encountered that implicitly for the intra-block data flow. It's only that we did not explicitly called it transfer function, instead the concept was formulated making use of kill and generate function. It turns out that many transfer functions can be formulated as we did via kill and transfer function (as many analysis domain are sets of information of interest), but not all. The general monotone framework simply requires a function that transform flow information on one end of a basic block to flow information at the other end. For a forward analysis, the flow information at the exit of a basic block is expressed as a function on the entry of the block, and for backward information, it's the other way around.

## Monotone frameworks

## direction of flow:

- forward analysis:
$-F=\operatorname{flow}\left(S_{*}\right)$
- Analysis。 for entry and Analysis. for exits
- assumption: isolated entries
- backward analysis: dually
$-F=$ flow $^{R}\left(S_{*}\right)$
- Analysis。 for exit and Analysis. for entry
- assumption: isolated exits


## sort of solution

- may analysis
- properties for some path
- smallest solution
- must analysis
- properties of /all paths
- greatest solution

Into which of the four categories a concrete analysis falls need to be thought through on a case-by-basis of course. However, it may not be a clean cut as it seems, resp. it may also be a matter of perspective. For example, live variable analysis. That one is a may (and a backward) analysis. We can switch perspective from concentrating on live variables to "dead" variables (those wich are not live), still with the same purpose of recylcing memory of variables with are not live $=$ dead. If the data flow analysis streams sets of deads variables throught its equations instead of live variables, the analysis will be a must analysis instead. After all, a variable is dead if it's not used in the future on all paths (which is the dual of being live, which refers to usage on some path). Consequently, one would be interested in the largest safe solution of dead variables.

In a way, both are the "same" analysis, or rather, dual to each other but ultimately equivalent. It's only that convetionally, it's referred to as "live variable analysis" and not as the more morbid dual one.

This switching to the dual perspective is easily possible if we are dealing with finite domains in the analysis, as we often do. Like in live variable analysis, there are only finitely many sets of variables.

$$
\begin{align*}
\text { Analysis }_{\circ}(l)= & \iota_{E}^{l} \sqcup \sqcup\left\{\text { Analysis }_{\bullet}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in F\right\}  \tag{2.9}\\
& \text { where } \iota_{E}^{l}=\left\{\begin{array}{cc}
\iota & \text { if } l \in E \\
\perp & \text { if } l \notin E
\end{array}\right. \\
\text { Analysis }_{\bullet}(l)= & f_{l}\left(\text { Analysis }_{\circ}(l)\right)
\end{align*}
$$

where $l \sqcup \perp=l$

## Explanation

Let's compare it to equation (2.8), where we did it for isolated entries: First remember that $E$ ("extremal") is an initial block (or a final one). Remember also that isolated entries does not mean that the is only one intial/final block, only that there is no loop-back. Let's consider the forward case. In this case, equation (2.8) makes sense. The distinguishing case is the one for inter-block flow, which says something for the entry of a block (in the forward case). And there are exactly two separate cases: if $l$ refers to the initial block, then there is only the initial information $\iota$. Note that this does not mean that there is only one initial label. Otherwise, if it is a non-initial label, then the intial flow is not mentioned in the equation/constraint. Note that it seems possible, that a non-initial block has no inflowing arc. Probably for the while-language that is not the case, however, the definitions seem to allow it. In this case the $\searrow$ as $\cup$ gives the empty set, which probably makes sense. Obviously, (2.8) makes no sense without isolated entries, because for initial labels, the equations overlooks the flowing-back information. The new equation (2.9) repairs that in
that it adds to the initial labels also the combined information from the posts of the connected nodes. However, it must of course not inject $\iota$ into non-initial nodes, hence the definition of $\iota_{E}^{l}$.\}

## Basic definitions: property space

- property space $L$, often complete lattice
- combination operator: $\sqcup: 2^{L} \rightarrow L$, $\sqcup$ : binary case
- $\perp=\bigsqcup \emptyset$
- often: ascending chain condition (stabilization)

The property space (here called $L$ ) captures the "information of interest" (sets of variables ...). Technically, it needs to be some form of lattice (see the corresponding section later). In good approximation, the lattices and its laws resemble closely the situation with sets of information (with $\cup, \cap, \subseteq, \supseteq, \emptyset$ $\ldots$...). Indeed, the power set of some set is a special case of a lattice (and an important one in the context of the lecture).

## Transfer functions

$$
f_{l}: L \rightarrow L
$$

with $l \in \mathbf{L a b}_{*}$

- associated with the blocks
- requirement: monotone
- $\mathcal{F}$ : monotone functions over $L$ :
- containing all transfer functions
- containing identity
- closed under composition

The transfer functions, as defined above, are attached to the elementary blocks (which here contain one single statement or expression, but in general may contain staight-line code). In other accounts, the control flow graphs and/or the transfer functions may be differently represented, without changing anything relevant. For instance, here, the nodes of the CFG contain assigmments and expression (i.e., relevant pieces of abstract syntax). Also the transfer functions are attached to the nodes. One can see it like the transfer function is the semantics of the corresponding piece of syntax. Not the "real" semantics, but on the chosen abstraction level of the analysis, i.e., on the level of the property space $L$.

Some authors prefer to attach the pieces of syntax and/or the transfer functions to edges of a control-flow graph (which therefore is of a slighty different format than the one we operate with). But it's only a different reprentation of the same principles.

## Summary

- complete lattice $L$, ascending chain condition
- $\mathcal{F}$ monotone functions, closed as stated
- distributive framework

$$
f\left(l_{1} \sqcup l_{2}\right)=f\left(l_{1}\right) \sqcup f\left(l_{2}\right)
$$

Instead of the above condition, one might require

$$
f\left(l_{1} \sqcup l_{2}\right) \sqsubseteq f\left(l_{1}\right) \sqcup f\left(l_{2}\right) .
$$

This weaker condition is enough as the other way around $f\left(l_{1} \sqcup l_{2}\right) \sqsupseteq f\left(l_{1}\right) \sqcup$ $f\left(l_{2}\right)$ follows by monotonicity of $f$ and the fact that $\sqcup$ is the least upper bound.

## The 4 classical examples

- for a label consistent program $S_{*}$, all are instances of a monotone, distributive, framework:
- conditions:
- lattice of properties: immediate (subset/superset)
- ascending chain condition: finite set of syntactic entities
- closure conditions on $\mathcal{F}$
* monotone
* closure under identity and composition
- distributivity: assured by using the kill- and generate-formulation


## Overview over the 4 examples

|  | avail. epxr. | reach. def's | very busy expr. | live var's |
| :---: | :---: | :---: | :---: | :---: |
| $L$ | $2^{\mathbf{A E x p}_{*}}$ | $2^{\mathbf{V a r}_{*} \times \mathbf{L a b}_{*}^{\prime}}$ | $2^{\mathbf{A E x p}_{*}}$ | $2^{\mathbf{V a r}_{*}}$ |
| $\sqsubseteq$ | $\supseteq$ | $\subseteq$ | $\supseteq$ | $\subseteq$ |
| $\sqcup$ | $\cap$ | $\bigcup$ | $\cap$ | $\bigcup$ |
| $\perp$ | $\mathbf{A E x p}_{*}$ | $\emptyset$ | AExp $_{*}$ | $\emptyset$ |
| $\iota$ | $\emptyset$ | $\left\{(x, ?) \mid x \in \operatorname{fv}\left(S_{*}\right)\right\}$ | $\emptyset$ | $\emptyset$ |
| $E$ | $\left\{\operatorname{init}\left(S_{*}\right)\right\}$ | $\left\{\operatorname{init}\left(S_{*}\right)\right\}$ | final $\left(S_{*}\right)$ | final $\left(S_{*}\right)$ |
| $F$ | $\operatorname{flow}\left(S_{*}\right)$ | flow $\left(S_{*}\right)$ | flow $^{R}\left(S_{*}\right)$ | flow $^{R}\left(S_{*}\right)$ |
| $\mathcal{F}$ | $f: L \rightarrow L \mid \exists l_{k}, l_{g .} f(l)=\left(l \backslash l_{k}\right) \cup l_{g}$ |  |  |  |
| $f_{l}$ | $f_{l}(l)=\left(l \backslash \operatorname{kill}\left([B]^{l}\right) \cup \operatorname{gen}\left([B]^{l}\right)\right)$ where $[B]^{l} \in \operatorname{blocks}\left(S_{*}\right)$ |  |  |  |

### 2.5 Equation solving

## Solving the analyses

- given: set of equations (or constraints) over finite sets of variables
- domain of variables: complete lattices + ascending chain condition
- 2 solutions for the monotone frameworks
- MFP: "maximal fix point"
- MOP: "meet over all paths"

Remarks Finally, we come to to how to solve the equations. We have seen two glimpses to the problem. One was at the introduction, the chaotic iteration, the other one was the "theory" related to the fixpoints. We will shortly revisit the chaotic iteration. What was lacking there was the concrete (=deterministic) realization.

## MFP

- terminology: historically "MFP" stands for maximal fix point (not minimal)
- iterative worklist algorithm:
- central data structure: worklist
- list (or container/set) of pairs
- related to chaotic iteration


## Chaotic iteration

```
Input: equations for reaching defs
    for the given program
Output: least solution: R\vec{D}=(\mp@subsup{RDD}{1}{},\ldots,\mp@subsup{R}{12}{})
```

Initialization:

$$
\mathrm{RD}_{1}:=\emptyset ; \ldots ; \mathrm{RD}_{12}:=\emptyset
$$

Iteration:
while $\mathrm{RD}_{j} \neq F_{j}\left(\mathrm{RD}_{1}, \ldots, \mathrm{RD}_{12}\right)$ for some $j$
do

$$
\mathrm{RD}_{j}:=F_{j}\left(\mathrm{RD}_{1}, \ldots, \mathrm{RD}_{12}\right)
$$

## Worklist algorithms

- fixpoint iteration algorithm
- general kind of algorithms, for DFA, CFA, ...
- same for equational and /constraint systems
- "specialization" i.e., determinization of chaotic iteration
$\Rightarrow$ worklist: central data structure, "container" containing "the work still to be done"
- for more details (different traversal strategies): see Chap. 6 from [8]


## WL-algo for DFA

- WL-algo for monotone frameworks
$\Rightarrow$ input: instance of monotone framework
- two central data structures
- worklist: /flow-edges yet to be (re-)considered:

1. removed when effect of transfer function has been taken care of
2. (re-) added, when point 1 endangers satisfaction of (in-)equations - array to store the "current state" of Analysis。

- one central control structure (after initialization): loop until worklist empty

Remember that the result of the analysis is a mapping from the entry and the exit points for each block. Here, only the entry blocks are stored. An array is of course a good representation of a finite function.

Why do we need only the "entry" of the blocks (assuming forward)? In the chaotic iteration we clearly see pre- and post-states. First we have to remember the chaotic iteration. There, an $\mathrm{RD}_{i}$ depends via $F$ on all $\mathrm{RD}_{j}$. Of course, in reality that's not the case, and moreover, we should distinguish between entry and exit points. For the exit points, of course, they only depend on the entry-point and nothing else. The worklist algorithm actually considers only the relation from the post-condition to the predondition, more precisely one pre-condition. That means, that only the inter-flow is actually checked. So, in some sense, the post-conditions are represented, but only implicitely in that they are calculated on the fly from the given pre-condition, when needed. That can be seen also in step 3 .

## Code

```
Input: ( }L,\mathcal{F},F,E,\iota,f
Output: MFP,,MFP.
Method: step 1: initialization
    W:= nil;
    for all (I,I') \inF do W:= (I, I'):: W;
    for all }l\inF\mathrm{ or }\inE\mathrm{ do
        if }l\inE\mathrm{ then Analysis[I]:= 
            else Analysis[/] := + L;
    step 2: iteration
        while W\not= nil do
        (I,I') := ( fst(head(W)), snd(head(W)));
        W := tail W;
        if f}\mp@subsup{f}{l}{(Analysis[/]) & Analysis[/']
        then Analysis[I'] := Analysis[㣙] }\sqcup\mp@subsup{f}{l}{\prime}(\mathrm{ Analysis[/]);
            for all I'' with (I', I') \inF do
                    W:=(I', I') ::W;
    step 3: presenting the result:
        for all }l\inF\mathrm{ or }\inE\mathrm{ do
            MFP黾(I) := Analysis[/];
            MFP.(I) := fl/(Analysis[/])
```


## ML Code

```
Let rec solve (wl1 : edge list) : unit =
    match wl1 with
    [] -> ()
        (1, -> () (* wl done *)
        let ana_pre: var list = lookx (ana,l) (* extract `.states *)
        and ana-post : var list = lookx (ana, l')
        in let ana_exitpre: var list = f_trans(ana_pre, l)
        in
        if not (subset (ana_exitpre, ana_post))
        then
            (enter (ana, l', union(ana_post, ana_exitpre))
                let (new_edges : edge list) =
                    (let (preds : node list) = Flow.Graph.pred (l')
                    in List.map (fun n m (1', n)) preds)
            in solve (new_edges @ wl')
        @\mp@code{O}
            (* Nothing to do here. *)
in
un (x: node) -> lookx (ana, x)
```

; ;

## MFP: properties

Lemma 2.5.1. The algo

- terminates and
- calculates the least solution

Proof. - termination: ascending chain condition \& loop is enlarging

- least FP:
- invariant: array always below Analysiso
- at loop exit: array "solves" (in-)equations


## Time complexity

- estimation of upper bound of number basic steps
- at most $b$ different labels in $E$
- at most $e \geq b$ pairs in the flow $F$
- height of the lattice: at most $h$
- non-loop steps: $O(b+e)$
- loop: at most $h$ times addition to the WL
$\Rightarrow$

$$
\begin{equation*}
O(e \cdot h) \tag{2.10}
\end{equation*}
$$

or $\leq O\left(b^{2} h\right)$

### 2.6 Interprocedural analysis

### 2.6.1 Introduction

## Adding procedures

- so far: very simplified language:
- minimalistic imperative language
- reading and writing to variables plus
- simple controlflow, given as flow graph
- now: procedures: interprocedural analysis
- complications:
- calls/return (control flow)
- parameter passing (call-by-value vs. call-by-reference)
- scopes
- potential aliasing (with call-by-reference)
- higher-order functions/procedures
- here: top-level procedures, mutual recursion, call-by-value parameter + call-by-result


## Syntax

- begin $D_{*} S_{*}$ end

$$
D::=\operatorname{proc} p(\operatorname{val} x, \operatorname{res} y) \text { is } S \text { end }_{l_{n}}^{l_{x}} \mid D D
$$

- procedure names $p$
- statements

$$
S::=\ldots[\operatorname{call} p(a, z)]_{l_{r}}^{l_{c}}
$$

- note: call statement with 2 labels
- statically scoped language, CBV parameter passing (1st parameter), and CBN for second
- mutual recursion possible
- assumption: unique labelling, only declared procedures are called, all procedures have different names.


