Programming Language Theory and its Implementation

Applicative and Imperative Paradigms

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To~Avra

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Preface

Formal methods are becoming an increasingly important part of the design of computer systems. This book provides elementary introductions to two of the mathematical theories upon which these methods are based:

- (i) Floyd-Hoare logic, a formal system for proving the correctness of imperative programs.
- (ii) The λ -calculus and combinators, the mathematical theory underlying functional programming.

The book is organised so that (i) and (ii) can be studied independently. The two theories are illustrated with working programs written in LISP, to which an introduction is provided. It is hoped that the programs will both clarify the theoretical material and show how it can be applied in practice. They also provide a starting point for student programming projects in some interesting and rapidly expanding areas of non-numerical computation.

Floyd-Hoare logic is a theory of reasoning about programs that are written in conventional imperative programming languages like Fortran, Algol, Pascal, Ada, Modula-2 etc. Imperative programming consists in writing commands that modify the state of a machine. Floyd-Hoare logic can be used to establish the correctness of programs already written, or (better) as the foundation for rigorous software development methodologies in which programs are constructed in parallel with their verifications (an example of such a methodology is VDM [37]). Furthermore, thinking about the logical properties of programs provides a useful perspective, even if one is not going to verify them in detail. Floyd-Hoare logic has influenced the design of several programming languages, notably Euclid [46]. It is also the basis for axiomatic semantics, in which the meaning of a programming language is specified by requiring that all programs written in it satisfy the rules and axioms of a formal logic. Since Hoare's first paper [32] was published, there have been many reformulations of his ideas, e.g. by Dijkstra [16]. We do not describe these developments here, partly because Hoare's original formulation is still the simplest and (in my opinion) the best to learn first, and partly because it is the basis for commercial program verifiers (e.g. Gypsy [20]). The principles underlying such verifiers are described in Chapter 3 xii Preface

and the implementation of an example system is presented in Chapter 11. Good introductions to the recent developments in verification theory are the books by Gries [26] and Backhouse [3].

The λ -calculus is a theory of higher-order functions, i.e. functions that take functions as arguments or return functions as results. It has inspired the design of functional programming languages including LISP [53], ML [55], Miranda [70] and Ponder [17]. These languages provide notations for defining functions that are based directly on the λ -calculus; they differ in the 'mathematical purity' of the other features provided. Closely related to the λ -calculus is the theory of combinators. This provides an elegant 'machine code' into which functional languages can be compiled and which is simple to interpret by firmware or hardware. It is straightforward to prove the correctness of the algorithm for compiling functional programs to combinators (see Section 8.2); proving the correctness of compiling algorithms for imperative languages is usually extremely difficult [13, 54, 62].

Although functional programs execute more slowly than imperative ones on current computers, it is possible that this situation will be reversed in the future. Functional languages are well suited to exploit the multiprocessor architectures that are beginning to emerge from research laboratories.

Both the λ -calculus and the theory of combinators were originally developed as foundations for mathematics before digital computers were invented. They languished as obscure branches of mathematical logic until rediscovered by computer scientists. It is remarkable that a theory developed by logicians has inspired the design of both the hardware and software for a new generation of computers. There is an important lesson here for people who advocate reducing support for 'pure' research: the pure research of today defines the applied research of tomorrow.

If one is forced to use an imperative language (as most programmers are), then Floyd-Hoare logic provides a tool for establishing program correctness. However, many people feel that imperative programs are intrinsically difficult to reason about and that functional programming is a better basis for formal correctness analysis [6]. One reason for this is that functions are well-understood mathematical objects and thus do not call for a special logic; ordinary mathematics suffices. However, this view is not universally held; an eloquent case for the mathematical simplicity of imperative programming can be found in recent work by Dijkstra [16], Hehner [27] and Hoare [34]. Furthermore, the functions arising in functional programming are often unlike traditional mathematical functions, and special logics, e.g. LCF [25, 60], have had to be devised for reasoning about them.

My own view is that both imperative and functional programming have their place, but that it is likely that functional languages will gradually replace imperative ones for general purpose use. This is already beginning Preface xiii

to happen; for example, ML replaced Pascal in 1987 as the first language that computer science students are taught at Cambridge University. There is growing evidence that

- (i) programmers can solve problems more quickly if they use a functional language, and
- (ii) the resulting solutions are more likely to be correct.

Because of (ii), and the relative ease of verifying the correctness of compilers for functional languages, functional programming is likely to have importance in safety-critical applications.

Parts I and II of this book provide an introduction to the theory underlying both imperative and functional programming. Part III contains some working programs which illustrate the material described in the first two parts. Floyd-Hoare logic is illustrated by an implementation of a complete program verifier. This consists of a verification condition generator and a simple theorem prover. The λ -calculus and combinators are illustrated by a toolkit for experimenting with the interpretation and compilation of functional programs. The example systems in Part III are implemented in LISP and are short enough that the reader can easily type them in and run them. A tutorial on the subset of LISP used is given in Chapter 9.

This book has evolved from the lecture notes for two undergraduate courses at Cambridge University which I have taught for the last few years. It should be possible to cover all the material in about thirty hours. There are no mathematical prerequisites besides school algebra. Familiarity with simple imperative programming (e.g. a first course in Pascal) would be useful, but is not essential.

I am indebted to the various students and colleagues at Cambridge who have provided me with feedback on the courses mentioned above, and to Graham Birtwistle of the University of Calgary and Elsa Gunter of the University of Pennsylvania, who pointed out errors in the notes and gave much advice and help on revising them into book form. In addition, Graham Birtwistle provided detailed and penetrating comments on the final draft of the manuscript, as did Jan van Eijck of SRI International, Mike Fourman of Brunel University and Avra Cohn of Cambridge University. These four people discovered various errors and made many helpful suggestions. Martin Hyland of Cambridge University explained to me a number of subtle points concerning the relationship between the λ -calculus and the theory of combinators.

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done using IATEX [41]. I am grateful to the versatile Elsa Gunter who, whilst simultaneously working as a researcher in the Cambridge Computer Laboratory and writing up her Ph.D. on group theory, was also employed by me to typeset this book. I would also like to thank Prentice-Hall's copy editors for helping to counteract excesses of informality in my writing style.

Finally, this book would not exist without the encouragement and patience of Helen Martin, Prentice-Hall's Acquisitions Editor, and Tony Hoare, the Series Editor.

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Part I Proving Programs Correct

Program Specification

A simple programming language containing assignments, conditionals, blocks, WHILE-commands and FOR-commands is introduced. This language is then used to illustrate Hoare's notation for specifying the partial correctness of programs. Hoare's notation uses predicate calculus to express conditions on the values of program variables. A fragment of predicate calculus is introduced and illustrated with examples.

1.1 Introduction

In order to prove mathematically the correctness of a program one must first specify what it means for it to be correct. In this chapter a notation for specifying the desired behaviour of *imperative* programs is described. This notation is due to C.A.R. Hoare.

Executing an imperative program has the effect of changing the *state*, i.e. the values of program variables¹. To use such a program, one first establishes an initial state by setting the values of some variables to values of interest. One then executes the program. This transforms the initial state into a final one. One then inspects (using print commands etc.) the values of variables in the final state to get the desired results. For example, to compute the result of dividing **y** into **x** one might load **x** and **y** into program variables **X** and **Y**, respectively. One might then execute a suitable program (see Example 7 in Section 1.4) to transform the initial state into a final state in which the variables QUOT and REM hold the quotient and remainder, respectively.

The programming language used in this book is described in the next section.

¹ For languages more complex than those described in this book, the state may consist of other things besides the values of variables [23].

1.2 A little programming language

Programs are built out of *commands* like assignments, conditionals etc. The terms 'program' and 'command' are really synonymous; the former will only be used for commands representing complete algorithms. Here the term 'statement' is used for conditions on program variables that occur in correctness specifications (see Section 1.3). There is a potential for confusion here because some writers use this word for commands (as in 'for-statement' [33]).

We now describe the *syntax* (i.e. form) and *semantics* (i.e. meaning) of the various commands in our little programming language. The following conventions are used:

- 1. The symbols V, V_1, \ldots, V_n stand for arbitrary variables. Examples of particular variables are X, REM, QUOT etc.
- 2. The symbols E, E_1, \ldots, E_n stand for arbitrary expressions (or terms). These are things like X + 1, $\sqrt{2}$ etc. which denote values (usually numbers).
- 3. The symbols S, S_1, \ldots, S_n stand for arbitrary statements. These are conditions like $X < Y, X^2 = 1$ etc. which are either true or false.
- 4. The symbols C, C_1, \ldots, C_n stand for arbitrary commands of our programming language; these are described in the rest of this section.

Terms and statements are described in more detail in Section 1.5.

1.2.1 Assignments

Syntax: V := E

Semantics: The state is changed by assigning the value of the term E to the variable V.

Example: X:=X+1

This adds one to the value of the variable X.

1.2.2 Sequences

Syntax: C_1 ; \cdots ; C_n

Semantics: The commands C_1, \dots, C_n are executed in that order.

Example: R:=X; X:=Y; Y:=R

The values of X and Y are swapped using R as a temporary variable. This command has the *side effect* of changing the value of the variable R to the old value of the variable X.

1.2.3 Blocks

Syntax: BEGIN VAR V_1 ; \cdots VAR V_n ; C END

Semantics: The command C is executed, and then the values of V_1, \dots, V_n are restored to the values they had before the block was entered. The initial values of V_1, \dots, V_n inside the block are unspecified.

Example: BEGIN VAR R; R:=X; X:=Y; Y:=R END

The values of X and Y are swapped using R as a temporary variable. This command does *not* have a side effect on the variable R.

1.2.4 One-armed conditionals

Syntax: IF S THEN C

Semantics: If the statement S is true in the current state, then C is executed. If S is false, then nothing is done.

Example: IF $\neg(X=0)$ THEN R:= Y DIV X

If the value X is not zero, then R is assigned the result of dividing the value of Y by the value of X.

1.2.5 Two-armed conditionals

Syntax: IF S THEN C_1 ELSE C_2

Semantics: If the statement S is true in the current state, then C_1 is executed. If S is false, then C_2 is executed.

Example: IF X<Y THEN MAX:=Y ELSE MAX:=X

The value of the variable MAX it set to the maximum of the values of X and Y.

1.2.6 WHILE-commands

 \mathbf{Syntax} : WHILE S DO C

Semantics: If the statement S is true in the current state, then C is executed and the WHILE-command is then repeated. If S is false, then nothing is done. Thus C is repeatedly executed until the value of S becomes true. If S never becomes true, then the execution of the command never terminates.

Example: WHILE $\neg(X=0)$ DO X:= X-2

If the value of X is non-zero, then its value is decreased by 2 and then process is repeated. This WHILE-command will terminate (with X having value 0) if the value of X is an even non-negative number. In all other states it will not terminate.

1.2.7 FOR-commands

Syntax: FOR $V := E_1$ UNTIL E_2 DO C

Semantics: If the values of terms E_1 and E_2 are positive numbers e_1 and e_2 respectively, and if $e_1 \leq e_2$, then C is executed $(e_2-e_1)+1$ times with the variable V taking on the sequence of values e_1, e_1+1, \ldots, e_2 in succession. For any other values, the FOR-command has no effect. A more precise description of this semantics is given in Section 2.1.11.

Example: FOR N:=1 UNTIL M DO X:=X+N

If the value of the variable M is m and $m \ge 1$, then the command X:=X+N is repeatedly executed with N taking the sequence of values 1, ..., m. If m < 1 then the FOR-command does nothing.

1.2.8 Summary of syntax

The syntax of our little language can be summarized with the following specification in BNF notation²

²BNF stands for Backus-Naur form; it is a well-known notation for specifying syntax.

Note that:

- Variables, terms and statements are as described in Section 1.5.
- Only declarations of the form 'VAR < variable>' are needed. The types of variables need not be declared (unlike in Pascal).
- Sequences C_1 ; ... C_n are valid commands; they are equivalent to BEGIN C_1 ; ... C_n END (i.e. blocks without any local variables).
- \bullet The BNF syntax is ambiguous: it does not specify, for example, whether IF S_1 THEN IF S_2 THEN C_1 ELSE C_2 means

IF
$$S_1$$
 THEN (IF S_2 THEN C_1 ELSE C_2)

or

IF
$$S_1$$
 THEN (IF S_2 THEN C_1) ELSE C_2

We will clarify, whenever necessary, using brackets.

1.3 Hoare's notation

In a seminal paper [32] C.A.R. Hoare introduced the following notation for specifying what a program does³:

$$\{P\}\ C\ \{Q\}$$

where:

• C is a program from the programming language whose programs are being specified (the language in Section 1.2 in our case).

 $^{^3}$ Actually, Hoare's original notation was P $\{C\}$ Q not $\{P\}$ C $\{Q\},$ but the latter form is now more widely used.

• P and Q are conditions on the program variables used in C.

Conditions on program variables will be written using standard mathematical notations together with *logical operators* like \land ('and'), \lor ('or'), \neg ('not') and \Rightarrow ('implies'). These are described further in Section 1.5.

We say $\{P\}$ C $\{Q\}$ is true, if whenever C is executed in a state satisfying P and if the execution of C terminates, then the state in which C's execution terminates satisfies Q.

Example: $\{X = 1\}$ X := X+1 $\{X = 2\}$. Here P is the condition that the value of X is 1, Q is the condition that the value of X is 2 and C is the assignment command X := X+1 (i.e. 'X becomes X+1'). $\{X = 1\}$ X := X+1 $\{X = 2\}$ is clearly true. \Box

An expression $\{P\}$ C $\{Q\}$ is called a partial correctness specification; P is called its precondition and Q its postcondition.

These specifications are 'partial' because for $\{P\}$ C $\{Q\}$ to be true it is not necessary for the execution of C to terminate when started in a state satisfying P. It is only required that if the execution terminates, then Q holds.

A stronger kind of specification is a total correctness specification. There is no standard notation for such specifications. We shall use $[P] \subset [Q]$.

A total correctness specification $[P] \subset [Q]$ is true if and only if the following conditions apply:

- (i) Whenever C is executed in a state satisfying P, then the execution of C terminates.
- (ii) After termination Q holds.

The relationship between partial and total correctness can be informally expressed by the equation:

Total correctness = Termination + Partial correctness.

Total correctness is what we are ultimately interested in, but it is usually easier to prove it by establishing partial correctness and termination separately.

Termination is often straightforward to establish, but there are some well-known examples where it is not. For example⁴, no one knows whether the program below terminates for all values if X:

```
WHILE X>1 DO
IF ODD(X) THEN X := (3\times X)+1 ELSE X := X DIV 2
```

⁴This example is taken from Exercise 2 on page 17 of Reynolds's book [63].

(The expression X DIV 2 evaluates to the result of rounding down X/2 to a whole number.)

Exercise 1

Write a specification which is true if and only if the program above terminates. \Box

In Part I of this book Floyd-Hoare logic is described; this only deals with partial correctness. Theories of total correctness can be found in the texts by Dijkstra [16] and Gries [26].

1.4 Some examples

The examples below illustrate various aspects of partial correctness specification.

In Examples 5, 6 and 7 below, **T** (for 'true') is the condition that is always true. In Examples 3, 4 and 7, \wedge is the logical operator 'and', i.e. if P_1 and P_2 are conditions, then $P_1 \wedge P_2$ is the condition that is true whenever both P_1 and P_2 hold.

1.
$$\{X = 1\}$$
 $Y := X$ $\{Y = 1\}$

This says that if the command Y:=X is executed in a state satisfying the condition X=1 (i.e. a state in which the value of X is 1), then, if the execution terminates (which it does), then the condition Y=1 will hold. Clearly this specification is true.

2.
$$\{X = 1\} Y := X \{Y = 2\}$$

This says that if the execution of Y:=X terminates when started in a state satisfying X=1, then Y=2 will hold. This is clearly false.

3.
$$\{X=x \land Y=y\}$$
 BEGIN R:=X; X:=Y; Y:=R END $\{X=y \land Y=x\}$

This says that if the execution of BEGIN R:=X; X:=Y; Y:=R END terminates (which it does), then the values of X and Y are exchanged. The variables x and y, which don't occur in the command and are used to name the initial values of program variables X and Y, are called auxiliary variables (or ghost variables).

4.
$$\{X=x \land Y=y\}$$
 BEGIN $X:=Y$; $Y:=X$ END $\{X=y \land Y=x\}$

This says that BEGIN X:=Y; Y:=X END exchanges the values of X and Y. This is not true.

5.
$$\{T\} \subset \{Q\}$$

This says that whenever C halts, Q holds.

```
6. \{P\} \ C \ \{\mathtt{T}\}
```

This specification is true for every condition P and every command C (because T is always true).

```
7. {T}
 \begin{array}{c} \text{BEGIN} \\ \text{R:=X;} \\ \text{Q:=0;} \\ \text{WHILE Y$\leq$R DO$} \\ \text{BEGIN R:=R-Y; Q:=Q+1 END} \\ \text{END} \\ \{\text{R} < \text{Y} \land \text{X} = \text{R} + (\text{Y} \times \text{Q})\} \end{array} \right\} C
```

This is $\{T\}$ C $\{R < Y \land X = R + (Y \times Q)\}$ where C is the command indicated by the braces above. The specification is true if whenever the execution of C halts, then Q is quotient and R is the remainder resulting from dividing Y into X. It is true (even if X is initially negative!)

In this example a program variable $\mathbb Q$ is used. This should not be confused with the Q used in 5 above. The program variable $\mathbb Q$ (notice the font) ranges over numbers, whereas the postcondition Q (notice the font) ranges over statements. In general, we use **typewriter font** for particular program variables and *italic font* for variables ranging over statements. Although this subtle use of fonts might appear confusing at first, once you get the hang of things the difference between the two kinds of 'Q' will be clear (indeed you should be able to disambiguate things from context without even having to look at the font).

Exercise 2

Let C be as in Example 7 above. Find a condition P such that:

$$[P] \ C \ [R < Y \land X = R + (Y \times Q)]$$

is true. □

Exercise 3

When is [T] C [T] true? \square

Exercise 4

Write a partial correctness specification which is true if and only if the command C has the effect of multiplying the values of X and Y and storing the result in X. \square

Exercise 5

Write a specification which is true if the execution of C always halts when execution is started in a state satisfying P. \square

1.5 Terms and statements

The notation used here for expressing pre- and postconditions is based on a language called *first-order logic* invented by logicians around the turn of this century. For simplicity, only a fragment of this language will be used. Things like:

$$T$$
, F , $X = 1$, $R < Y$, $X = R + (Y \times Q)$

are examples of atomic statements. Statements are either true or false. The statement T is always true and the statement F is always false. The statement F is true if the value of F is equal to 1. The statement F is true if the value of F is less than the value of F. The statement F is true if the value of F is equal to the sum of the value of F with the product of F and F and F.

Statements are built out of terms like:

$$X$$
, 1, R , Y , $R+(Y\times Q)$, $Y\times Q$

Terms denote values such as numbers and strings, unlike statements which are either true or false. Some terms, like 1 and 4+5, denote a fixed value, whilst other terms contain variables like X, Y, Z etc. whose value can vary. We will use conventional mathematical notation for terms, as illustrated by the examples below:

$$X, Y, Z,$$
 $1, 2, 325,$
 $-X, -(X+1), (X\times Y)+Z,$
 $\sqrt{(1+X^2)}, X!, \sin(X), \operatorname{rem}(X,Y)$

T and F are atomic statements that are always true and false respectively. Other atomic statements are built from terms using *predicates*. Here are some more examples:

ODD(X), PRIME(3),
$$X = 1$$
, $(X+1)^2 \ge X^2$

ODD and PRIME are examples of predicates and = and \geq are examples of infixed predicates. The expressions X, 1, 3, X+1, $(X+1)^2$, X^2 are examples of terms.

Compound statements are built up from atomic statements using the following logical operators:

$$\begin{array}{ll}
\neg & (not) \\
\land & (and) \\
\lor & (or) \\
\Rightarrow & (implies) \\
\Leftrightarrow & (if and only if)
\end{array}$$

The single arrow \rightarrow is commonly used for implication instead of \Rightarrow . We use \Rightarrow to avoid possible confusion with the use of \rightarrow for λ -conversion in Part II.

Suppose P and Q are statements, then:

- $\neg P$ is true if P is false, and false if P is true.
- $P \wedge Q$ is true whenever both P and Q are true.
- $P \lor Q$ is true if either P or Q (or both) are true.
- $P\Rightarrow Q$ is true if whenever P is true, then Q is true also. By convention we regard $P\Rightarrow Q$ as being true if P is false. In fact, it is common to regard $P\Rightarrow Q$ as equivalent to $\neg P\vee Q$; however, some philosophers called intuitionists disagree with this treatment of implication.
- $P \Leftrightarrow Q$ is true if P and Q are either both true or both false. In fact $P \Leftrightarrow Q$ is equivalent to $(P \Rightarrow Q) \land (Q \Rightarrow P)$.

Examples of statements built using the connectives are:

```
\begin{array}{ll} \mathtt{ODD}(\mathtt{X}) \vee \mathtt{EVEN}(\mathtt{X}) & \mathtt{X} \text{ is odd or even.} \\ \\ \neg (\mathtt{PRIME}(\mathtt{X}) \Rightarrow \mathtt{ODD}(\mathtt{X})) & \text{It is not the case that if $\mathtt{X}$ is prime, then $\mathtt{X}$ is odd.} \\ \\ \mathtt{X} \leq \mathtt{Y} \Rightarrow \mathtt{X} \leq \mathtt{Y}^2 & \text{If $\mathtt{X}$ is less than or equal to $\mathtt{Y}$, then $\mathtt{X}$ is less than or equal to $\mathtt{Y}^2$.} \end{array}
```

To reduce the need for brackets it is assumed that \neg is more binding than \land and \lor , which in turn are more binding than \Rightarrow and \Leftrightarrow . For example:

```
\begin{array}{lll} \neg P \wedge Q & \text{is equivalent to} & (\neg P) \wedge Q \\ P \wedge Q \Rightarrow R & \text{is equivalent to} & (P \wedge Q) \Rightarrow R \\ P \wedge Q \Leftrightarrow \neg R \vee S & \text{is equivalent to} & (P \wedge Q) \Leftrightarrow ((\neg R) \vee S) \end{array}
```

Floyd-Hoare Logic

The idea of formal proof is discussed. Floyd-Hoare logic is then introduced as a method for reasoning formally about programs.

In the last chapter three kinds of expressions that could be true or false were introduced:

- (i) Partial correctness specifications $\{P\}$ C $\{Q\}$.
- (ii) Total correctness specifications [P] C [Q].
- (iii) Statements of mathematics (e.g. $(X + 1)^2 = X^2 + 2 \times X + 1$).

It is assumed that the reader knows how to prove simple mathematical statements like the one in (iii) above. Here, for example, is a proof of this fact.

1.	$(X + 1)^2$	$= (X+1) \times (X+1)$	Definition of $()^2$.
2.	$(X+1)\times(X+1)$	$= (\mathtt{X} + \mathtt{1}) \times \mathtt{X} + (\mathtt{X} + \mathtt{1}) \times \mathtt{1}$	Left distributive law
			of \times over $+$.
3.	$(X + 1)^2$	$= (X+1) \times X + (X+1) \times 1$	Substituting line 2
			into line 1.
4.	$(X + 1) \times 1$	= X + 1	Identity law for 1.
5.	$(X + 1) \times X$	$= X \times X + 1 \times X$	Right distributive law
			of \times over $+$.
6.	$(X + 1)^2$	$= \texttt{X} \times \texttt{X} + \texttt{1} \times \texttt{X} + \texttt{X} + \texttt{1}$	Substituting lines 4
			and 5 into line 3.
7.	$1 \times X$	= X	Identity law for 1.
8.	$(X + 1)^2$	$= X \times X + X + X + 1$	Substituting line 7
			into line 6.
9.	${\tt X} \times {\tt X}$	$= X^2$	Definition of $()^2$.
10.	X + X	$=2 \times X$	2=1+1, distributive law.
11.	$(X + 1)^2$	$= X^2 + 2 \times X + 1$	Substituting lines 9
	, ,		and 10 into line 8.

This proof consists of a sequence of lines, each of which is an instance of an axiom (like the definition of ()²) or follows from previous lines by a

rule of inference (like the substitution of equals for equals). The statement occurring on the last line of a proof is the statement proved by it (thus $(X+1)^2 = X^2 + 2 \times X + 1$ is proved by the proof above).

To construct formal proofs of partial correctness specifications axioms and rules of inference are needed. This is what Floyd-Hoare logic provides. The formulation of the deductive system is due to Hoare [32], but some of the underlying ideas originated with Floyd [18].

A proof in Floyd-Hoare logic is a sequence of lines, each of which is either an *axiom* of the logic or follows from earlier lines by a *rule of inference* of the logic.

The reason for constructing formal proofs is to try to ensure that only sound methods of deduction are used. With sound axioms and rules of inference, one can be confident that the conclusions are true. On the other hand, if any axioms or rules of inference are unsound then it may be possible to deduce false conclusions; for example¹

```
\begin{array}{lll} 1. & \sqrt{-1\times -1} & = \sqrt{-1\times -1} & \text{Reflexivity of } =. \\ 2. & \sqrt{-1\times -1} & = (\sqrt{-1})\times (\sqrt{-1}) & \text{Distributive law of } \sqrt{\text{ over }} \times. \\ 3. & \sqrt{-1\times -1} & = (\sqrt{-1})^2 & \text{Definition of } ()^2. \\ 4. & \sqrt{-1\times -1} & = -1 & \text{definition of } \sqrt{.} \\ 5. & \sqrt{1} & = -1 & \text{As } -1\times -1 = 1. \\ 6. & 1 & = -1 & \text{As } \sqrt{1} = 1. \end{array}
```

A formal proof makes explicit what axioms and rules of inference are used to arrive at a conclusion. It is quite easy to come up with plausible rules for reasoning about programs that are actually unsound (some examples for FOR-commands can be found in Section 2.1.11). Proofs of correctness of computer programs are often very intricate and formal methods are needed to ensure that they are valid. It is thus important to make fully explicit the reasoning principles being used, so that their soundness can be analysed.

Exercise 6

Find the flaw in the 'proof' of 1 = -1 above. \square

For some applications, correctness is especially important. Examples include life-critical systems such as nuclear reactor controllers, car breaking systems, fly-by-wire aircraft and software controlled medical equipment. At the time of writing, there is a legal action in progress resulting from the death of several people due to radiation overdoses by a cancer treatment machine that had a software bug [38]. Formal proof of correctness provides a way of establishing the absence of bugs when exhaustive testing is impossible (as it almost always is).

¹ This example was shown to me by Sylva Cohn.

The Floyd-Hoare deductive system for reasoning about programs will be explained and illustrated, but the mathematical analysis of the soundness and completeness of the system is beyond the scope of this book (however, there is a brief discussion of what is involved in Section 2.2).

2.1 Axioms and rules of Floyd-Hoare logic

As discussed at the beginning of this chapter, a formal proof of a statement is a sequence of lines ending with the statement and such that each line is either an instance of an axiom or follows from previous lines by a rule of inference. If S is a statement (of either ordinary mathematics or Floyd-Hoare logic) then we write $\vdash S$ to mean that S has a proof. The statements that have proofs are called theorems. As discussed earlier, in this book only the axioms and rules of inference for Floyd-Hoare logic are described; we will thus simply assert $\vdash S$ if S is a theorem of mathematics without giving any formal justification. Of course, to achieve complete rigour such assertions must be proved, but for details of this the reader will have to consult a book (such as [10, 47, 49]) on formal logic.

The axioms of Floyd-Hoare logic are specified below by *schemas* which can be *instantiated* to get particular partial correctness specifications. The inference rules of Floyd-Hoare logic will be specified with a notation of the form:

$$\frac{\vdash S_1, \ldots, \vdash S_n}{\vdash S}$$

This means the *conclusion* $\vdash S$ may be deduced from the *hypotheses* $\vdash S_1, \ldots, \vdash S_n$. The hypotheses can either all be theorems of Floyd-Hoare logic (as in the sequencing rule below), or a mixture of theorems of Floyd-Hoare logic and theorems of mathematics (as in the rule of preconditioning strengthening described in Section 2.1.2).

2.1.1 The assignment axiom

The assignment axiom represents the fact that the value of a variable V after executing an assignment command V := E equals the value of the expression E in the state before executing it. To formalize this, observe that if a statement P is to be true after the assignment, then the statement obtained by substituting E for V in P must be true before executing it.