## Example: Fibonacci

### 2.6.2 Semantics

Explanation Next comes the adaptation of the definition of the flow graph. To do so, we need to adapt and extend the definitions of flow, block, etc. The new part deals, obviously, with the procedures. The basic trick is that we introduce new kinds of edges to deal with the procedures. The definition/presentation proceeds (in the slides) in two steps, first the call sites, afterwards the procedures themselves.

## Block, labels, etc.

$$
\begin{aligned}
\operatorname{init}\left([\operatorname{call} p(a, z)]_{l_{c}}^{l_{c}}\right. & =l_{c} \\
\text { final }\left([\operatorname{call} p(a, z)]_{l_{r}}^{l_{c}}\right. & =\left\{l_{r}\right\} \\
\operatorname{blocks}\left([\operatorname{call} p(a, z)]_{l_{c}}^{l_{c}}\right. & =\left\{[\operatorname{call} p(a, z)]_{l_{r}}^{l_{c}}\right\} \\
\operatorname{labels}([\operatorname{call} p(a, z)]]_{c_{c}}^{l_{c}} & =\left\{l_{c}, l_{r}\right\} \\
\text { flow }\left([\operatorname{call} p(a, z)]_{l_{r}}^{l_{c}}\right) & =\left\{\left(l_{\mathrm{c}} ; \mathbf{1}_{\mathrm{n}}\right),\left(\mathbf{l}_{\mathbf{x}} ; \mathbf{1}_{\mathbf{r}}\right)\right\}
\end{aligned}
$$

where $\operatorname{proc} p(\operatorname{val} x$, res $y)$ is $^{l_{n}} S$ end $^{l_{x}}$ is in $D_{*}$.

- two new kinds of flows (written slightly different(!)): calling and returning
- static dispatch only


## For procedure declaration

$$
\begin{aligned}
\operatorname{init}(p) & =l_{n} \\
\operatorname{final}(p) & =\left\{l_{x}\right\} \\
\operatorname{blocks}(p) & =\left\{\text { is }^{l_{n}}, \operatorname{end}^{l_{x}}\right\} \cup \operatorname{blocks}(S) \\
\operatorname{labels}(p) & =\left\{l_{n}, l_{x}\right\} \cup \operatorname{labels}(S) \\
\operatorname{flow}(p) & =\left\{\left(l_{n}, \operatorname{init}(S)\right)\right\} \cup \operatorname{flow}(S) \cup\left\{\left(l, l_{x}\right) \mid l \in \operatorname{final}(S)\right\}
\end{aligned}
$$

## "Standard" flow of complete program

not yet interprocedural flow (IF)

```
    init \(_{*}=\operatorname{init}\left(S_{*}\right)\)
final \(_{*}=\operatorname{final}\left(S_{*}\right)\)
blocks* \(=\bigcup\left\{\right.\) blocks \((p) \mid \operatorname{proc} p(\operatorname{val} x\), res \(y)\) is \(^{l_{n}} S\) end \(\left.^{l_{x}} \in D_{*}\right\}\)
    \(\cup\) blocks \(\left(S_{*}\right)\)
labels \(_{*}=\bigcup\left\{\operatorname{labels}(p) \mid \operatorname{proc} p(\operatorname{val} x\right.\), res \(y)\) is \(^{l_{n}} S\) end \(\left.^{l_{x}} \in D_{*}\right\}\)
        \(\cup\) labels \(\left(S_{*}\right)\)
flow \(_{*}=\bigcup\left\{\right.\) flow \((p) \mid \operatorname{proc} p(\operatorname{val} x\), res \(y)\) is \(^{l_{n}} S\) end \(\left.^{l_{x}} \in D_{*}\right\}\)
    \(\cup\) flow \(\left(S_{*}\right)\)
```

side remark: $S_{*}$ : notation for complete program "of interest"

## New kind of edges: Interprocedural flow (IF)

- inter-procedural: from call-site to procedure, and back: $\left(l_{c} ; l_{n}\right)$ and $\left(l_{x} ; l_{r}\right)$.
- more precise (= better) capture of flow
- abbreviation: IF for inter-flow * or inter-flow ${ }_{*}^{R}$


## IF

$$
\left.\begin{array}{rl}
\text { inter-flow }_{*}=\left\{\left(l_{c}, l_{n}, l_{x}, l_{r}\right) \mid P_{*}\right. \text { contains } & {\left[\begin{array}{c}
[\operatorname{call} p(a, z)]]_{l_{r}}^{l_{c}} \text { and } \\
\\
\\
\text { proc(val } x, \text { res } y) \text { is }^{l_{n}} S \text { end }^{l_{x}}
\end{array}\right\}}
\end{array}\right\}
$$

## Example: fibonacci flow

Example: fibonacci flow


## Semantics: stores, locations,. . .

- not only new syntax
- new semantical concept: local data!
- different "incarnations" of a variable $\Rightarrow$ locations
- remember: $\sigma \in$ State $=\mathbf{V a r}_{*} \rightarrow \mathbf{Z}$


## Representation of "memory"

$$
\begin{array}{lll}
\xi \in \mathbf{L o c} & \text { locations } \\
\rho \in \mathbf{E n v}^{\prime}=\mathbf{V a r}_{*} \rightarrow \mathbf{L o c} & \text { environment } \\
\varsigma \in \mathbf{S t o r e}=\mathbf{L o c} \rightarrow_{\text {fin }} \mathbf{Z} & \text { store }
\end{array}
$$

- $\sigma=\varsigma \circ \rho$ : total $\Rightarrow \operatorname{ran}(\rho) \subseteq \operatorname{dom}(\varsigma)$
- top-level environment: $\rho_{*}$ : all var's are mapped to unique locations (no aliasing !!!!)

Explanations $\rightarrow_{f i n}$ represents finite partial functions.

## SOS steps

- steps relative to environment $\rho$

$$
\rho \vdash_{*}\langle S, \varsigma\rangle \rightarrow\langle\dot{S}, \dot{\varsigma}\rangle
$$

or

$$
\rho \vdash_{*}\langle S, \varsigma\rangle \rightarrow \zeta
$$

- old rules needs to be adapted
- "global" environment $\rho_{*}$


## Call-rule

$$
\begin{gathered}
\xi_{1}, \xi_{2} \notin \operatorname{dom}(\varsigma) \quad v \in \mathbf{Z} \\
\operatorname{proc} p(\operatorname{val} x, \operatorname{res} y) \text { is }^{l_{n}} S \text { end }^{l_{x}} \in D_{*} \\
\dot{\varsigma}=\varsigma\left[\xi_{1} \mapsto[a]_{\mathrm{Sop}}^{4}\right]\left[\xi_{2} \mapsto v\right] \\
\frac{\rho \vdash_{*}\left\langle[\operatorname{call} p(a, z)]_{l_{r}}^{l_{c}}, \varsigma\right\rangle \rightarrow\left\langle\operatorname{bind} \rho_{*}\left[x \mapsto \xi_{1}\right]\left[y \mapsto \xi_{2}\right] \text { in } S \text { then } z:=y, \dot{\varsigma}\right\rangle}{l^{\prime}} \mathrm{CALL}
\end{gathered}
$$

## Bind-construct

$$
\begin{gathered}
\frac{\dot{\rho} \vdash_{*}\langle S, \varsigma\rangle \rightarrow\langle\dot{S}, \dot{\varsigma}\rangle}{\rho \vdash_{*}\langle\text { bind } \hat{\rho} \text { in } S \text { then } z:=y, \varsigma\rangle \rightarrow\langle\text { bind } \rho \text { in } \dot{S} \text { then } z:=y, \dot{\varsigma}\rangle} \text { Bind }_{1} \\
\frac{\rho \dot{\rho} \vdash_{*}\langle S, \varsigma\rangle \rightarrow \dot{\varsigma}}{\rho \vdash_{*}\langle\text { bind } \hat{\rho} \text { in } S \text { then } z:=y, \varsigma\rangle \rightarrow \dot{\varsigma}[\rho(z) \mapsto \dot{\varsigma}(\hat{\rho}(y))]} \text { Bind }_{2}
\end{gathered}
$$

- bind-syntax: "runtime syntax"
$\Rightarrow$ formulation of correctness must be adapted, too (Chap. 3) ${ }^{1}$


### 2.6.3 Analysis

## Transfer function: Naive formulation

- first attempt
- assumptions:
- for each proc. call: 2 transfer functions: $f_{l_{c}}$ (call) and $f_{l_{r}}$ (return)

[^1]- for each proc. definition: 2 transfer functions: $f_{l_{n}}$ (enter) and $f_{l_{x}}$ (exit)
- given: mon. framework $(L, \mathcal{F}, F, E, \iota, f)$


## Naive

- treat IF edges $\left(l_{c} ; l_{n}\right)$ and $\left(l_{x} ; l_{r}\right)$ as ordinary flow edges $\left(l_{1}, l_{2}\right)$
- ignore parameter passing: transfer functions for proc. calls and proc definitions are identity


## Equation system ("naive" version")

$$
\begin{aligned}
& A_{\bullet}(l)=f_{l}\left(A_{\circ}(l)\right) \\
& A_{\circ}(l)=\sqcup\left\{A_{\bullet}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in F \text { or }\left(l^{\prime} ; l\right) \in F\right\} \sqcup \iota_{E}^{l}
\end{aligned}
$$

with

$$
\iota_{E}^{l}= \begin{cases}\iota & \text { if } l \in E \\ \perp & \text { if } l \notin E\end{cases}
$$

- analysis: safe
- unnecessarily imprecise, too abstract

The equational system here is / without/ the assumption of isolated entries/exits. That corresponds to page 64 in the book.

### 2.6.4 Paths

## Paths

- remember: "MFP"
- historically: MOP stands for meet over all paths
- here: dually mosty joins
- 2 "versions" of a path:
- path to entry of a block: blocks traversed from the "extremal block" of the program, but not including it
- path to exit of a block


## Paths

$$
\begin{aligned}
& \text { path }_{\circ}(l)=\left\{\left[l_{1}, \ldots \mathbf{l}_{\mathbf{n}-\mathbf{1}}\right] \mid l_{i} \rightarrow_{\text {fow }} l_{i+1} \wedge l_{n}=l \wedge l_{1} \in E\right\} \\
& \text { path }_{\bullet}(l)=\left\{\left[l_{1}, \ldots \mathbf{l}_{\mathbf{n}}\right] \mid l_{i} \rightarrow_{\text {foow }} l_{i+1} \wedge l_{n}=l \wedge l_{1} \in E\right\}
\end{aligned}
$$

- transfer function for paths $\vec{l}$

$$
f_{\vec{l}}=f_{l_{n}} \circ \ldots f_{l_{1}} \circ i d
$$

## Meet over all paths

- paths:
- forward: paths from init block to entry of a block
- backwards: paths from exits of a block to a final block
- two versions for the MOP solution (for given $l$ ):
- up-to but not including $l$
- up-to including $l$


## MOP

$$
\begin{aligned}
& \operatorname{MOP}_{\circ}(l)=\bigsqcup\left\{f_{\vec{l}}(\iota) \mid \vec{l} \in \text { path }_{\circ}(l)\right\} \\
& M O P_{\bullet}(l)=\bigsqcup\left\{f_{\vec{l}}(\iota) \mid \vec{l} \in \operatorname{path}_{\bullet}(l)\right\}
\end{aligned}
$$

## MOP vs. MFP

- MOP: can be undecidable
- MFP approximates MOP ("MFP $\sqsupseteq M O P ")$


## Lemma 2.6.1.

$$
\begin{equation*}
M F P_{\circ} \sqsupseteq M O P_{\circ} \text { and } M F P_{\bullet} \sqsupseteq M O P_{\bullet} \tag{2.11}
\end{equation*}
$$

In case of a distributive framework

$$
\begin{equation*}
M F P_{\circ}=M O P_{\circ} \text { and } M F P_{\bullet}=M O P_{\bullet} \tag{2.12}
\end{equation*}
$$

If the transfer function is given by kill and generate as shown, the analysis is distributive.