In order to say this formally, define P[E/V] to mean the result of replacing all occurrences of V in P by E. Read P[E/V] as 'P with E for

V'. For example,

$$(X+1 > X)[Y+Z/X] = ((Y+Z)+1 > Y+Z)$$

The way to remember this notation is to remember the 'cancellation law'

$$V[E/V] = E$$

which is analogous to the cancellation property of fractions

$$v \times (e/v) = e$$

The assignment axiom

$$\vdash \{P [E/V]\} V := E \{P\}$$

Where V is any variable, E is any expression, P is any statement and the notation P[E/V] denotes the result of substituting the term E for all occurrences of the variable V in the statement P.

Instances of the assignment axiom are:

1.
$$\vdash \{Y = 2\} \ X := 2 \ \{Y = X\}$$

$$2. \quad \vdash \ \{\mathtt{X}+\mathtt{1}=\mathtt{n}+\mathtt{1}\} \ \mathtt{X} := \mathtt{X}+\mathtt{1} \ \{\mathtt{X}=\mathtt{n}+\mathtt{1}\}$$

3.
$$\vdash \{E = E\} \ \mathbf{X} := E \ \{\mathbf{X} = E\}$$

Many people feel the assignment axiom is 'backwards' from what they would expect. Two common erroneous intuitions are that it should be as follows:

(i) $\vdash \{P\} \ V := E \{P[V/E]\}.$

Where the notation P[V/E] denotes the result of substituting V for E in P.

This has the clearly false consequence that $\vdash \{x=0\} \ x:=1 \ \{x=0\}$, since the (x=0)[x/1] is equal to (x=0) as 1 doesn't occur in (x=0).

(ii)
$$\vdash \{P\} \ V := E \{P [E/V]\}.$$

This has the clearly false consequence $\vdash \{x=0\} \ x:=1 \{1=0\}$ which follows by taking P to be X=0, V to be X=0 and E to be 1.

The fact that it is easy to have wrong intuitions about the assignment axiom shows that it is important to have rigorous means of establishing the validity of axioms and rules. We will not go into this topic here aside from remarking that it is possible to give a formal semantics of our little programming language and then to prove that the axioms and rules of inference of Floyd-Hoare logic are sound. Of course, this process will only increase our confidence in the axioms and rules to the extent that we believe the correctness of the formal semantics. The simple assignment axiom above is not valid for 'real' programming languages. For example, work by G. Ligler [44] shows that it can fail to hold in six different ways for the language Algol 60.

One way that our little programming language differs from real languages is that the evaluation of expressions on the right of assignment commands cannot 'side effect' the state. The validity of the assignment axiom depends on this property. To see this, suppose that our language were extended so that it contained the 'block expression'

This expression, E say, has value 2, but its evaluation also 'side effects' the variable Y by storing 1 in it. If the assignment axiom applied to expressions like E, then it could be used to deduce:

$$\vdash$$
 {Y=0} X:=BEGIN Y:=1; 2 END {Y=0}

(since (Y=0)[E/X] = (Y=0) as X does not occur in (Y=0)). This is clearly false, as after the assignment Y will have the value 1.

2.1.2 Precondition strengthening

The next rule of Floyd-Hoare logic enables the preconditions of (i) and (ii) on page 16 to be simplified. Recall that

$$\frac{\vdash S_1, \ldots, \vdash S_n}{\vdash S}$$

means that $\vdash S$ can be deduced from $\vdash S_1, \ldots, \vdash S_n$.

Using this notation, the rule of precondition strengthening is

Precondition strengthening

$$\frac{\vdash \ P \Rightarrow P', \quad \vdash \ \{P'\} \ C \ \{Q\}}{\vdash \ \{P\} \ C \ \{Q\}}$$

Examples

1. From the arithmetic fact \vdash X+1=n+1 \Rightarrow X=n, and 2 on page 16 it follows by precondition strengthening that

$$\vdash \{X = n\} \ X := X + 1 \ \{X = n + 1\}.$$

The variable \mathbf{n} is an example of an auxiliary (or ghost) variable. As described earlier (see page 9), auxiliary variables are variables occurring in a partial correctness specification $\{P\}$ C $\{Q\}$ which do not occur in the command C. Such variables are used to relate values in the state before and after C is executed. For example, the specification above says that if the value of \mathbf{X} is \mathbf{n} , then after executing the assignment $\mathbf{X}:=\mathbf{X}+\mathbf{1}$ its value will be $\mathbf{n}+\mathbf{1}$.

2. From the logical truth \vdash T \Rightarrow (E=E), and 3 on page 16 one can deduce²:

$$\vdash \{T\} X := E \{X = E\}$$

2.1.3 Postcondition weakening

Just as the previous rule allows the precondition of a partial correctness specification to be strengthened, the following one allows us to weaken the postcondition.

Postcondition weakening

$$\frac{\vdash \ \{P\} \ C \ \{Q'\}, \qquad \vdash \ Q' \Rightarrow Q}{\vdash \ \{P\} \ C \ \{Q\}}$$

Example: Here is a little formal proof.

- 1. $\vdash \{R=X \land 0=0\} \ Q:=0 \ \{R=X \land Q=0\}$ By the assignment axiom.
- 2. $\vdash R=X \Rightarrow R=X \land 0=0$ By pure logic.
- By precondition strengthening $\{R=X \land Q=0\}$
- $\vdash R=X \land Q=0 \Rightarrow R=X+(Y \times Q)$ By laws of arithmetic.
- 5. $\vdash \{R=X\} \ Q:=0 \ \{R=X+(Y\times Q)\}$ By postcondition weakening.

² If it is not obvious that \vdash T \Rightarrow (E=E) is a logical truth, then you should read an elementary introduction to formal logic, e.g. [10, 19, 47, 49].

The rules precondition strengthening and postcondition weakening are sometimes called the *rules of consequence*.

2.1.4 Specification conjunction and disjunction

The following two rules provide a method of combining different specifications about the same command.

Specification conjunction

$$\frac{\vdash \{P_1\} \ C \ \{Q_1\}, \qquad \vdash \ \{P_2\} \ C \ \{Q_2\}}{\vdash \ \{P_1 \land P_2\} \ C \ \{Q_1 \land Q_2\}}$$

Specification disjunction

$$\frac{\vdash \ \{P_1\} \ C \ \{Q_1\}, \qquad \vdash \ \{P_2\} \ C \ \{Q_2\}}{\vdash \ \{P_1 \lor P_2\} \ C \ \{Q_1 \lor Q_2\}}$$

These rules are useful for splitting a proof into independent bits. For example, they enable $\vdash \{P\} \ C \ \{Q_1 \land Q_2\}$ to be proved by proving separately that both $\vdash \{P\} \ C \ \{Q_1\}$ and $\vdash \{P\} \ C \ \{Q_2\}$.

The rest of the rules allow the deduction of properties of compound commands from properties of their components.

2.1.5 The sequencing rule

The next rule enables a partial correctness specification for a sequence C_1 ; C_2 to be derived from specifications for C_1 and C_2 .

The sequencing rule

$$\frac{\vdash \{P\} \ C_1 \ \{Q\}, \qquad \vdash \ \{Q\} \ C_2 \ \{R\}}{\vdash \ \{P\} \ C_1; C_2 \ \{R\}}$$

Example: By the assignment axiom:

- $(i) \vdash \{X=x \land Y=y\} R := X \{R=x \land Y=y\}$
- (ii) $\vdash \{R=x \land Y=y\} \ X:=Y \ \{R=x \land X=y\}$
- (iii) $\vdash \{R=x \land X=y\} \ Y:=R \ \{Y=x \land X=y\}$

Hence by (i), (ii) and the sequencing rule

$$(iv) \vdash \{X=x \land Y=y\} R:=X; X:=Y \{R=x \land X=y\}$$

Hence by (iv) and (iii) and the sequencing rule

$$(v) \vdash \{X=x \land Y=y\} \ R:=X; \ X:=Y; \ Y:=R \ \{Y=x \land X=y\}$$

2.1.6 The derived sequencing rule

The following rule is derivable from the sequencing and consequence rules.

The derived sequencing rule

The derived sequencing rule enables (v) in the previous example to be deduced directly from (i), (ii) and (iii) in one step.

2.1.7 The block rule

The block rule is like the sequencing rule, but it also takes care of local variables.

The block rule

$$\frac{ \vdash \{P\} \ C \ \{Q\}}{\vdash \ \{P\} \ \mathtt{BEGIN} \ \mathtt{VAR} \ V_1; \ \ldots; \ \mathtt{VAR} \ V_n; \ C \ \mathtt{END} \ \{Q\}}$$

where none of the variables V_1, \ldots, V_n occur in P or Q.

The syntactic condition that none of the variables V_1, \ldots, V_n occur in P or Q is an example of a *side condition*. It is a syntactic condition that must hold whenever the rule is used. Without this condition the rule is invalid; this is illustrated in the example below.

Note that the block rule is regarded as including the case when there are no local variables (the 'n = 0' case).

Example: From \vdash {X=x \land Y=y} R:=X; X:=Y; Y:=R {Y=x \land X=y} (see page 20) it follows by the block rule that

$$\vdash \{X=x \land Y=y\} \text{ BEGIN VAR R}; R:=X; X:=Y; Y:=R \text{ END } \{Y=x \land X=y\}$$

since R does not occur in $X=x \land Y=y$ or $X=y \land Y=x$. Notice that from

$$\vdash \{X=x \land Y=y\} R:=X; X:=Y \{R=x \land X=y\}$$

one cannot deduce

$$\vdash \{X=x \land Y=y\}$$
 BEGIN VAR R; R:=X; X:=Y END $\{R=x \land X=y\}$

since R occurs in $\{R=x \land X=y\}$. This is as required, because assignments to local variables of blocks should not be felt outside the block body. Notice, however, that it is possible to deduce:

$$\vdash \{X=x \land Y=y\} \text{ BEGIN } R:=X; X:=Y \text{ END } \{R=x \land X=y\}.$$

This is correct because R is no longer a local variable. \square

The following exercise addresses the question of whether one can show that changes to local variables inside a block are invisible outside it.

Exercise 7

Consider the specification

$$\{X=x\}$$
 BEGIN VAR X; X:=1 END $\{X=x\}$

Can this be deduced from the rules given so far?

- (i) If so, give a proof of it.
- (ii) If not, explain why not and suggest additional rules and/or axioms to enable it to be deduced.

2.1.8 The derived block rule

From the derived sequencing rule and the block rule the following rule for blocks can be derived.

The derived block rule

where none of the variables V_1, \ldots, V_n occur in P or Q.

Using this rule, it can be deduced $in\ one\ step$ from (i), (ii) and (iii) on page 20 that:

$$\vdash$$
 {X=x \land Y=y} BEGIN VAR R; R:=X; X:=Y; Y:=R END {Y=x \land X=y}

Exercise 8

$$\mathrm{Show} \vdash \ \{\mathtt{X=x} \ \land \ \mathtt{Y=y}\} \ \mathtt{X:=X+Y} \,; \ \mathtt{Y:=X-Y} \,; \ \mathtt{X:=X-Y} \, \left\{\mathtt{Y=x} \ \land \ \mathtt{X=y}\right\}$$

Exercise 9

```
Show \vdash {X=R+(Y×Q)} BEGIN R:=R-Y; Q:=Q+1 END {X=R+(Y×Q)} \sqcap
```

2.1.9 The conditional rules

There are two kinds of conditional commands: one-armed conditionals and two-armed conditionals. There are thus two rules for conditionals.

The conditional rules

$$\frac{\vdash \ \{P \land S\} \ C \ \{Q\}, \qquad \vdash \ P \land \neg S \Rightarrow Q}{\vdash \ \{P\} \ \text{If } S \ \text{THEN} \ C \ \{Q\}}$$

$$\frac{\vdash \ \{P \land S\} \ C_1 \ \{Q\}, \qquad \vdash \ \{P \land \neg S\} \ C_2 \ \{Q\}}{\vdash \ \{P\} \ \mathtt{IF} \ S \ \mathtt{THEN} \ C_1 \ \mathtt{ELSE} \ C_2 \ \{Q\}}$$

Example: Suppose we are given that

- (i) $\vdash X > Y \Rightarrow max(X,Y) = X$
- (ii) $\vdash Y \ge X \Rightarrow max(X,Y) = Y$

Then by the conditional rules (and others) it follows that

$$\vdash \ \{\mathtt{T}\} \ \mathtt{IF} \ \mathtt{X} \underline{>} \mathtt{Y} \ \mathtt{THEN} \ \mathtt{MAX} := \mathtt{X} \ \mathtt{ELSE} \ \mathtt{MAX} := \mathtt{Y} \ \big\{ \mathtt{MAX} = \mathtt{max}(\mathtt{X}, \mathtt{Y}) \big\}$$

Exercise 10

Give a detailed formal proof that the specification in the previous example follows from hypotheses (i) and (ii). \Box

Exercise 11

Devise an axiom and/or rule of inference for a command SKIP that has no effect. Show that if IF S THEN C is regarded as an abbreviation for IF S THEN C ELSE SKIP, then the rule for one-armed conditionals is derivable from the rule for two-armed conditionals and your axiom/rule for SWAP. \square

Exercise 12

Suppose we add to our little programming language commands of the form:

CASE
$$E$$
 OF BEGIN C_1 ; ...; C_n END

These are evaluated as follows:

(i) First E is evaluated to get a value x.

- (ii) If x is not a number between 1 and n, then the CASE-command has no effect.
- (iii) If x = i where $1 \le i \le n$, then command C_i is executed.

Why is the following rule for CASE-commands wrong?

```
\frac{\vdash \{P \land E=1\} \ C_1 \ \{Q\}, \ \dots \ , \vdash \{P \land E=n\} \ C_n \ \{Q\}}{\vdash \{P\} \ \texttt{CASE} \ E \ \texttt{OF} \ \texttt{BEGIN} \ C_1; \ \dots \ ; \ C_n \ \texttt{END} \ \{Q\}}
```

Hint: Consider the case when P is 'X=0', E is 'X', C_1 is 'Y:=0' and Q is 'Y=0'. \square

Exercise 13

Devise a proof rule for the CASE-commands in the previous exercise and use it to show:

Exercise 14

Show that if $\vdash \{P \land S\}$ C_1 $\{Q\}$ and $\vdash \{P \land \neg S\}$ C_2 $\{Q\}$, then it is possible to deduce:

```
\vdash {P} IF S THEN C<sub>1</sub> ELSE IF \negS THEN C<sub>2</sub> {Q}.
```

2.1.10 The WHILE-rule

If $\vdash \{P \land S\}$ C $\{P\}$, we say: P is an invariant of C whenever S holds. The WHILE-rule says that if P is an invariant of the body of a WHILE-command whenever the test condition holds, then P is an invariant of the whole WHILE-command. In other words, if executing C once preserves the truth of P, then executing C any number of times also preserves the truth of P.

The WHILE-rule also expresses the fact that after a WHILE-command has terminated, the test must be false (otherwise, it wouldn't have terminated).

The WHILE-rule

$$\frac{\vdash \ \{P \land S\} \ C \ \{P\}}{\vdash \ \{P\} \ \mathtt{WHILE} \ S \ \mathtt{DO} \ C \ \{P \land \neg S\}}$$

Example: By Exercise 9 on page 22

```
\vdash \{X=R+(Y\times Q)\} \text{ BEGIN } R:=R-Y; Q:=Q+1 \text{ END } \{X=R+(Y\times Q)\}
```

Hence by precondition strengthening

```
\vdash \{X=R+(Y\times Q)\wedge Y\leq R\} \text{ BEGIN } R:=R-Y; \ Q:=Q+1 \text{ END } \{X=R+(Y\times Q)\}
```

Hence by the WHILE-rule (with $P = (X=R+(Y\times Q))$)

```
 \begin{array}{ll} (i) \; \vdash \; \big\{ \, X = R + (\Upsilon \times \mathbb{Q}) \, \big\} \\ & \quad \text{WHILE } \; \Upsilon \leq R \; \; DO \\ & \quad \text{BEGIN } \; R := R - \Upsilon ; \; \; \mathbb{Q} := \mathbb{Q} + 1 \; \; \text{END} \\ & \quad \big\{ \, X = R + (\Upsilon \times \mathbb{Q}) \wedge \neg \; (\Upsilon \leq R) \, \big\} \end{array}
```

It is easy to deduce that

```
(ii) \{T\} R:=X; Q:=0 \{X=R+(Y\times Q)\}
```

Hence by (i) and (ii), the sequencing rule and postcondition weakening

```
\begin{array}{l} \vdash \ \left\{T\right\} \\ \text{R:=X;} \\ \text{Q:=0;} \\ \text{WHILE Y$\leq$R DO$} \\ \text{BEGIN R:=R-Y; Q:=Q+1 END} \\ \left\{R{<}Y{\wedge}X{=}R{+}(Y{\times}Q)\right\} \end{array}
```

With the exception of the WHILE-rule, all the axioms and rules described so far are sound for total correctness as well as partial correctness. This is because the only commands in our little language that might not terminate are WHILE-commands. Consider now the following proof:

```
1. \vdash {T} X:=0 {T} (assignment axiom)
2. \vdash {T \land T} X:=0 {T} (precondition strengthening)
3. \vdash {T} WHILE T DO X:=0 {T \land \neg T} (2 and the WHILE-rule)
```

If the WHILE-rule were true for total correctness, then the proof above would show that:

$$\vdash \ [\texttt{T}] \ \texttt{WHILE} \ \texttt{T} \ \texttt{DO} \ \texttt{X} \colon \texttt{=0} \ [\texttt{T} \land \neg \texttt{T}]$$

but this is clearly false since WHILE T DO X:=0 does not terminate, and even if it did then $T \land \neg T$ could not hold in the resulting state.

Extending Floyd-Hoare logic to deal with termination is quite tricky. One approach can be found in Dijkstra [16].

2.1.11 The FOR-rule

It is quite hard to capture accurately the intended semantics of FOR-commands in Floyd-Hoare logic. Axioms and rules are given here that appear to be sound, but they are not necessarily complete (see Section 2.2). An early reference on the logic of FOR-commands is Hoare's 1972 paper [33]; a comprehensive treatment can be found in Reynolds [63].

The intention here in presenting the FOR-rule is to show that Floyd-Hoare logic can get very tricky. All the other axioms and rules were quite straightforward and may have given a false sense of simplicity: it is very difficult to give adequate rules for anything other than very simple programming constructs. This is an important incentive for using simple languages.

One problem with FOR-commands is that there are many subtly different versions of them. Thus before describing the FOR-rule, the intended semantics of FOR-commands must be described carefully. In this book, the semantics of

FOR
$$V := E_1$$
 UNTIL E_2 DO C

is as follows:

- (i) The expressions E_1 and E_2 are evaluated once to get values e_1 and e_2 , respectively.
- (ii) If either e_1 or e_2 is not a number, or if $e_1 > e_2$, then nothing is done.

(iii) If $e_1 < e_2$ the FOR-command is equivalent to:

BEGIN VAR
$$V$$
;
$$V:=e_1\;;\;C\;;\;V:=e_1+1\;;\;C\;;\;\ldots\;;\;V:=e_2\;;\;C$$
 END

i.e. C is executed $(e_2-e_1)+1$ times with V taking on the sequence of values e_1, e_1+1, \ldots, e_2 in succession. Note that this description is not rigorous: ' e_1 ' and ' e_2 ' have been used both as numbers and as expressions of our little language; the semantics of FOR-commands should be clear despite this.

FOR-rules in different languages can differ in subtle ways from the one here. For example, the expressions E_1 and E_2 could be evaluated at each iteration and the controlled variable V could be treated as global rather than local. Note that with the semantics presented here, FOR-commands cannot go into infinite loops (unless, of course, they contain non-terminating WHILE-commands).

To see how the FOR-rule works, suppose that

$$\vdash \{P\} \ C \ \{P[V+1/V]\}$$

Suppose also that C does not contain any assignments to the variable V. If this is the case, then it is intuitively clear (and can be rigorously proved) that

$$\vdash \{(V = v)\}\ C\ \{(V = v)\}\$$

hence by specification conjunction

$$\vdash \{P \land (V = v)\} \ C \ \{P [V+1/V] \land (V = v)\}$$

Now consider a sequence V := v; C. By Example 2 on page 18,

$$\vdash \{P \llbracket v/V \rrbracket\} \ V := v \{P \land (V = v)\}$$

Hence by the sequencing rule

$$\vdash \{P[V/v]\}\ V := v;\ C\{P[V+1/V]\ \land (V=v)\}$$

Now it is a truth of logic alone that

$$\vdash P[V+1/V] \land (V=v) \Rightarrow P[v+1/V]$$

hence by postcondition weakening

$$\vdash \{P[v/V]\}\ V := v;\ C\{P[v+1/V]\}$$

Taking v to be e_1, e_1+1, \ldots, e_2 and using the derived sequencing rule we can thus deduce

$$\{P[e_1/V]\}\ V:=e_1;\ C;\ V:=e_1+1;\ \dots;\ V:=e_2;\ C\ \{P[e_2/V]\}$$

This suggests that a FOR-rule could be:

$$\frac{ \vdash \{P\} \ C \ \{P \ [V+1/V]\}}{\vdash \{P \ [E_1/V]\} \ \text{FOR} \ V := E_1 \ \text{UNTIL} \ E_2 \ \text{DO} \ C \ \{P \ [E_2+1/V]\}}$$

Unfortunately, this rule is unsound. To see this, first note that:

- $\begin{array}{lll} 1. & \vdash & \{ \texttt{Y+1=Y+1} \} & \texttt{X:=Y+1} & \{ \texttt{X=Y+1} \} \\ 2. & \vdash & \{ \texttt{T} \} & \texttt{X:=Y+1} & \{ \texttt{X=Y+1} \} \end{array} \qquad \text{$(1$ and precondition strengthening)}$
- 3. $\vdash X=Y \Rightarrow T$ (logic: 'anything implies true') 4. $\vdash \{X=Y\} X:=Y+1 \{X=Y+1\}$ (2 and precondition strengthening)

Thus if P is 'X=Y' then:

$$\vdash \{P\} \ X := Y+1 \{P[Y+1/Y]\}$$

and so by the FOR-rule above, if we take V to be Y, E_1 to be 3 and E_2 to be 1, then

$$\vdash \ \{ \ \ \underbrace{\text{X=3}}_{P \, \text{[3/Y]}} \ \ \} \ \text{FOR} \ \ \text{Y:=3 UNTIL 1 DO X:=Y+1} \ \ \{ \ \ \underbrace{\text{X=2}}_{P \, \text{[1+1/Y]}} \ \ \}$$

This is clearly false: it was specified that if the value of E_1 were greater than the value of E_2 then the FOR-command should have no effect, but in this example it changes the value of X from 3 to 2.

To solve this problem, the FOR-rule can be modified to

$$\frac{ \vdash \{P\} \ C \ \{P \ [V+1/V]\}}{\vdash \{P \ [E_1/V] \ \land \ E_1 \leq E_2\} \ \text{FOR} \ V := E_1 \ \text{UNTIL} \ E_2 \ \text{DO} \ C \ \{P \ [E_2+1/V]\}}$$

If this rule is used on the example above all that can be deduced is

$$\vdash \ \{ \, \texttt{X=3} \ \, \land \quad \underbrace{3 \leq 1}_{\text{never true!}} \, \} \ \, \texttt{FOR} \ \, \texttt{Y:=3} \ \, \texttt{UNTIL 1 DO X:=Y+1} \ \, \{ \, \texttt{X=2} \, \}$$

This conclusion is harmless since it only asserts that X will be changed if the FOR-command is executed in an impossible starting state.

Unfortunately, there is still a bug in our FOR-rule. Suppose we take P to be 'Y=1', then it is straightforward to show that:

$$\vdash \ \underbrace{\{\underbrace{\mathtt{Y=1}}_{P}\}} \ \mathtt{Y:=Y-1} \ \underbrace{\{\underbrace{\mathtt{Y+1=1}}_{P}\ [\mathtt{Y+1/Y}]} \}$$

so by our latest FOR-rule

$$\vdash \{ \underbrace{1=1}_{P \text{ [1/Y]}} \land 1 \leq 1 \} \text{ FOR } \text{ Y} := 1 \text{ UNTIL 1 DO Y} := \text{Y-1} \ \{ \underbrace{2=1}_{P \text{ [1+1/Y]}}]$$

Whatever the command does, it doesn't lead to a state in which 2=1. The problem is that the body of the FOR-command modifies the controlled variable. It is not surprising that this causes problems, since it was explicitly assumed that the body didn't modify the controlled variable when we motivated the FOR-rule. It turns out that problems also arise if any variables in the expressions E_1 and E_2 (which specify the upper and lower bounds) are modified. For example, taking P to be Z=Y, then it is straightforward to show

$$\vdash \{\underbrace{Z=Y}_{P}\} \ Z:=Z+1 \{\underbrace{Z=Y+1}_{P \ [Y+1/Y]}\}$$

hence the rule allows us the following to be derived:

$$\vdash \ \{ \ \ \, \underbrace{\mathbf{Z=1}}_{P \ \texttt{[1/Y]}} \ \ \, \land \ \, \mathbf{1} \leq \mathbf{Z} \} \ \, \texttt{FOR} \ \ \, \mathbf{Y:=1} \ \, \texttt{UNTIL Z DO Z:=Z+1} \ \ \, \{ \ \ \, \underbrace{\mathbf{Z=Z+1}}_{P \ \texttt{[Z+1/Y]}} \ \ \, \}$$

This is clearly wrong as one can never have Z=Z+1 (subtracting Z from both sides would give 0=1). One might think that this is not a problem because the FOR-command would never terminate. In some languages this might be the case, but the semantics of our language were carefully defined in such a way that FOR-commands always terminate (see the beginning of this section).

To rule out the problems that arise when the controlled variable or variables in the bounds expressions, are changed by the body, we simply impose a side condition on the rule that stipulates that the rule cannot be used in these situations. The final rule is thus:

The FOR-rule

$$\frac{\vdash \{P \land (E_1 \leq V) \land (V \leq E_2)\} \ C \ \{P \lceil V + 1/V \rceil\}}{\vdash \{P \lceil E_1/V \rceil \land (E_1 \leq E_2)\} \ \text{FOR} \ V := E_1 \ \text{UNTIL} \ E_2 \ \text{DO} \ C \ \{P \lceil E_2 + 1/V \rceil\}}$$

where neither V, nor any variable occurring in E_1 or E_2 , is assigned to in the command C.

This rule does not enable anything to be deduced about FOR-commands whose body assigns to variables in the bounds expressions. This precludes

such assignments being used if commands are to be reasoned about. The strategy of only defining rules of inference for non-tricky uses of constructs helps ensure that programs are written in a perspicuous manner. It is possible to devise a rule that does cope with assignments to variables in bounds expressions, but it is not clear whether it is a good idea to have such a rule.

The FOR-axiom

To cover the case when $E_2 < E_1$, we need the FOR-axiom below.

The FOR-axiom

$$\vdash \ \{P \land (E_2 < E_1)\} \ \text{FOR} \ V := E_1 \ \text{UNTIL} \ E_2 \ \text{DO} \ C \ \{P\}$$

This says that when E_2 is less than E_1 the FOR-command has no effect.

Example: By the assignment axiom and precondition strengthening

$$\vdash \{X = ((N-1)\times N) \text{ DIV } 2\} X := X+N \{X=(N\times(N+1)) \text{ DIV } 2\}$$

Strengthening the precondition of this again yields

$$\vdash \{(X=((N-1\times N) \ DIV \ 2) \land (1 \le N) \land (N \le M) \ X:=X+N \ \{X=(N\times (N+1)) \ DIV \ 2\}$$

Hence by the FOR-rule

$$\vdash \{(X=((1-1)\times 1) \text{ DIV } 2) \land (1\leq M)\}$$
FOR N:=1 UNTIL M DO X:=X+N
$$\{X=(M\times (M+1)) \text{ DIV } 2\}$$

Hence

$$\vdash \{(\texttt{X=0}) \land (\texttt{1} \leq \texttt{M}) \texttt{ FOR N:=1 UNTIL M DO X:=X+N } \{\texttt{X=(M} \times (\texttt{M+1})) \texttt{ DIV 2}\}$$

Note that if

- (i) $\vdash \{P\} \ C \{P[V+1/V]\}, \text{ or }$
- (ii) $\vdash \{P \land (E_1 < V)\} \ C \{P [V+1/V]\}, \text{ or }$
- (iii) $\vdash \{P \land (V < E_2)\} \ C \{P [V+1/V]\}$

then by precondition strengthening one can infer

$$\vdash \{P \land (E_1 < V) \land (V < E_2)\} \ C \ \{P[V+1/V]\}$$

Exercise 15

Show that

```
⊢ {M≥1}
BEGIN
X:=0;
FOR N:=1 UNTIL M DO X:=X+N
END
{X=(M×(M+1)) DIV 2}
```

2.1.12 Arrays

Floyd-Hoare logic can be extended to cope with arrays so that, for example, the correctness of inplace sorting programs can be verified. However, it is not as straightforward as one might expect to do this. The main problem is that the assignment axiom does not apply to array assignments of the form $A(E_1) := E_2$ (where A is an array variable and E_1 and E_2 are expressions).

One might think that the axiom in Section 2.1.1 could be generalized to

$$\vdash \{P[E_2/A(E_1)]\} \ A(E_1) := E_2 \ \{P\}$$

where ' $P[E_2/A(E_1)]$ ' denotes the result of substituting E_2 for all occurrences of $A(E_1)$ throughout P. Alas, this does not work. Consider the following case:

$$P \equiv \text{`A(Y)=0'}, \qquad E_1 \equiv \text{`X'}, \qquad E_2 \equiv \text{`1'}$$

Since A(X) does not occur in P, it follows that P[1/A(X)] = P, and hence the generalized axiom yields

$$\vdash \{A(Y)=0\} A(X):=1 \{A(Y)=0\}$$

This specification is clearly false if X=Y. To avoid this, the array assignment axiom must take into account the possibility that changes to A(X) may also change A(Y), A(Z), ... (since X might equal Y, Z, ...).

We will not go into details of the Floyd-Hoare logic of arrays here, but a thorough treatment can be found in more advanced books (e.g. [63, 1, 26]).

2.2 Soundness and completeness

It is clear from the discussion of the FOR-rule in Section 2.1.11 that it is not always straightforward to devise correct rules of inference. As discussed at the beginning of Chapter 2, it is very important that the axioms and rules be sound. There are two approaches to ensure this:

- (i) Define the language by the axioms and rules of the logic.
- (ii) Prove that the logic fits the language.

Approach (i) is called axiomatic semantics. The idea is to define the semantics of the language by requiring that it make the axioms and rules of inference true. It is then up to implementers to ensure that the logic matches the language. One snag with this approach is that most existing languages have already been defined in some other way (usually by informal and ambiguous natural language statements). An example of a language defined axiomatically is Euclid [46]. The other snag with axiomatic semantics is that it is known to be impossible to devise complete Floyd-Hoare logics for certain constructs (this is discussed further below). It could be argued that this is not a snag at all but an advantage, because it forces programming languages to be made logically tractable. I have some sympathy for this latter view; it is clearly not the position taken by the designers of Ada

Approach (ii) requires that the axioms and rules of the logic be proved valid. To do this, a mathematical model of states is constructed and then a function, **Meaning** say, is defined which takes an arbitrary command C to a function **Meaning** (C) from states to states. Thus **Meaning** (C) (s) denotes the state resulting from executing command C in state s. The specification $\{P\}C\{Q\}$ is then defined to be true if whenever P is true in a state s and **Meaning** (C) (s) = s' then Q is true in state s'. It is then possible to attempt to prove rigorously that all the axioms are true and that the rules of inference lead from true premisses to true conclusions. Actually carrying out this proof is likely to be quite tedious, especially if the programming language is at all complicated, and there are various technical details which require care (e.g. defining **Meaning** to correctly model non-termination). The precise formulation of such soundness proofs is beyond the scope of this book, but details can be found in the text by Loeckx and Sieber [45].

Even if we are sure that our logic is sound, how can we be sure that every true specification can be proved? It might be the case that for some particular P, Q and C the specification $\{P\}C\{Q\}$ was true, but the rules of our logic were too weak to prove it (see Exercise 7 on page 21 for an example). A logic is said to be *complete* if every true statement in it is provable.

2.3. Some exercises 33

There are various subtle technical problems in formulating precisely what it means for a Floyd-Hoare logic to be complete. For example, it is necessary to distinguish incompleteness arising due to incompleteness in the assertion language (e.g. arithmetic) from incompleteness due to inadequate axioms and rules for programming language constructs. The completeness of a Floyd-Hoare logic must thus be defined independently of that of its assertion language. Good introductions to this area can be found in Loeckx and Sieber [45] and Clarke's paper [11]. Clarke's paper also contains a discussion of his important results showing the impossibility of giving complete inference systems for certain combinations of programming language constructs. For example, he proves that it is impossible to give a sound and complete system for any language combining procedures as parameters of procedure calls, recursion, static scopes, global variables and internal procedures as parameters of procedure calls. These features are found in Algol 60, which thus cannot have a sound and complete Floyd-Hoare logic.

2.3 Some exercises

The exercises in this section have been taken from various sources, including Alagić and Arbib's book [1] and Cambridge University Tripos examinations.

Exercise 16

The exponentiation function exp satisfies:

```
exp(m,0) = 1

exp(m,n+1) = m \times exp(m,n)
```

Devise a command C that uses repeated multiplication to achieve the following partial correctness specification:

```
{X = x \land Y = y \land Y \ge 0} \ C \ {Z = exp(x, y) \land X = x \land Y = y}
```

Prove that your command C meets this specification. \square

Exercise 17

Show that

```
├ {M≥0}
BEGIN
X:=0;
FOR N:=1 UNTIL M DO X:=X+N
END
{X=(M×(M+1)) DIV 2}
```

WHILE ¬ODD(X) DO

S := S+Y;X := X-1

 $\begin{array}{l} \mathtt{END} \\ \{\mathtt{S} = \mathtt{x} \! \times \! \mathtt{y}\} \end{array}$

BEGIN $Y:=2\times Y$; X:=X DIV 2 END;

Exercise 20

Deduce:

```
H {X=x \( \times Y=y\)}
BEGIN
S:=0;
WHILE ¬(X=0) DO
BEGIN
WHILE ¬ODD(X) DO
BEGIN Y:=2×Y; X:=X DIV 2 END;
S:=S+Y;
X:=X-1
END
END
{S = x×y}
```

2.3. Some exercises

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Exercise 21

Prove the following invariant property.

```
├ {S = (x-X) × y}
BEGIN
VAR R;
R:=0;
WHILE ¬(R=Y) DO
BEGIN S:=S+1; R:=R+1 END;
X:=X-1
END
{S = (x-X) × y}
```

Hint: Show that $S = (x-X) \times y + R$ is an invariant for S:=S+1; R:=R+1. \square

Exercise 22

Deduce:

Exercise 23

Using $(P \times X^{\mathbb{N}} = x^n) \wedge \neg (X=0) \wedge (N>0)$ as an invariant, deduce:

END

```
\vdash \{X=x \land N=n\}
        BEGIN
           P := 1;
           IF \neg(X=0)
             THEN
                WHILE \neg(N=0) DO
                  BEGIN
                     IF ODD(N) THEN P := P \times X;
                     N:=N DIV 2;
                     X := X \times X
                   END
             ELSE P:=0
        END
         \{P = x^n\}
Exercise 24
Prove that the command
      BEGIN
        Z := 0;
        WHILE \neg(X=0) DO
           BEGIN
             IF ODD(X) THEN Z:=Z+Y;
             Y:=Y\times 2;
             X:=X DIV 2
           END
      END
computes the product of the initial values of X and Y and leaves the result
in Z. \square
Exercise 25
Prove that the command
      BEGIN
        Z := 1;
        WHILE N>O DO
           BEGIN
             IF ODD(N) THEN Z := Z \times X;
             N:=N DIV 2;
             \mathtt{X}:=\mathtt{X}\times\mathtt{X}
           END
```

2.3. Some exercises

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assigns x^n to \mathbf{Z} , where x and n are the initial values of \mathbf{X} and \mathbf{N} respectively and we assume $n \geq 0$. \square

Exercise 26

Devise a proof rule for a command

```
REPEAT command UNTIL statement
```

The meaning of REPEAT C UNTIL S is that C is executed and then S is tested; if the result is true, then nothing more is done, otherwise the whole REPEAT command is repeated. Thus REPEAT C UNTIL S is equivalent to C; WHILE S DO C. \Box

Exercise 27

Use your REPEAT rule to deduce:

```
\label{eq:continuous_state} \begin{array}{ll} \vdash & \{ \texttt{S} = \texttt{C} + \texttt{R} \ \land \ \texttt{R} < \texttt{Y} \} \\ & \texttt{REPEAT} \\ & \texttt{S} : = \texttt{S} + \texttt{1} \ ; \ \texttt{R} : = \texttt{R} + \texttt{1} \\ & \texttt{UNTIL} \ \texttt{R} = \texttt{Y} \\ & \{ \texttt{S} = \texttt{C} + \texttt{Y} \} \end{array}
```

Exercise 28

Use your REPEAT rule to deduce:

```
├ {X=x ∧ Y=y}
BEGIN
S:=0;
REPEAT
R:=0;
REPEAT
S:=S+1; R:=R+1
UNTIL R=Y;
X:=X-1
UNTIL X=0
END
{S = x×y}
```

Exercise 29

Assume gcd(X,Y) satisfies:

```
⊢ (X>Y) ⇒ gcd(X,Y)=gcd(X-Y,Y)
⊢ gcd(X,Y)=gcd(Y,X)
⊢ gcd(X,X)=X
```

```
Prove:
     \vdash \{(A>0) \land (B>0) \land (gcd(A,B)=gcd(X,Y))\}
        WHILE A>B DO A:=A-B;
        WHILE B>A DO B:=B-A
        \{(0 < B) \land (B \le A) \land (gcd(A,B)=gcd(X,Y))\}
   Hence, or otherwise, use your rule for REPEAT commands to prove:
     \vdash {A=a \land B=b}
        REPEAT
          WHILE A>B DO A:=A-B;
          WHILE B>A DO B:=B-A
        UNTIL A=B
        \{A=B \land A=gcd(a,b)\}
Exercise 30
Prove:
     \vdash \{N \ge 1\}
        BEGIN
          PROD=0;
          FOR X:=1 UNTIL N DO PROD := PROD+M
        \{PROD = M \times N\}
Exercise 31
Prove:
     \vdash \{X>0 \land Y>0\}
        BEGIN
          S:=0;
          FOR I:=1 UNTIL X DO
             FOR J:=I UNTIL Y DO
               S:=S+1
        END
        \{S = X \times Y\}
```

Mechanizing Program Verification

The architecture of a simple program verifier is described. Its operation is justified with respect to the rules of Floyd-Hoare logic.

After doing only a few exercises, the following two things will be painfully clear:

- (i) Proofs are typically long and boring (even if the program being verified is quite simple).
- (ii) There are lots of fiddly little details to get right, many of which are trivial (e.g. proving $\vdash (R=X \land Q=0) \Rightarrow (X=R+Y\times Q)$).

Many attempts have been made (and are still being made) to automate proof of correctness by designing systems to do the boring and tricky bits of generating formal proofs in Floyd-Hoare logic. Unfortunately logicians have shown that it is impossible in principle to design a decision procedure to decide automatically the truth or falsehood of an arbitrary mathematical statement [58]. However, this does not mean that one cannot have procedures that will prove many useful theorems. The non-existence of a general decision procedure merely shows that one cannot hope to prove everything automatically. In practice, it is quite possible to build a system that will mechanize many of the boring and routine aspects of verification. This chapter describes one commonly taken approach to doing this.

Although it is impossible to decide automatically the truth or falsity of arbitrary statements, it is possible to check whether an arbitrary formal proof is valid. This consists in checking that the results occurring on each line of the proof are indeed either axioms or consequences of previous lines. Since proofs of correctness of programs are typically very long and boring,

they often contain mistakes when generated manually. It is thus useful to check proofs mechanically, even if they can only be generated with human assistance.

3.1 Overview

In the previous chapter it was shown how to prove $\{P\}C\{Q\}$ by proving properties of the components of C and then putting these together (with the appropriate proof rule) to get the desired property of C itself. For example, to prove $\vdash \{P\}C_1; C_2\{Q\}$ first prove $\vdash \{P\}C_1\{R\}$ and $\vdash \{R\}C_2\{Q\}$ (for suitable R), and then deduce $\vdash \{P\}C_1; C_2\{Q\}$ by the sequencing rule.

This process is called forward proof because one moves forward from axioms via rules to conclusions. In practice, it is more natural to work backwards: starting from the goal of showing $\{P\}C\{Q\}$ one generates subgoals, subsubgoals etc. until the problem is solved. For example, suppose one wants to show:

$$\{X=x \land Y=y\} R:=X; X:=Y; Y:=R \{Y=x \land X=y\}$$

then by the assignment axiom and sequencing rule it is sufficient to show the subgoal

$$\{X=x \land Y=y\} R:=X; X:=Y \{R=x \land X=y\}$$

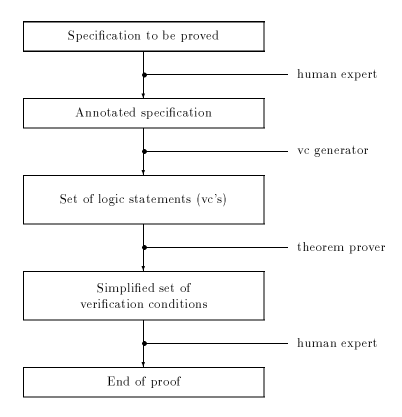
(because \vdash {R=x \land X=y} Y:=R {Y=x \land X=y}). By a similar argument this subgoal can be reduced to

$$\{ \hspace{.05cm} \texttt{X=x} \hspace{.2cm} \wedge \hspace{.2cm} \texttt{Y=y} \} \hspace{.2cm} \texttt{R:=X} \hspace{.2cm} \{ \hspace{.05cm} \texttt{R=x} \hspace{.2cm} \wedge \hspace{.2cm} \texttt{Y=y} \}$$

which clearly follows from the assignment axiom.

This chapter describes how such a *goal oriented* method of proof can be formalized; in Chapter 11 a complete LISP program verifier is given to illustrate how it can be mechanized.

The verification system described here can be viewed as a proof checker that also provides some help with generating proofs. The following diagram gives an overview of the system. 3.1. Overview 41



The system takes as input a partial correctness specification annotated with mathematical statements describing relationships between variables. From the annotated specification the system generates a set of purely mathematical statements, called *verification conditions* (or ve's). In Section 3.5 it is shown that if these verification conditions are provable, then the original specification can be deduced from the axioms and rules of Floyd-Hoare logic.

The verification conditions are passed to a theorem prover program which attempts to prove them automatically; if it fails, advice is sought from the user. We will concentrate on those aspects pertaining to Floyd-Hoare logic and say very little about theorem proving here. Chapter 10 contains the description of a very simple theorem prover based on rewriting and implemented in LISP. It is powerful enough to do most of the examples discussed in this chapter automatically.

The aim of much current research is to build systems which reduce the role of the slow and expensive human expert to a minimum. This can be

achieved by:

- reducing the number and complexity of the annotations required, and
- increasing the power of the theorem prover.

The next section explains how verification conditions work. In Section 3.5 their use is justified in terms of the axioms and rules of Floyd-Hoare logic. Besides being the basis for mechanical verification systems, verification conditions are a useful way of doing proofs by hand.

3.2 Verification conditions

The following sections describe how a goal oriented proof style can be formalized. To prove a goal $\{P\}C\{Q\}$, three things must be done. These will be explained in detail later, but here is a quick overview:

- (i) The program C is annotated by inserting into it statements (often called assertions) expressing conditions that are meant to hold at various intermediate points. This step is tricky and needs intelligence and a good understanding of how the program works. Automating it is a problem of artificial intelligence.
- (ii) A set of logic statements called *verification conditions* (vc's for short) is then generated from the annotated specification. This process is purely mechanical and easily done by a program.
- (iii) The verification conditions are proved. Automating this is also a problem of artificial intelligence.

It will be shown that if one can prove all the verification conditions generated from $\{P\}C\{Q\}$ (where C is suitably annotated), then $\vdash \{P\}C\{Q\}$.

Since verification conditions are just mathematical statements, one can think of step 2 above as the 'compilation', or translation, of a verification problem into a conventional mathematical problem.

The following example will give a preliminary feel for the use of verification conditions.

Suppose the goal is to prove (see the example on page 25)

```
 \begin{cases} T \\ BEGIN \\ R:=X; \\ Q:=0; \\ WHILE \ Y \leq R \ DO \\ BEGIN \ R:=R-Y; \ Q:=Q+1 \ END \\ END \\ \{X=R+Y \times Q \ \land \ R < Y \} \end{cases}
```

This first step (1 above) is to insert annotations. A suitable annotated specification is:

```
 \begin{array}{l} \{T\} \\ \text{BEGIN} \\ \text{R:=X;} \\ \text{Q:=0;} \ \{\text{R=X} \ \land \ \text{Q=0}\} \ \longleftarrow P_1 \\ \text{WHILE } \text{Y$\le$R DO } \{\text{X} = \text{R+Y}{\times}\text{Q}\} \ \longleftarrow P_2 \\ \text{BEGIN } \text{R:=R-Y;} \ \text{Q:=Q+1 END} \\ \text{END} \\ \{\text{X} = \text{R+Y}{\times}\text{Q} \ \land \ \text{R<Y}\} \end{array}
```

The annotations P_1 and P_2 state conditions which are intended to hold whenever control reaches them. Control only reaches the point at which P_1 is placed once, but it reaches P_2 each time the WHILE body is executed and whenever this happens P_2 (i.e. $X=R+Y\times Q$) holds, even though the values of R and Q vary. P_2 is an invariant of the WHILE-command.

The second step (2 above), which has yet to be explained, will generate the following four verification conditions:

```
(i) T \Rightarrow (X=X \land 0=0)

(ii) (R=X \land Q=0) \Rightarrow (X = R+(Y\times Q))

(iii) (X = R+(Y\times Q)) \land Y \leq R \Rightarrow (X = (R-Y)+(Y\times (Q+1)))

(iv) (X = R+(Y\times Q)) \land \neg (Y\leq R) \Rightarrow (X = R+(Y\times Q) \land R\leq Y)
```

Notice that these are statements of arithmetic; the constructs of our programming language have been 'compiled away'.

The third step (3 above) consists in proving these four verification conditions. They are all easy and are proved automatically by the theorem prover described in Chapter 11 (see page 212). The steps are now explained in detail.

3.3 Annotation

An annotated command is a command with statements (called assertions) embedded within it. A command is said to be properly annotated if statements have been inserted at the following places:

- (i) Before each command C_i (where i > 1) in a sequence $C_1; C_2; \ldots; C_n$ which is *not* an assignment command,
- (ii) After the word DO in WHILE and FOR commands.

Intuitively, the inserted assertions should express the conditions one expects to hold *whenever* control reaches the point at which the assertion occurs.

A properly annotated specification is a specification $\{P\}C\{Q\}$ where C is a properly annotated command.

Example: To be properly annotated, assertions should be at points ① and ② of the specification below:

```
 \begin{cases} X = n \\ BEGIN \\ Y := 1; & \longleftarrow (1) \\ WHILE & X \neq 0 & DO & \longleftarrow (2) \\ BEGIN & Y := Y \times X; & X := X-1 & END \\ END \\ X = 0 & \land & Y = n ! \end{cases}
```

Suitable statements would be:

```
at ①: \{Y = 1 \land X = n\}
at ②: \{Y \times X! = n!\}
```

The verification conditions generated from an annotated specification $\{P\}C\{Q\}$ are described by considering the various possibilities for C in turn. This process is justified in Section 3.5 by showing that $\vdash \{P\}C\{Q\}$ if all the verification conditions can be proved.

3.4 Verification condition generation

In this section a procedure is described for generating verification conditions for an annotated partial correctness specification $\{P\}C\{Q\}$. This procedure is recursive on C.

Assignment commands

The single verification condition generated by

$$\{P\}\ V := E\ \{Q\}$$

is

$$P \Rightarrow Q[E/V]$$

Example: The verification condition for

$$\{X=0\}\ X:=X+1\ \{X=1\}$$

is

$$X=0 \Rightarrow (X+1)=1$$

(which is clearly true). □

One-armed conditional

The verification conditions generated by

$$\{P\}$$
 IF S THEN C $\{Q\}$

are

- (i) $(P \land \neg S) \Rightarrow Q$
- (ii) the verifications generated by

$${P \land S} C {Q}$$

Example: The verification conditions for

$$\{T\}$$
 IF X<0 THEN X:=-X $\{X \ge 0\}$

are $T \land \neg (X<0) \Rightarrow X \ge 0$ together with the verification conditions for $\{T \land (X<0)\}\ X:=-X\ \{X\ge 0\}$, i.e. $T \land (X<0) \Rightarrow -X\ge 0$. The two vc's are thus:

- (i) $T \land \neg(X>0) \Rightarrow X \ge 0$
- (ii) T \wedge (X<0) \Rightarrow -X \geq 0

These are equivalent to $X \ge 0 \implies X \ge 0$ and $X < 0 \implies -X \ge 0$, respectively, which are both clearly true. \square

Two-armed conditional

The verification conditions generated from

$$\{P\}$$
 IF S THEN C_1 ELSE C_2 $\{Q\}$

are

(i) the verification conditions generated by

$$\{P \land S\} C_1 \{Q\}$$

(ii) the verifications generated by

$$\{P \land \neg S\} C_2 \{Q\}$$

Exercise 32

What are the verification conditions for the following specification?

$$\{T\}$$
 IF $X \ge Y$ THEN MAX:=X ELSE MAX:=Y $\{MAX=max(X,Y)\}$

Do they follow from the assumptions about max(X,Y) given in the example on page 23?

If C_1 ; ...; C_n is properly annotated, then (see page 44) it must be of one of the two forms:

1.
$$C_1$$
; ...; C_{n-1} ; $\{R\}C_n$, or

2.
$$C_1$$
; ...; C_{n-1} ; $V := E$.

where, in both cases, C_1 ; ...; C_{n-1} is a properly annotated command.

Sequences

1. The verification conditions generated by

$$\{P\}\ C_1;\ldots;C_{n-1};\ \{R\}\ C_n\ \{Q\}$$

(where C_n is not an assignment) are:

(a) the verification conditions generated by

$$\{P\}\ C_1;\ldots;C_{n-1}\ \{R\}$$

(b) the verifications generated by

$$\{R\}\ C_n\ \{Q\}$$

2. The verification conditions generated by

$$\{P\}\ C_1; \ldots; C_{n-1}; I := E\ \{Q\}$$

are the verification conditions generated by

$$\{P\}\ C_1; \ldots; C_{n-1}\ \{Q[E/V]\}$$

Example: The verification conditions generated from

$${X=x \land Y=y} R:=X; X:=Y; Y:=R {X=y \land Y=x}$$

are those generated by

$${X=x \land Y=y} R:=X; X:=Y {(X=y \land Y=x)[R/Y]}$$

which, after doing the substitution, simplifies to

$${X=x \land Y=y} R:=X; X:=Y {X=y \land R=x}$$

The verification conditions generated by this are those generated by

$${X=x \land Y=y} R:=X {(X=y \land R=x)[Y/X]}$$

which, after doing the substitution, simplifies to

$$\left\{ \begin{array}{llll} \mathtt{X=x} & \wedge & \mathtt{Y=y} \end{array} \right\} \ \mathtt{R:=X} \ \left\{ \begin{array}{llll} \mathtt{Y=y} & \wedge & \mathtt{R=x} \end{array} \right\}.$$

The only verification condition generated by this is

$$X=x \land Y=y \Rightarrow (Y=y \land R=x)[X/R]$$

which, after doing the substitution, simplifies to

$$X=x \land Y=y \Rightarrow Y=y \land X=x$$

which is obviously true. \Box

The procedure for generating verification conditions from blocks involves checking the syntactic condition that the local variables of the block do not occur in the precondition or postcondition. The need for this is clear from the side condition in the block rule (see page 21); this will be explained in more detail when the procedure for generating verification conditions is justified in Section 3.5.

Blocks

The verification conditions generated by

$$\{P\}$$
 BEGIN VAR V_1 ; ...; VAR V_n ; C END $\{Q\}$

are

- (i) the verification conditions generated by $\{P\}C\{Q\}$, and
- (ii) the syntactic condition that none of V_1, \ldots, V_n occur in either P or Q.

Example: The verification conditions for

```
\{X=x \land Y=y\} BEGIN VAR R; R:=X; X:=Y; Y:=R END \{X=y \land Y=x\}
```

are those generated by $\{X=x \land Y=y\}$ R:=X; X:=Y; Y:=R $\{X=y \land Y=x\}$ (since R does not occur in $\{X=x \land Y=y\}$ or $\{X=y \land Y=x\}$). See the previous example for the verification conditions generated by this. \square

Exercise 33

What are the verification conditions for the following specification?

```
\{X = R+(Y\times Q)\}\ BEGIN\ R:=R-Y;\ Q:=Q+1\ END\ \{X = R+(Y\times Q)\}
```

Exercise 34

What are the verification conditions for the following specification?

$$\{X=x\}$$
 BEGIN VAR X; X:=1 END $\{X=x\}$

Relate your answer to this exercise to your answer to Exercise 7 on page 21.

A correctly annotated specification of a WHILE-command has the form

$$\{P\}$$
 WHILE S DO $\{R\}$ C $\{Q\}$

Following the usage on page 25, the annotation R is called an invariant.

WHILE-commands

The verification conditions generated from

$$\{P\}$$
 WHILE S DO $\{R\}$ C $\{Q\}$

- $\begin{array}{ccc} \text{(i)} & P & \Rightarrow & R \\ \\ \text{(ii)} & R & \wedge & \neg S & \Rightarrow & Q \end{array}$
- (iii) the verification conditions generated by $\{R \land S\}$ $C\{R\}$.

Example: The verification conditions for

are:

- (i) $R=X \land Q=0 \Rightarrow (X = R+(Y\times Q))$
- (ii) $X = R+Y\times Q \land \neg (Y\leq R) \Rightarrow (X = R+(Y\times Q) \land R<Y)$

together with the verification condition for

$$\begin{array}{lll} \left\{ \textbf{X} &= \textbf{R} + (\textbf{Y} \times \textbf{Q}) \ \land \ (\textbf{Y} \leq \textbf{R}) \right\} \\ & \textbf{BEGIN R} := \textbf{R} - \textbf{Y}; \ \textbf{Q} := \textbf{Q} + \textbf{1} \ \textbf{END} \\ \left\{ \textbf{X} = \textbf{R} + (\textbf{Y} \times \textbf{Q}) \right\} \end{array}$$

which (see Exercise 33) consists of the single condition

(iii)
$$X = R+(Y\times Q) \wedge (Y\leq R) \Rightarrow X = (R-Y)+(Y\times (Q+1))$$

The WHILE-command specification is thus true if (i), (ii) and (iii) hold, i.e.

$$\vdash \{R=X \ \land \ Q=0\}$$
 WHILE $Y \le R$ DO
 BEGIN $R:=R-Y$; $Q:=Q+1$ END
 $\{X = R+(Y \times Q) \ \land \ R < Y\}$

if

$$\vdash$$
 R=X \land Q=0 \Rightarrow (X = R+(Y \times Q))

and

$$\vdash X = R + (Y \times Q) \land \neg (Y \leq R) \Rightarrow (X = R + (Y \times Q) \land R < Y)$$

and

$$\vdash X = R + (Y \times Q) \land (Y \leq R) \Rightarrow X = (R - Y) + (Y \times (Q + 1))$$

Exercise 35

What are the verification conditions generated by the annotated program for computing n! (the factorial of n) given in the example on page 44?

A correctly annotated specification of a FOR-command has the form

$$\{P\}$$
 FOR $V:=E_1$ UNTIL E_2 DO $\{R\}$ C $\{Q\}$

FOR-commands

The verification conditions generated from

$$\{P\}$$
 FOR $V:=E_1$ UNTIL E_2 DO $\{R\}$ C $\{Q\}$

are

- (i) $P \Rightarrow R[E_1/V]$
- (ii) $R[E_2+1/V] \Rightarrow Q$
- (iii) $P \wedge E_2 < E_1 \Rightarrow Q$
- (iv) the verification conditions generated by

$$\{R \land E_1 \leq V \land V \leq E_2\} \ C \ \{R[V+1/V]\}$$

(v) the syntactic condition that neither V, nor any variable occurring in E_1 or E_2 , is assigned to inside C.