## MVP

- take calls and returns (IF) serious
- restrict attention to valid ("possible") paths
$\Rightarrow$ capture the nesting structure
- from MOP to MVP: "meet over all valid paths"
- complete path:
- appropriate call-nesting
- all calls are answered


## Complete paths

- given $P_{*}=\operatorname{begin} D_{*} S_{*}$ end
- $C P_{l_{1}, l_{2}}$ : complete paths from $l_{1}$ to $l_{2}$
- generated by the following productions (l's are the terminals) (we assume forward analysis here)
- basically a context-free grammar

$$
\begin{aligned}
& \overline{C P_{l, l} \longrightarrow l} \\
& \frac{\left(l_{1}, l_{2}\right) \in F}{C P_{l_{1}, l_{3}} \longrightarrow l_{1}, C P_{l_{2}, l_{3}}} \\
& \frac{\left(l_{c}, l_{n}, l_{x}, l_{r}\right) \in I F}{C P_{l_{c}, l} \longrightarrow l_{c}, C P_{l_{n}, l_{x}}, C P_{l_{r}, l}}
\end{aligned}
$$

The notion of complete path is rather straightforward. It informally says that each call is answered by one corresponding return, and also that each return is matched by one corresponding call. It directly corresponds to the prototypical context-free parenthetic languages, except that we have an arbitrary number of different "parentheses" namely the different calls. The calls are not being identified by the name of the function being called, but by the call-sites and the identity of the function being called, more precisely by the two labels at the call site plus the two labels of the entry and the exit of the procedure. That is visible in the third rule.

The definition is given in a rule-like manner. They are not really like derivation rule, though. It's more like a family of grammar productions, namely for each label (first rule), for pairs of labels (second rule), resp. quadupel of labels (last rule). As the premises of the last two rules show, not for all tuples or all quadruples, of course, only those as given by the control flow graph, in particular taking care of the inter-procedural flow in the last rule.

The interpretation is as follows. There is only one complete path from a label to itself, that's the trivial path. The second rule just splits off one label on the left (one could do also differently). Also in the third rule, there is a split-off of the first step. A terminating execution will have a complete path. There are only a finite number of productions.

As a side remark: being a complete path, in some way, is not a safety property, whereas being a valid path, is.

## Example: Fibonacci

- concrete grammar for fibonacci program:

$$
\begin{aligned}
C P_{9,10} & \longrightarrow 9, C P_{1,8}, C P_{10,10} \\
C P_{10,10} & \longrightarrow 10 \\
C P_{1,8} & \longrightarrow 1, C P_{2,8} \\
C P_{2,8} & \longrightarrow 2, C P_{3,8} \\
C P_{2,8} & \longrightarrow 2, C P_{4,8} \\
C P_{3,8} & \longrightarrow 3, C P_{8,8} \\
C P_{8,8} & \longrightarrow 8 \\
C P_{4,8} & \longrightarrow 4, C P_{1,8}, C P_{5,8} \\
C P_{5,8} & \longrightarrow 5, C P_{6,8} \\
C P_{6,8} & \longrightarrow 6, C P_{1,8}, C P_{7,8} \\
C P_{7,8} & \longrightarrow 7, C P_{8,8}
\end{aligned}
$$

## Valid paths (context-free grammar)

## Valid path (generated from non-terminal $V P_{*}$ ):

- start at extremal node $(E)$,
- all proc exits have matching entries

$$
\begin{aligned}
& \frac{l_{1} \in E \quad l_{2} \in \mathbf{L a b}_{*}}{V P_{*} \longrightarrow V P_{l_{1}, l_{2}}} \\
& \frac{\left(l_{1}, l_{2}\right) \in F}{V P_{l, l} \longrightarrow l} \\
& \frac{V P_{l_{1}, l_{3}} \longrightarrow l_{1}, V P_{l_{2}, l_{3}}}{} \\
& \frac{\left(l_{c}, l_{n}, l_{x}, l_{r}\right) \in I F}{V P_{l_{c}, l} \longrightarrow l_{c}, C P_{l_{n}, l_{x}}, V P_{l_{r}, l}}
\end{aligned} \quad \frac{\left(l_{c}, l_{n}, l_{x}, l_{r}\right) \in I F}{V P_{l_{c}, l} \longrightarrow l_{c}, V P_{l_{n}, l}} .
$$

The grammar for valid paths is slightly more complex than the one for complete paths. There is an easy explanation what a valid path is: a valid path is a prefix of a complete path. One could leave it at that. The definition shows, basically, that also that property is a context-free property (namely by giving the corresponding (family of) productions.

## MVP

- adapt the definition of paths

$$
\begin{aligned}
\text { vpath }_{\circ}(l) & =\left\{\left[l_{1}, \ldots l_{\mathbf{n}-\mathbf{1}}\right] \mid l_{n}=l \wedge\left[l_{1}, \ldots, l_{n}\right] \text { valid }\right\} \\
\text { vpath }_{\bullet}(l) & =\left\{\left[l_{1}, \ldots l_{\mathbf{n}}\right] \mid l_{n}=l \wedge\left[l_{1}, \ldots, l_{n}\right] \text { valid }\right\}
\end{aligned}
$$

- MVP solution:

$$
\begin{aligned}
& M V P_{\circ}(l)=\sqcup\left\{f_{\vec{l}}(l) \mid \vec{l} \in \operatorname{vpath}_{\circ}(l)\right\} \\
& M V P_{\bullet}(l)=\sqcup\left\{f_{\vec{l}}(l) \mid \vec{l} \in \operatorname{vpath}_{\bullet}(l)\right\}
\end{aligned}
$$

- but still: "meets over paths" is impractical

Fixpoint calculations next: how to reconcile the path approach with MFP

### 2.6.5 Context-sensitive analysis

## Contexts

- MVP/MOP / undecidable( but more precise than basic MFP
$\Rightarrow$ instead of MVP: "embellish" MFP

$$
\begin{equation*}
\delta \in \Delta \tag{2.13}
\end{equation*}
$$

- $\delta$ : context information
- for instance: representing/recording of the path taken
$\Rightarrow$ "embellishment": adding contexts
embellished monotone framework

$$
(\hat{L}, \hat{\mathcal{F}}, F, E, \hat{\iota}, \hat{f})
$$

- intra-procedural (no change of embellishment $\Delta$ )
- inter-procedural

Embellishment, notationally, is indicated by a hat on top. The following will proceed in two stages. The intra-procedural part and the interprocedural part. The first part is (or course) simpler. One might ask, why we need to consider the first part at all? Well, we change the framework slightly by embellishing it (indicated by the hatted syntax). That will involve a change in the lattice and other concomitant changes. Consequently, also the intraprocedual part needs to be adapted, basically taking care of the embellishement, i.e., taking care of the contexts. Taking care basically is rather trivial and consists of "ignoring"
the context: as long as one deals with data-flow within one function body, the context remains the same. Nonetheless, the additional context-component has to be mentioned as being unchanged when dealing with the embelished transfer functions and other parts of the definition of the monotone framework. But the keyword is: the intra-procedural part is "basically unchanged".

## Intra-procedural: basically unchanged

- this part: "independent" of $\Delta$
- property lattice $\hat{L}=\Delta \rightarrow L$
- mononote functions $\hat{\mathcal{F}}$
- transfer functions: pointwise

$$
\begin{equation*}
\hat{f}_{l}(\hat{l})(\delta)=f_{l}(\hat{l}(\delta)) \tag{2.14}
\end{equation*}
$$

- flow equations: "unchanged" for intra-proc. part

$$
\begin{align*}
& A_{\bullet}(l)=\hat{f}_{l}\left(A_{\circ}(l)\right)  \tag{2.15}\\
& \left.A_{\circ}(l)=\sqcup\left\{A_{\bullet}\left(l^{\prime}\right) \mid\left(l^{\prime}, l\right) \in F \text { or }\left(l^{\prime} ; l\right) \in F\right)\right\} \sqcup \hat{\iota_{E}}
\end{align*}
$$

- in equation for $A_{\bullet}$ : except for labels $l$ for proc. calls (i.e., not $l_{c}$ and $l_{r}$ )

There is an unfortunate notational collision: Lattice $L$ with its elements on the one hand and labels/nodes in the CFG $l$ from $\mathbf{L a b}_{*}$

Apart from that: The above definitions define $\hat{L}$ as a function of type $\Delta \rightarrow L$. That gives raise to some "higher-order" explanations of what the embellished framework means (see already equation (2.14). That is a clean explanation of what is going on, but one may also see it as follows.

Remember that in the unembellished framework, a solution of a problem is a mapping from the nodes of the cfg to elements of the lattice. Let's use $N$ for the nodes or labels (not $L \ldots$ ). To be very precise, we are interested not in a mapping from nodes to the lattice but from the entries and the exits of the nodes to the lattice, but let's ignore that for now.

That means, that an unembelished solution is of the type $N \rightarrow L$, whereas now, the solution is of type

$$
N \rightarrow \Delta \rightarrow L
$$

That is the same as

$$
(N \times \Delta) \rightarrow L .
$$

Seen like that, the context is simply "paired" with the location or node in the control-flow graph and represents relevant information of the call-site where the function was called.

## Sign analysis (unembellished)

- $\boldsymbol{\operatorname { S i g n }}=\{-, 0,+\}, L_{\text {sign }}=2^{\mathbf{V a r}_{*} \rightarrow \mathbf{S i g n}}$
- abstract states $\sigma^{\text {sign }} \in L_{\text {sign }}$
- for expressions: []$_{-}^{\mathcal{A}_{\text {sign }}}: \mathbf{A E x p} \rightarrow\left(\mathbf{V a r}_{*} \rightarrow \mathbf{S i g n}\right) \rightarrow 2^{\text {Sign }}$

Transfer function for $[x:=a]^{l}$

$$
\begin{equation*}
f_{l}^{s i g n}(Y)=\bigcup\left\{\phi_{l}^{s i g n}\left(\sigma^{s i g n}\right) \mid \sigma^{s i g n} \in Y\right\} \tag{2.16}
\end{equation*}
$$

where $Y \subseteq$ Var $_{*} \rightarrow$ Sign and

$$
\begin{equation*}
\phi_{l}^{s i g n}\left(\sigma^{s i g n}\right)=\left\{\sigma^{s i g n}[x \mapsto s] \mid s \in[a]_{\sigma^{s i g n}}^{\mathcal{A}_{s i n}}\right\} \tag{2.17}
\end{equation*}
$$

We start with the unembellished part, i.e., without even considering contexts. For that basic setting, the lattice we start with is a set of functions; we can think of it as a set of states.

As a side remark: for what is called sign-analysis, that's not the only possible choice. An alternative to $L_{\text {sign }}$ would be to use a function Var $\rightarrow 2^{\text {Sign }}$. It would be a "state with abstract values" (where an abstract value is a set of concrete values) as opposed to an "abstract state" consististing of a set of concrete states. The alternative interpretation would be "weaker", i.e., more abstract.

Anyway: The above definition proceeds in 3 steps: At the core is the semantic function [-]. This function is for expressions, and is already non-deterministic. Eq. (2.17) reflects the effect of an assignment for one abstract state and (2.16) is the transfer function (lifted pointwise).
Why does []$_{-}^{\mathcal{A}_{s i g n}}$ give back a set? Clearly, because of the non-determinism due to abstraction.

The sign-analysis is not yet embellished here (embellished $=$ adding context). This means, there is not even a mentioning of $\Delta$ here. The real work is done in $\phi$ : the overall input to that function is $Y$, which is a set of states, and $\phi_{l}^{\text {sign }}$ just applies it pointwise, interpreting the expression on the right-hand side of the assignment and updating the state accordingly.

On the next slides, we will embellish the analysis, but since we are not yet in the inter-procedural part, the embellishment is not very interesting, just a "lifting" to the embellished setting.

## Sign analysis: embellished

$$
\begin{align*}
\hat{L}_{\text {sign }} & =\Delta \rightarrow L_{\text {sign }}  \tag{2.18}\\
& =\Delta \rightarrow 2^{\text {Var }_{*} \rightarrow \text { Sign }} \simeq 2^{\Delta \times\left(\mathbf{V a r}_{*} \rightarrow \mathbf{S i g n}\right)}
\end{align*}
$$

Transfer function for $[x:=a]^{l}$

$$
\begin{equation*}
\hat{f}_{l}^{s i g n}(Z)=\bigcup\left\{\{\delta\} \times \phi_{l}^{s i g n}\left(\sigma^{s i g n}\right) \mid\left(\delta, \sigma^{s i g n}\right) \in Z\right\} \tag{2.19}
\end{equation*}
$$

The unembellished one so far was a simple instance of the monotone framework. The transfer function just "joins" all possible outcomes, where it is assumed that we have as function that calculates the set of signs for expression. That was completely standard. Now, it does not get really more complex: equation (2.19) just does nothing with the $\delta$, since we are still within a single process. In the following we go to the inter-procedural fragment and there things get more complex, since for dealing with calls and returns we have to connect the contexts of the caller and the callee. It's a bit like parameter passing.

## Inter-procedural

- procedure definition proc(val $x$, res $y)$ is ${ }^{l_{n}} S$ end $^{l_{x}}$ :

$$
\hat{f}_{l_{n}}, \hat{f}_{l_{x}}:(\Delta \rightarrow L) \rightarrow(\Delta \rightarrow L)=i d
$$

- procedure call: $\left(l_{c}, l_{n}, l_{x}, l_{r}\right) \in I F$
- here: forward analysis
- call: 2 transfer functions $/ 2$ sets of equations, i.e., for all $\left(l_{c}, l_{n}, l_{x}, l_{r}\right) \in I F$


## 2 transfer functions

1. for calls: $\hat{f}^{1}{ }_{l_{c}}:(\Delta \rightarrow L) \rightarrow(\Delta \rightarrow L)$

$$
\begin{equation*}
A_{\bullet}\left(l_{c}\right)=\hat{f}_{l_{c}}\left(A_{\circ}\left(l_{c}\right)\right) \tag{2.20}
\end{equation*}
$$

1. for returns: $\hat{f}^{2}{ }_{l_{c}, l_{r}}:(\Delta \rightarrow L) \times(\Delta \rightarrow L) \rightarrow(\Delta \rightarrow L)$

$$
\begin{equation*}
\left.A_{\bullet}\left(l_{r}\right)=\hat{f}^{2}{ }_{l_{c}, l_{r}}\left(A_{\circ}\left(l_{c}\right), A_{\circ}\left(l_{r}\right)\right)\right) \tag{2.21}
\end{equation*}
$$

## Procedure call



Note again the unfortunate notational collision: $\hat{l}$ : element of embellished lattice (abstract value), $l_{c}$ etc: nodes/labels in the control flow graph. The situation may become even more confusing for analyses like RD: there labels (which are nodes in the control-flow graph) are part of the values of interest and thus also elements of the lattice.

Next come two different simplifications for $f^{2}$. However, one way to understand the 2 arguments for the return is that often one wants to match the return with the call (via the context).

Ignoring the call context

$$
\hat{f}_{l_{c}, l_{r}}^{2}\left(\hat{l}, \hat{l}^{\prime}\right)=\hat{f}_{l_{r}}^{2}\left(\hat{l}^{\prime}\right)
$$



## Merging call contexts

$$
\hat{f}_{l_{c}, l_{r}}^{2}\left(\hat{l}, \hat{l}^{\prime}\right)=\hat{f}_{l_{c}, l_{r}}^{2 A}(\hat{l}) \sqcup \hat{f}_{l_{c}, l_{r}}^{2 B}\left(\hat{l}^{\prime}\right)
$$



## Context sensitivity

- IF-edges: allow to relate returns to matching calls
- context insensitive: proc-body analysed combining flow information from all call-sites.
- contexts: used to distinguish different call-sites
$\Rightarrow$ context sensitive analysis $\Rightarrow$ more precision + more effort

In the following: 2 specializations:

1. control ("call strings")
2. data
(combinations of course possible) Combinations of the two approaches are not covered in the lecture. The call-strings corresponds more or less to the previously sketched MVP approach.

## Call strings

- context $=$ path
- call-string = sequence of currently "active" calls
- concentrating on calls: flow-edges $\left(l_{c}, l_{n}\right)$, where just $l_{c}$ is recorded

$$
\Delta=\text { Lab }^{*} \quad \text { call strings }
$$

- extremal value (from $\hat{L}=\Delta \rightarrow L$ )

$$
\hat{\iota}(\delta)= \begin{cases}\iota & \text { if } \delta=\epsilon \\ \perp & \text { otherwise }\end{cases}
$$

The definition of $\hat{\iota}$ should be clear: at the beginning of the program, there are no calls, and that is what we recording, hence the call string must be empty. Note again the higher-order approach. The $\hat{\iota}$ is somehow defined again point-wise. The initial value, an element from the lattice $\Delta \rightarrow L$, which here contains those functions.

## Fibonacci flow



## Fibonacci call strings

some call strings:

$$
\epsilon,[9],[9,4],[9,6],[9,4,4],[9,4,6],[9,6,4],[9,6,6], \ldots
$$

similar, but not same as valid paths

Explanations The call strings is not the same as the valid paths. It's related, though. The difference is the treatment of the returns. In the valid (or complete) path description, the returns are part of the paths, and the paths "never forget", they only grow longer. Here, when dealing with a return, the path does not get longer, it gets shorter be removing the the previous call. The call string only tracks the currently open calls. It corresponds to the current depth in the call-stack. That is also the way to match the contexts of the callee and the caller.

Note that if a function body calls another function 2 times, then the two call sites may still be confused!

## Transfer functions for call strings

- here: forward analysis
- 2 cases: define $\hat{f}_{l_{c}}^{1}$ and $\hat{f}_{l_{c}, l_{r}}^{2}$


## Transfer functions

- calls (basically: check that the path ends with $l_{c}$ ):

$$
\begin{align*}
\hat{f}_{l_{c}}^{1}(\hat{l})\left(\left[\delta, l_{c}\right]\right) & =f_{l_{c}}^{1}(\hat{l}(\delta))  \tag{2.22}\\
\hat{f}_{l_{c}}^{1}(-) & =\perp
\end{align*}
$$

- returns (basically: match return with the call)

$$
\begin{equation*}
\hat{f}_{l_{c}, l_{r}}^{2}\left(\hat{l}, \hat{l}^{\prime}\right)(\delta)=f_{l_{c}, l_{r}}^{2}\left(\hat{l}(\delta), \hat{l}^{\prime}\left(\left[\delta, l_{c}\right]\right)\right) \tag{2.23}
\end{equation*}
$$

- rather "higher-order" way of connecting the flows, using the call-strings as contexts
- connection between the arguments (via $\delta$ ) of $f_{l_{c}, l_{r}}$
- given: underlying $f_{l_{c}}^{1}$ and $f_{l_{c}, l_{r}}^{2}$.
- Notation: $\left[\delta, l_{c}\right]$ : concatenation of calls string
- $l^{\prime}$ : at procedure exit.


## Sign analysis (continued)

- so far: "unconcrete", i.e.,
- given some underlying analysis: how to make it context-sensitive
- call-strings as context
- now: apply to some simple case: signs
- remember: $\hat{L} \simeq 2^{\Delta \times\left(\mathbf{V a r}_{*} \rightarrow \mathbf{S i g n}\right)}$ (see Eq. (2.18))
- before: standard embellished $\hat{f}_{l}^{\text {Sign }}$ (with the help of $\phi_{l}^{\text {Sign }}$ )
- now: inter-procedural


## Sign analysis: aux. functions $\phi$

still unembellished
calls: abstract parameter-passing

$$
\phi_{l_{c}}^{s i g n 1}\left(\sigma^{s i g n}\right)=\left\{\sigma^{s i g n}[x \mapsto s]\left[y \mapsto s^{\prime}\right] \mid s \in[a]_{\sigma^{s i g n}}^{\mathcal{A}_{s i g n}}, s^{\prime} \in\{-, 0,+\}\right\}
$$

returns (analogously)

$$
\phi_{l_{c}, l_{r}}^{s i n 2}\left(\sigma_{1}^{s i g n}, \sigma_{2}^{s i g n}\right)=\left\{\sigma_{2}^{s i g n}\left[x, y, z \mapsto \sigma_{1}^{s i g n}(x), \sigma_{1}^{s i g n}(y), \sigma_{2}^{s i g n}(y)\right]\right\}
$$

(formal params: $x, y$, where $y$ is the result parameter, call-site return variable $z$ )

- non-det "assignment" to $y$
- remember: operational semantics,


## Sign analysis

calls: abstract parameter-passing + glueing calls-returns

$$
\left.\hat{f}_{l_{c}}^{s i g n 1}(Z)=\bigcup\left\{\left\{\delta^{\prime}\right\} \times \phi_{l_{c}}^{s i g n 1}\left(\sigma^{s i g n}\right) \mid\left(\delta^{\prime}, \sigma^{s i g n}\right) \in Z, \delta^{\prime}=\left[\delta, l_{c}\right]\right)\right\}
$$

## Returns: analogously

$$
\begin{aligned}
\hat{f}_{l_{c}, l_{r}}^{s i g n}\left(Z, Z^{\prime}\right)=\bigcup\left\{\{\delta\} \times \phi_{l_{c}, l_{r}}^{s i g n 2}\left(\sigma_{1}^{s i g n}, \sigma_{2}^{s i g n}\right) \mid\right. & \left(\delta, \sigma_{1}^{s i g n}\right) \in Z \\
& \left(\delta^{\prime}, \sigma_{2}^{\text {sign }}\right) \in Z^{\prime} \\
& \delta^{\prime}=\left[\delta, l_{c}\right]
\end{aligned}
$$

(formal params: $x, y$, call-site return variable $z$ )
The sign analysis was introduced before. The start was unembellished, just the context-non-sensitive case. There, the $\phi$ was done, as a pre-step for the unembellished transfer functions. The underlying lattice was not a mapping from variables to sets of signs (which would have been possible) but the more precise ets of such
mappings. Those were called abstract states. A that point we had already the embellished transfer function, but only for the non-procedure case. Note also: in this particular setting: the embellished lattice, in general, is a mapping from contexts to the old lattice. Here, the lattice is isomorphic to sets of pairs (see equation (2.18) for that).

How does this work? It works as before, i.e., as for the intra-procedural analysis. The auxiliary function $\phi$ (for signs) in (2.17) when the example sign-example was introduced which was already used in the unembellished setting to define the (unembellished) $f_{l}^{\text {sign }}$. In the embellished setting in the old intra-procedural part, the $\phi$ is also not used for touching the paths $\delta$.

Here we now see, that the auxiliary $\phi_{l}^{\text {sign }}$ is split into 2 functions $\phi_{l_{c}}^{s i g n 1}$ and $\phi_{l_{c}, l_{r}}^{s i g n 2}$. This is done analogous to the splitting of the transfer functions (the $\phi$ 's are just auxiliary constructions to it anyway). As before, the $\phi$ 's have nothing to do with the paths. But they have to be different, because of the parameter passing (for $x$ and $y$ e.g., in the call). Note that the value for $y$ is set arbitrarily.

The interesting coupling is in $\delta$ and $\delta^{\prime}$ (resp. $Z$ and $Z^{\prime}$ )!
The definition can be best understood into two states, both for calls and for returns. As before, we assume that the abstract denotational semantics for expression is given (which is already non-deterministic and not repeated here). The two stages are 1) defined " f " for one abstract state (resp. for a pair, in case of the return) and then 2) lift it to sets of such.

1. that one is for parameter passing on the abstract level. The $s^{\prime}$ is just because of the call-by-result semantics for the result parameter, it's just the same as in the semantics (remember the SOS).
2. The second stage, the real transfer function, lifts it to sets. Cf. also the unembellished lifting function (for signs) in equation (2.17). The important change is that now we have the contexts (as call strings) $\delta$ in the lattice

## Call strings of bounded length

- recursion $\Rightarrow$ call-strings of unbounded length
$\Rightarrow$ restrict the length

$$
\Delta=\mathbf{L a b} \quad{ }^{\leq k} \quad \text { for some } k \geq 0
$$

- for $k=0$ context-insensitive $(\Delta=\{\epsilon\})$


## Assumption sets

- alternative to call strings
- not tracking the path, but assumption about the state
- assume here: lattice
$L=2^{D}$
$\Rightarrow \hat{L}=\Delta \rightarrow L \simeq 2^{\Delta \times D}$
restrict to only the last call
dependency on data only $\Rightarrow$

$$
\Delta=2^{D}
$$

- $\hat{\iota}=\{(\{\iota\}, \iota)\}$ extremal value


## Transfer functions

- calls

$$
\left.\begin{array}{rl}
\hat{f}_{l_{c}}^{1}(Z)=\bigcup\left\{\left\{\delta^{\prime}\right\} \times \phi_{l_{c}}^{1}(d) \mid\right. & (\delta, d) \in Z \wedge \\
& \delta^{\prime}=\left\{d^{\prime \prime} \mid\left(\delta, d^{\prime \prime}\right) \in Z\right\}
\end{array}\right\}
$$

where $\phi_{l_{c}}^{1}: D \rightarrow 2^{D}$

- note: new context $\delta^{\prime}$ for the procedure body
- "caller-callee" connection via the context (= data) $\delta$
- return

$$
\begin{aligned}
\hat{f}_{l_{c}, l_{r}}^{2}\left(Z, Z^{\prime}\right)=\bigcup\left\{\{\delta\} \times \phi_{l_{c}, l_{r}}^{2}\left(d, d^{\prime}\right) \mid\right. & (\delta, d) \in Z \wedge \\
& \left(\delta^{\prime}, d^{\prime}\right) \in Z^{\prime} \wedge \\
& \delta^{\prime}=\left\{d^{\prime \prime} \mid\left(\delta, d^{\prime \prime}\right) \in Z\right\}
\end{aligned}
$$

## Small assumption sets

- throw away even more information.

$$
\Delta=D
$$

- instead of $2^{D} \times D$ : now only $D \times D$.
- transfer functions simplified
- call

$$
\hat{f}_{l_{c}}^{1}(Z)=\bigcup\left\{\{\delta\} \times \phi_{l_{c}}^{1}(d) \mid(\delta, d) \in Z\right\}
$$

- return

$$
\begin{aligned}
\hat{f}_{l_{c}, l_{r}}^{2}\left(Z, Z^{\prime}\right)=\bigcup\left\{\{\delta\} \times \phi_{l_{c}, l_{r}}^{2}\left(d, d^{\prime}\right) \mid\right. & (\delta, d) \in Z \wedge\} \\
& \left(\delta, d^{\prime}\right) \in Z^{\prime}
\end{aligned}
$$

## Flow-(in-)sensitivity

- "execution order" influences result of the analysis:

$$
S_{1} ; S_{2} \quad \text { vs. } \quad S_{2} ; S_{1}
$$

- flow in-sensitivity: order is irrelevant
- less precise (but "cheaper")
- for instance: kill is empty
- sometimes useful in combination with inter-proc. analysis


## Set of assigned variables

- for procedure $p$ : determine

$$
\operatorname{IAV}(p)
$$

global variables that may be assigned to (also indirectly) when $p$ is called

- two aux. definitions (straightforwardly defined, obviously flow-insensitive)
- AV $(S)$ : assigned variables in $S$
- $\mathrm{CP}(S)$ : called procedures in $S$

$$
\begin{equation*}
\operatorname{IAV}(p)=(\operatorname{AV}(S) \backslash\{x\}) \cup \bigcup\left\{\operatorname{IAV}\left(p^{\prime}\right) \mid p^{\prime} \in C P(S)\right\} \tag{2.24}
\end{equation*}
$$

where $\operatorname{proc} p(\operatorname{val} x$, res $y)$ is $^{l_{n}} S$ end $^{l_{x}} \in D_{*}$

- $\mathrm{CP} \Rightarrow$ procedure call graph (which procedure calls which one; see example)


## Example

```
begin procfib(val z) is
            if [z<3]
            then [call add(a)]
            else [call fib(z-1)];
                                    [call fib(z-2)]
            end;
            proc add(val u) is(y := y+1;u:= 0)
            end
            y:= 0; [call fib(x)]
end
```


## Example



$$
\begin{aligned}
\operatorname{IAV}(f i b) & =(\emptyset \backslash\{z\}) \cup \operatorname{IAV}(f i b) \cup \operatorname{IAV}(a d d) \\
\operatorname{IAV}(a d d) & =\{y, u\} \backslash\{u\}
\end{aligned}
$$

$\Rightarrow$ smallest solution

$$
\operatorname{IAV}(f i b)=\{y\}
$$

### 2.7 Static single assignment

## Introduction

This section is not covered by the book Nielson et al. [8]. It's added this year as it's an interesting and important angle on data flow analysis. The book covers so-called definition-use or use-definition chains (aka du- and ud-chains), which are left out instead this time. Only so much: du- and du-chains are basically a generalization of reaching definitions, one version working forward (like rd) one backward. And SSA can be seen as a generalization, in turn.