Example: The verification conditions generated by

are

- (i) X=0 \wedge 1 \leq M \Rightarrow X=((1-1) \times 1) DIV 2
- (ii) $X = (((M+1)-1)\times(M+1))$ DIV $2 \Rightarrow X = (M\times(M+1))$ DIV 2
- (iii) X=0 \land 1 \le M \land M<1 \Rightarrow X = (M \times (M+1)) DIV 2
- (iv) The verification condition generated by

which, after some simplification, is

which is true since

$$\frac{(N-1)\times N}{2} + N = \frac{2N + (N-1)\times N}{2}$$

$$= \frac{2N + N^2 - N}{2}$$

$$= \frac{N + N^2}{2}$$

$$= \frac{N \times (N+1)}{2}$$

(Exercise: justify this calculation in the light of the fact that

$$(x + y)$$
 DIV $z \neq (x$ DIV $z) + (y$ DIV $z)$

as is easily seen by taking x, y and z to be 3, 5 and 8, respectively.)

(v) Neither N or M is assigned to in X:=X+N

3.5 Justification of verification conditions

It will be shown in this section that an annotated specification $\{P\}C\{Q\}$ is provable in Floyd-Hoare logic (i.e. $\vdash \{P\}C\{Q\}$) if the verification conditions generated by it are provable. This shows that the verification conditions are *sufficient*, but not that they are necessary. In fact, the verification conditions are the weakest sufficient conditions, but we will neither make this more precise nor go into details here. An in-depth study of preconditions can be found in Dijkstra's book [16].

It is easy to show (see the exercise below) that the verification conditions are not necessary, i.e. that the verification conditions for $\{P\}C\{Q\}$ not being provable doesn't imply that $\vdash \{P\}C\{Q\}$ cannot be deduced.

Exercise 36

Show that

(i) The verification conditions from the annotated specification

$$\{T\}$$
 WHILE F DO $\{F\}$ X:=0 $\{T\}$

are not provable.

(ii) \vdash {T} WHILE F DO X:=0 {T}

The argument that the verification conditions are sufficient will be by induction on the structure of C. Such inductive arguments have two parts. First, it is shown that the result holds for assignment commands. Second, it is shown that when C is not an assignment command, then if the result holds for the constituent commands of C (this is called the induction hypothesis), then it holds also for C. The first of these parts is called the basis of the induction and the second is called the step. From the basis and the step it follows that the result holds for all commands.

Assignments

The only verification condition for $\{P\}V := E\{Q\}$ is $P \Rightarrow Q[E/V]$. If this is provable, then as $\vdash \{Q[E/V]\}V := E\{Q\}$ (by the assignment axiom on page 16) it follows by precondition strengthening (page 17) that $\vdash \{P\}V := E\{Q\}$.

One-armed conditionals

If the verification conditions for $\{P\}$ IF S THEN C $\{Q\}$ are provable, then $\vdash P \land \neg S \Rightarrow Q$ and all the verification conditions for $\{P \land S\}$ C $\{Q\}$ are provable. Hence by the induction hypothesis $\vdash \{P \land S\}$ C $\{Q\}$ and hence by the one-armed conditional rule (page 23) it follows that $\vdash \{P\}$ IF S THEN C $\{Q\}$.

Two-armed conditionals

If the verification conditions for $\{P\}$ IF S THEN C_1 ELSE C_2 $\{Q\}$ are provable, then the verification conditions for both $\{P \land S\}$ C_1 $\{Q\}$ and $\{P \land \neg S\}$ C_2 $\{Q\}$ are provable. By the induction hypothesis we can assume that $\vdash \{P \land S\}$ C_1 $\{Q\}$ and $\vdash \{P \land \neg S\}$ C_2 $\{Q\}$. Hence by the two-armed conditional rule (page 23) $\vdash \{P\}$ IF S THEN C_1 ELSE C_2 $\{Q\}$.

Sequences

There are two cases to consider:

(i) If the verification conditions for $\{P\}$ C_1 ; ...; C_{n-1} ; $\{R\}$ C_n $\{Q\}$ are provable, then the verification conditions for $\{P\}$ C_1 ; ...; C_{n-1} $\{R\}$ and $\{R\}$ C_n $\{Q\}$ must both be provable and hence by induction we

have $\vdash \{P\}$ $C_1; \ldots; C_{n-1}$ $\{R\}$ and $\vdash \{R\}$ C_n $\{Q\}$. Hence by the sequencing rule (page 19) $\vdash \{P\}$ $C_1; \ldots; C_{n-1}; C_n$ $\{Q\}$.

(ii) If the verification conditions for $\{P\}$ $C_1; \ldots; C_{n-1}; V := E$ $\{Q\}$ are provable, then it must be the case that the verification conditions for $\{P\}$ $C_1; \ldots; C_{n-1}$ $\{Q \mid E/V\}$ are also provable and hence by induction we have \vdash $\{P\}$ $C_1; \ldots; C_{n-1}$ $\{Q \mid E/V\}$. It then follows by the assignment axiom that \vdash $\{Q \mid E/I \mid V := E \mid Q\}$, hence by the sequencing rule \vdash $\{P\}$ $C_1; \ldots; C_{n-1}; V := E \mid Q\}$.

Blocks

If the verification conditions for $\{P\}$ BEGIN VAR V_1 ; ...; VAR V_n ; C END $\{Q\}$ are provable, then the verification conditions for $\{P\}$ C $\{Q\}$ are provable and V_1, \ldots, V_n do not occur in P or Q. By induction $\vdash \{P\}$ C $\{Q\}$ hence by the block rule (page 21) $\vdash \{P\}$ BEGIN VAR V_1 ; ...; VAR V_n ; C END $\{Q\}$.

WHILE-commands

If the verification conditions for $\{P\}$ WHILE S DO $\{R\}$ C $\{Q\}$ are provable, then $\vdash P \Rightarrow R$, $\vdash (R \land \neg S) \Rightarrow Q$ and the verification conditions for $\{R \land S\}$ C $\{R\}$ are provable. By induction $\vdash \{R \land S\}$ C $\{R\}$, hence by the WHILE-rule (page 25) $\vdash \{R\}$ WHILE S DO C $\{R \land \neg S\}$, hence by the consequence rules (see page 19) $\vdash \{P\}$ WHILE S DO C $\{Q\}$.

FOR-commands

Finally, if the verification conditions for

$$\{P\}$$
 FOR $V:=E_1$ UNTIL E_2 DO $\{R\}$ C $\{Q\}$

are provable, then

- (i) $\vdash P \Rightarrow R[E_1/V]$
- (ii) $\vdash R[E_1/V] \Rightarrow Q$
- (iii) $\vdash P \land E_2 < E_1 \Rightarrow Q$
- (iv) The verification conditions for

$$\{R \land E_1 \leq V \land V \leq E_2\} \ C \ \{R[V+1/V]\}$$

are provable.

(v) Neither V, nor any variable in E_1 or E_2 , is assigned to in C.

By induction $\vdash \{R \land E_1 \leq V \land V \leq E_2\} \ C \ \{R[V+1/V]\}$, hence by the FOR-rule

$$\vdash \ \{R \llbracket E_1/V \rrbracket \ \land \ E_1 \leq E_2\} \ \text{FOR} \ V := E_1 \ \text{UNTIL} \ E_2 \ \text{DO} \ C \ \{R \llbracket E_2 + 1/V \rrbracket \}$$

hence by (i), (ii) and the consequence rules

(vi)
$$\vdash \{P \land E_1 \leq E_2\} \text{ FOR } V := E_1 \text{ UNTIL } E_2 \text{ DO } C \{Q\}.$$

Now by the FOR-axiom (page 30)

$$\vdash \{(P \land E_2 < E_1) \land E\} \text{ FOR } V := E_1 \text{ UNTIL } E_2 \text{ DO } C \{P \land E_2 < E_1\},$$

hence by the consequence rules and (iii)

$$\vdash \{P \land E_2 < E_1\} \text{ FOR } V := E_1 \text{ UNTIL } E_2 \text{ DO } C \{Q\}.$$

Combining this last specification with (vi) using specification disjunction (page 19) yields

$$\vdash \{P \land E_2 < E_1) \lor (P \land E_1 \leq E_2)\} \text{ FOR } V := E_1 \text{ UNTIL } E_2 \text{ DO } C \{Q \lor Q\}$$

Now $\vdash Q \lor Q \Rightarrow Q$ and

$$\vdash (P \land E_2 < E_1) \lor (P \land E_1 \leq E_2) \Rightarrow P \land (E_2 < E_1 \lor E_1 \leq E_2)$$

but $\vdash E_2 < E_1 \lor E_1 \leq E_2$, hence

$$\vdash (P \land E_2 < E_1) \lor (P \land E_1 \leq E_2)$$

and so one can conclude:

$$\vdash \{P\} \text{ FOR } V := E_1 \text{ UNTIL } E_2 \text{ DO } C \{Q\}$$

Thus the verification conditions for the FOR-command are sufficient.

Exercise 37

Annotate the specifications in Exercises 17 to 25 (they start on page 33) and then generate the corresponding verification conditions. \Box

Exercise 38

Devise verification conditions for commands of the form

REPEAT
$$C$$
 UNTIL S

(See Exercise 26, page 37.) □

Exercise 39

Do Exercises 27–31 using verification conditions. □

Exercise 40

Show that if no variable occurring in P is assigned to in C, then \vdash $\{P\}$ $C\{P\}$. Hint: Use induction on the structure of C, see page 52. \square

Part II The λ -calculus and Combinators

Introduction to the λ -calculus

The λ -calculus notation for specifying functions is introduced. Various technical definitions are explained and motivated, including the rules of α -, β - and η -conversion.

The λ -calculus (or lambda-calculus) is a theory of functions that was originally developed by the logician Alonzo Church as a foundation for mathematics. This work was done in the 1930s, several years before digital computers were invented. A little earlier (in the 1920s) Moses Schönfinkel developed another theory of functions based on what are now called 'combinators'. In the 1930s, Haskell Curry rediscovered and extended Schönfinkel's theory and showed that it was equivalent to the λ -calculus. About this time Kleene showed that the λ -calculus was a universal computing system; it was one of the first such systems to be rigorously analysed. In the 1950s John McCarthy was inspired by the λ -calculus to invent the programming language LISP. In the early 1960s Peter Landin showed how the meaning of imperative programming languages could be specified by translating them into the λ -calculus. He also invented an influential prototype programming language called ISWIM [42]. This introduced the main notations of functional programming and influenced the design of both functional and imperative languages. Building on this work, Christopher Strachey laid the foundations for the important area of denotational semantics [23, 67]. Technical questions concerning Strachey's work inspired the mathematical logician Dana Scott to invent the theory of domains, which is now one of the most important parts of theoretical computer science. During the 1970s Peter Henderson and Jim Morris took up Landin's work and wrote a number of influential papers arguing that functional programming had important advantages for software engineering [29, 28]. At about the same time David Turner proposed that Schönfinkel and Curry's combinators could be used as the machine code of computers for executing functional programming languages. Such computers could exploit mathematical properties of the

 λ -calculus for the parallel evaluation of programs. During the 1980s several research groups took up Henderson's and Turner's ideas and started working on making functional programming practical by designing special architectures to support it, some of them with many processors.

We thus see that an obscure branch of mathematical logic underlies important developments in programming language theory, such as:

- (i) The study of fundamental questions of computation.
- (ii) The design of programming languages.
- (iii) The semantics of programming languages.
- (iv) The architecture of computers.

4.1 Syntax and semantics of the λ -calculus

The λ -calculus is a notation for defining functions. The expressions of the notation are called λ -expressions and each such expression denotes a function. It will be seen later how functions can be used to represent a wide variety of data and data-structures including numbers, pairs, lists etc. For example, it will be demonstrated how an arbitrary pair of numbers (x,y) can be represented as a λ -expression. As a notational convention, mnemonic names are assigned in **bold** or <u>underlined</u> to particular λ -expressions; for example $\underline{1}$ is the λ -expression (defined in Section 5.3) which is used to represent the number one.

There are just three kinds of λ -expressions:

- (i) Variables: x, y, z etc. The functions denoted by variables are determined by what the variables are bound to in the *environment*. Binding is done by abstractions (see 3 below). We use V, V_1 , V_2 etc. for arbitrary variables.
- (ii) Function applications or combinations: if E_1 and E_2 are λ -expressions, then so is $(E_1 \ E_2)$; it denotes the result of applying the function denoted by E_1 to the function denoted by E_2 . E_1 is called the rator (from 'operator') and E_2 is called the rand (from 'operand'). For example, if $(\underline{m},\underline{n})$ denotes a function representing the pair of numbers m and n (see Section 5.2) and sum denotes the addition function $1 \ \lambda$ -calculus (see Section 5.5), then the application $(\mathbf{sum}(\underline{m},\underline{n}))$ denotes $\underline{m}+\underline{n}$.

¹Note that sum is a λ -expression, whereas + is a mathematical symbol in the 'metalanguage' (i.e. English) that we are using for talking about the λ -calculus.

(iii) Abstractions: if V is a variable and E is a λ -expression, then λV . E is an abstraction with bound variable V and body E. Such an abstraction denotes the function that takes an argument a and returns as result the function denoted by E in an environment in which the bound variable V denotes a. More specifically, the abstraction λV . E denotes a function which takes an argument E' and transforms it into the thing denoted by E[E'/V] (the result of substituting E' for V in E, see Section 4.8). For example, λx . $\mathbf{sum}(x,\underline{1})$ denotes the add-one function.

Using BNF, the syntax of λ -expressions is just:

$$< \lambda \text{-expression} > ::= < \text{variable} >$$

$$| (< \lambda \text{-expression} > < \lambda \text{-expression} >)$$

$$| (\lambda < \text{variable} > . < \lambda \text{-expression} >)$$

If V ranges over the syntax class < variable > and E, E_1, E_2, \dots etc. range over the syntax class $< \lambda$ -expression >, then the BNF simplifies to:

$$E ::= V \mid \underbrace{(E_1 \ E_2)}_{\text{variables}} \mid \underbrace{\lambda V. \ E}_{\text{abstractions}}$$

$$\underset{\text{(combinations)}}{\text{abstractions}}$$

The description of the meaning of λ -expressions just given above is vague and intuitive. It took about 40 years for logicians (Dana Scott, in fact [66]) to make it rigorous in a useful way. We shall not be going into details of this.

Example: $(\lambda x. x)$ denotes the 'identity function': $((\lambda x. x) E) = E. \square$

Example: $(\lambda x. (\lambda f. (f x)))$ denotes the function which when applied to E yields $(\lambda f. (f x))[E/x]$, i.e. $(\lambda f. (f E))$. This is the function which when applied to E' yields (f E)[E'/f] i.e. (E' E). Thus

$$((\lambda x. (\lambda f. (f x))) E) = (\lambda f. (f E))$$

and

$$((\lambda f. (f E)) E') = (E' E)$$

Exercise 41

Describe the function denoted by $(\lambda x. (\lambda y. y))$. \square

Example: Section 5.3 describes how numbers can be represented by λ -expressions. Assume that this has been done and that $\underline{0}$, $\underline{1}$, $\underline{2}$, ... are λ -expressions which represent 0, 1, 2, ..., respectively. Assume also that add is a λ -expression denoting a function satisfying:

$$((\mathbf{add} \ \underline{m}) \ \underline{n}) = m + n.$$

Then $(\lambda x. ((\mathbf{add} \ \underline{1}) \ x))$ is a λ -expression denoting the function that transforms \underline{n} to $\underline{1+n}$, and $(\lambda x. (\lambda y. ((\mathbf{add} \ x)y)))$ is a λ -expression denoting the function that transforms \underline{m} to the function which when applied to \underline{n} yields $\underline{m+n}$, namely $\lambda y. ((\mathbf{add} \ \underline{m})y))$. \square

The relationship between the function **sum** in (ii) at the beginning of this section (page 60) and the function **add** in the previous example is explained in Section 5.5.

4.2 Notational conventions

The following conventions help minimize the number of brackets one has to write.

1. Function application associates to the left, i.e. $E_1 \ E_2 \ \cdots \ E_n$ means $((\cdots (E_1 \ E_2) \ \cdots) \ E_n)$. For example:

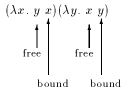
```
\begin{array}{lll} E_1 \ E_2 & \text{means} & (E_1 \ E_2) \\ E_1 \ E_2 \ E_3 & \text{means} & ((E_1 \ E_2) E_3) \\ E_1 \ E_2 \ E_3 \ E_4 & \text{means} & (((E_1 \ E_2) E_3) E_4) \end{array}
```

- 2. $\lambda V. E_1 E_2 \ldots E_n$ means $(\lambda V. (E_1 E_2 \ldots E_n))$. Thus the scope of ' λV ' extends as far to the right as possible.
- 3. $\lambda V_1 \cdots V_n$. E means $(\lambda V_1 \cdot (\cdots \cdot (\lambda V_n \cdot E) \cdots))$. For example:

Example: $\lambda x \ y$. add $y \ x$ means $(\lambda x. \ (\lambda y. \ ((add \ y) \ x)))$.

4.3 Free and bound variables

An occurrence of a variable V in a λ -expression is free if it is not within the scope of a ' λV ', otherwise it is bound. For example



4.4 Conversion rules

In Chapter 5 it is explained how λ -expressions can be used to represent data objects like numbers, strings etc. For example, an arithmetic expression like $(2+3)\times 5$ can be represented as a λ -expression and its 'value' 25 can also be represented as a λ -expression. The process of 'simplifying' $(2+3)\times 5$ to 25 will be represented by a process called *conversion* (or *reduction*). The rules of λ -conversion described below are very general, yet when they are applied to λ -expressions representing arithmetic expressions they simulate arithmetical evaluation.

There are three kinds of λ -conversion called α -conversion, β -conversion and η -conversion (the original motivation for these names is not clear). In stating the conversion rules the notation E[E'/V] is used to mean the result of substituting E' for each *free* occurrence of V in E. The substitution is called *valid* if and only if no free variable in E' becomes bound in E[E'/V]. Substitution is described in more detail in Section 4.8.

The rules of λ -conversion

• α -conversion.

Any abstraction of the form λV . E can be converted to $\lambda V'$. E[V'/V] provided the substitution of V' for V in E is valid.

β-conversion.

Any application of the form $(\lambda V. E_1)$ E_2 can be converted to $E_1 [E_2/V]$, provided the substitution of E_2 for V in E_1 is valid.

• η -conversion.

Any abstraction of the form λV . (E V) in which V has no free occurrence in E can be reduced to E.

The following notation will be used:

- $E_1 \longrightarrow E_2$ means E_1 α -converts to E_2 .
- $E_1 \xrightarrow{\beta} E_2$ means $E_1 \beta$ -converts to E_2 .
- $E_1 \xrightarrow{\eta} E_2$ means $E_1 \eta$ -converts to E_2 .

In Section 4.4.4 below this notation is extended.

The most important kind of conversion is β -conversion; it is the one that can be used to simulate arbitrary evaluation mechanisms. α -conversion is to do with the technical manipulation of bound variables and η -conversion expresses the fact that two functions that always give the same results on the same arguments are equal (see Section 4.7). The next three subsections give further explanation and examples of the three kinds of conversion (note that 'conversion' and 'reduction' are used below as synonyms).

4.4.1 α -conversion

A λ -expression (necessarily an abstraction) to which α -reduction can be applied is called an α -redex. The term 'redex' abbreviates 'reducible expression'. The rule of α -conversion just says that bound variables can be renamed provided no 'name-clashes' occur.

Examples

$$\lambda x \cdot x \xrightarrow{\alpha} \lambda y \cdot y$$

$$\lambda x \cdot f \ x \xrightarrow{\alpha} \lambda y \cdot f \ y$$

It is *not* the case that

$$\lambda x \cdot \lambda y \cdot \mathbf{add} \ x \ y \xrightarrow{\alpha} \lambda y \cdot \lambda y \cdot \mathbf{add} \ y \ y$$

because the substitution $(\lambda y. \text{ add } x y)[y/x]$ is not valid since the y that replaces x becomes bound. \square

4.4.2 β -conversion

A λ -expression (necessarily an application) to which β -reduction can be applied is called a β -redex. The rule of β -conversion is like the evaluation of a function call in a programming language: the body E_1 of the function λV . E_1 is evaluated in an environment in which the 'formal parameter' V is bound to the 'actual parameter' E_2 .

Examples

It is *not* the case that

$$(\lambda x.\ (\lambda y.\ \mathbf{add}\ x\ y))\ (\mathbf{square}\ y) \xrightarrow{\ \beta\ } \lambda y.\ \mathbf{add}\ (\mathbf{square}\ y)\ y$$

because the substition $(\lambda y. \ \mathbf{add} \ x \ y) \lceil (\mathbf{square} \ y)/x \rceil$ is not valid, since y is free in $(\mathbf{square} \ y)$ but becomes bound after substitution for x in $(\lambda y. \ \mathbf{add} \ x \ y)$. \square

It takes some practice to parse λ -expressions according to the conventions of Section 4.2 so as to identify the β -redexes. For example, consider the application:

$$(\lambda x. \lambda y. \mathbf{add} \ x \ y) \ \underline{3} \ \underline{4}.$$

Putting in brackets according to the conventions expands this to:

$$(((\lambda x. (\lambda y. ((\mathbf{add} x) y))) \underline{3}) \underline{4})$$

which has the form:

$$((\lambda x. E) 3) 4$$

where

$$E = (\lambda y. \text{ add } x y)$$

 $(\lambda x. E) \ \underline{3}$ is a β -redex and could be reduced to $E[\underline{3}/x]$.

4.4.3 η -conversion

A λ -expression (necessarily an abstraction) to which η -reduction can be applied is called an η -redex. The rule of η -conversion expresses the property that two functions are equal if they give the same results when applied to the same arguments. This property is called extensionality and is discussed further in Section 4.7. For example, η -conversion ensures that λx . (sin x) and sin denote the same function. More generally, λV . (EV) denotes the function which when applied to an argument E' returns (EV)[E'/V]. If V does not occur free in E then (EV)[E'/V] = (EV). Thus V0 and V1 and V2 both yield the same result, namely V3 when applied to the same arguments and hence they denote the same function.

Examples

$$\lambda x. \text{ add } x \xrightarrow{\eta} \text{add}$$

$$\lambda y. \text{ add } x \text{ } y \xrightarrow{\eta} \text{ add } x$$

It is not the case that

$$\lambda x. \ \mathbf{add} \ x \ x \xrightarrow{\eta} \mathbf{add} \ x$$

because x is free in add x. \square

4.4.4 Generalized conversions

The definitions of $\xrightarrow{\alpha}$, $\xrightarrow{\beta}$ and $\xrightarrow{\eta}$ can be generalized as follows:

• $E_1 \xrightarrow{\alpha} E_2$ if E_2 can be got from E_1 by α -converting any subterm.

- $E_1 \xrightarrow{\beta} E_2$ if E_2 can be got from E_1 by β -converting any subterm.
- $E_1 \xrightarrow{\eta} E_2$ if E_2 can be got from E_1 by η -converting any subterm.

Examples

$$((\lambda x.\ \lambda y.\ \mathbf{add}\ x\ y)\ \underline{3})\ \underline{4} \xrightarrow{\beta} (\lambda y.\ \mathbf{add}\ \underline{3}\ y)\ \underline{4}$$

$$(\lambda y. \ \mathbf{add} \ \underline{3} \ y) \ \underline{4} \xrightarrow{\beta} \mathbf{add} \ \underline{3} \ \underline{4}$$

The first of these is a β -conversion in the generalized sense because $(\lambda y. \ \mathbf{add} \ \underline{3} \ y)\underline{4}$ is obtained from $((\lambda x. \ \lambda y. \ \mathbf{add} \ x \ y)\underline{3})\underline{4}$ (which is not itself a β -redex) by reducing the subexpression $(\lambda x. \ \lambda y. \ \mathbf{add} \ x \ y)\underline{3}$. We will sometimes write a sequence of conversions like the two above as:

$$((\lambda x\,.\,\lambda y.\ \mathbf{add}\ x\ y)\ \underline{3})\ \underline{4} \xrightarrow[\beta]{} (\lambda y.\ \mathbf{add}\ \underline{3}\ y)\ \underline{4} \xrightarrow[\beta]{} \mathbf{add}\ \underline{3}\ \underline{4}$$

Exercise 42

Which of the three β -reductions below are generalized conversions (i.e. reductions of subexpressions) and which are conversions in the sense defined on page 63? \square

(i)
$$(\lambda x. x) \xrightarrow{\underline{1} \xrightarrow{\beta} \underline{1}}$$

(ii)
$$(\lambda y.\ y)\ ((\lambda x.\ x)\ \underline{1}) \xrightarrow{\beta} (\lambda y.\ y)\underline{1} \xrightarrow{\beta} \underline{1}$$

(iii)
$$(\lambda y. \ y) \ ((\lambda x. \ x) \ \underline{1}) \xrightarrow{\beta} (\lambda x. \ x) \ \underline{1} \xrightarrow{\beta} \underline{1}$$

In reductions (ii) and (iii) in the exercise above one starts with the same λ -expression, but reduce redexes in different orders.

An important property of β -reductions is that no matter in which order one does them, one always ends up with equivalent results. If there are several disjoint redexes in an expression, one can reduce them in parallel. Note, however, that some reduction sequences may never terminate. This is discussed further in connection with the normalization theorem of Chapter 7. It is a current hot research topic in 'fifth-generation computing' to design processors which exploit parallel evaluation to speed up the execution of functional programs.

4.5 Equality of λ -expressions

The three conversion rules preserve the meaning of λ -expressions, i.e. if E_1 can be converted to E_2 then E_1 and E_2 denote the same function. This property of conversion should be intuitively clear. It is possible to give a mathematical definition of the function denoted by a λ -expression and then to prove that this function is unchanged by α -, β - or η -conversion. Doing this is surprisingly difficult [67] and is beyond the scope of this book.

We will simply define two λ -expressions to be equal if they can be transformed into each other by a sequence of (forwards or backwards) λ -conversions. It is important to be clear about the difference between equality and identity. Two λ -expressions are identical if they consist of exactly the same sequence of characters; they are equal if one can be converted to the other. For example, λx . x is equal to λy . y, but not identical to it. The following notation is used:

- $E_1 \equiv E_2$ means E_1 and E_2 are identical.
- $E_1 = E_2$ means E_1 and E_2 are equal.

Equality (=) is defined in terms of identity (\equiv) and conversion ($\xrightarrow{\alpha}$, $\xrightarrow{\beta}$ and $\xrightarrow{\eta}$) as follows.

Equality of λ -expressions

If E and E' are λ -expressions then E=E' if $E\equiv E'$ or there exist expressions E_1, E_2, \ldots, E_n such that:

- 1. $E \equiv E_1$
- 2. $E' \equiv E_n$
- 3. For each i either
 - (a) $E_i \xrightarrow{\alpha} E_{i+1}$ or $E_i \xrightarrow{\beta} E_{i+1}$ or $E_i \xrightarrow{\eta} E_{i+1}$ or
 - (b) $E_{i+1} \xrightarrow{\alpha} E_i$ or $E_{i+1} \xrightarrow{\beta} E_i$ or $E_{i+1} \xrightarrow{\eta} E_i$.

Examples

$$(\lambda x. x) \underline{1} = \underline{1}$$

$$(\lambda x. x) ((\lambda y. y) \underline{1}) = \underline{1}$$
$$(\lambda x. \lambda y. \mathbf{add} x y) \underline{3} \underline{4} = \mathbf{add} \underline{3} \underline{4}$$

From the definition of = it follows that:

- (i) For any E it is the case that E = E (equality is reflexive).
- (ii) If E = E', then E' = E (equality is symmetric).
- (iii) If E = E' and E' = E'', then E = E'' (equality is transitive).

If a relation is reflexive, symmetric and transitive then it is called an equivalence relation. Thus = is an equivalence relation.

Another important property of = is that if $E_1 = E_2$ and if E_1' and E_2' are two λ -expressions that only differ in that where one contains E_1 the other contains E_2 , then $E_1' = E_2'$. This property is called *Leibnitz's law*. It holds because the same sequence of reduction for getting from E_1 to E_2 can be used for getting from E_1' to E_2' . For example, if $E_1 = E_2$, then by Leibnitz's law λV . $E_1 = \lambda V$. E_2 .