More than one ever wants to know can about SSA can be found in [6].

Origins Early citations are Rosen et al. [11] Alpern et al. [1], and Cytron et al. [4]. So also historically, it's connected to value numbering.

## Intro

- improvement on def-use chains, connecting "definitions" of variables to their uses
- important intermediate representation
- used in many compilers (gcc, go, swift, llvm, ...)
- referential transparency

SSA A program is in SSA form if each variable is a target of exactly one assignment in the program text.

Referential transparency refers to a in generally welcome property of expressions (in this case variables). It means that a value of an expression (or here a variable) is independent of where the expression appears.

The statement should perhaps be qualified in that the value of a variable is always the same (unless there is no value, i.e., the variable is undefined). If we ignore the "being undefined" as special status, then a referentially transparent value means indeed the value of the variable is "always the same" (and that means the value is immutable and the variable is single assignment, and there might be other characterizations as well). It's also related to functional or declarative programming, If one deals with referentially transparent variables, many things become more easy, the concept of variable becomes more logical. Variables in imperative programming languages have the flavor of being names to memory addresses, whereas variables in mathematical text books, equations, or in logical formulas do not give that feeling. The reason being that expressions and variables are intuitvely understood as being referentially transparent (not that any math text would ever point that out, it simply does not cross anybody's mind what a variable has to do with a mutable memory cell ...) Also in the lecture, we carefully try to separate assigmnents $x:=e$ from equations $x=e$, the latter being understood as referentially transparent (or declarative ...). The opposite of referentially transpatent is also known as referentially opaque.

Anyway, one way of understanding the general motivation of SSA is that it's a format that is referentially transparent. One nice property is, that if one has a unique assignment $\mathrm{x}:=\mathrm{e} ; \mathrm{S}$ which is referentially transparent. which we might also write $\mathrm{x}=\mathrm{e} ; \mathrm{S}$ or actually let $\mathrm{x}=\mathrm{e}$ in S . The semantics of the construct is, that first, $e$ is evaluated to some value, then that value is stored in $x$, and finally, the rest $S$ is evaluated. Now, in a single-assignment setting, once $x$ has gotten its value (say $v$ ), then that value of $x$ won't change any more. As a consequence, the variable and the value are "synonymous", $x$ "is" $v$. That captures the most fundamental property what it means to be "equal", the standard mathematical, logical (referentially transparent, declarative ...) meaning of equality:

## two things being equal can be used interchangably.

After evaluating $e$ to $v$, the let $x=v$ in $S$ can be explained as $S[v / x]$, replacing or substituting $x$ by $v$ in $S$. Obviously, such a substitution explanation does not work for general (non-single) assignments $x:=v ; S$ (but for single-assignments it holds).

What relevance does that have for static analysis? Basically, static analysis in the form of this lecture is an automatic logical analysis of (abstract) properties of programs (like variables being live, etc.). The cleaner the program, the easier the analysis, and if the variables of the program behave like logical variables, that may help in a correct analysis. Note that ultimately, of course, the "logical, singleassignment variables" must be mapped to mutable memory cells of a standard vonNeuman architecture or variations thereof. Nonetheless, during the semantic phase, the analysis may profit from a logically clean intermediate representation.

All that may be a bit philosophical. More down to earth is SSA as intermediate representation: it's a format which has some data flow analysis already "built in". If there is only one assignment to each variable, then when using a variable, it's already clear by the identity of the variable "where it comes from", where it was "defined". Because of that, SSA can be seen as a generalization of so-called defuse chains. Since this valuable piece of data flow information is already built into the variables of the SSA, it's a good general starting point for all kinds of more specialized, subsequent analyses. For instance, reaching definitions.

## Example in SLC

## 3AC

```
a := x + y
b}:=\textrm{a}-
a := y + b
b}:=\textrm{x}*
a := a + b
```


## 3AC in SSA

- $x$ and $y$ : input variables, "read only"
- assigned to via initialization, "before" the program

```
a}1:= x + y
b
a}2:=y+\mp@subsup{b}{2}{
\mp@subsup{b}{2}{}}:=\textrm{x}*
a}\mp@subsup{3}{3}{:= a}\mp@subsup{\textrm{a}}{2}{}+\mp@subsup{\textrm{b}}{2}{
```

The transformation for straight-line code is straightforward. Appel [2] states that it's a form of value numbering.

## Basis idea (for SLC)

- later more complex
- for straight line code: simple "rename" the variables, like: use different versions $x_{1}, x_{2}, x_{3}$ for $x$ on the left-hand side.
- some easy data flow analysis needed to get a fitting "versioning" for left-hand sides


## Compare: 3AC (here for expressions)

$2 * a+(b-3)$


## Three-address code

```
t1 = 2 * a
t2 = b - 3
t3= t1 + t2
```


## Compare to 3AC and temporaries

- code generation of 3 AC for straight-line code
- three-address code: linear IR, close to machine code
- restricted right-hand sides
- temporaries: to store intermediate results
- often
- temporaries $=$ abstract form of register
- unboundedly many asssumed
$\Rightarrow$ each one assigned to only once
For the participants of the compiler construction course (INF5110), this comparison with the generation of 3 AC should be familiar. But also without having participated, the idea is simple. Complex expressions are not supported by standard 3AC (as it's supposed to be close to machine code). That means they need to be broken into pieces and intermediate results need to be stored somewhere. For that purpose the compiler (at that stage) introduces "special" variables, special at least in the sense, that they don't show up in the source code. Otherwise, they are not too special anyway.

Since the compile can generate a new temporary when needed and since no upper bound on their number is assumed, that leads to a code in static single assignment form, at least as far as assignment to temporaries is concerned.

Static single assignment just pushes that idea a bit further and makes sure that also ordinary variable are assigned to once, only (at least "statically").

If we were sticking to straight-line code, without conditional jumps, conditionals, loops etc., everything would be rather simplistic and straightforward. The problem starts with conditionals. That will be treated next, and leads to the introduction of so-called $\Phi$-functions.

## Join points and phony functions

- simple illustration: two "definitions" of $x$


1. Phony functions $\Phi$ Assignments using functions like $\Phi\left(x_{1}, x_{2}\right)$ placed judiciously at (join) nodes to assure SSA format.

## SSA in a nutshell

## Transformation to SSA

- $\operatorname{SSA}=\Phi+$ variable (re)naming scheme


## Phony functions

- "non-standard" function
- encodes "control flow": value depends on if program "came from the left or from the right" in the last step
- $\Phi$
- "virtual", for purpose of analysis only, or
- ultimately "real", i.e., code for $\Phi$ 's will be generated


## 2 phase algorithm(s), in this order

1. strategical placement of $\Phi$-functions
2. renaming of variables

- main challenge: placement of $\Phi$


## Brainless SSA form

- place $\Phi$ "everywhere"


## Maximal SSA recipe

Placement: For all variables, at the beginning of each join block add

$$
x \leftarrow \Phi(x, \ldots, x)
$$

where number of $x$ 's is the number of predecessors of the node $\geq 2$

Renaming: rename variables consistently (making use of reaching definition analysis)

- note: over-generous placement
- guarantees single-assignment format
- is sound


## Room for improvements

- phony functions everywhere: sound but wasteful and generally undesirable
- costly extra computations
- subsequent analyses may suffer loss of precision
- phony function unnessessary
- result not live
- no real choice: $x_{i} \leftarrow \Phi\left(x_{j}, x_{j}\right)$
- in the following: efficient general algo for $\Phi$-placement based on
dominance


## Improvement

Core idea Assume assignment $x:=e$ in $n_{1}$. If all paths from $n_{0}$ to $n_{2}$ must go through $n_{1}$, then $n_{1}$ 's assignment to $x$ does not need to be covered by a phony function for $x$ at $n_{2}$.

## Rest



## Domination

- CFG: directed graph with 1 entry (and 1 exit)
- "content" of the nodes / basic blocks irrelevant right now

Domination Node $n_{1}$ dominates $n_{2}$, if all paths from entry $n_{0}$ to $n_{2}$ must pass through $n_{1}$.

- $\operatorname{dom}(n)$ :dominators of $n$ (" $n$ being dominated")
- $\operatorname{dom}(n)$ : determined by a simple data flow analysis (must and forward)
- also: strict dominance


## Dominance frontier

- we know, where not to put $\Phi$ for a given assignment at $n=$ all nodes being dominated by $n$
- danger zone: "undominated", but where to put there?
- solution: as "early" as possible in the danger zone (thinking backwards)

Dominance frontier The dominance frontier $d f(n)$ of a node is the collection of $m$ s.t.:

1. $n$ dominates a predecessor of $m(q \rightarrow m$ and $q \in \operatorname{dom}(n))$, and
2. $n$ does not strictly dominate $m$.

- "strict" (non-)dominance condition \& loops
- dominance frontier contains join nodes only


## Dominator trees

- $n$ 's dominators: $\operatorname{dom}(n)$
- $n$ 's immediate dominator $\operatorname{idom}(n)$ (the dominator closest to $n$, if exists)



## Dominance frontier

```
for all nodes }n\mathrm{ in the CFG
    DF(n) :=\emptyset
for all nodes n
    if }\quadn\mathrm{ has multiple predecessors
    then for all predecessors p of n
        runner := p
        while runner }\not=i\operatorname{dom}(n
        do }\quad\textrm{DF}(\mathrm{ runner ) := DF(runner) }\cup{n
        runner := idom(runner)
```

The algo calculates, for each node, the dominance frontier $d f(n)$. A node dominating another is a node "in the past", but the dominance frontier works forwardly, "to the future", calculating for each node, those other in $\rightarrow{ }^{*}$ that are being dominated, or at least almost so. So, the nodes of $d f(n)$ are not being dominated by $n$, i.e., it's not $n_{2} \in d f\left(n_{1}\right)$, then $n_{1}$ dominates $n_{2}$, it's that $n_{1}$ does not dominate $n_{2}$, but almost does, in that its (immediate) predecessor is being dominated. So, the dominator front describes some form of "front" for $n$ but just one step beyond the front into the undominated territory. Remember that being dominated is the "safe" zone in that it does not need $\Phi$-functions, whereas the undominated territory is the "danger zone" where $\Phi$ functions are necessary (for join nodes). And the frontier, one step beyond the border, is the place of the earliest convenience, where to put the $\Phi$.

Now to the algorithm. The frontier is calculated for all nodes which have more than one predecessor ( $\Phi$-functions are needed for join-nodes only). The nodes are treated one by one.

The algo is simple enough, and needs both the original graph as well as the immediate domintor tree. First one takes all predecessors in the CFG of the chosen node (and there has to be more than one, as we concentrate on join nodes). From each of those predecessors, we walk up the dominator tree. That walk is "deterministic", as we use the dominator tree upwards. Note that we are not only following the tree upwards, it's also like following the original graph backwards, only not step by step, but one dominator-edge may correspond to $\rightarrow^{+}$. In case of a loop in the program, the predecessor of $n$ used as a starting point is, of course, both a predecessor as well as a successor of $n$ (that's what makes it a loop). Therefore, following the dominator tree upwards ends up where the runner is immediatedly domminated by $n$ (that's the while-loops exit condition). In that loop-situation, runner is a successor of $n$ as well as a predecessor to $n$. Then we add $n$ to all the "runners" (not the other way around). Note also that we climb up the tree until the runner equals the immidate dominator. In case of a loop, that will lead to the situation that the entry-point of a loop dominates itself (which is consistent with the definition, in particular the second part which mentions strict dominance. That may be seen in the algo applied to $n_{1}$. Note again that applying the inner loop of the algo to a node do not contribute by calculating the dominator front for $n$, but rather the other way around: adding $n$ to other nodes (represented by runner).

## Example

$$
\begin{array}{cccccccccc} 
& n_{0} & n_{1} & n_{2} & n_{3} & n_{4} & n_{5} & n_{6} & n_{7} & n_{8} \\
D F & \emptyset & \left\{n_{1}\right\} & \left\{n_{3}\right\} & \left\{n_{1}\right\} & \emptyset & \left\{n_{3}\right\} & \left\{n_{7}\right\} & \left\{n_{3}\right\} & \left\{n_{7}\right\}
\end{array}
$$

## Further improvement

- left out here:
- how to actually place the $\Phi$
- how to then rename the variables
- Basically: problem solved: place it at the dominance frontier
- we forgot: adding $x_{i} \leftarrow \Phi\left(x_{j}, x_{j}\right)$ is another assignment ...
- further improvements possible (liveness: focus on "global" names)


## Renaming

```
rename (b)
    for each }\Phi\mathrm{ -function `` }x\leftarrow\Phi(...) in 
        rewrite x as newname(x)
    for each operation }x\leftarrowy\oplus
        rewrite y with subscript top(stack[y])
        rewrite z with subscript top(stack[z])
        rewrite }x\mathrm{ as newname(x)
    for each successor of b in the CFG
        full in \Phi-function parameters
    for each successor s of b in the dominator tree
        rename s
    for each operation '` }x\leftarrowy\oplus\mp@subsup{z}{}{\prime\prime}\mathrm{ ' in }
        and each }\Phi\mathrm{ -function ``}x\leftarrow\Phi(\ldots)''
            pop(stack[x])
```


## Learning Targets of this Chapter

type systems
effects
functional languages
type inference and unification

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### 3.1 Introduction

In this part, we cover various type systems, but focussing clearly on the effects parts. In the introduction, we used the while-language to illustrate type and effect systems or annotated type systems in a very simple setting, where the typing part was trivial. In this section, we deal mainly with functional languages, i.e., $\lambda$-calculi.
in 2107 , we only came up-to the standard type inference part.

### 3.2 Control flow analysis

## Syntax

$$
\begin{aligned}
& e::= c|x| \mathrm{fn}_{\pi} x \Rightarrow e\left|\mathrm{fun}_{\pi} f x \Rightarrow e\right| e e \quad \text { terms } \\
& \quad \mid \quad \text { if } e \text { then } e \text { else } e \mid \text { let } x=e \text { in } e \mid e \text { op } e
\end{aligned}
$$

Table 3.1: Abstract syntax

| $\pi$ | $\in \mathbf{P n t}$ | program points |
| ---: | :--- | :--- | :--- |
| $e$ | $\in$ Expr | expressions |
| $c$ | $\in$ Const | constants |
| op | $\in \mathbf{O p}$ | operators |
| $f, x$ | $\in$ Var | variables |

The syntax is a a variation of an (untyped) $\lambda$-calculus, the prototypical "functional language". Instead of " $\lambda$ " as symbol for functional abstraction, the language here uses fn and fun. The language distinguishes syntactivally between the "standard" abstraction $f n$ and abstraction for recursive functions. Standard representations of the untyped $\lambda$-calculus would not bother to make that distinction: the untyped $\lambda$-calculus is Turing complete and therefore expressive enough to encode recursion. Likewise it could encode conditionals etc. However, the lecture uses the more elaborate syntax for two reasons. We are interested in illustrating program analysis, where control flow constructs like conditionals are most likely to be part of the abstract syntax. Even if conditional would be encodable in a more basic syntax, that will obscure the analysis (and make it less "realistic"). Likewise, recursion will pose specific problems, the treatment of which would be obscured if relying on an encoding. Finally, we do make use of typed versions of the language, and for the typed $\lambda$-calculus, things change: without adding recursion explicitly, the calculus is not Turing complete (which make it hard to argue that it's a simple form of a standard functional programming language).

What does not belong to the standard calculus are the labels. Those are needed (as before) for the intended analysis. It's worthwile to remember the CFA from the introduction, where the functional calculus was labelled, as well. Unlike here, the labelling in the introduction was for all constructs whereas here, we label only the two forms abstractions. The difference are motivated by the fact that here we are interested in a slightly different analysis.

## Examples

Example 3.2.1 (Application).

$$
\left(\mathrm{fn}_{X} x \Rightarrow x\right)\left(\mathrm{fn}_{Y} y \Rightarrow y\right)
$$

Example 3.2.2.

$$
\begin{array}{ll}
\text { let } g & =\left(\operatorname{fun}_{F} f x \Rightarrow f\left(\mathrm{fn}_{Y} y \Rightarrow y\right)\right) \\
\text { in } & g\left(\mathrm{fn}_{Z} x \Rightarrow x\right)
\end{array}
$$

It's worthwile to think about the latter example and especially the role of fun (the "recursive" function abstraction). Its

## Types

- Curry-style typing

$\tau \in$ Type types<br>$\Gamma \in$ TEnv type environment

## Types

$$
\tau::=\text { int } \mid \text { bool } \mid \tau \rightarrow \tau
$$

- base types:
- bool and int
- standard constants and operators assumed (true, $5,+, \leq, \ldots$ )
- each constant has a base type $\tau_{c}$
- type environments (finite mappings)

$$
\Gamma::=[] \mid \Gamma, x: \tau
$$

"Curry-style" means that abstractions like $\mathrm{fn} x . e$ don't mention a type for the formal parameter. The alternative with a syntax like $\mathrm{fn} x: \tau . e$ is called "Church style". Both variants exist in real programming languages. Also, leaving it up to a user whether mentioning a type or leaving it out on a case by case basis is possible. Some programmers may prefer to add types for being more explicit and documenting the type of expected arguments and prefer to leave it out, for compactness or convenience In Curry-style typing, it's a problem to figure out what the type of an argument actually is (if any). The process of figuring that out is commonly known as type inference; some people prefer the word type reconstruction for it. Some other interpretations of the terminology of "type inference" also exists, but they are not so common. In case giving the types is optional, the problem is called partial type inference (or partial type reconstruction).

The base types won't play a prominent role in the development, the calculus simply picks some common ones, where Booleans are needed for the conditionals. There is one restriction for the constants here, namely they are typed by one of the base types. That means there are no functional constants. More precisely: the operators can be seen as built-in functional constants, but their arguments are base types, which means, the operators are not higher-order. That may well be different in real languages, and is done to simplify the representation here.

As for type environments $\Gamma$. They play the role of "contexts" (also in the technical sense of dealing with context-sensitive analyses). It is assumed that they work as finite mappings, basically as a representation of the symbol table. Three fundamental things one can do with such a "table" is: creating an empty one, adding a new binding ("enter"), and looking up an entry ("lookup"). The first two operations are part of the definition from above, for the lookup of $x$ in $\Gamma$ we write $\Gamma(x)$. We may also write $\Gamma[x \mapsto \tau]$, and use $\operatorname{dom}(\Gamma)$. $\Gamma$ in general acts like a stack.

## Judgments and derivation system

## Type judgments

$$
\begin{equation*}
\Gamma \vdash \vdash_{\mathrm{UL}} e: \tau \tag{3.1}
\end{equation*}
$$

- derivation system:
- Curry-style formulation
$\Rightarrow$ non-deterministic
- nonetheless: monomorphic let
- type reconstruction/type inference

The remark about monomorphic let is relevent at that point only for the ones who know, that a very famous contribution in type inference is the treatment of the socalled "polymorphic let". Polymorphism and monymorphim relate to the character of the type system for a language. A language is monomorphic if each program has (at most) one type. It's at most and not exactly one as a program may fail to be well-typed. Alternatively can can say, a language is monomorphic, if every well-typed program has exactly one type. A language is polymorphic, if it's not the case, i.e., if it's not monomorphic.

There are various forms of polymorphism, according to a classical classification by Cardelli and Wegner [3], there are 4 main variants. Arguably the two most important or most interesting are parametric polymorphism and inclusion polymorphism (both together called in [3] also as universal polymorphism). Inclusion polymorphism is also known as subtype polymorphism (as made popular by object-oriented languages).

Let-polymoprhism is of the parametric kind, and we might encounter it later. But as said: not right now. Subtype or inclusion polymorphism will probably not be covered in this lecture, at least not as far as the underlying type system is concerned. On the other hand: when dealing with the "non-standard" part of the type system (the annotations, the effects etc.) then there will be "inclusion polymorphism". That is related to the "lattice treatment" for instance when dealing with data flow information as in the monotone frameworks (except that it was not formulated making use of type-theoretic notions).

## Underlying type system (1) (Curry style)

$$
\begin{aligned}
& \Gamma \vdash c: \tau_{c} \quad \text { Con } \quad \frac{\Gamma(x)=\tau}{\Gamma \vdash x: \tau} \mathrm{VAR} \\
& \frac{\Gamma \vdash e_{1}: \tau_{\mathrm{op}}^{1} \quad \Gamma \vdash e_{2}: \tau_{\mathrm{op}}^{2}}{\Gamma \vdash e_{1} \mathrm{op} e_{2}: \tau_{\mathrm{op}}} \mathrm{OP}
\end{aligned}
$$

## Underlying type system (2) (Curry style)

$$
\begin{aligned}
& \frac{\Gamma, x: \tau_{1} \vdash e: \tau_{2}}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e: \tau_{1} \rightarrow \tau_{2}} \mathrm{FN} \quad \frac{\Gamma, x: \tau_{1}, f: \tau_{1} \rightarrow \tau_{2} \vdash e: \tau_{2}}{\Gamma \vdash \mathrm{fun}_{\pi} x \Rightarrow e: \tau_{1} \rightarrow \tau_{2}} \mathrm{FUN} \\
& \frac{\Gamma \vdash e_{1}: \tau_{1} \rightarrow \tau_{2} \quad \Gamma \vdash e_{2}: \tau_{1}}{\Gamma \vdash e_{1} e_{2}: \tau_{2}} \mathrm{APP} \\
& \frac{\Gamma \vdash e_{0}: \text { bool } \quad \Gamma \vdash e_{1}: \tau \quad \Gamma \vdash e_{2}: \tau}{\Gamma \vdash \text { if } e_{0} \text { then } e_{1} \text { else } e_{2}: \tau} \mathrm{IF} \\
& \frac{\Gamma \vdash e_{1}: \tau_{1} \quad \Gamma, x: \tau_{1} \vdash e_{2}: \tau_{2}}{\Gamma \vdash \text { let } x=e_{1} \text { in } e_{2}: \tau_{2}} \text { LET }
\end{aligned}
$$

### 3.2.1 Control flow analysis

## Intro

This part was not covered in the lecture 2017. We directly jumped to the unification and standard type inference part. A flavor of similar analysis, though, was covered in the introduction.

## Control flow analysis

- remember introduction: CFA touched 2 times
- constraint analysis
- as effect-system ("call-tracking")
- goal CFA (general): "which functions may be applied in an application" more precisely:

CFA to which function abstractions may an expression evaluate to

- augment or annotate the type system with effects
- note: data "=" control here

It is worthwhile to compare the control flow analysis presented next with the treatments in the introduction. Also in the simple while-language, to determine the edges of the control-flow graph, we called a simple form of control flow analysis, but normally, no-one calls it like that, as it's so simple and it does not require any complex techniques.

More seriously, control flow analysis in the introduction was touched upon 2 times in the context of a functional langauge. At one point, the problem of control flow analysis (in the section about constaint based analysis) was defined as the question "for each function application, which functions are potentially applied". That's of course related to the formulation of CFA from above.

The second time an analysis related to control-flow analysis was touched upon in the introduction was the call-tracking analysis. There the question was for each expression "which function abstraction may be applied during execution". That was formulated as effect analysis. It's worthwhile to compare the rules presented next to the ones from earlier. They are quite similar, as call tracking analysis is not much different to the cfa problem here. Note, however, the analysis here is not an effect system, we are not interested what happens during evaluation, as is done for the call-tracking analysis.

## Annotations

labelled abstraction $\mathrm{fn}_{\pi}$ and fun $_{\pi}$
annotations: set of function names,

$$
\begin{array}{rlll}
\varphi & \in & \text { Ann } & \text { annotations } \\
\hat{\tau} & \in & \text { Type } & \text { annotated types } \\
\hat{\Gamma} & \in & \text { TÊnv } & \text { ann. type environments } \\
& & \varphi::=\{\pi\}|\varphi \cup \varphi| \emptyset \\
& \hat{\tau} \quad::= & \text { int } \mid \text { bool } \mid \hat{\tau} \xrightarrow{\varphi} \hat{\tau} \\
& & \hat{\Gamma}::= & {[] \mid \hat{\Gamma}, x: \hat{\tau}}
\end{array}
$$

Erasure to underlying type system:

$$
\lfloor\hat{\tau}\rfloor \quad\lfloor\hat{\Gamma}\rfloor
$$

As for the call-tracking analysis in the introduction, the functions, i.e., the abstractions, get labelled or annotated to make them identifiable and the analysis will deal with sets $\varphi$ of such labels. The arrow types are annotated accordingly (as was the case for the call tracking analysis from before).

So, both the call tracking analysis and the control flow analysis here are operating with annotated types of the form

$$
\hat{\tau}_{1} \xrightarrow{\varphi} \hat{\tau}_{2} .
$$

The interpretation, however, of these types are different! Here, the $\varphi$ on the arrow constructor approximates the set of functions that the corresponding expression can evaluate to. That refers to the values at the end. For the call tracking analysis, $\varphi$ on the arrow type was called latent effect and approximated the set of functions
potentially being called during an evaluations of that function. The judgments of the call-tracking analysis where more complex as well, as they contained also an effect component (the "typing part" being of the form $\hat{\tau}:: \varphi$ ). The analysis here is not an effect analysis.