It is essential for the substitutions in the α - and β -reductions to be valid. The validity requirement disallows, for example, λx . $(\lambda y. x)$ being α -reduced to λy . $(\lambda y. y)$ (since y becomes bound after substitution for x in $\lambda y. x$). If this invalid substitution were permitted, then it would follow by the definition of = that:

$$\lambda x \cdot \lambda y \cdot x = \lambda y \cdot \lambda y \cdot y$$

But then since:

$$(\lambda x . \ (\lambda y \ .x)) \ \underline{1} \ \underline{2} \xrightarrow{\beta} (\lambda y . \ \underline{1}) \ \underline{2} \xrightarrow{\beta} \underline{1}$$

and

$$(\lambda y. \ (\lambda y. \ y)) \ \underline{1} \ \underline{2} \xrightarrow{\beta} (\lambda y. \ y) \ \underline{2} \xrightarrow{\beta} \underline{2}$$

one would be forced to conclude that $\underline{1} = \underline{2}$. More generally by replacing $\underline{1}$ and $\underline{2}$ by any two expressions, it could be shown that any two expressions are equal!

Exercise 43

Find an example which shows that if substitutions in β -reductions are allowed to be invalid, then it follows that any two λ -expressions are equal. \Box

Example: If V_1, V_2, \ldots, V_n are all distinct and none of them occur free in any of E_1, E_2, \ldots, E_n , then

$$(\lambda V_1 \ V_2 \cdots V_n \cdot E) \ E_1 \ E_2 \cdots E_n$$

$$= ((\lambda V_1 \cdot (\lambda V_2 \cdots V_n \cdot E)) E_1) \ E_2 \cdots E_n$$

$$\xrightarrow{\beta} ((\lambda V_2 \cdots V_n \cdot E) [E_1/V_1]) \ E_2 \cdots E_n$$

$$= (\lambda V_2 \cdots V_n \cdot E [E_1/V_1]) E_2 \cdots E_n$$

$$\vdots$$

$$= E [E_1/V_1] [E_2/V_2] \cdots [E_n/V_n]$$

Exercise 44

In the last example, where was the assumption used that V_1, V_2, \ldots, V_n are all distinct and that none of them occur free in any of E_1, E_2, \ldots, E_n ?

Exercise 45

Find an example to show that if $V_1 = V_2$, then even if V_2 is not free in E_1 , it is not necessarily the case that:

$$(\lambda V_1 V_2 . E) E_1 E_2 = E [E_1/V_1] [E_2/V_2]$$

Exercise 46

Find an example to show that if $V_1 \neq V_2$, but V_2 occurs free in E_1 , then it is not necessarily the case that:

$$(\lambda V_1 V_2 . E) E_1 E_2 = E[E_1/V_1][E_2/V_2]$$

4.6 The \longrightarrow relation

In the previous section $E_1 = E_2$ was defined to mean that E_2 could be obtained from E_1 by a sequence of forwards or backwards conversions. A special case of this is when E_2 is got from E_1 using only forwards conversions. This is written $E_1 \longrightarrow E_2$.

Definition of \longrightarrow

If E and E' are λ -expressions, then $E \longrightarrow E'$ if $E \equiv E'$ or there exist expressions E_1, E_2, \ldots, E_n such that:

- 1. $E \equiv E_1$ 2. $E' \equiv E_n$ 3. For each i either $E_i \xrightarrow{\alpha} E_{i+1}$ or $E_i \xrightarrow{\beta} E_{i+1}$ or $E_i \xrightarrow{\eta} E_{i+1}$.

Notice that the definition of \longrightarrow is just like the definition of = on page 68 except that part (b) of 3 is missing.

Exercise 47

Find E, E' such that E = E' but it is not the case that $E \longrightarrow E'$. \square

Exercise 48

[very hard!] Show that if $E_1 = E_2$, then there exists E such that $E_1 \longrightarrow E$ and $E_2 \longrightarrow E$. (This property is called the Church-Rosser theorem. Some of its consequences are discussed in Chapter 7.) \square

4.7 Extensionality

Suppose V does not occur free in E_1 or E_2 and

$$E_1 V = E_2 V$$

Then by Leibnitz's law (see page 69)

$$\lambda V. E_1 V = \lambda V. E_2 V$$

so by η -reduction applied to both sides

$$E_1 = E_2$$

It is often convenient to prove that two λ -expressions are equal using this property, i.e. to prove $E_1 = E_2$ by proving $E_1 V = E_2 V$ for some V not occurring free in E_1 or E_2 . We will refer to such proofs as being by extensionality.

Exercise 49

Show that

$$(\lambda f g x. f x (g x)) (\lambda x y. x) (\lambda x y. x) = \lambda x. x$$

4.8 Substitution

At the beginning of Section 4.4 E[E'/V] was defined to mean the result of substituting E' for each free occurrence of V in E. The substitution was said to be valid if no free variable in E' became bound in E[E'/I]. In the definitions of α - and β -conversion, it was stipulated that the substitutions involved must be valid. Thus, for example, it was only the case that

$$(\lambda V. E_1) E_2 \xrightarrow{\beta} E_1 [E_2/V]$$

as long as the substitution $E_1[E_2/V]$ was valid.

It is very convenient to extend the meaning of E[E'/V] so that we don't have to worry about validity. This is achieved by the definition below which has the property that for all expressions E, E_1 and E_2 and all variables V and V':

$$(\lambda V. E_1) E_2 \longrightarrow E_1[E_2/V]$$
 and $\lambda V. E \longrightarrow \lambda V'. E[V'/V]$

To ensure this property holds, E[E'/V] is defined recursively on the structure of E as follows:

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E	$E\left[E'/V ight]$
V	E'
V' (where $V \neq V'$)	V'
E_1 E_2	$E_1[E'/V]$ $E_2[E'/V]$
$\lambda V. E_1$	$\lambda V. E_1$
$\lambda V'$. E_1 (where $V \neq V'$ and V' is not free in E')	$\lambda V'$. $E_1[E'/V]$
$\lambda V'$. E_1 (where $V \neq V'$ and V' is free in E')	$\lambda V''$. $E_1[V''/V'][E'/V]$ where V'' is a variable not free in E' or E_1

This particular definition of E[E'/V] is based on (but not identical to) the one in Appendix C of [4]. A LISP implementation of it is given in Chapter 12 on page 228.

To illustrate how this works consider $(\lambda y. y. x)[y/x]$. Since y is free in y x we must use the last case of the table above. Since z does not occur in y x or y,

$$(\lambda y. \ y. \ x) [y/x] \equiv \lambda z. \ (y. x) [z/y] [y/x] \equiv \lambda z. \ (z. x) [y/x] \equiv \lambda z. \ z. y$$

In the last line of the table above, the particular choice of V'' is not specified. Any variable not occurring in E' or E_1 will do. In Chapter 12 an implementation of substitution in LISP is given.

A good discussion of substitution can be found in the book by Hindley and Seldin [31] where various technical properties are stated and proved. The following exercise is taken from that book.

Exercise 50

Use the table above to work out

- (i) $(\lambda y. \ x \ (\lambda x. \ x)) [(\lambda y. \ y \ x)/x].$
- (ii) $(y (\lambda z. x z)) [(\lambda y. z y)/x].$

It is straightforward, but rather tedious, to prove from the definition of E[E'/V] just given that indeed

$$(\lambda V. E_1) E_2 \longrightarrow E_1[E_2/V]$$
 and $\lambda V. E \longrightarrow \lambda V'. E[V'/V]$

for all expressions E, E_1 and E_2 and all variables V and V'.

In Chapter 8 it will be shown how the theory of combinators can be used to decompose the complexities of substitution into simpler operations. Instead of combinators it is possible to use the so-called nameless terms of De Bruijn [8]. De Bruijn's idea is that variables can be thought of as 'pointers' to the λ s that bind them. Instead of 'labelling' λ s with names (i.e. bound variables) and then pointing to them via these names, one can point to the appropriate λ by giving the number of levels 'upwards' needed to reach it. For example, λx . λy . x y would be represented by $\lambda \lambda 2$ 1. As a more complicated example, consider the expression below in which we indicate the number of levels separating a variable from the λ that binds it.

$$\lambda x. \lambda y. x y (\lambda y. x y y)$$

In De Bruijn's notation this is $\lambda\lambda 2$ 1 $\lambda 3$ 1 1.

A free variable in an expression is represented by a number bigger than the depth of λ s above it; different free variables being assigned different numbers. For example,

$$\lambda x. (\lambda y. y. x. z) x. y. w$$

would be represented by

$$\lambda(\lambda 1\ 2\ 3)\ 1\ 2\ 4$$

Since there are only two λ s above the occurrence of 3, this number must denote a free variable; similarly there is only one λ above the second occurrence of 2 and the occurrence of 4, so these too must be free variables. Note that 2 could not be used to represent w since this had already been used to represent the free y; we thus chose the first available number bigger than 2 (3 was already in use representing z).

Care must be taken to assign big enough numbers to free variables. For example, the first occurrence of z in λx . z (λy . z) could be represented by 2, but the second occurrence requires 3; since they are the same variable we must use 3.

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Example: With De Bruijn's scheme λx . x (λy . x y y) would be represented by $\lambda 1(\lambda 2 \ 1 \ 1)$. \square

Exercise 51

What λ -expression is represented by $\lambda 2(\lambda 2)$? \square

Exercise 52

Describe an algorithm for computing the De Bruijn representation of the expression E[E'/V] from the representations of E and E'. \square

Representing Things in the λ -calculus

The representation in the λ -calculus of various data objects (e.g. numbers), data-structures (e.g. pairs) and useful functions (e.g. addition) is described. Definition by recursion using the fixed-point operator \mathbf{Y} is explained. It is shown that all the recursive functions can be represented by suitable λ -expressions.

The λ -calculus appears at first sight to be a very primitive language. However, it can be used to represent most of the objects and structures needed for modern programming. The idea is to code these objects and structures in such a way that they have the required properties. For example, to represent the truth values true and false and the Boolean function \neg ('not'), λ -expressions true, false and not are devised with the properties that:

```
not true = false
not false = true
```

To represent the Boolean function \wedge ('and') a λ -expression **and** is devised such that:

```
and true true = true
and true false = false
and false true = false
and false false = false
```

and to represent \vee ('or') an expression or such that:

```
or true true = true
or true false = true
or false true = true
or false false = false
```

The λ -expressions used to represent things may appear completely unmotivated at first. However, the definitions are chosen so that they work together in unison.

We will write

LET
$$\sim = \lambda$$
-expression

to introduce \sim as a new notation. Usually \sim will just be a name such as **true** or **and**. Such names are written in **bold** face, or underlined, to distinguish them from variables. Thus, for example, *true* is a variable but **true** is the λ -expression λx . λy . x (see Section 5.1 below) and 2 is a number but $\underline{2}$ is the λ -expression λf x. f(f x) (see Section 5.3).

Sometimes \sim will be a more complicated form like the conditional notation $(E \to E_1 \mid E_2)$.

5.1 Truth-values and the conditional

This section defines λ -expressions **true**, **false**, **not** and $(E \to E_1 \mid E_2)$ with the following properties:

$$not true = false$$

 $not false = true$

$$(\mathbf{true} \to E_1 \mid E_2) = E_1$$
$$(\mathbf{false} \to E_1 \mid E_2) = E_2$$

The λ -expressions **true** and **false** represent the truth-values true and false, **not** represents the negation function \neg and $(E \rightarrow E_1 \mid E_2)$ represents the conditional 'if E then E_1 else E_2 '.

There are infinitely many different ways of representing the truth-values and negation that work; the ones used here are traditional and have been developed over the years by logicians.

LET
$$\mathbf{true} = \lambda x$$
. λy . x
LET $\mathbf{false} = \lambda x$. λy . y
LET $\mathbf{not} = \lambda t$. t \mathbf{false} \mathbf{true}

It is easy to use the rules of λ -conversion to show that these definitions have the desired properties. For example:

not true =
$$(\lambda t. t$$
 false true $)$ **true** (definition of **not**)

= true false true (
$$\beta$$
-conversion)
= $(\lambda x. \lambda y. x)$ false true (definition of true)
= $(\lambda y.$ false) true (β -conversion)
= false (β -conversion)

Similarly not false = true.

Conditional expressions $(E \to E_1 \mid E_2)$ can be defined as follows:

LET
$$(E \to E_1 \mid E_2) = (E \ E_1 \ E_2)$$

This means that for any λ -expressions E, E_1 and E_2 , $(E \to E_1 \mid E_2)$ stands for $(E \ E_1 \ E_2)$.

The conditional notation behaves as it should:

$$\begin{aligned} (\mathbf{true} \rightarrow E_1 \mid E_2) &= \mathbf{true} \ E_1 \ E_2 \\ &= (\lambda x \ y. \ x) \ E_1 \ E_2 \\ &= E_1 \end{aligned}$$

and

(false
$$\rightarrow E_1 \mid E_2$$
) = false $E_1 \mid E_2$
= $(\lambda x \mid y. \mid y) \mid E_1 \mid E_2$
= E_2

Exercise 53

Let **and** be the λ -expression $\lambda x \ y$. $(x \to y \mid \mathbf{false})$. Show that:

and true true = true and true false = false and false true = false and false false = false

Exercise 54

Devise a λ -expression or such that:

```
or true true = true
or true false = true
or false true = true
or false false = false
```

5.2 Pairs and tuples

The following abbreviations represent pairs and n-tuples in the λ -calculus.

LET
$$\mathbf{fst} = \lambda p.\ p\ \mathbf{true}$$

LET $\mathbf{snd} = \lambda p.\ p\ \mathbf{false}$
LET $(E_1,E_2) = \lambda f.\ f\ E_1\ E_2$

 (E_1, E_2) is a λ -expression representing an ordered pair whose first component (i.e. E_1) is accessed with the function **fst** and whose second component (i.e. E_2) is accessed with **snd**. The following calculation shows how the various definitions co-operate together to give the right results.

$$\begin{aligned} \textbf{fst} \ (E_1, E_2) &= (\lambda p. \ p \ \textbf{true}) \ (E_1, E_2) \\ &= (E_1, E_2) \ \textbf{true} \\ &= (\lambda f. \ f \ E_1 \ E_2) \ \textbf{true} \\ &= \textbf{true} \ E_1 \ E_2 \\ &= (\lambda x \ y. \ x) \ E_1 \ E_2 \\ &= E_1 \end{aligned}$$

Exercise 55 Show that $\operatorname{\mathbf{snd}}(E_1,E_2)=E_2$. \square

A pair is a data-structure with two components. The generalization to n components is called an n-tuple and is easily defined in terms of pairs.

LET
$$(E_1, E_2, \dots, E_n) = (E_1, (E_2, (\dots (E_{n-1}, E_n) \dots)))$$

 (E_1, \ldots, E_n) is an *n*-tuple with components E_1, \ldots, E_n and length n. Pairs are 2-tuples. The abbreviations defined next provide a way of extracting the components of n-tuples.

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LET
$$E \downarrow^n 1 = \mathbf{fst} \ E$$

LET $E \downarrow^n 2 = \mathbf{fst}(\mathbf{snd} \ E)$

$$\vdots$$

LET $E \downarrow^n i = \mathbf{fst}(\underbrace{\mathbf{snd}(\mathbf{snd}(\cdots(\mathbf{snd} \ E) \cdots)))}_{i-1 \ \mathbf{snds}} E) \cdots))) (if $i < n$)

$$\vdots$$

LET $E \downarrow^n n = \underbrace{\mathbf{snd}(\mathbf{snd}(\cdots(\mathbf{snd} \ E) \cdots)))}_{n-1 \ \mathbf{snds}}$$

It is easy to see that these definitions work, for example:

$$(E_1, E_2, \dots, E_n) \stackrel{n}{\downarrow} 1 = (E_1, (E_2, (\dots))) \stackrel{n}{\downarrow} 1$$

$$= \mathbf{fst} (E_1, (E_2, (\dots)))$$

$$= E_1$$

$$(E_1, E_2, \dots, E_n) \stackrel{n}{\downarrow} 2 = (E_1, (E_2, (\dots))) \stackrel{n}{\downarrow} 2$$

$$= \mathbf{fst} \ (\mathbf{snd} \ (E_1, (E_2, (\dots))))$$

$$= \mathbf{fst} \ (E_2, (\dots))$$

$$= E_2$$

In general $(E_1, E_2, \dots, E_n) \stackrel{n}{\downarrow} i = E_i$ for all i such that $1 \leq i \leq n$.

Convention

We will usually just write $E \downarrow i$ instead of $E \downarrow^n i$ when it is clear from the context what n should be. For example,

$$(E_1, \ldots, E_n) \downarrow i = E_i$$
 (where $1 \le i \le n$)

5.3 Numbers

There are many ways to represent numbers by λ -expressions, each with their own advantages and disadvantages [72, 40]. The goal is to define for each number n a λ -expression \underline{n} that represents it. We also want to define λ -expressions to represent the primitive arithmetical operations. For

example, we will need λ -expressions **suc**, **pre**, **add** and **iszero** representing the successor function $(n \mapsto n+1)$, the predecessor function $(n \mapsto n-1)$, addition and a test for zero, respectively. These λ -expressions will represent the numbers correctly if they have the following properties:

suc
$$\underline{n} = \underline{n+1}$$
 (for all numbers n)

pre $\underline{n} = \underline{n-1}$ (for all numbers n)

add $\underline{m} \ \underline{n} = \underline{m+n}$ (for all numbers m and n)

iszero $\underline{0} = \mathbf{true}$

iszero (suc \underline{n}) = false

The representation of numbers described here is the original one due to Church. In order to explain this it is convenient to define f^n x to mean n applications of f to x. For example,

$$f^5 x = f(f(f(f(f(x)))))$$

By convention f^0 x is defined to mean x. More generally:

LET
$$E^0$$
 E' = E'

LET E^n E' = $\underbrace{E(E(\cdots(E) E')\cdots)}_{n Es}$

Note that $E^n(EE') = E^{n+1} E' = E(E^n E')$; we will use the fact later.

Example:

$$f^4x = f(f(f(f(x)))) = f(f^3x) = f^3(f(x))$$

Using the notation just introduced we can now define Church's numerals. Notice how the definition of the λ -expression \underline{n} below encodes a unary representation of n.

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```
LET \underline{0} = \lambda f \ x. \ x

LET \underline{1} = \lambda f \ x. \ f \ x

LET \underline{2} = \lambda f \ x. \ f(f \ x)

\vdots

LET \underline{n} = \lambda f \ x. \ f^n \ x

\vdots
```

The representations of **suc**, **add** and **iszero** are now magically pulled out of a hat. The best way to see how they work is to think of them as operating on unary representations of numbers. The exercises that follow should help.

LET
$$\mathbf{suc} = \lambda n \ f \ x. \ n \ f(f \ x)$$

LET $\mathbf{add} = \lambda m \ n \ f \ x. \ m \ f \ (n \ f \ x)$
LET $\mathbf{iszero} = \lambda n. \ n \ (\lambda x. \ \mathbf{false}) \ \mathbf{true}$

Exercise 56

Show:

- (i) **suc** 0 = 1
- (ii) **suc** 5 = 6
- (iii) iszero $\underline{0} = \mathbf{true}$
- (iv) iszero $\underline{5} = \mathbf{false}$
- (v) **add** 0 1 = 1
- (vi) **add** $\underline{2} \ \underline{3} = \underline{5}$

Exercise 57

Show for all numbers m and n:

(i) **suc** $\underline{n} = \underline{n+1}$

- (ii) iszero (suc \underline{n}) = false
- (iii) add $\underline{0} \ \underline{n} = \underline{n}$
- (iv) add $\underline{m} \ \underline{0} = \underline{m}$
- (v) add $\underline{m} \underline{n} = \underline{m+n}$

The predecesor function is harder to define than the other primitive functions. The idea is that the predecessor of \underline{n} is defined by using $\lambda f(x)$. $f^n(x)$ (i.e. \underline{n}) to obtain a function that applies f only n-1 times. The trick is to 'throw away' the first application of f in f^n . To achieve this, we first define a function **prefn** that operates on pairs and has the property that:

- (i) **prefn** f (**true**, x) = (**false**, x)
- (ii) **prefn** f (**false**, x) = (**false**, f x)

From this it follows that:

- (iii) (**prefn** f)ⁿ (**false**, x) = (**false**, fⁿ x)
- (iv) $(\mathbf{prefn}\ f)^n\ (\mathbf{true}, x) = (\mathbf{false}, f^{n-1}\ x)$ (if n > 0)

Thus n applications of **prefn** to (\mathbf{true}, x) result in n-1 applications of f to x. With this idea, the definition of the predecessor function \mathbf{pre} is straightforward. Before giving it, here is the definition of \mathbf{prefn} :

LET prefn =
$$\lambda f$$
 p. (false, (fst $p \to \text{snd } p \mid (f(\text{snd } p))))$

Exercise 58

Show **prefn** $f(b, x) = (\mathbf{false}, (b \to x \mid f(x)))$ and hence:

- (i) **prefn** f (**true**, x) = (**false**, x)
- (ii) **prefn** f (**false**, x) = (**false**, f x)
- (iii) (**prefn** f)ⁿ (**false**, x) = (**false**, fⁿ x)
- (iv) (**prefn** f)ⁿ (**true**, x) = (**false**, f^{n-1} x) (if n > 0)

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The predecessor function **pre** can now be defined.

LET pre =
$$\lambda n \ f \ x. \ \mathbf{snd} \ (n \ (\mathbf{prefn} \ f) \ (\mathbf{true}, x))$$

It follows that if n > 0

$$\mathbf{pre} \ \underline{n} \ f \ x = \mathbf{snd} \ (\underline{n} \ (\mathbf{prefn} \ f) \ (\mathbf{true}, x)) \quad \text{(definition of } \mathbf{pre})$$

$$= \mathbf{snd} \ ((\mathbf{prefn} \ f)^n \ (\mathbf{true}, x)) \quad \text{(definition of } \underline{n})$$

$$= \mathbf{snd} (\mathbf{false}, f^{n-1} \ x) \quad \text{(by (v) above)}$$

$$= f^{n-1} \ x$$

hence by extensionality (Section 4.7 on page 71)

$$\begin{array}{rcl} \mathbf{pre} \; \underline{n} & = & \lambda f \; x \cdot f^{n-1} \; x \\ & = & \underline{n-1} & & \text{(definition of } \underline{n-1} \text{)} \end{array}$$

Exercise 59

Using the results of the previous exercise (or otherwise) show that

- (i) **pre** (**suc** \underline{n}) = \underline{n}
- (ii) **pre** 0 = 0

The numeral system in the next exercise is the one used in [4] and has some advantages over Church's (e.g. the predecessor function is easier to define).

Exercise 60

$$\begin{aligned} & \text{LET } \underline{\widehat{0}} = \lambda x.x \\ & \text{LET } \underline{\widehat{1}} = (\mathbf{false}, \underline{\widehat{0}}) \\ & \text{LET } \underline{\widehat{2}} = (\mathbf{false}, \underline{\widehat{1}}) \\ & \vdots \\ & \text{LET } \underline{\widehat{n+1}} = (\mathbf{false}, \underline{\widehat{n}}) \\ & \vdots \end{aligned}$$

Devise λ -expressions $\widehat{\mathbf{suc}}$, $\widehat{\mathbf{iszero}}$, $\widehat{\mathbf{pre}}$ such that for all n:

- (i) $\widehat{\mathbf{suc}} \ \widehat{\underline{n}} = \widehat{n+1}$
- (ii) $\widehat{\mathbf{iszero}} \ \widehat{\underline{0}} = \mathbf{true}$
- (iii) $\widehat{\mathbf{iszero}}$ ($\widehat{\mathbf{suc}}$ $\widehat{\underline{n}}$) = \mathbf{false}
- (iv) $\widehat{\mathbf{pre}}$ ($\widehat{\mathbf{suc}}$ $\widehat{\underline{n}}$) = $\widehat{\underline{n}}$

5.4 Definition by recursion

To represent the multiplication function in the λ -calculus we would like to define a λ -expression, **mult** say, such that:

$$\mathbf{mult}\ m\ n = \underbrace{\mathbf{add}\ n\ (\mathbf{add}\ n\ (\cdots\ (\mathbf{add}\ n\ \underline{0})\ \cdots))}_{m\ \mathbf{add}\,s}$$

This would be achieved if **mult** could be defined to satisfy the equation:

$$\mathbf{mult}\ m\ n = (\mathbf{iszero}\ m \to \underline{0}\ |\ \mathbf{add}\ n\ (\mathbf{mult}\ (\mathbf{pre}\ m)\ n))$$

If this held then, for example,

mult
$$\underline{2} \ \underline{3} = (\mathbf{iszero} \ \underline{2} \to \underline{0} \ | \ \mathbf{add} \ \underline{3} \ (\mathbf{mult} \ (\mathbf{pre} \ \underline{2}) \ \underline{3}))$$
(by the equation)
$$= \mathbf{add} \ \underline{3} \ (\mathbf{mult} \ \underline{1} \ \underline{3})$$
(by properties of \mathbf{iszero} , the conditional and \mathbf{pre})
$$= \mathbf{add} \ \underline{3} \ (\mathbf{iszero} \ \underline{1} \to \underline{0} \ | \ \mathbf{add} \ \underline{3} \ (\mathbf{mult} \ (\mathbf{pre} \ \underline{1}) \ \underline{3}))$$
(by the equation)
$$= \mathbf{add} \ \underline{3} \ (\mathbf{add} \ \underline{3} \ (\mathbf{mult} \ \underline{0} \ \underline{3}))$$
(by properties of \mathbf{iszero} , the conditional and \mathbf{pre})
$$= \mathbf{add} \ \underline{3} \ (\mathbf{add} \ \underline{3} \ (\mathbf{iszero} \ \underline{0} \to \underline{0} \ | \ \mathbf{add} \ \underline{3} \ (\mathbf{mult} \ (\mathbf{pre} \ \underline{0}) \ \underline{3})))$$
(by the equation)
$$= \mathbf{add} \ \underline{3} \ (\mathbf{add} \ \underline{3} \ (\mathbf{add} \ \underline{3} \ \underline{0})$$
(by properties of \mathbf{iszero} and the conditional)

The equation above suggests that **mult** be defined by:

$$\mathbf{mult} = \lambda m \ n. \ (\mathbf{iszero} \ m \to \underline{0} \mid \mathbf{add} \ n \ (\mathbf{mult} \ (\mathbf{pre} \ m) \ n))$$

$$\uparrow_{\mathrm{N.B.}}$$

Unfortunately, this cannot be used to define **mult** because, as indicated by the arrow, **mult** must already be defined for the λ -expression to the right of the equals to make sense.

Fortunately, there is a technique for constructing λ -expressions that satisfy arbitrary equations. When this technique is applied to the equation above it gives the desired definition of **mult**. First define a λ -expression **Y** that, for any expression E, has the following odd property:

$$\mathbf{Y} E = E (\mathbf{Y} E)$$

This says that \mathbf{Y} E is unchanged when the function E is applied to it. In general, if E E' = E' then E' is called a fixed point of E. A λ -expression \mathbf{Fix} with the property that \mathbf{Fix} E = $E(\mathbf{Fix}$ E) for any E is called a fixed-point operator. There are known to be infinitely many different fixed-point operators [57]; \mathbf{Y} is the most famous one, and its definition is:

Let
$$\mathbf{Y} = \lambda f_{\cdot} \; (\lambda x_{\cdot} \; f(x \; x)) \; (\lambda x_{\cdot} \; f(x \; x))$$

It is straightforward to show that Y is indeed a fixed-point operator:

$$\mathbf{Y} E = (\lambda f. (\lambda x. f(x x)) (\lambda x. f(x x))) E$$
 (definition of \mathbf{Y})

$$= (\lambda x. E(x x)) (\lambda x. E(x x))$$
 (β -conversion)

$$= E ((\lambda x. E(x x)) (\lambda x. E(x x)))$$
 (β -conversion)

$$= E (\mathbf{Y} E)$$
 (the line before last)

This calculation shows that every λ -expression E has a fixed point (namely \mathbf{Y} E); this is sometimes referred to as the first fixed-point theorem. The second fixed-point theorem is introduced in Section 7.1.

Armed with **Y**, we can now return to the problem of solving the equation for **mult**. Suppose **multfn** is defined by

LET multfn =
$$\lambda f \ m \ n$$
. (iszero $m \to \underline{0} \mid \mathbf{add} \ n \ (f \ (\mathbf{pre} \ m) \ n))$

and then define mult by:

$$LET$$
 $mult = Y$ $mult fn$

Then:

$$\mathbf{mult} \ m \ n = (\mathbf{Y} \ \mathbf{multfn}) \ m \ n$$
 (definition of \mathbf{mult})

= multfn (Y multfn)
$$m$$
 n (fixed-point property of Y)
= multfn mult m n (definition of mult)
= $(\lambda f \ m \ n. \ (\text{iszero} \ m \to \underline{0} \mid \text{add} \ n \ (f \ (\text{pre} \ m) \ n)))$ mult m n (definition of multfn)
= $(\text{iszero} \ m \to \underline{0} \mid \text{add} \ n \ (\text{mult} \ (\text{pre} \ m) \ n))$ (β -conversion)

An equation of the form $f x_1 \cdots x_n = E$ is called *recursive* if f occurs free in E. **Y** provides a general way of solving such equations. Start with an equation of the form:

$$\mathbf{f} x_1 \dots x_n = \mathbf{f}$$

where $\frown \mathbf{f} \frown$ is some λ -expression containing \mathbf{f} . To obtain an \mathbf{f} so that this equation holds define:

LET
$$\mathbf{f} = \mathbf{Y} (\lambda f \ x_1 \dots x_n - f - f)$$

The fact that the equation is satisfied can be shown as follows:

$$\mathbf{f} \ x_1 \dots x_n = \mathbf{Y} \ (\lambda f \ x_1 \dots x_n \dots f \longrightarrow) \ x_1 \dots x_n \qquad \text{(definition of } \mathbf{f})$$

$$= (\lambda f \ x_1 \dots x_n \dots f \longrightarrow) \ (\mathbf{Y} \ (\lambda f \ x_1 \dots x_n \dots f \longrightarrow)) \ x_1 \dots x_n \qquad \text{(fixed-point property)}$$

$$= (\lambda f \ x_1 \dots x_n \dots f \longrightarrow) \ \mathbf{f} \ x_1 \dots x_n \qquad \text{(definition of } \mathbf{f})$$

$$= \dots f \longrightarrow \qquad (\beta\text{-conversion})$$

Exercise 61

Construct a λ -expression eq such that

eq
$$m \ n = ($$
iszero $m \rightarrow$ iszero $n \mid$ $($ iszero $n \rightarrow$ false \mid eq $($ pre $m) ($ pre $n)))$

Exercise 62

Show that if \mathbf{Y}_1 is defined by:

LET
$$\mathbf{Y}_1 = \mathbf{Y} (\lambda y \ f. \ f(y \ f))$$

then \mathbf{Y}_1 is a fixed-point operator, i.e. for any E:

$$\mathbf{Y}_1 \ E = E \ (\mathbf{Y}_1 \ E)$$

The fixed-point operator in the next exercise is due to Turing (Barendregt [4], page 132).

Exercise 63

Show that $(\lambda x \ y. \ y \ (x \ x \ y)) \ (\lambda x \ y. \ y \ (x \ x \ y))$ is a fixed-point operator. \Box

The next exercise also comes from Barendregt's book, where it is attributed to Klop.

Exercise 64

Show that \mathbf{Y}_2 is a fixed-point operator, where:

Exercise 65

Is it the case that $\mathbf{Y} f \longrightarrow f(\mathbf{Y} f)$? If so prove it; if not find a λ -expression $\widehat{\mathbf{Y}}$ such that $\widehat{\mathbf{Y}} f \longrightarrow f(\widehat{\mathbf{Y}} f)$. \square

In the pure λ -calculus as defined on page 60, λ -expressions could only be applied to a single argument; however, this argument could be a *tuple* (see page 80). Thus one can write:

$$E(E_1,\ldots,E_n)$$

which actually abbreviates:

$$E(E_1, (E_2, (\cdots (E_{n-1}, E_n) \cdots)))$$

For example, $E(E_1, E_2)$ abbreviates $E(\lambda f, f E_1, E_2)$.

5.5 Functions with several arguments

In conventional mathematical usage, the application of an n-argument function f to arguments x_1, \ldots, x_n would be written as $f(x_1, \ldots, x_n)$. There are two ways of representing such applications in the λ -calculus:

- (i) as $(f x_1 ... x_n)$, or
- (ii) as the application of f to an n-tuple (x_1, \ldots, x_n) .

In case (i), f expects its arguments 'one at a time' and is said to be curried after a logician called Curry (the idea of currying was actually invented by Schönfinkel [65]). The functions **and**, **or** and **add** defined earlier were all curried. One advantage of curried functions is that they can be 'partially applied'; for example, **add** $\underline{1}$ is the result of partially applying **add** to $\underline{1}$ and denotes the function $n \mapsto n+1$.

Although it is often convenient to represent n-argument functions as curried, it is also useful to be able to represent them, as in case (ii) above, by λ -expressions expecting a single n-tuple argument. For example, instead of representing + and \times by λ -expressions add and mult such that

add
$$\underline{m} \ \underline{n} = \underline{m+n}$$

mult $\underline{m} \ \underline{n} = \underline{m \times n}$

it might be more convenient to represent them by functions, \mathbf{sum} and \mathbf{prod} say, such that

$$\mathbf{sum} \ (\underline{m}, \underline{n}) = \underline{m+n}$$

$$\mathbf{prod} \ (\underline{m}, \underline{n}) = m \times n$$

This is nearer to conventional mathematical usage and has applications that will be encountered later. One might say that **sum** and **prod** are uncurried versions of **add** and **mult** respectively.

Define:

LET
$$\mathbf{curry} = \lambda f \ x_1 \ x_2 . \ f \ (x_1, x_2)$$

LET $\mathbf{uncurry} = \lambda f \ p . \ f \ (\mathbf{fst} \ p) \ (\mathbf{snd} \ p)$

then defining

results in **sum** and **prod** having the desired properties; for example:

```
\begin{array}{ll} \mathbf{sum} \ (\underline{m},\underline{n}) &= \mathbf{uncurry} \ \mathbf{add} \ (\underline{m},\underline{n}) \\ &= (\lambda f \ p. \ f \ (\mathbf{fst} \ p) \ (\mathbf{snd} \ p)) \mathbf{add} \ (\underline{m},\underline{n}) \\ &= \mathbf{add} \ (\mathbf{fst} \ (\underline{m},\underline{n})) \ (\mathbf{snd} \ (\underline{m},\underline{n})) \\ &= \mathbf{add} \ \underline{m} \ \underline{n} \\ &= \underline{m+n} \end{array}
```

Exercise 66

Show that for any E:

curry (uncurry
$$E$$
) = E
uncurry (curry E) = E

hence show that:

We can define n-ary functions for currying and uncurrying. For n>0 define:

LET curry_n =
$$\lambda f x_1 \cdots x_n \cdot f (x_1, \dots, x_n)$$

Let
$$\mathbf{uncurry}_n = \lambda f \ p. \ f \ (p \stackrel{n}{\downarrow} 1) \ \cdots \ (p \stackrel{n}{\downarrow} n)$$

If E represents a function expecting an n-tuple argument, then $\operatorname{\mathbf{curry}}_n E$ represents the curried function which takes its arguments one at a time. If E represents a curried function of n arguments, then $\operatorname{\mathbf{uncurry}}_n E$ represents the uncurried version which expects a single n-tuple as argument.

Exercise 67

Show that:

- (i) $\operatorname{\mathbf{curry}}_n (\operatorname{\mathbf{uncurry}}_n E) = E$
- (ii) $\mathbf{uncurry}_n \ (\mathbf{curry}_n \ E) = E$

Exercise 68

Devise λ -expressions E_1^n and E_2^n built out of **curry** and **uncurry** such that $\mathbf{curry}_n = E_1^n$ and $\mathbf{uncurry}_n = E_2^n$. \square

The following notation provides a convenient way to write λ -expressions which expect tuples as arguments.

Generalized λ -abstractions

LET
$$\lambda(V_1, \ldots, V_n)$$
. $E = \mathbf{uncurry}_n \ (\lambda V_1 \ \ldots \ V_n . \ E)$

Example: $\lambda(x,y)$. **mult** x y abbreviates:

$$\mathbf{uncurry}_2 \ (\lambda x \ y. \ \mathbf{mult} \ x \ y) = (\lambda f \ p. \ f \ (p \downarrow 1) \ (p \downarrow 2)) \ (\lambda x \ y. \ \mathbf{mult} \ x \ y)$$
$$= (\lambda f \ p. \ f \ (\mathbf{fst} \ p) \ (\mathbf{snd} \ p)) \ (\lambda x \ y. \ \mathbf{mult} \ x \ y)$$
$$= \lambda p. \ \mathbf{mult} \ (\mathbf{fst} \ p) (\mathbf{snd} \ p)$$

Thus:

$$(\lambda(x,y).$$
 mult $x y)$ $(E_1, E_2) = (\lambda p.$ mult $($ fst $p)$ $($ snd $p))$ (E_1, E_2)
= mult $($ fst $(E_1, E_2))$ $($ snd $(E_1, E_2))$
= mult $E_1 E_2$

This example illustrates the rule of generalized β -conversion in the box below. This rule can be derived from ordinary β -conversion and the definitions of tuples and generalized λ -abstractions. The idea is that a tuple of arguments is passed to each argument position in the body of the generalized abstraction; then each individual argument can be extracted from the tuple without affecting the others.

Generalized β -conversion

$$(\lambda(V_1,\ldots,V_n),E)$$
 $(E_1,\ldots,E_n)=E[E_1,\ldots,E_n/V_1,\ldots,V_n]$

where $E[E_1, \ldots, E_n/V_1, \ldots, V_n]$ is the *simultaneous substitution* of E_1, \ldots, E_n for V_1, \ldots, V_n respectively and none of these variables occur free in any of E_1, \ldots, E_n .

It is convenient to extend the notation $\lambda V_1 \ V_2 \dots V_n$. E described on page 62 so that each V_i can either be an identifier or a tuple of identifiers. The meaning of $\lambda V_1 \ V_2 \dots V_n$. E is still $\lambda V_1 . (\lambda V_2 . (\dots (\lambda V_n . E) \dots))$, but now if V_i is a tuple of identifiers then the expression is a generalized abstraction.

Example: $\lambda f(x,y)$. f(x,y) means λf . $(\lambda(x,y), f(x,y))$ which in turn means λf . **uncurry** $(\lambda x, y, f(x,y))$ which equals λf . $(\lambda p, f(\mathbf{fst}, p), (\mathbf{snd}, p))$. \square

Exercise 69

Show that if the only free variables in E are x_1, \ldots, x_n and f, then if:

$$\mathbf{f} = \mathbf{Y} (\lambda f (x_1, \dots, x_n), E)$$

then

$$\mathbf{f}(x_1,\ldots,x_n) = E[\mathbf{f}/f]$$

Exercise 70

Define a λ -expression **div** with the property that:

$$\mathbf{div}\ (\underline{m},\underline{n})=(q,\underline{r})$$

where q and r are the quotient and remainder of dividing n into m. \square

5.6 Mutual recursion

To solve a set of mutually recursive equations like:

$$\mathbf{f}_1 = F_1 \ \mathbf{f}_1 \cdots \mathbf{f}_n$$

$$\mathbf{f}_2 = F_2 \ \mathbf{f}_1 \cdots \mathbf{f}_n$$

$$\vdots$$

$$\mathbf{f}_n = F_n \ \mathbf{f}_1 \cdots \mathbf{f}_n$$

we simply define for $1 \le i \le n$

$$\mathbf{f}_i = \mathbf{Y} \left(\lambda(f_1, \dots f_n) \cdot (F_1 \ f_1 \cdots f_n, \dots, F_n \ f_1 \cdots f_n) \right) \downarrow i$$

This works because if

$$\vec{\mathbf{f}} = \mathbf{Y} \left(\lambda(f_1, \dots f_n) \cdot (F_1 \ f_1 \cdots f_n, \dots, F_n \ f_1 \cdots f_n) \right)$$

then $\mathbf{f}_i = \vec{\mathbf{f}} \downarrow i$ and hence:

$$\vec{\mathbf{f}} = (\lambda(f_1, \dots, f_n) \cdot (F_1 \ f_1 \cdots f_n, \dots, F_n \ f_1 \cdots f_n))\vec{\mathbf{f}}$$

$$= (F_1(\vec{\mathbf{f}} \downarrow 1) \cdots (\vec{\mathbf{f}} \downarrow n), \dots, F_n(\vec{\mathbf{f}} \downarrow 1) \cdots (\vec{\mathbf{f}} \downarrow n))$$

$$= (F_1 \ \mathbf{f}_1 \cdots \mathbf{f}_n, \dots, F_n \ \mathbf{f}_1 \cdots \mathbf{f}_n) \qquad (\text{since } \vec{\mathbf{f}} \downarrow i = \mathbf{f}_i).$$

Hence:

$$\mathbf{f}_i = F_i \ \mathbf{f}_1 \cdots \mathbf{f}_n$$

5.7 Representing the recursive functions

The recursive functions form an important class of numerical functions. Shortly after Church invented the λ -calculus, Kleene proved that every

recursive function could be represented in it [39]. This provided evidence for *Church's thesis*, the hypothesis that any intuitively computable function could be represented in the λ -calculus. It has been shown that many other models of compution define the same class of functions that can be defined in the λ -calculus [56].

In this section it is described what it means for an arithmetic function to be represented in the λ -calculus. Two classes of functions, the *primitive* recursive functions and the recursive functions, are defined and it is shown that all the functions in these classes can be represented in the λ -calculus.

In Section 5.3 it was explained how a number n is represented by the λ -expression \underline{n} . A λ -expression \underline{f} is said to represent a mathematical function f if for all numbers x_1, \ldots, x_n :

$$\underline{f}(\underline{x_1}, \dots, \underline{x_n}) = \underline{y}$$
 if $f(x_1, \dots, x_n) = \underline{y}$

5.7.1 The primitive recursive functions

A function is called *primitive recursive* if it can be constructed from 0 and the functions S and U_n^i (defined below) by a finite sequence of applications of the operations of substitution and primitive recursion (also defined below).

The successor function S and projection functions U_n^i (where n and i are numbers) are defined by:

- (i) S(x) = x + 1
- (ii) $U_n^i(x_1, x_2, ..., x_n) = x_i$

Substitution

Suppose g is a function of r arguments and h_1, \ldots, h_r are r functions each of n arguments. We say f is defined from g and h_1, \ldots, h_r by substitution if:

$$f(x_1, \ldots, x_n) = q(h_1(x_1, \ldots, x_n), \ldots, h_r(x_1, \ldots, x_n))$$

Primitive recursion

Suppose g is a function of n-1 arguments and h is a function of n+1 arguments. We say f is defined from g and h by primitive recursion if:

$$f(0, x_2, ..., x_n) = g(x_2, ..., x_n)$$

$$f(S(x_1), x_2, ..., x_n) = h(f(x_1, x_2, ..., x_n), x_1, x_2, ..., x_n)$$

g is called the *base function* and h is called the *step function*. It can proved that for any base and step function there always exists a unique function

defined from them by primitive recursion. This result is called the primitive recursion theorem; proofs of it can be found in textbooks on mathematical logic (e.g. [64]).

Example: The addition function sum is primitive recursive because:

$$sum(0, x_2) = x_2$$

 $sum(S(x_1), x_2) = S(sum(x_1, x_2))$

It is now shown that every primitive recursive function can be represented by λ -expressions.

It is obvious that the λ -expressions $\underline{0}$, $\operatorname{\mathbf{suc}}$, λp . $p \downarrow^n i$ represent the initial functions 0, S and U_n^i respectively.

Suppose function g of r variables is represented by \mathbf{g} and functions h_i $(1 \le i \le r)$ of n variables are represented by \mathbf{h}_i . Then if a function f of n variables is defined by substitution by:

$$f(x_1, ..., x_n) = g(h_1(x_1, ..., x_n), ..., h_r(x_1, ..., x_n))$$

then f is represented by \mathbf{f} where:

$$\mathbf{f} = \lambda(x_1, \ldots, x_n) \cdot \mathbf{g}(\mathbf{h}_1(x_1, \ldots, x_n), \ldots, \mathbf{h}_r(x_1, \ldots, x_n))$$

Suppose function f of n variables is defined inductively from a base function g of n-1 variables and an inductive step function h of n+1 variables. Then

$$f(0, x_2, \dots, x_n) = g(x_2, \dots, x_n)$$

$$f(S(x_1), x_2, \dots, x_n) = h(f(x_1, x_2, \dots, x_n), x_1, x_2, \dots, x_n)$$

Thus if \mathbf{g} represents g and \mathbf{h} represents h then \mathbf{f} will represent f if

$$\begin{aligned} \mathbf{f} \; & (x_1, x_2, \dots, x_n) = \\ & (\mathbf{iszero} \; x_1 \rightarrow \mathbf{g}(x_2, \dots, x_n) \mid \\ & \quad \quad \mathbf{h}(\mathbf{f} \; (\mathbf{pre} \; x_1, x_2, \dots, x_n), \mathbf{pre} \; x_1, x_2, \dots, x_n)) \end{aligned}$$

Using the fixed-point trick, an \mathbf{f} can be constructed to satisfy this equation defining \mathbf{f} to be¹:

$$\begin{aligned} \mathbf{Y}(\lambda f. \ \lambda(x_1, x_2, \dots, x_n). \\ (\mathbf{iszero} \ x_1 \rightarrow \mathbf{g}(x_2, \dots, x_n) \mid \\ \mathbf{h}(f(\mathbf{pre} \ x_1, x_2, \dots, x_n), \mathbf{pre} \ x_1, x_2, \dots, x_n))) \end{aligned}$$

Thus any primitive recursive function can be represented by a λ -expression.

¹See Exercise 69 on page 92.

5.7.2 The recursive functions

A function is called *recursive* if it can be constructed from 0, the successor function and the projection functions (see page 94) by a sequence of substitutions, primitive recursions and *minimizations*.

Minimization

Suppose g is a function of n arguments. We say f is defined from g by minimization if:

$$f(x_1, x_2, \dots, x_n) =$$
 'the smallest y such that $g(y, x_2, \dots, x_n) = x_1$ '

The notation MIN(f) is used to denote the minimization of f. Functions defined by minimization may be undefined for some arguments. For example, if one is the function that always returns 1, i.e. one(x) = 1 for every x, then MIN(one) is only defined for arguments with value 1. This is obvious because if f(x) = MIN(one)(x), then:

$$f(x)$$
 = 'the smallest y such that $one(y)=x$ '

and clearly this is only defined if x = 1. Thus

$$MIN(one)(x) = \begin{cases} \underline{0} & \text{if } x = 1\\ & \text{undefined} & \text{otherwise} \end{cases}$$

To show that any recursive function can be represented in the λ -calculus it is necessary to show how to represent the minimization of an arbitrary function. Suppose g represents a function g of n variables and f is defined by:

$$f = MIN(g)$$

Then if a λ -expression **min** can be devised such that **min** \underline{x} **f** $(\underline{x}_1, \dots, \underline{x}_n)$ represents the least number y greater than x such that

$$f(y, x_2, \ldots, x_n) = x_1$$

then g will represent g where:

$$\mathbf{g} = \lambda(x_1, x_2, \dots, x_n) \cdot \min \underline{0} \mathbf{f}(x_1, x_2, \dots, x_n)$$

min will clearly have the desired property if:

where eq \underline{m} \underline{n} is equal to true if m = n and false otherwise (a suitable definition of eq occurs on page 88). Thus min can simply be defined to be:

$$\mathbf{Y}(\lambda m. \\ \lambda x \ f \ (x_1, x_2, \dots, x_n). \\ (\mathbf{eq} \ (f(x, x_2, \dots, x_n)) \ x_1 \ \to x \mid m \ (\mathbf{suc} \ x) \ f \ (x_1, x_2, \dots, x_n)))$$

Thus any recursive function can be represented by a λ -expression.

Higher-order primitive recursion

There are functions which are recursive but not primitive recursive. An example given in Barendregt ([4], page 569) is the version of Ackermann's function, ψ , defined by:

$$\psi(0,n) = n+1
\psi(m+1,0) = \psi(m,1)
\psi(m+1,n+1) = \psi(m,\psi(m+1,n))$$

However, if one allows functions as arguments, then any recursive function can be defined by a primitive recursion. For example, if the higher-order function *rec* is defined by primitive recursion as follows:

$$rec(0, x_2, x_3) = x_2$$

 $rec(S(x_1), x_2, x_3) = x_3(rec(x_1, x_2, x_3))$

then ψ can be defined by:

$$\psi(m,n) = rec (m, S, f \mapsto (x \mapsto rec(x, f(1), f))) (n)$$

where $x \mapsto \theta(x)$ denotes the function² that maps x to $\theta(x)$. Notice that the third argument of rec, viz. x_3 , must be a function. In the definition of ψ we also took x_1 to be a function, viz. S.

Exercise 71

Show that the definition of ψ in terms of rec works, i.e. that with ψ defined as above:

$$\begin{array}{lll} \psi(0,n) &=& n+1 \\ \psi(m+1,0) &=& \psi(m,1) \\ \psi(m+1,n+1) &=& \psi(m,\psi(m+1,n)) \end{array}$$

² Note that λx . $\theta(x)$ is an expression of the λ -calculus whereas $x \mapsto \theta(x)$ is a notation of informal mathematics.

A function which takes another function as an argument, or returns another function as a result, is called *higher-order*. The example ψ shows that higher-order primitive recursion is more powerful than ordinary primitive recursion³. The use of operators like *rec* is one of the things that makes functional programming very powerful. The functions **map** and **lit** described in Section 6.5 illustrate this further.

5.7.3 The partial recursive functions

A partial function is one that is not defined for all arguments. For example, the function MIN(one) described above is partial. Another example is the division function, since division by 0 is not defined. Functions that are defined for all arguments are called total.

A partial function is called partial recursive if it can be constructed from 0, the successor function and the projection functions by a sequence of substitutions, primitive recursions and minimizations. Thus the recursive functions are just the partial recursive functions which happen to be total. It can be shown that every partial recursive function f can be represented by a λ -expression f in the sense that

- (i) $\underline{f}(\underline{x_1}, \dots, \underline{x_n}) = \underline{y}$ if $f(x_1, \dots, x_n) = \underline{y}$
- (ii) If $f(x_1, \ldots, x_n)$ is undefined then $\underline{f(x_1, \ldots, x_n)}$ has no normal form.

Note that despite (ii) above, it is not in general correct to regard expressions with no normal form as being 'undefined'; this subtle point is discussed further on page 120 (see also pages 41 and 42 of Barendregt [4]).

Exercise 72

Write down the λ -expression that represents MIN(f), where f(x) = 0 for all x. \square

5.8 Representing lists (LISP S-expressions)

The data-structures manipulated by LISP (see Chapter 9) are called symbolic expressions or S-expressions. The LISP primitives for operating on these include car, cdr and cons. A first approximation to representing S-expressions is to represent the formation of LISP's dotted-pairs (i.e. conscells) by the λ -calculus pairing defined in Section 5.2 (see page 80).

³The kind of primitive recursion defined in Section 5.7.1 is *first-order* primitive recursion.

This appears to work at first, since by defining $\mathbf{car} = \mathbf{fst}$ and $\mathbf{cdr} = \mathbf{snd}$ then, since $(E_1, E_2, \dots, E_n) = (E_1, (E_2, \dots, E_n))$, it follows that

$$\operatorname{car} (E_1, E_2, \dots, E_n) = E_1$$

 $\operatorname{cdr} (E_1, E_2, \dots, E_n) = (E_2, \dots, E_n)$

as in LISP. The only thing that is missing is the empty list **nil**. Unfortunately, we cannot simply devise some λ -expression E to represent **nil** because there would be no way (see Exercise 87 on page 127) of defining a λ -expression **null** such that:

$$\mathbf{null} \ E = \left\{ \begin{array}{ll} \mathbf{true} & E = \mathbf{nil} \\ \\ \mathbf{false} & \mathrm{otherwise} \end{array} \right.$$

In LISP implementations, machine words are marked to indicate whether they represent atoms (e.g. **nil**) or dotted pairs. One can adopt the same approach in the λ -calculus and represent a list by a pair (b, E) where b represents a boolean which indicates whether the list is empty or not. Then a **null** test can easily be represented by inspecting the first component:

$$LET null = fst$$

To avoid confusion between λ -expressions representing pairs and those representing S-expressions (defined below) we will use $[E_1; \dots; E_n]$ to represent an S-expression list whose components are E_1, \dots, E_n . This notation is similar to the M-expression syntax of LISP 1.5 [53].

The empty list, for which we will use the notation [], is represented by a pair of the form (**true**, \frown). Thus ensuring that:

$$\mathbf{null}$$
 [] = \mathbf{true}

It doesn't really matter what \sim is, but we will use a special λ -expression called \perp (pronounced 'bottom'). We can also use \perp as the head (**car**) and tail (**cdr**) of the empty list. The λ -expression \perp is supposed to represent an 'undefined function'. Intuitively, the result of applying such a function to any argument is also 'undefined', so it is natural to require for every E that:

$$\perp E = \perp$$

Using Y we can solve the equation $\perp x = \perp$.

$$\mathtt{LET} \perp = \mathbf{Y} \ (\lambda f \ x. \ f)$$

Now define

```
LET [] = (\mathbf{true}, \bot)

LET [E] = (\mathbf{false}, (E, []))

LET [E_1; E_2] = (\mathbf{false}, (E_1, [E_2]))

\vdots

LET [E_1; \cdots; E_n] = (\mathbf{false}, (E_1, [E_2; \cdots; E_n]))
```

The primitive functions can then be defined by:

```
LET \mathbf{hd} = \lambda l. (\mathbf{null}\ l \to \bot \mid \ \mathbf{fst}(\mathbf{snd}\ l))

LET \mathbf{tl} = \lambda l. (\mathbf{null}\ l \to \bot \mid \ \mathbf{snd}(\mathbf{snd}\ l))

LET \mathbf{cons} = \lambda x\ l. (\mathbf{false}, (x, l))
```

Exercise 73

Show that the following properties hold:

```
hd (cons x l) = x

tl (cons x l) = l

null (cons x l) = false

[E_1; \dots; E_n] = cons E_1 (cons E_2 ( \dots (cons E_n []) \dots))
```

Example: A function for appending one list to another can be defined by constructing a λ -expression **append** such that:

append
$$x \ y = (\text{null } x \to y \mid \text{ cons } (\text{hd } x) \ (\text{append}(\text{tl } x)y))$$

i.e.