One may compare the rule for applications in both analyses.

## Annotated type system

$$
\begin{aligned}
& \hat{\Gamma} \vdash c: \tau_{c} \quad \operatorname{CoN} \quad \frac{\hat{\Gamma}(x)=\hat{\tau}}{\hat{\Gamma} \vdash x: \hat{\tau}} \mathrm{VAR} \\
& \frac{\hat{\Gamma} \vdash e_{1}: \tau_{\mathrm{op}}^{1} \quad \hat{\Gamma} \vdash e_{2}: \tau_{\mathrm{op}}^{2}}{\hat{\Gamma} \vdash e_{1} \text { op } e_{2}: \tau_{\mathrm{op}}} \mathrm{OP} \\
& \frac{\hat{\Gamma} \vdash e_{0}: \text { bool } \quad \hat{\Gamma} \vdash e_{1}: \hat{\tau} \quad \hat{\Gamma} \vdash e_{2}: \hat{\tau}}{\Gamma \vdash \text { if } e_{0} \text { then } e_{1} \text { else } e_{2}: \hat{\tau}} \mathrm{IF} \\
& \frac{\hat{\Gamma} \vdash e_{1}: \hat{\tau}_{1} \quad \Gamma, x: \hat{\tau}_{1} \vdash e_{2}: \hat{\tau}_{2}}{\hat{\Gamma} \vdash \text { let } x=e_{1} \text { in } e_{2}: \hat{\tau}_{2}} \text { LET } \quad \hat{\hat{\Gamma} \vdash \mathrm{fn}_{\pi} x \Rightarrow e: \hat{\tau}_{1} \xrightarrow{\{\pi\} \cup \varphi} \hat{\tau}_{2}} \text { FN } \\
& \frac{\Gamma, x: \hat{\tau}_{1}, f: \hat{\tau}_{1} \xrightarrow{\{\pi\} \cup \varphi} \hat{\tau}_{2} \vdash e: \hat{\tau}_{2}}{\Gamma \vdash \mathrm{fun}_{\pi} x \Rightarrow e: \hat{\tau}_{1} \stackrel{\{\pi\} \cup \varphi}{\rightarrow} \hat{\tau}_{2}} \text { FUN } \\
& \hat{\Gamma} \vdash e_{1}: \hat{\tau}_{1} \rightarrow \hat{\tau}_{2} \quad \hat{\Gamma} \vdash e_{2}: \hat{\tau}_{1} \\
& \frac{\hat{\Gamma} \vdash e_{1} e_{2}: \hat{\tau}_{2}}{} \text { APP }
\end{aligned}
$$

The interesting rules are the ones for functions, i.e., for dealing with the two forms of abstractions and with function application.

For Fn, for instance: the rule involves "guessing", insofar that in the premise an arbitrary $\varphi$ is mentioned! As a consquence, the rules are not interpretable as a straightforward as algorithm. Another aspect which makes the rules non-algorithmic has to do with the fact that the abstractions don't mention the type $\hat{\tau}_{1}$ of the formal parameter (as we are dealing with a Curry-type formulation). This has nothing directly to do with the control-flow analysis, but was already the case for the underlying type system (without annotations).

Anyway, the rules of this (and similar) systems cannot be seen as type checking algorithm, but rather as specification of the analysis. How to algorithmically check and analyse a given program is a separate question.

## Example

$$
\frac{\frac{x: \hat{\tau}_{Y} \vdash x: \hat{\tau}_{Y}}{\vdash\left(\mathrm{fn}_{X} x \Rightarrow x\right): \hat{\tau}_{Y} \xrightarrow{\{X\}} \hat{\tau}_{Y}} \quad \frac{y: \text { int } \vdash y: \text { int }}{\vdash\left(\mathrm{fn}_{Y} y \Rightarrow y\right): \hat{\tau}_{Y}}}{\vdash\left(\mathrm{fn}_{X} x \Rightarrow x\right)\left(\mathrm{fn}_{Y} y \Rightarrow y\right): \hat{\tau}_{Y}}
$$

with

$$
\hat{\tau}_{Y}=\operatorname{int} \xrightarrow{\{Y\}} \text { int }
$$

## Equivalence of annotations

- annotations $\varphi$ are considered as sets
- one could axiomatise this (UCAI)
- i.e., one could import equality on sets into equality on types:

$$
\begin{gathered}
\hat{\tau}_{1}=\hat{\tau}_{1}^{\prime} \quad \hat{\tau}_{2}=\hat{\tau}_{2}^{\prime} \quad \varphi=\varphi^{\prime} \\
\hat{\tau}_{1} \xrightarrow{\varphi} \hat{\tau}_{2}=\hat{\tau}_{1}^{\prime} \xrightarrow{\varphi^{\prime}} \hat{\tau}_{2}^{\prime}
\end{gathered}
$$

- types (and $\hat{\Gamma}$ 's) are considered only modulo this equality

Equivalences seems like a minor or obvious point, but: it can give serious technical headaches when we go for an algorithm (type reconstruction or inference)

The discussion around the treatment of $\varphi$ seems slightly obscure. The $\varphi$ s are intended as finite sets and it seems clear enough, when two sets are equal (namely when they have the same elements), end of story.

That would be indeed ok, especially for now. Later, however, we will deal with techniques to algorithmically do type checking resp. type inference and also do type inference in combination with annotated types. One core technique there will involve unification. That works straightforward if one unifies "terms" (the standard unification is understood as "term unification"). However, if one wants to use unification on things that are more complex than terms, things can get hairy. Sets are more complex as non-trivial equivalences hold, like $\{1,2\}=\{2,1\}$. Sets are still comparatively simple (a few well-understood, simple equivalences: commutativity, associativity, identity, unit), so that unification still works fine ("unification modulo set equivalences"), it even has a name: UCAI-unification. But as said, the sets as the information of interest is quite simple, if one tries to figure out more complex stuff in the analysis, corresponding unification may no longer work (one may even end up with an undecidable problem).

Therefore, the book follows a slightly different route (later), not relying on unification modulo some equivalences.

## Underlying vs. annotated type system

- desired relationship between the original type system and the annotated one:
$\Rightarrow$ The annotation does not "disturb" the original one
- conservative extension
- note:
- type systems reject programs
- flow analysis and similar: typically don't reject, just analyse


## Fact

- if $\hat{\Gamma} \vdash_{\text {CFA }}: e: \hat{\tau}$ then $\lfloor\hat{\Gamma}\rfloor \vdash_{\mathrm{UL}} e:\lfloor\hat{\tau}\rfloor$
- if $\Gamma \vdash_{\mathrm{UL}} e: \tau$, then $\hat{\Gamma} \vdash_{\mathrm{CFA}} e: \hat{\tau}$ for some $\hat{\Gamma}$ and $\hat{\tau}$ s.t. $\tau=\lfloor\hat{\tau}\rfloor$ and $\Gamma=\lfloor\hat{\Gamma}\rfloor$.


### 3.2.2 Correctness

Also this section is left out 2017, at least the technicalities. But it's good to have a feeling of what guarantees a typing judgment $\Gamma \vdash e: \tau$ is intended to mean (i.e., what type safety means).

## Semantic correctness

- as always: the analysis as (over)approximation
- correctness formulated here as subject reduction
- assume : typing for op is properly given

Theorem 3.2.3. If [] $\vdash_{\text {CFA }} e: \hat{\tau}$ and $\vdash e \longrightarrow v$, then []$\vdash_{\text {CFA }} v: \hat{\tau}$

The name "subject reduction" is often used for this kind of results (in type systems). There is a (historical?) reason why such results are called subject reduction, but it does not matter for us and the word does not provide much inside.

A better term for the theorem would be preservation of well-typedness under reduction. Indeed, sometimes the result is also called like that, or just preservation. Anyway, it's a very central lemma, basically justifying the term "static typing". Why is that? The initial the state of a program is well-typed by definition, as the compiler allows only well-typed programs to run. If one has "subject reduction" it means, well-typedness is preserved when doing one step (well, in our case it's a "big step"). Both facts together immediately imply that all configurations reachable from the initial one are well-typed.

Since, once established at the beginning, well-typedness is preserved, there's no need to check it over and over again when running the program. In other word: there is no need for run-time type checking! Hence the word static typing.

While being central for static type systems, subject reduction in isolation does not guarantee "correctness" or that in general the type system does anything meaningfull. For illustration, a trivial analysis, where all programs are well-typed (the type would correspond to the logical "true") would enjoy subject reduction. The same holds for the type system that rejects all programs (corresponding to "false"). The latter one lead to the fact that the initial program is ill-typed as well, therefore the "base case" for all reachable configurations is not met. Anyway, both are cases of rather useless type systems that do enjoy subject reduction.

The missing piece that make a type system "meaningfull" wrt. to a semantics is to connected it to the absence of the errors one intends to target (or the things one wishes to analyse). Informally speaking the connection between the type system and the freedom-of-error is something like "a well-typed program does not exhibit an error right now". Now if all reachable programs are indeed well-typed, then obviously the program is error-free.

The "right-now" could mean, the next step does not raise an error (resp., if there is non-determinism: none of the possible next steps raise an error) or alternatively: the current configuration is not "erroneous".

By speaking of error-freeness, of course we don't mean the program is absolutely error free. Free of errors refers only to the errors the type system is designed to catch. For standard type systems (not effects, or other fancy stuff), the type system refers to values at the end of a computation, and consequently, errors are of the kind "wrong data value" (like a string where an int is expected etc.)

In some contexts and for traditional reasons mostly: the absence-of-error-now result is sometimes called progress. The intuition being the following: The operational semantics describes error-free behavior, there are no rules describing what happens if there are run-time type errors (or other errors). So, one way of dealing with "errors" would be simple not to put in a rule that describes what happens in that case. The operational semantics simply concentrates on the "positive" side of things. As an example, the rules don't describe what happens of we apply an integer to a boolean value:

## 5 true

In reality, probably some run-time error would occur (hopefully at least, that would be better than some random behavior). However, there is not SOS rule applying to that situation, which means the above application is stuck (there is no "progress"). Of course, values are "stuck" in that sense too, but legitimately so: values are those expressions which are intended to be the end of a computation. The above application is not a value, and lack of progress there is an indication of an error. So, connecting the type system to the intended absence of errors would be something like: "a well-typed expression is either a value or can done one more step (progress)". In the type system, it is immediately clear 5 true is ill-typed ("now"). Less clear is wether an well-typed expression can ever reach such an ill-typed (and stuck)
expression, so it refers to a future error, not an error now. For that, one needs subject reduction.

That describes in some detail the connection between (absence of) errors and the type system for standard type systems. For the non-standard type system here (i.e., the annotation part dealing with control-flow information), the question we want to ask is: given $\vdash e: \hat{\tau}_{1} \xrightarrow{\varphi} \hat{\tau}_{2}$ and $e$ reduces to an abstraction, then it is immediate to see that the label of the abstraction must be in $\varphi$.

One key lemma for subject reduction is preservation under substitution.
Lemma 3.2.4 (Substitution). Assume [] $\vdash_{\text {CFA }} e_{0}: \hat{\tau}_{0}$ and $\hat{\Gamma}\left[x \mapsto \hat{\tau}_{0}\right] \vdash_{\text {CFA }} e: \hat{\tau}$. Then $\hat{\Gamma} \vdash_{\text {CFA }} e\left[x \mapsto e_{0}\right]: \hat{\tau}$

## Semantics correctness \& subject reduction

- subject reduction: standard name for key to correctess (aka type safety) in static type systems (here type and effects)

Goal ("Milner's dictum") A well-typed program cannot go wrong.

- goal a bit more technically: no "erroneous" state is reachable, starting from a/the initial state
- erroneous state: a state where a run-time type error "manifests" itself
- wrong arguments to a function
- data stored in variable not declared/dimensioned to hold that kind of data
- "method not supported" error
- ...


## Type safety

## Type safety: 3 easy pieces

- Induction: all reachable "states" are well-typed
base case initial state is well-typed
induction step Well-typedness is preserved under doing a step (= subject reduction)
- a well-typed state is not erroneous at that point
- base case trivial/by assumption: only well-typed programs are run
- since well-typing is preserved: no run-time type checks needed (efficiency, static typing)
- with effects: subject reduction $=$ simulation (however: CFA here no effects yet)

As mentioned earlier, the key to correctness of a type system, i.e., type safety, is subject reduction. Type safety is a statement over all reachable states of a program, no matter whereto the program runs, nothing bad will happen. That leads to an inductive proof, where the induction step is known as subject reduction (or with a more understandable word preservation of well-typedness under reduction.

## Complete lattice of annotated types

- to assure existence of solutions

$$
(\mathrm{Ann}, \sqsubseteq) \quad\left(\simeq\left(2^{\mathrm{Pnt}}, \subseteq\right)\right)
$$

- write $\operatorname{Type}(\tau)$ : set of $\hat{\tau}$ 's s.t. $\lfloor\hat{\tau}\rfloor=\tau$

$$
\begin{array}{cc}
\frac{}{\hat{\tau} \sqsubseteq \hat{\tau}} \quad \frac{\hat{\tau}_{1} \sqsubseteq \hat{\tau}_{1}^{\prime} \quad \hat{\tau}_{2} \sqsubseteq \hat{\tau}_{2}^{\prime} \quad \varphi \sqsubseteq \varphi^{\prime}}{\hat{\tau}_{1} \xrightarrow{\varphi} \hat{\tau}_{2} \sqsubseteq \hat{\tau}_{1}^{\prime} \xrightarrow{\varphi^{\prime}} \hat{\tau}_{2}^{\prime}} \\
\Rightarrow \quad(\mathbf{T y p e}(\tau), \sqsubseteq) \text { is complete lattice }
\end{array}
$$

### 3.2.3 Type inference

That was one of the sections covered 2017. Basically, we left out all effect or flow parts, and covered only the basics here: type inference or reconstruction and the role unification plays in that.

## Inference algorithms

- take care of terminology
- so far: no algorithm! (price of laxness)
- foresight needed
- guessing wrong $\Rightarrow$ backtracking (and we seriously don't want that)
$\Rightarrow$ required: mechanism to make
- tentative guesses
- refine guesses
- we start first: with the underlying system

Next we tackle what is called type inference. The terminology of type inference is standard (it goes back to Milner etc), but not all like it. The point why it's not the best name is: the type systems we are dealing with are given in the form of inference systems i.e. with the help of inference rules. Therefore the word "type inference" may be confusing. Consequently, the word type reconstruction is used sometimes.

No matter how it's called, it's here about algorithms. The type system so far did not correspond directly to any algorithm for type checking (or flow or effect analysis)

It is useful here to revisit the previous rules of the "Curry-style" type system and reflect on why those rules are not directly an algorithm. The basic culprit is the rule for abstraction FN. In the formulation here, where we have also a rule for recusive function abstraction, also that rule is not "algorithmic" (rule Fun). Concentrating here on Fn: the culprit is the fact that the premise of the rule has to guess type $\tau_{1}$. What is the status of $\tau_{1}$ ? Well, $\tau_{1}$ is a symbol we use to represent types (a non-terminal in the grammar for types). It's not a type itself. Another terminology is that it's a meta-variable representing types. That $\tau$ is a meta-variable for types means, we use that symbol to "speak about" the type system, but it's of course not itself a type. Especially it's not a variable in or of the type system.

Another way of seeing it is: assuming that the type rules where an inmpementation in the sense of a (non-deterministic) recursive procecure. The conclusion would be a call to a function, say tcheck (for do-a-type-check), with input a context, an expression, and a type) and a boolean return. That's a simplifying assumption, as we typically want to have the context $\Gamma$ and expression $e$ as input, and the type as output, but let's use the simpler "boolean" yes/no problem for illustration. That would mean, the type-check procedure could look like (in some unspecified programming language):

```
tcheck(gamma : Context, exp : Exp, type : Type) =
    if exp = app (exp1 exp2 ) ....
```

Now, type is a variable in the unspecified programming language, the formal parameter of the procedure, and is of type Type, which, one the one hand, is some concrete type in that said programming language and, on the other hand, is used to implement the types of the language whose type system we intend to implement. At any rate, type is a variable, but not of the language we are implementing, but of the language we use to implement it. Thus it's a meta variable, a variable of the language use to implement the target language (or the language used to "speak about" the target language).

One core "enabler" to make type inference work is to internalize the notion of "variable" from the meta-level to the level of the language we are dealing with. That means that the concept of types needs to be extended to include type variables. That will be called augmented types.

## Augmented types

fancy name for: "we have added type variables"

$$
\begin{aligned}
\tau & \in \text { AType } \\
\alpha & \in \text { TVar } \quad \text { augmented types } \\
\tau & ::=\text { int } \mid \text { bool }|\tau \rightarrow \tau| \alpha \\
\alpha & ::={ }^{\prime} \mathrm{a} \mid \text { 'b } \mid \ldots
\end{aligned}
$$

## Substitutions

Substitution (in general) mapping from variables to "terms"

- "syntactic mapping" here:
- "terms" are (augmented) types
- variables: type variables

$$
\theta: \text { TVar } \rightarrow_{f i n} \text { AType }
$$

- considered as finite functions: we write $\operatorname{dom}(\theta)$.
- ground substitution: mapping to ordinary types (no variables)
- substitutions: lifted to types in the standard manner
- composition of substitutions: $\theta_{1} \circ \theta_{2}$ (or just $\theta_{2} \theta_{1}$ )

Substitutions will play an important role in the following (and are an important concept in general). They are called "syntactic mapping" above, since terms (here types) are considered syntax (as opposed to values or similar, which is considered "semantics"). So, a state as a mapping from variables to values is considered a semantic thing and different from a substitution. Terms normally contains variables (otherwise there would be no point of substitutions anyhow ...) and terms without variables are generally called ground terms. Consequently, a substitution where the result does not contain variables is called a ground substitution.

By lifting, one simply means; if one knows the effect of a substitution on variables, then it's straightforward to use the substitution also as a mapping from terms to terms (here types to types), simply by replacing all the variables inside a term one by one. It's a simple recursive algorithm.

Above we defined substitutions as finite functions, i.e., function with a finite domain of variables. Sometimes, $\theta$ 's are also considered as total functions (over "all" variables), setting $\theta(\alpha)=\alpha$ when $\alpha \notin \operatorname{dom}(\theta)$.

## Algorithm: basic idea

- instead of guessing type /now/ $\Rightarrow$ postpone the decision
$\Rightarrow$ use of *type variables*
replace:

$$
\frac{\Gamma, x: \tau_{1} \vdash e: \tau_{2}}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e: \tau_{1} \rightarrow \tau_{2}} \mathrm{FN}
$$

by

$$
\frac{\Gamma, x: \alpha \vdash e: \tau_{2}}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e: \alpha \rightarrow \tau_{2}} \mathrm{FN}
$$

- $x: \alpha$ when $\alpha$ is fresh (otherwise unused) means: type of $x$ is completely arbitrary.
- syntax-directed now?
- $\tau_{1}$ : meta-variable for concrete types
- $\alpha$ : (still meta variable for) type variables
$\alpha$ 's completely arbitrary?

Consider body

$$
e=x g
$$

for $\mathrm{fn}_{\pi} x \Rightarrow e$
$\Rightarrow$

- a function type: $\alpha=\beta \rightarrow \gamma$
- fit together with type of $g \Rightarrow$ condition or constraint on $\beta$
- judments "give back" not just the type, but also "restrictions" on type variables.
- represented as substitutions ${ }^{1}$
- $\Rightarrow$

$$
\Gamma \vdash e:(\tau, \theta)
$$

Under the assumptions $\Gamma$ (which might "assign" to (program) variables: type variables), program $e$ possesses type $\tau$ (again potentially containing type variables) and imposes the restrictions "embodied" by $\theta$ on the type variables.

$$
\frac{\Gamma, x: \alpha \vdash e_{0}:\left(\tau_{0}, \theta_{0}\right)}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e:\left(\left(\theta_{0} \alpha\right) \rightarrow \tau_{0}, \theta_{0}\right)} \mathrm{FN}_{\mathrm{N}}
$$

[^2]
## Unification

$\overline{\Gamma \vdash c:\left(\tau_{c}, i d\right)}$ T-Const $\overline{\Gamma \vdash x:(\Gamma(x), i d)}$ T-VAR

$$
\frac{\alpha \text { fresh } \quad \Gamma, x: \alpha \vdash e_{0}:\left(\tau_{0}, \theta_{0}\right)}{\Gamma \vdash \mathrm{fn}_{\pi} x \Rightarrow e_{0}:\left(\theta_{0} \alpha \rightarrow \tau_{0}, \theta_{0}\right)} \text { T-FN }
$$

$$
\frac{\alpha, \alpha_{0} \text { fresh } \Gamma, f: \alpha \rightarrow \alpha_{0}, x: \alpha \vdash e_{0}:\left(\tau_{0}, \theta_{0}\right) \quad \theta_{1}=\mathcal{U}\left(\tau_{0}, \theta_{0} \alpha_{0}\right)}{\Gamma \vdash \mathrm{fun}_{\pi} f x \Rightarrow e_{0}:\left(\theta_{1} \theta_{0} \alpha \rightarrow \theta_{1}\left(\tau_{0}\right), \theta_{1} \circ \theta_{0}\right)} \text { T-FUN }
$$

$$
\frac{\Gamma \vdash e_{1}:\left(\tau_{1}, \theta_{1}\right) \quad \theta_{1} \Gamma \vdash e_{2}:\left(\tau_{2}, \theta_{2}\right) \quad \alpha \text { fresh } \quad \theta_{3}=\mathcal{U}\left(\theta_{2} \tau_{1}, \tau_{2} \rightarrow \alpha\right)}{\Gamma \vdash e_{1} e_{2}:\left(\theta_{3} \alpha, \theta_{3} \theta_{2} \theta_{1}\right)} \text { T-APP }
$$

$$
\Gamma \vdash e_{0}:\left(\tau_{0}, \theta_{0}\right) \quad \theta_{0} \Gamma \vdash e_{1}:\left(\tau_{1}, \theta_{1}\right) \quad \theta_{1} \theta_{0} \Gamma \vdash e_{2}:\left(\tau_{2}, \theta_{2}\right)
$$

$$
\frac{\theta_{3}=\mathcal{U}\left(\theta_{2} \theta_{0} \tau_{1}, \text { bool }\right) \quad \theta_{4}=\mathcal{U}\left(\theta_{3} \tau_{2}, \theta_{3} \theta_{2} \tau_{1}\right)}{\Gamma \vdash \text { if } e_{0} \text { then } e_{1} \text { else } e_{2}:\left(\theta_{4} \theta_{3} \tau_{2}, \theta_{4} \theta_{3} \theta_{2} \theta_{1} \theta_{0}\right)} \text { IF }
$$

$$
\frac{\Gamma \vdash e_{1}:\left(\tau_{1}, \theta_{1}\right) \quad \theta_{1} \Gamma, x: \tau_{1} \vdash e_{2}:\left(\tau_{2}, \theta_{2}\right)}{\Gamma \vdash \operatorname{let} x=e_{1} \text { in } e_{2}:\left(\tau_{2}, \theta_{2} \theta_{1}\right)} \text { LET }
$$

$$
\Gamma \vdash e_{1}:\left(\tau_{1}, \theta_{1}\right) \quad \theta_{2} \Gamma \vdash e_{2}:\left(\tau_{2}, \theta_{2}\right)
$$

$$
\frac{\theta_{3}=\mathcal{U}\left(\theta_{2} \tau_{1}, \tau_{\mathrm{op}}^{1}\right) \quad \theta_{3}=\mathcal{U}\left(\theta_{3} \tau_{2}, \tau_{\mathrm{op}}^{2}\right)}{\Gamma \vdash e_{1} \text { op } e_{2}:\left(\tau_{\mathrm{op}}, \theta_{4} \theta_{3} \theta_{2} \theta_{1}\right)} \text { OP }
$$

Remarks Note again: we do not have the full famous Damas Milner type system with let-polymorphism. To do that we would need to add type schemes (see later).

## Unification

- "classical" algorithm ([10])
- many applications (theorem proving, Prolog etc.)
- unifier of two types $\tau_{1}$ and $\tau_{2}$ : a substitution $\theta$ such that

$$
\theta\left(\tau_{1}\right)=\theta\left(\tau_{2}\right)
$$

- unfication problem given $\tau_{1}$ and $\tau_{2}$, determine a unifier for them, if it exists
- better formulation of unfication problem: given $\tau_{1}$ and $\tau_{2}$, determine the best $=$ most general unifier for them (if they are unifiable).

Remarks in other areas: formulas, terms ...

## Unification algorithm for underlying types

$$
\begin{aligned}
& \mathcal{U}(\text { int, int }))=i d \\
& \mathcal{U}(\text { bool, bool }))=i d \\
& \mathcal{U}\left(\tau_{1} \rightarrow \tau_{2}, \tau_{1}^{\prime} \rightarrow \tau_{2}^{\prime}\right)=\text { let } \theta_{1}=\mathcal{U}\left(\tau_{1}, \tau_{1}^{\prime}\right) \\
& \theta_{2}=\mathcal{U}\left(\theta_{1} \tau_{2}, \theta_{1} \tau_{2}^{\prime}\right) \\
& \mathcal{U}(\tau, \alpha)= \begin{cases}\text { in } \theta_{2} \circ \theta_{1} \\
{[\alpha \mapsto \tau]} & \text { if } \alpha \text { does not occur in } \tau \\
\text { fail } & \text { or if } \alpha=\tau\end{cases} \\
& \mathcal{U}(\alpha, \tau)=\text { symmetrically } \\
& \mathcal{U}\left(\tau_{1}, \tau_{2}\right)=\text { fail in all other cases }
\end{aligned}
$$

## Type inference algorithm

- formulated here as rule system
- immediate correspondence to a recursive function:

$$
\mathcal{W}(\Gamma, e)=(\tau, \theta)
$$

instead of

$$
\Gamma \vdash e:(\tau, \theta)
$$

## "Classic" type inference

- we did not look at the full well-known Hindley-Damas-Milner type inference algorithm
- missing here: polymorphic let
- monomoprhic let: "almost useless" polymorphism
- Note the fine line
- polymorphic let: yes
- polymorphic functions as function arguments: no!


## the classical type "inference" algo

- higher-order functions,
- polymorphic functions,
- but no "higher-order polymorphic functions"
- dropping the last restriction: type inference undecidable
- no type variables in the underlying type system (the "specification"), the type inference algo does
- types (with variables) and type schemes $\forall \alpha . \tau$


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[^0]:    ${ }^{1}$ source code, intermediate code at various levels
    ${ }^{2}$ Playing with words, one could call full-scale (hand?) verification "static" analysis, and likewise call lexical analysis a static analysis.

[^1]:    ${ }^{1}$ Not covered in the lecture.

[^2]:    ${ }^{1}$ One could also collect the constraints/restrictions as a set of equations and solve them at the very end.