```
\texttt{LET append} = \mathbf{Y} \ (\lambda f \ x \ y. \ (\mathbf{null} \ x \to y \ | \ \mathbf{cons} \ (\mathbf{hd} \ x) \ (f \ (\mathbf{tl} \ x) \ y)))
```

Then, for example,

```
\begin{aligned} & \mathbf{append}[E_1; E_2] \ [E_3; E_4] \\ &= (\mathbf{null}[E_1; E_2] \to [E_3; E_4] \ | \\ &\qquad \qquad \mathbf{cons} \ (\mathbf{hd}[E_1; E_2]) \ (\mathbf{append}(\mathbf{tl}[E_1; E_2])[E_3; E_4])) \\ &= \mathbf{cons} \ E_1 \ (\mathbf{append}[E_2] \ [E_3; E_4]) \\ &= \mathbf{cons} \ E_1 \ (\mathbf{cons} \ E_2 \ (\mathbf{append}[\ ] \ [E_3; E_4])) \\ &= \mathbf{cons} \ E_1 \ (\mathbf{cons} \ E_2 \ [E_3; E_4]) \\ &= [E_1; E_2; E_3; E_4]) \end{aligned}
```

In general:

```
\mathbf{append}[E_1; \dots; E_n][E'_1; \dots; E'_m] = [E_1; \dots; E_n; E'_1; \dots; E'_m]
```

Exercise 74

Define a λ -expression **reverse** such that:

```
reverse[E_1; E_2; \dots; E_{n-1}; E_n] = [E_n; E_{n-1}; \dots; E_2; E_1]
```

There is an important difference between the lists we have just represented in the λ -calculus and LISP lists: the former lists can be *infinite*. To see this consider the equation:

```
from n = cons \ n \ (from \ (suc \ n))
```

Then

```
from \underline{0} = \mathbf{cons} \ \underline{0} \ (\mathbf{from} \ \underline{1})
= \mathbf{cons} \ \underline{0} \ (\mathbf{from} \ \underline{1})
= \mathbf{cons} \ \underline{0} \ (\mathbf{cons} \ \underline{1} \ (\mathbf{from} \ \underline{2}))
= \mathbf{cons} \ \underline{0} \ (\mathbf{cons} \ \underline{1} \ (\mathbf{cons} \ \underline{2} \ (\mathbf{from} \ \underline{3})))
\vdots
= [0; 1; 2; 3; \cdots]
```

If one attempts to define **from** in LISP, then because the LISP evaluator does *call by value* (i.e. it always evaluates arguments before entering functions), expressions like **hd** (**from** $\underline{0}$) would generate non-terminating evaluations. However, from the definitions above it is straightforward to show that, for example,

$$\mathbf{hd} \ (\mathbf{from} \ \underline{0}) = \underline{0}$$

Languages with this behaviour (i.e. **from** works) are said to employ *lazy* or *normal order* evaluation. There is more on this topic in the next chapter (e.g. see the normalization theorem on page 121).

5.9 Extending the λ -calculus

Although it is possible to represent data-objects and data-structures with λ -expressions, it is often inefficient to do so. For example, most computers have hardware for arithmetic and it is reasonable to use this, rather than λ -conversion, to compute with numbers. A mathematically clean way of 'interfacing' computation rules to the λ -calculus is via so called δ -rules. The idea is to add a new constant, \mathbf{c} say, and then to specify an algorithm, called a δ -rule, for reducing applications \mathbf{c} $E_1 \ldots E_n$. For example, one might add + as a new constant together with the δ -rule:

$$+ \underline{m} \underline{n} \xrightarrow{\delta} \underline{m+n}$$

 $(E_1 \xrightarrow{\delta} E_2 \text{ means } E_2 \text{ results by applying a } \delta\text{-rule to some subexpression of } E_1).$

When adding such constants and rules to the λ -calculus one must be careful not to destroy its nice properties, e.g. the Church-Rosser theorem (see page 118). For example, suppose the following rule were added:

$$+ E_1 E_2 \xrightarrow{\delta} \begin{cases} \underline{m+n} & \text{if } E_1 \equiv \underline{m} \text{ and } E_2 \equiv \underline{n} \\ \bot & \text{otherwise} \end{cases}$$

Then

$$+ ((\lambda x.x)\underline{0}) \underline{0} \longrightarrow \bot$$

because although $(\lambda x.x)\underline{0} = \underline{0}$, it is not the case that $(\lambda x.x)\underline{0} \equiv \underline{m}$ for any number m. It would also follow that

$$+ ((\lambda x.x)\underline{0}) \ \underline{0} \ \xrightarrow{\beta} \ + \ \underline{0} \ \underline{0} \ \xrightarrow{\delta} \ \underline{0}$$

This shows that $\underline{0} = \bot$ if the above rule for + is added (and the definition of = on page 68 is extended to include δ -reduction).

The following condition due to Mitschke (see Barendregt [4], page 401) guarantees that a δ -rule preserves the Church-Rosser theorem.

Suppose the following three things hold:

- 1. $R_1(E_1, \ldots, E_n)$, $R_2(E_1, \ldots, E_n)$, ..., $R_m(E_1, \ldots, E_n)$ are disjoint relations. This means that:
 - (a) $R_i(E_1, \ldots, E_n)$ is either true or false.
 - (b) For a given E_1, \ldots, E_n at most one of $R_i(E_1, \ldots, E_n)$ is true.
- 2. The relations R_i are closed under λ -conversion. This means that if $R_i(E_1, \ldots, E_j, \ldots, E_n)$ is true and also if $E_j \longrightarrow E'_j$ then $R_i(E_1, \ldots, E'_j, \ldots, E_n)$ is also true.
- 3. The relations R_i are closed under substitution. This means that if $R_i(E_1, \ldots, E_j, \ldots, E_n)$ is true, E is any expression and V is any variable then $R_i(E_1, \ldots, E_i[E/V], \ldots, E_n)$ is also true.

If 1, 2 and 3 hold, then adding a constant c together with the $\delta\text{-reductions}\colon$

$$\mathbf{c} \ E_1 \ \dots \ E_n \xrightarrow{\delta} E^1 \quad \text{If } R_1(E_1, \dots, E_n) \text{ holds}$$
 \vdots

$$\mathbf{c} \ E_1 \ \dots \ E_n \ \xrightarrow{\delta} \ E^m \quad \text{If } R_m(E_1, \dots, E_n) \text{ holds}$$

(where E^1, \ldots, E^n are arbitrary λ -expressions) results in an extension of the λ -calculus for which the Church-Rosser theorem still holds.

Exercise 75

Devise δ -rules for arithmetic and show that they satisfy Mitschke's conditions. \square

Functional Programs

Functional programming is briefly described and illustrated with some simple examples. A little functional programming language is defined. It is shown how it can be translated into the λ -calculus.

Functional programming languages all contain some subset equivalent to the λ -calculus. They differ in the particular notation used and in the 'impure' features provided. A feature is impure if it cannot be viewed as an abbreviation for some λ -calculus construct (and hence cannot be analysed using λ -calculus theory). For example, older functional languages like LISP contain imperative features such as assignment and goto commands.

In the rest of this chapter some typical functional programming notations are described; these descend from Landin's pioneering work in the 1960s [42]. It is shown that these notations are 'pure', i.e. that they can be viewed as abbreviations for λ -calculus constructs.

6.1 Functional notation

Let let $x = E_1$ in E_2 be written to mean $(\lambda x. E_2) E_1$. Then, for example, let $n = \underline{0}$ in suc n means $(\lambda n. \mathbf{suc} n) \underline{0}$. By the rules of λ -conversion this reduces to $\underline{1}$.

Here is another example in which there is a let inside a let.

$$\begin{array}{ll} \text{let } m = \underline{0} \text{ in} \\ \text{let } n = \underline{1} \text{ in} \\ \text{add } m \ n \end{array}$$

This means:

$$(\lambda m. \ \text{let} \ n = \underline{1} \ \text{in add} \ m \ n) \ \underline{0}$$

By the rules of λ -conversion

One can think of let x = E as a kind of local declaration of x with value E. The scope of the declaration is written after in.

The basic idea can be extended in various ways:

let
$$(x_1,\ldots,x_n)=E_1$$
 in E_2

means:

$$(\lambda(x_1,\ldots,x_n),E_2)$$
 E_1

Example:

$$let(x,y) = (\underline{2},\underline{3})$$
 in add $x y$

means

$$(\lambda(x,y))$$
 add $x(y)$ $(\underline{2},\underline{3})$

This is equal to $\underline{5}$ by λ -conversion. \square

The notation:

let
$$x_1 = E_1$$
 and $x_2 = E_2 \cdots$ and $x_n = E_n$ in E

is defined to mean:

let
$$(x_1, x_2, \dots, x_n) = (E_1, E_2, \dots, E_n)$$
 in E

i.e.

$$(\lambda(x_1, x_2, \ldots, x_n), E) (E_1, E_2, \ldots, E_n)$$

Note that:

$$\begin{array}{lll} \text{let} & x_1=E_1 & \text{in} \\ \\ \text{let} & x_2=E_2 & \text{in} \\ \\ & & \vdots \\ \\ \text{let} & x_n=E_n & \text{in} & E \end{array}$$

is different from:

$$\begin{array}{ll} \text{let} \ x_1 = E_1 \\ \text{and} \ x_2 = E_2 \\ & \vdots \\ \text{and} \ x_n = E_n \ \text{in} \ E \end{array}$$

In the former, the scope of each let $x_i = E_i$ is

let
$$x_{i+1}=E_{i+1}$$
 in let $x_{i+2}=E_{i+2}$ in \cdots let $x_n=E_n$ in E

In the latter, the declarations let $x_i = E_i$ are done 'in parallel'. For example,

let
$$x = \underline{1}$$
 in let $y = \operatorname{suc} x$ in y

is equal to $\underline{2}$, but

let
$$x = \underline{1}$$
 and $y = \mathbf{suc}\ x$ in y

is equal to suc x, since the declaration let $x = \underline{1}$ isn't felt by the x in suc x.

In general:

$$\begin{array}{lll} \text{let} \ x_1=E_1 \ \text{in} \\ \text{let} \ x_2=E_2 \ \text{in} \\ & \vdots \\ \text{let} \ x_n=E_n \ \text{in} \ E \end{array}$$

means $E[E_n/x_n] \cdots [E_2/x_2][E_1/x_1]$, whereas:

$$\begin{array}{ll} \text{let } x_1=E_1\\ \text{and } x_2=E_2\\ & \vdots\\ \text{and } x_n=E_n \text{ in } E \end{array}$$

means $E[E_1, E_2, ..., E_n/x_1, x_2, ..., x_n]$.

To make the definitions of functions look nicer, the notation:

let
$$f x_1 \cdots x_n = E$$

is defined to mean:

let
$$f = \lambda x_1 \cdots x_n$$
. E

and

let
$$f(x_1,\ldots,x_n)=E$$

is defined to mean:

let
$$f = \lambda(x_1, \ldots, x_n)$$
. E

Example

let
$$\operatorname{suc} n = \lambda f \ x \cdot n \ f \ (f \ x)$$
 in $\operatorname{suc} \ \underline{0}$

means

let
$$suc = \lambda n \cdot \lambda f x \cdot n f (f x)$$
 in $suc 0$

By λ -conversion it follows that:

$$\begin{array}{ll} (\lambda suc.\ suc\ \underline{0})\ (\lambda n.\ \lambda f\ x.\ n\ f\ (f\ x)) &= (\lambda n.\ \lambda f\ x.\ n\ f\ (f\ x))\ \underline{0} \\ &= \lambda f\ x.\ \underline{0}\ f\ (f\ x) \\ &= \lambda f\ x.\ f\ x \\ &= 1 \end{array}$$

See the description of \mathbf{suc} on page 83 for more explanation. \square

Example

let sum
$$(m,n) = \lambda f x \cdot m (n f x)$$
 in E

means:

let sum =
$$\lambda(m, n)$$
. $\lambda f x$. $m (n f x)$ in E

which is:

$$(\lambda sum \cdot E) (\lambda (m, n) \cdot \lambda f x \cdot (n f x))$$

i.e.

$$E[(\lambda(m,n), \lambda f \ x. (n \ f \ x))/sum]$$

See the description of **sum** on page 90 for more explanation. \square A common notation for recursive definition is:

letrec
$$f = E$$

This is defined to mean:

let
$$f = \mathbf{Y} (\lambda f. E)$$

For simultaneous recursive definitions, the notation

letrec
$$f_1=E_1$$
 and \cdots and $f_n=E_n$

is defined to mean:

let
$$(f_1,\ldots,f_n)=\mathbf{Y}\;(\lambda(f_1,\ldots,f_n)\,.\,(E_1,\ldots E_n))$$

The function-defining notation described above can also be used with letrec as well as with let, thus:

letrec
$$f x_1 \cdots x_n = E$$

means:

letrec
$$f = \lambda x_1 \cdots x_n \cdot E$$

and

letrec
$$f(x_1 \cdots x_n) = E$$

means:

letrec
$$f = \lambda(x_1, \ldots, x_n)$$
. E

Example: The declaration:

letrec mult
$$m \ n = (\mathbf{iszero} \ m \to \underline{0} \mid \mathbf{add} \ n \ (\mathbf{mult} \ (\mathbf{pre} \ m) \ n))$$

is equivalent to:

let
$$\mathbf{mult} = \mathbf{Y} \ (\lambda \ mult \ m \ n. \ (\mathbf{iszero} \ m \to \underline{0} \ | \ \mathbf{add} \ n \ (mult \ (\mathbf{pre} \ m) \ n)))$$

6.2 Combining declarations

So far declarations have just been things of the form let B or letrec B. One can introduce some operators for combining such declarations. For example, suppose D_1 and D_2 are declarations, then:

1. D_1 ; D_2 is a declaration whose meaning is defined by;

$$(D_1; D_2)$$
 in $E = D_1$ in $(D_2$ in $E)$

2. D₁ ins D₂ is a declaration in which the bindings in D₁ only apply to the expressions in D₂ (i.e. the scope of D₁ is D₂) and these bindings are not exported as bindings of the compound declaration D₁ ins D₂. This is clarified by the examples below.

Examples

- (i) let $x = \underline{1}$; let $y = x + \underline{1}$ is equivalent to let $x = \underline{1}$ and $y = \underline{2}$.
- (ii) let $x = \underline{1}$ ins let $y = x + \underline{1}$ is equivalent to let $y = \underline{2}$.

In general, if D_2 is equivalent to:

let
$$X_1 = E_1$$
 and \cdots and $X_n = E_n$

then D_1 ins D_2 is equivalent to:

let
$$X_1=(D_1 \text{ in } E_1) \text{ and } \cdots \text{ and } X_n=(D_1 \text{ in } E_n)$$

If D_2 is equivalent to:

letrec
$$X_1 = E_1$$
 and \cdots and $X_n = E_n$

then D_1 ins D_2 is equivalent to:

letrec
$$X_1=(D_1 \text{ in } E_1)$$
 and \cdots and $X_n=(D_1 \text{ in } E_n)$

The name ins derives from 'inside'.

6.3 Predeclared definitions

The names $\underline{0}, \underline{1}, \underline{2}, \ldots$, suc , add , cond , true , ... are part of the language used to talk about the λ -calculus, i.e. they are part of the *metalanguage*. In a real functional programming language these things will be predefined variables; the definitions being done automatically for the user. One way to represent this is to imagine a big declaration $D_{\operatorname{initial}}$ which binds the representations of standard objects like numbers, truth values, etc. to their usual names. So $D_{\operatorname{initial}}$ might start out:

```
let 0 = \lambda f \ x. \ x;

\vdots

let n = \lambda f \ x. \ f^n \ x;

\vdots

let true = \lambda x \ y. \ x;

\vdots

let hd = \lambda l. \ (null \ l \to \bot \mid fst(snd \ l));

\vdots
```

Any expressions E will then be assumed to be in the scope of D_{initial} . (i.e. E will really mean D_{initial} in E).

6.4 A compiling algorithm

In this section the notations just described are gathered together into a language. The syntax of declarations is;

$$D ::=$$
let $B \mid$ letrec $B \mid D_1; D_2 \mid D_1$ ins D_2
 $B ::= X = E \mid X_1 = E_1$ and \cdots and $X_n = E_n$
 $X ::= V \mid (X_1, \dots, X_n) \mid V \mid F_1 \dots F_n \mid V(F_1, \dots, F_n)$
 $F ::= V \mid (V_1, \dots, V_n)$

Here E, E_1, \ldots, E_n range over expressions; these are either ordinary λ -expressions, or things of the form D in E where D is a declaration. Thus:

$$E ::= V \mid (E_1 \ E_2) \mid \lambda V. \ E \mid D \ \textbf{in} \ E$$

V ranges over variables. The Bs used in defining declarations are called bindings. The Xs are called variable structures and the Fs are called parameter specifications.

The language just described includes things that have not been discussed, for example:

let
$$((f x, y), (z_1, z_2)) = E$$
 in E'

There follows a description of a 'compiler' from this language to simple λ -expressions as defined on page 60. For example, the expression above will compile to:

$$(\lambda(f, y, z_1, z_2), E')$$
 $((\lambda x, E \downarrow 1 \downarrow 1), E \downarrow 1 \downarrow 2, E \downarrow 2 \downarrow 1, E \downarrow 2 \downarrow 2))$

This denotes a simple λ -expression via the definitions of $\lambda(x_1, \ldots, x_n)$. E and \downarrow .

The compilation alogorithm will be described via a sequence of transformations (passes). The notation ' $X \rightsquigarrow Y$ ' will mean 'replace X by Y'.

Expressions of the form D in E are compiled by the following steps (which should be performed in the order given):

Step 1: Remove bindings of the form $(X_1, \ldots, X_n) = E$.

$$(X_1,\ldots,X_n)=E \quad \leadsto \quad X_1=E_1 \stackrel{n}{\downarrow} 1 \text{ and } \cdots \text{ and } X_n=E_n \stackrel{n}{\downarrow} n$$

Example: Applying Step 1 to:

let
$$((f x, y), (z_1, z_2)) = E$$
 in E'

yields:

let
$$(f \ x, \ y) = E \downarrow 1$$
 and $(z_1, z_2) = E \downarrow 2$ in E'

Applying Step 1 again yields:

$$\begin{array}{lll} \text{let } f \ x &= E \downarrow 1 \downarrow 1 \\ \text{and } y &= E \downarrow 1 \downarrow 2 \\ \text{and } z_1 &= E \downarrow 2 \downarrow 1 \\ \text{and } z_2 &= E \downarrow 2 \downarrow 2 \\ \text{in } E' \end{array}$$

Step 2: Put bindings in the form V = E.

$$V X_1 \dots X_n = E \quad \leadsto \quad V = \lambda X_1 \dots X_n \cdot E$$

$$V(X_1, \ldots, X_n) = E \quad \rightsquigarrow \quad V = \lambda(X_1, \ldots, X_n) \cdot E$$

Example: Applying Step 2 to the result of the previous example yields:

let
$$f = \lambda x. E \downarrow 1 \downarrow 1$$

and $y = E \downarrow 1 \downarrow 2$
and $z_1 = E \downarrow 2 \downarrow 1$
and $z_2 = E \downarrow 2 \downarrow 2$
in E'

Step 3: Remove ins.

$$D$$
 ins let $V_1=E_1$ and \cdots and $V_n=E_n$
 \Rightarrow let $V_1=(D \text{ in } E_1)$ and \cdots and let $V_n=(D \text{ in } E_n)$

Example:

let
$$x=\underline{1}$$
 ins let $y=x+\underline{1}$ \leadsto let $y=(\text{let }x=\underline{1} \text{ in }x+\underline{1})$

Step 4: Remove in.

let
$$V_1=E_1$$
 and \cdots and $V_n=E_n$ in E \rightarrow $(\lambda(V_1,\ldots,V_n),E)(E_1,\ldots,E_n)$ letrec $V_1=E_1$ and \cdots and $V_n=E_n$ in E

 $\rightsquigarrow (\lambda(V_1,\ldots,V_n).E)(\mathbf{Y}(\lambda(V_1,\ldots,V_n)(E_1,\ldots,E_n)))$

Example: Applying Step 4 to the result of Step 2 in the example before last yields:

```
(\lambda(f, y, z_1, z_2). E') ((\lambda x. E \downarrow 1 \downarrow 1), E \downarrow 1 \downarrow 2, E \downarrow 2 \downarrow 1, E \downarrow 2 \downarrow 2)
```

Exercise 76

Compile:

```
letrec fact n = (\mathbf{iszero} \ n \to \underline{0} \mid \mathbf{mult} \ n \ (\mathbf{fact} \ (\mathbf{pre} \ n))) in fact 6
```

6.5 Example functional programs

A functional program consists of a sequence of function definitions followed by an expression (involving the functions) which evaluates to the result one is trying to compute.

For example, the functional program below computes the average of the squares of the numbers 64, 23, 104, 8, 72 and 20. It has the form

```
letrec sumlist l = \cdots and letrec length l = \cdots and letrec map f \mid l = \cdots ins let avsq l = \cdots in avsq [\underline{64}; \ \underline{23}; \ \underline{104}; \ \underline{8}; \ \underline{72}; \ \underline{20}]
```

where the three functions **sumlist**, **length** and **map** are local to the declaration of **avsq**, which is the function for computing the average of the sum of squares. The functions **sumlist** and **length** compute the sum of a list of numbers and its length, respectively. The higher-order function **map** takes as arguments a function f and a list $[E_1; \ldots; E_n]$ and returns as result the list $[f(f, E_1); \ldots; f(f, E_n)]$. For example,

map
$$(\lambda n. \mathbf{mult} \ n \ n) \ l$$

evaluates to the list obtained from l by squaring each element. Here is the complete program:

```
letrec sumlist l=(\operatorname{null} l \to \underline{0} \mid \operatorname{add} \ (\operatorname{hd} \ l) \ (\operatorname{sumlist} \ (\operatorname{tl} \ l))) and letrec length l=(\operatorname{null} \ l \to \underline{0} \mid \operatorname{add} \ \underline{1} \ (\operatorname{length} \ (\operatorname{tl} \ l))) and letrec map f \ l=(\operatorname{null} \ l \to \underline{\mathbb{I}} \ | \ (\operatorname{cons} \ (f \ (\operatorname{hd} \ l)) \ (\operatorname{map} \ f \ (\operatorname{tl} \ l))) ins let avsq l=\operatorname{fst} \ (\operatorname{div} \ (\operatorname{sumlist} \ (\operatorname{map} \ (\lambda n. \ \operatorname{mult} \ n \ n) \ l)) \ (\operatorname{length} \ l)) in avsq [\underline{64}; \ \underline{23}; \ \underline{104}; \ \underline{8}; \ \underline{72}; \ \underline{20}]
```

Higher-order 'iteration functions' like \mathbf{map} are very powerful and are used frequently in functional programming. Another example is \mathbf{lit}^1

lit
$$f [x_1; x_2; ...; x_n] x = f x_1 (f x_2 ... (f x_n x) ...)$$

lit is a generalization of the standard mathematical notation for iterated sums and products:

$$\sum_{i=1}^{n} x_i = \mathbf{lit} \ \mathbf{add} \ [x_1; x_2; \dots; x_n] \ \underline{0}$$

$$\prod_{i=1}^{n} x_i = \text{lit mult } [x_1; x_2; \dots; x_n] \ \underline{1}$$

Thus sumlist l is just lit add $l \underline{0}$.

Many simple list processing functions can be defined using **lit** and no other recursion. For example:

This works because:

```
\begin{array}{l} \mathbf{append} \ [x_1; \ \dots \ ; x_m] \ [y_1; \ \dots \ ; y_n] \\ = \mathbf{lit} \ \mathbf{cons} \ [x_1; \ \dots \ ; x_m] \ [y_1; \ \dots \ ; y_n] \\ = \mathbf{cons} \ x_1 \ (\mathbf{cons} \ x_2 \ (\ \dots \ (\mathbf{cons} \ x_m \ [y_1; \ \dots \ ; y_n]) \ \dots)) \\ = \ [x_1; \ \dots \ ; x_m; \ y_1; \ \dots \ ; \ y_n] \end{array}
```

The list-reversing function rev, where:

$$\mathbf{rev} [x_1; \ldots; x_m] = [x_m; \ldots; x_1]$$

can then be defined by:

rev
$$l = \mathbf{lit} (\lambda x \ l_1. \mathbf{append} \ l_1 \ [x]) \ l \ []$$

Here are some more examples:

¹ The function **lit** is sometimes called **reduce** or **itlist**: it is similar to operators in the programming language APL. An early occurrence of it is in Barron and Strachey's article [5].

1. If **member** x l is **true** if x is a member of list l and **false** otherwise, then:

```
 \begin{array}{l} \mathbf{member} \ x \ l \\ = \mathbf{lit} \ (\lambda x_1 \ b. \ (\mathbf{eq} \ b \ \mathbf{true} \rightarrow \mathbf{true} \ | \ (\mathbf{eq} \ x_1 \ x \rightarrow \mathbf{true} \ | \ \mathbf{false}))) \\ l \\ \mathbf{false} \end{array}
```

For example, **member** $\underline{2}$ [$\underline{5}$; $\underline{2}$; $\underline{3}$; $\underline{4}$; $\underline{5}$; $\underline{6}$; $\underline{7}$] = **true** and **member** $\underline{8}$ [$\underline{1}$; $\underline{2}$; $\underline{3}$; $\underline{4}$; $\underline{5}$; $\underline{6}$; $\underline{7}$] = **false**.

2. If union l_1 l_2 is the set-theoretic union of l_1 and l_2 then:

```
union l_1 l_2
= lit (\lambda x \ l_3. (member x \ l_2 \rightarrow l_3 \mid \mathbf{cons} \ x \ l_3)) l_1 l_2
```

For example, union $[\underline{1}; \underline{2}; \underline{3}]$ $[\underline{2}; \underline{4}; \underline{5}] = [\underline{1}; \underline{3}; \underline{2}; \underline{4}; \underline{5}]$.

3. If **intersection** l_1 l_2 is the set-theoretic union of l_1 and l_2 then:

```
intersection l_1 l_2
= lit (\lambda x \ l_3. (member x \ l_2 \to \mathbf{cons} \ x \ l_3 \ | \ l_3)) l_1 []
```

For example, **intersection** $[\underline{1}; \underline{2}; \underline{3}]$ $[\underline{2}; \underline{4}; \underline{5}] = [\underline{2}]$.

These examples are from the paper 'On the power of list iteration' [21] where it is also shown that both the set of all sublists of a list and the cartesian product of a list of lists can be computed by expressions built up out of lit. It is also shown that any primitive recursive function can be computed in this way, but that the equality of lists cannot.

Exercise 77

Program the function **length** for computing the length of a list using **lit**. \Box

An excellent book containing lots of examples of functional programs is Recursive Programming Techniques by W. Burge [9].

Theorems about the λ -calculus

Several important theorems about the λ -calculus are stated without proof. These include the Church-Rosser theorem, the normalization theorem, and the undecidability of the halting problem. The significance of these is explained and various applications are described.

If $E_1 \longrightarrow E_2$ then E_2 can be thought of as having been got from E_1 by 'evaluation'. If there are no $(\beta$ - or η -) redexes in E_2 then it can be thought of as 'fully evaluated'.

A λ -expression is said to be in normal form if it contains no β - or η -redexes (i.e. if the only conversion rule that can be applied is α -conversion). Thus a λ -expression in normal form is 'fully evaluated'.

Examples

- (i) The representations of numbers are all in normal form.
- (ii) $(\lambda x. x)$ 0 is not in normal form.

Suppose an expression E is 'evaluated' in two different ways by applying two different sequences of reductions until two normal forms E_1 and E_2 are obtained. The Church-Rosser theorem stated below shows that E_1 and E_2 will be the same except for having possibly different names of bound variables.

Because the results of reductions do not depend on the order in which they are done, separate redexes can be evaluated in parallel. Various research projects are currently trying to exploit this fact by designing multiprocessor architectures for evaluating λ -expressions. It is too early to tell how successful this work will be. There is a possibility that the communication overhead of distributing redexes to different processors and then

collecting together the results will cancel out the theoretical advantages of the approach. Let us hope this pessimistic possibility can be avoided. It is a remarkable fact that the Church-Rosser theorem, an obscure mathematical result dating from before computers were invented, may underpin the design of the next generation of computing systems.

Here is the statement of the Church-Rosser theorem. It is an example of something that is intuitively obvious, but very hard to prove. Many properties of the λ -calculus share this property.

The Church-Rosser theorem

If $E_1 = E_2$ then there exists an E such that $E_1 \longrightarrow E$ and $E_2 \longrightarrow E$.

It is now possible to see why the Chuch-Rosser theorem shows that λ -expressions can be evaluated in any order. Suppose an expression E is 'evaluated' in two different ways by applying two different sequences of reductions until two normal forms E_1 and E_2 are obtained. Since E_1 and E_2 are obtained from E by sequences of conversions, it follows by the definition of E that E is an expression, E and E is an expression, E are in normal form, then the only redexes they can contain are α -redexes and so the only way that E and E can be reduced to E is by changing the names of bound variables. Thus E_1 and E_2 must be the same up to renaming of bound variables (i.e. α -conversion).

Another application of the Church-Rosser theorem is to show that if $m \neq n$ then the λ -expressions representing m and n are not equal, i.e. $\underline{m} \neq \underline{n}$. Suppose $m \neq n$ but $\underline{m} = \underline{n}$; by the Church-Rosser theorem $\underline{m} \longrightarrow E$ and $\underline{n} \longrightarrow E$ for some E. But it is obvious from the definitions of \underline{m} and \underline{n} , namely

$$\underline{m} = \lambda f \ x. \ f^m \ x$$
$$\underline{n} = \lambda f \ x. \ f^n \ x$$

that no such E can exist. The only conversions that are applicable to $\underline{\mathbf{m}}$ and $\underline{\mathbf{n}}$ are α -conversions and these cannot change the number of function applications in an expression (\underline{m} contains m applications and \underline{n} contains n applications).

A λ -expression E has a normal form if E=E' for some E' in normal form. The following corollary relates expressions in normal form to those that have a normal form; it summarizes some of the statements made above.

Corollary to the Church-Rosser theorem

- (i) If E has a normal form then $E \longrightarrow E'$ for some E' in normal form.
- (ii) If E has a normal form and E=E' then E' has a normal form.
- (iii) If E=E' and E and E' are both in normal form, then E and E' are identical up to α -conversion.

Proof

- (i) If E has a normal form then E=E' for some E' in normal form. By the Church-Rosser theorem there exists E'' such that $E\longrightarrow E'$ and $E\longrightarrow E''$. As E' is in normal form the only redexes it can have are α -redexes, so the reduction $E'\longrightarrow E''$ must consist of a sequence of α -conversions. Thus E'' must be identical to E' except for some renaming of bound variables; it must thus be in normal form as E' is.
- (ii) Suppose E has a normal form and E = E'. As E has a normal form, E = E'' where E'' is in normal form. Hence E' = E'' by the transitivity of = (see page 69) and so E' has a normal form.
- (iii) This was proved above.

Exercise 78

For each of the following λ -expressions *either* find its normal form or show that it has no normal form:

- (i) **add** <u>3</u>
- (ii) add 35
- (iii) $(\lambda x. x x) (\lambda x. x)$
- (iv) $(\lambda x. x x) (\lambda x. x x)$
- (v) **Y**
- (vi) $\mathbf{Y} (\lambda y. y)$
- (vii) **Y** ($\lambda f \ x$. (iszero $x \to \underline{0} \mid f \ (\mathbf{pre} \ x)$)) $\underline{7}$

Notice that a λ -expression E might have a normal form even if there exists an infinite sequence $E \longrightarrow E_1 \longrightarrow E_2 \cdots$. For example $(\lambda x. \underline{1})$ (**Y** f) has a normal form $\underline{1}$ even though:

$$(\lambda x. \underline{1}) (\mathbf{Y} f) \longrightarrow (\lambda x. \underline{1}) (f (\mathbf{Y} f)) \longrightarrow \cdots (\lambda x. \underline{1}) (f^n (\mathbf{Y} f)) \longrightarrow \cdots$$

The normalization theorem stated below tells us that such blind alleys can always be avoided by reducing the *leftmost* β - or η -redex, where by 'leftmost' is meant the redex whose beginning λ is as far to the left as possible.

Another important point to note is that E_1 may not have a normal form even though E_1 E_2 does have one. For example, \mathbf{Y} has no normal form, but \mathbf{Y} $(\lambda x. \underline{1}) \longrightarrow \underline{1}$. It is a common mistake to think of λ -expressions without a normal form as denoting 'undefined' functions; \mathbf{Y} has no normal form but it denotes a perfectly well defined function¹. Analysis beyond the scope of this book (see Wadsworth's paper [71]) shows that a λ -expression denotes an undefined function if and only if it cannot be converted to an expression in head normal form, where E is in head normal form if it has the form

$$\lambda V_1 \cdot \cdot \cdot \cdot V_m \cdot V E_1 \cdot \cdot \cdot \cdot E_n$$

where V_1, \ldots, V_m and V are variables and E_1, \ldots, E_n are λ -expressions (V can either be equal to V_i , for some i, or it can be distinct from all of them). It follows that the fixed-point operator \mathbf{Y} is not undefined because it can be converted to

$$\lambda f. f((\lambda x. f(x x)) (\lambda x. f(x x)))$$

which is in head normal form.

It can be shown that an expression E has a head normal form if and only if there exist expressions E_1, \ldots, E_n such that E E_1, \ldots, E_n has a normal form. This supports the interpretation of expressions without head normal forms as denoting undefined functions: E being undefined means that E E_1, \ldots, E_n never terminates for any E_1, \ldots, E_n . Full details on head normal forms and their relation to definedness can be found in Barendregt's book [4].

Exercise 79

In Section 5.7.3 a representation of a partial recursive function f was defined by a λ -expression f. Is it the case that with this representation, if

¹ The mathematical characterization of the function denoted by \mathbf{Y} can be found in Stoy's book [67].

 $f(x_1,\ldots,x_n)=y$ is undefined then $\underline{f}(\underline{x_1},\ldots,\underline{x_n})=\underline{y}$ does not have a head normal form? \square

The normalization theorem

If E has a normal form, then repeatedly reducing the leftmost β - or η -redex (possibly after an α -conversion to avoid invalid substitutions) will terminate with an expression in normal form.

The remark about α -conversion in the statement of the theorem is to cover cases like:

$$(\lambda x . \ (\lambda y . \ x \ y)) \ y \longrightarrow \lambda y' . \ y \ y'$$

where $\lambda y. \ x \ y \longrightarrow \lambda y'. \ x \ y'$ has been α -converted so as to avoid the invalid substitution $(\lambda y. \ x \ y) [y/x] = \lambda y. \ y. y$.

A sequence of reductions in which the leftmost redex is always reduced is called a *normal order reduction sequence*.

The normalization theorem says that if E has a normal form (i.e. for some E' in normal form E = E') then it can be found by normal order reduction. This, however, is not usually the 'most efficient' way to find it. For example, normal order reduction requires

$$(\lambda x. \longrightarrow x \longrightarrow x \longrightarrow) E$$

to be reduced to

$$\sim E \sim E \sim$$

If E is not in normal form then it would be more efficient to first reduce E to E' say (where E' is in normal form) and then to reduce

$$(\lambda x - x - x - x) E'$$

to

$$\sim E' \sim E' \sim$$

thereby avoiding having to reduce E twice.

Note, however, that this 'call-by-value' scheme is disastrous in cases like

$$(\lambda x \cdot \underline{1}) ((\lambda x \cdot x \cdot x) (\lambda x \cdot x \cdot x))$$

It is a difficult problem to find an optimal algorithm for choosing the next redex to reduce. For recent work in this area see Levy's paper [43].

Because normal order reduction appears so inefficient, some programming languages based on the λ -calculus, e.g. LISP, have used call by

value even though it doesn't always terminate. Actually, call by value has other advantages besides efficiency, especially when the language is 'impure', i.e. has constructs with side effects (e.g. assignments). On the other hand, recent research suggests that maybe normal order evaluation is not as inefficient as was originally thought if one uses cunning implementation tricks like graph reduction (see page 136). Whether functional programming languages should use normal order or call by value is still a controversial issue.

7.1 Some undecidability results

In Section 5.7 it was shown that the λ -calculus is a computing mechanism at least as powerful as the partial recursive functions. There follow some examples of things that cannot be computed.

Suppose a λ -expression **hasnf** (for 'has a normal form') could be devised with the property that for any E:

$$\mathbf{hasnf} \ E = \left\{ \begin{array}{ll} \mathbf{true} & \text{if } E \text{ has a normal form} \\ \\ \mathbf{false} & \text{if } E \text{ doesn't have a normal form} \end{array} \right.$$

Define another λ -expression **W** which satisfies the equation:

$$\mathbf{W} = (\mathbf{hasnf} \ \mathbf{W} \to \bot \mid \underline{0})$$

(where \perp is a λ -expression which doesn't have a normal form, e.g. the one defined on page 99). A suitable definition of **W** is:

$$\mathbf{W} = \mathbf{Y} \left(\lambda f. \left(\mathbf{hasnf} \ f \to \bot \mid \underline{0} \right) \right)$$

Does ${\bf W}$ have a normal form? There are two possible answers: 'yes' or 'no'.

- (i) If 'yes' then hasnf W = true and then by the equation for W and properties of conditionals (see page 78) it follows that W = ⊥. But ⊥ doesn't have a normal form which contradicts (ii) of the corollary to the Church-Rosser theorem on page 118.
- (ii) If 'no' then **hasnf W** = **false** and then by the equation for **W** and properties of conditionals (see page 78) it follows that **W** = <u>0</u>. But <u>0</u> has a normal form (indeed, it is in normal form) which again contradicts (ii) of the corollary to the Church-Rosser theorem.

This shows that the assumption that **hasnf** exists leads to a contradiction in all cases and thus it cannot exist.

Exercise 80

Show that there cannot exist a λ -expression equal with the property that:

equal
$$E_1$$
 $E_2 = \begin{cases} \mathbf{true} & \text{if } E_1 = E_2 \\ \mathbf{false} & \text{otherwise} \end{cases}$

Hint: Choose **W** to satisfy **W** = (equal **W** $\underline{0} \rightarrow \underline{1} \mid \underline{0}$) and then consider whether **W** = $\underline{0}$. \Box

Exercise 81

Why doesn't non-existence of **equal** in the previous exercise contradict the existence of **eq** in Exercise 61 on page 88? □

The result on the non-existence of **hasnf** is not very strong because all that the evaluation of **hasnf** could do is apply expressions to E, and E to expressions. Maybe if **hasnf** were given some kind of representation of the syntax of E then it could use some algorithm to compute whether E terminates. To investigate this possibility, suppose that for each λ -expression E another λ -expression E has been defined such that $E_1 = E_2$ if and only if E_1 is identical to E_2 (an outline of how to define E is given below). One can think of E as the parse tree of E represented as a λ -expression. Note that one cannot just take E to be E because then, for example, it would follow that $(\lambda x. x) = 1$ is not identical to 1.

It will be shown that there is no λ -expression, **halts** say, such that for all E:

$$\mathbf{halts}^{\mathsf{r}} E^{\mathsf{r}} = \begin{cases} \mathbf{true} & \text{if } E \text{ has a normal form} \\ \mathbf{false} & \text{if } E \text{ doesn't have a normal form} \end{cases}$$

The argument is similar to the one above for **hasnf**, but slighly more cunning. It makes use of the second fixed-point theorem stated below.

Defining [E] for arbitary E is straightforward, but a bit tedious. First it is necessary to define a representation for character strings. If $c_1 c_2 \ldots c_n$ is a character string, define $"c_1 c_2 \ldots c_n"$ be the λ -expression $[\mathbf{a}_1; \mathbf{a}_2; \cdots; \mathbf{a}_n]$ where \mathbf{a}_i is the ASCII code for \mathbf{c}_i (e.g. "fred" is [102; 114; 101; 100]).

Exercise 82

Using the definitions on pages 82 and 99 write out the λ -expression denoted by "def". \square

Exercise 83

Define a λ -expression **eqstr** such that:

$$\mathbf{eqstr}("\mathtt{c}_1 \dots \mathtt{c}_n", "\mathtt{c}_1' \dots \mathtt{c}_m'") = \left\{ \begin{array}{ll} \mathbf{true} & \text{if } n = m \text{ and} \\ & \mathtt{c}_i = \mathtt{c}_i' \ (1 \leq i \leq n) \\ \mathbf{false} & \text{otherwise} \end{array} \right.$$

If E is a λ -expression we define E by cases as follows:

- (i) If E is a variable I then: [E] = (0, "I")
- (ii) If E is a combination $(E_1 \ E_2)$ then: $E = (\underline{1}, E_1, E_2)$
- (iii) If E is an abstraction (λV . E') then: E' = (2, "V", E')

Think of E as an internal represention (i.e. parse tree) of E. The first component of the pair E indicates the sort of expression that E is.

Example:

$$\begin{split} \lceil \lambda x. \; (\lambda y. \; x \; y) \rceil &= (\underline{2}, "\mathbf{x}", \lceil \lambda y. \; x \; y \rceil) \\ &= (\underline{2}, "\mathbf{x}", (\underline{2}, "\mathbf{y}", \lceil x \; y \rceil)) \\ &= (\underline{2}, "\mathbf{x}", (\underline{2}, "\mathbf{y}", (\underline{1}, \lceil x \rceil, \lceil y \rceil))) \\ &= (\underline{2}, "\mathbf{x}", (\underline{2}, "\mathbf{y}", (\underline{1}, (\underline{0}, "\mathbf{x}"), (\underline{0}, "\mathbf{y}")))) \\ &= (\underline{2}, [\underline{120}], (\underline{2}, [\underline{121}], (\underline{1}, (\underline{0}, [\underline{120}]), (\underline{0}, [\underline{121}])))) \end{split}$$

The following λ -expressions are defined to represent abstract syntax constructors, selectors and tests.

Constructors

LET
$$\mathbf{mkvar} = \lambda x \cdot (\underline{0}, x)$$

LET $\mathbf{mkcomb} = \lambda(x, y) \cdot (\underline{1}, x, y)$
LET $\mathbf{mkabs} = \lambda(x, y) \cdot (\underline{2}, x, y)$

Selectors

LET rator =
$$\lambda e \cdot e \downarrow 2$$

LET rator = $\lambda e \cdot e \downarrow 3$
LET by = $\lambda e \cdot e \downarrow 3$

LET body =
$$\lambda e. e \downarrow 3$$

Tests

LET is
$$\mathbf{var} = \lambda e$$
. \mathbf{eq} (fst e) $\underline{0}$
LET is $\mathbf{comb} = \lambda e$. \mathbf{eq} (fst e) $\underline{1}$
LET is $\mathbf{abs} = \lambda e$. \mathbf{eq} (fst e) $\underline{2}$

See Exercise 61 on page 88 for eq.

Exercise 84

Show

- (ii) $\mathbf{b}\mathbf{v}^{\mathsf{T}}\lambda x$. $E^{\mathsf{T}} = \mathbf{v}\mathbf{x}^{\mathsf{T}}$
- (iii) body $\lceil \lambda x \cdot E \rceil = \lceil E \rceil$
- (iv) rator $E_1 E_2 = E_1$
- (v) rand $E_1 E_2 = E_2$
- (vii) $\mathbf{mkcomb}(E_1, E_2) = E_1 E_2$
- (viii) $\mathbf{mkabs}(\mathbf{"x"}, \mathbf{\Basis}) = \mathbf{\Basis}\lambda x \cdot \mathbf{\Basis}$

The second fixed-point theorem stated below should be compared with the first fixed-point theorem of Section 5.4 (see page 87). It is useful for showing the non-existence of **halts**.

The second fixed-point theorem

For any λ -expression E there exists a λ -expression E' such that:

$$E' = E'E'$$

 $Proof\ outline$

Suppose there is a λ -expression quote such that for every E

$$quote^{\Gamma}E^{\Gamma} = (\Gamma E^{\Gamma})^{\Gamma}$$

Then one can define $E' = \mathbf{F} \mathsf{\lceil} \mathbf{F} \mathsf{\rceil}$ where:

$$\mathbf{F} = \lambda x \cdot E \text{ (mkcomb } (x, \mathbf{quote } x))$$

This works because

$$\begin{split} E' &= \mathbf{F} \ \mathbf{\bar{F}} \\ &= E \ (\mathbf{mkcomb} \ (\mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \\ &= E \ (\mathbf{mkcomb} \ (\mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \\ &= E \ (\mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \\ &= E \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \ \mathbf{\bar{F}} \\ \end{split}$$

The definition of **quote** is straightforward; one just considers the various forms $^{r}E^{r}$ can take as described on page 124. \square

Exercise 85

This exercise outlines the details of the definition of quote.

(i) Define a λ -expression **quotenum** such that for all n:

quotenum
$$\underline{n} = \underline{n}$$

(ii) Define a λ -expression **quotestring** such that for all strings " $\mathbf{x}_1 \dots \mathbf{x}_n$ "

quotestring
$$\mathbf{x}_1 \dots \mathbf{x}_n = \mathbf{x}_1 \dots \mathbf{x}_n$$

(iii) Define a λ -expression **quote** such that for all E

quote
$$E' = E''$$

7.2 The halting problem

Suppose one could devise a λ -expression halts with the property that:

$$\mathbf{halts} \, \mathsf{T}E \, \mathsf{T} = \left\{ \begin{array}{ll} \mathbf{true} & \text{if } E \text{ has a normal form} \\ \\ \mathbf{false} & \text{if } E \text{ doesn't have a normal form.} \end{array} \right.$$

Define **foo** by

foo =
$$\lambda x$$
. (halts $x \to \bot \mid \underline{0}$)

By the second fixed-point theorem there is a λ -expression, **W** say, such that:

$$\mathbf{W} = \mathbf{foo} \mathbf{W} = (\mathbf{halts} \mathbf{W} \rightarrow \bot \mid \underline{0})$$

It can be seen that, in fact, such a **W** cannot exists by asking whether it has a normal form.

- (i) If W has a normal form then halts W = true, so by the definition of foo: W = ⊥. But this is impossible because if W = ⊥ then by the corollary to the Church-Rosser theorem (see page 118) ⊥ would have to have a normal form also.
- (ii) If **W** does not have a normal form then $\mathbf{halt}^{\mathsf{T}}\mathbf{W}^{\mathsf{T}} = \mathbf{false}$, so $\mathbf{W} = \underline{0}$. But this is also impossible as $\underline{0}$ is in normal form.

This contradiction was derived by assuming that **halts** exists, hence **halts** does not exist.

Exercise 86

Show that there cannot exist a λ -expression equiv such that

equiv(
$$[E_1], [E_2]$$
) =
$$\begin{cases} \mathbf{true} & \text{if } E_1 = E_2 \\ \mathbf{false} & \text{if } E_1 \neq E_2 \end{cases}$$

Hint: Replace halts by λx . equiv (E_1, x) in the argument above. \Box

Exercise 87

Suppose lists are represented as in Section 5.8 on page 98. Show that there cannot exist a λ -expression **null** such that:

$$\mathbf{null} \ E = \left\{ \begin{array}{ll} \mathbf{true} & \text{if } E = [\] \\ \\ \mathbf{false} & \text{otherwise} \end{array} \right.$$

Does this result hold if [] is replaced by an arbitrary λ -expression E?

Combinators

Combinators are motivated and described, and their relationship to the λ -calculus is discussed. Various algorithms for compiling λ -expressions into combinators are explained. A brief introduction to the combinator machines is given.

Combinators provide an alternative theory of functions to the λ -calculus. They were originally introduced by logicians as a way of studying the process of substitution. More recently, Turner has argued that combinators provide a good 'machine code' into which functional programs can be compiled [68]. Several experimental computers have been built based on Turner's ideas (see e.g. [12]) and the results are promising. How these machines work is explained in Section 8.3. Combinators also provide a good intermediate code for conventional machines; several of the best compilers for functional languages are based on them (e.g. [17, 2]).

There are two equivalent ways of formulating the theory of combinators:

- (i) within the λ -calculus, or
- (ii) as a completely separate theory.

The approach here is to adopt (i) here as it is slightly simpler, but (ii) was how it was done originally¹. It will be shown that $any \lambda$ -expression is equal to an expression built from variables and two particular expressions, \mathbf{K} and \mathbf{S} , using only function application. This is done by mimicking λ -abstractions using combinations of \mathbf{K} and \mathbf{S} . It will be demonstrated how β -reductions can be simulated by simpler operations involving \mathbf{K} and \mathbf{S} . It is these simpler operations that combinator machines implement directly in hardware. The definitions of \mathbf{K} and \mathbf{S} are

¹The two-volume treatise Combinatory Logic [14, 15] is the definitive reference, but the more recent textbooks [31, 4] are better places to start.

LET
$$\mathbf{K} = \lambda x \ y. \ x$$

 LET $\mathbf{S} = \lambda f \ g \ x. \ (f \ x) \ (g \ x)$

From these definitions it is clear by β -reduction that for all E_1 , E_2 and E_3 :

K
$$E_1$$
 $E_2 = E_1$
S E_1 E_2 $E_3 = (E_1$ $E_3)$ $(E_2$ $E_3)$

Any expression built by application (i.e. combination) from K and S is called a *combinator*; K and S are the *primitive combinators*.

In BNF, combinators have the following syntax:

$$< combinator > ::= K | S | (< combinator > < combinator >)$$

A combinatory expression is an expression built from K, S and zero or more variables. Thus a combinator is a combinatory expression not containing variables. In BNF, the syntax of combinatory expressions is:

Exercise 88

Define I by:

LET
$$I = \lambda x \cdot x$$

Show that $\mathbf{I} = \mathbf{S} \ \mathbf{K} \ \mathbf{K}$. \square

The identity function I defined in the last exercise is often taken as a primitive combinator, but as the exercise shows this is not necessary as it can be defined from K and S.

8.1 Combinator reduction

If E and E' are combinatory expressions then the notation $E \xrightarrow{c} E'$ is used if E = E' or if E' can be got from E by a sequence of rewritings of the form:

(i) **K**
$$E_1$$
 $E_2 \xrightarrow{c} E_1$

(ii) **S**
$$E_1 E_2 E_3 \xrightarrow{c} (E_1 E_3) (E_2 E_3)$$

(iii) I
$$E \xrightarrow{c} E$$

Note that the reduction I $E \xrightarrow{c} E$ is derivable from (i) and (ii).

Example

$$\mathbf{S} \mathbf{K} \mathbf{K} x \xrightarrow{c} \mathbf{K} x (\mathbf{K} x) \qquad \text{by (ii)}$$

$$\xrightarrow{c} x \qquad \text{by (i)}$$

This example shows that for any $E: \mathbf{I} \to E$.

Any sequence of combinatory reductions, i.e. reductions via \xrightarrow{c} , can be expanded into a sequence of β -conversions. This is clear because \mathbf{K} E_1 E_2 and \mathbf{S} E_1 E_2 E_3 reduce to E_1 and $(E_1$ $E_3)$ $(E_2$ $E_3)$, respectively, by sequences of β -conversions.

8.2 Functional completeness

A surprising fact is that any λ -expression can be translated to an equivalent combinatory expression. This result is called the functional completeness of combinators and is the basis for compilers for functional languages to the machine code of combinator machines.

The first step is to define, for an arbitrary variable V and combinatory expression E, another combinatory expression λ^*V . E that simulates λV . E in the sense that λ^*V . $E=\lambda V$. E. This provides a way of using $\mathbf K$ and $\mathbf S$ to simulate adding ' λV ' to an expression.

If V is a variable and E is a combinatory expression, then the combinatory expression λ^*V . E is defined inductively on the structure of E as follows:

(i)
$$\lambda^* V$$
. $V = \mathbf{I}$

(ii)
$$\lambda^* V$$
. $V' = \mathbf{K} V'$ (if $V \neq V'$)

(iii)
$$\lambda^* V$$
. $C = \mathbf{K} C$ (if C is a combinator)

(iv)
$$\lambda^* V$$
. $(E_1 \ E_2) = \mathbf{S} \ (\lambda^* V . E_1) \ (\lambda^* V . E_2)$

Note that $\lambda^* V$. E is a combinatory expression not containing V.

Example: If f and x are variables and $f \neq x$, then:

$$\lambda^* x$$
. $f x = \mathbf{S} (\lambda^* x \cdot f) (\lambda^* x \cdot x)$
= $\mathbf{S} (\mathbf{K} f) \mathbf{I}$

The following theorem shows that λ^*V . E simulates λ -abstraction.

Theorem
$$(\lambda^* V. E) = \lambda V. E$$

Proof

We show that $(\lambda^* V. E) V = E$. It then follows immediately that $\lambda V. (\lambda^* V. E) V = \lambda V. E$ and hence by η -reduction that $\lambda^* V. E = \lambda V. E$.

The proof that $(\lambda^* V. E) V = E$ is by mathematical induction on the 'size' of E. The argument goes as follows:

(i) If E = V then:

$$(\lambda^* V. E) V = \mathbf{I} V = (\lambda x. x) V = V = E$$

(ii) If E = V' where $V' \neq V$ then:

$$(\lambda^* V. E) V = \mathbf{K} V' V = (\lambda x y. x) V' V = V' = E$$

(iii) If E = C where C is a combinator, then:

$$(\lambda^* V. E) V = \mathbf{K} C = (\lambda x y. x) C V = C = E$$

(iv) If $E = (E_1 E_2)$ then we can assume by induction that:

$$(\lambda^* V. E_1) V = E_1$$
$$(\lambda^* V. E_2) V = E_2$$

and hence

$$(\lambda^* V. E) \ V = (\lambda^* V. (E_1 \ E_2)) \ V$$

$$= (\mathbf{S} \ (\lambda^* V. \ E_1) \ (\lambda^* V. \ E_2)) \ V$$

$$= (\lambda f \ g \ x. \ f \ x \ (g \ x)) \ (\lambda^* V. \ E_1) \ (\lambda^* V. \ E_2) \ V$$

$$= (\lambda^* V. \ E_1) \ V \ ((\lambda^* V. \ E_2) \ V)$$

$$= E_1 \ E_2$$
 (by induction assumption)
$$= E$$

The notation

$$\lambda^* V_1 \ V_2 \ \cdots \ V_n \ E$$

is used to mean

$$\lambda^* V_1 \cdot \lambda^* V_2 \cdot \cdots \lambda^* V_n \cdot E$$

Now define the translation of an arbitrary λ -expression E to a combinatory expression $(E)_{\mathtt{C}}$:

- (i) $(V)_{c} = x$
- (ii) $(E_1 \ E_2)_c = (E_1)_c \ (E_2)_c$
- (iii) $(\lambda V. E)_c = \lambda^* V. (E)_c$

Theorem For every λ -expression E we have: $E = (E)_{\mathbb{C}}$

Proof

The proof is by induction on the size of E.

- (i) If E = V then $(E)_{c} = (V)_{c} = V$
- (ii) If $E = (E_1 \ E_2)$ we can assume by induction that

$$E_1 = (E_1)_{\,\mathrm{C}}$$

$$E_2 = (E_2)_{\, C}$$

hence

$$(E)_{c} = (E_{1} E_{2})_{c} = (E_{1})_{c} (E_{2})_{c} = E_{1} E_{2} = E$$

(iii) If $E = \lambda V$. E' then we can assume by induction that

$$(E')_{\tt C} = E'$$

hence

$$(E)_{\texttt{C}} = (\lambda V. E')_{\texttt{C}}$$

$$= \lambda^* V. (E')_{\texttt{C}}$$
 (by translation rules)
$$= \lambda^* V. E'$$
 (by induction assumption)
$$= \lambda V. E'$$
 (by previous theorem)
$$= E$$

This theorem shows that any λ -expression is equal to a λ -expression built up from **K** and **S** and variables by application, i.e. the class of λ -expressions E defined by the BNF:

$$E ::= V \mid \mathbf{K} \mid \mathbf{S} \mid E_1 E_2$$

is equivalent to the full λ -calculus.

A collection of n combinators C_1, \ldots, C_n is called an n-element basis (Barendregt [4], Chapter 8) if every λ -expression E is equal to an expression built from C_i s and variables by function applications. The theorem above shows that \mathbf{K} and \mathbf{S} form a 2-element basis. The exercise below (from Section 8.1.5. of Barendregt) shows that there exists a 1-element basis.

Exercise 89

Find a combinator, \mathbf{X} say, such that any λ -expression is equal to an expression built from \mathbf{X} and variables by application. *Hint*: Consider $(\mathbf{K}, \mathbf{S}, \mathbf{K})$ $(\mathbf{K}, \mathbf{S}, \mathbf{K})$ $(\mathbf{K}, \mathbf{S}, \mathbf{K})$ and $(\mathbf{K}, \mathbf{S}, \mathbf{K})$ $((\mathbf{K}, \mathbf{S}, \mathbf{K}))$

Examples:

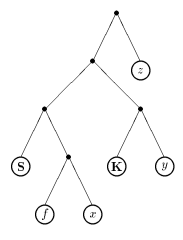
```
\begin{split} \lambda^*f.\ \lambda^*x.\ f\ (x\ x) &= \lambda^*f.\ (\lambda^*x.\ f\ (x\ x)) \\ &= \lambda^*f.\ (\mathbf{S}\ (\lambda^*x.\ f)\ (\lambda^*x.\ x\ x)) \\ &= \lambda^*f.\ (\mathbf{S}\ (\mathbf{K}f)\ (\mathbf{S}(\lambda^*x.\ x)\ (\lambda^*x.\ x))) \\ &= \lambda^*f.\ (\mathbf{S}\ (\mathbf{K}f)\ (\mathbf{S}\ \mathbf{I}\ \mathbf{I})) \\ &= \mathbf{S}\ (\lambda^*f.\ \mathbf{S}\ (\mathbf{K}f))\ (\lambda^*f.\ \mathbf{S}\ \mathbf{I}\ \mathbf{I}) \\ &= \mathbf{S}\ (\mathbf{S}\ (\lambda^*f.\ \mathbf{S})\ (\lambda^*f.\ \mathbf{K}\ f))\ (\mathbf{K}\ (\mathbf{S}\ \mathbf{I}\ \mathbf{I})) \\ &= \mathbf{S}\ (\mathbf{S}\ (\mathbf{K}\ \mathbf{S})\ (\mathbf{S}\ (\lambda^*f.\ \mathbf{K})\ (\lambda^*f.\ f)))\ (\mathbf{K}\ (\mathbf{S}\ \mathbf{I}\ \mathbf{I})) \\ &= \mathbf{S}\ (\mathbf{S}\ (\mathbf{K}\ \mathbf{S})\ (\mathbf{S}\ (\mathbf{K}\ \mathbf{K})\ \mathbf{I}))\ (\mathbf{K}\ (\mathbf{S}\ \mathbf{I}\ \mathbf{I})) \end{split}
```

```
\begin{split} (\mathbf{Y})_{\mathtt{C}} &= (\lambda f.\ (\lambda x.\ f(x\ x))\ (\lambda x.\ f(x\ x)))_{\mathtt{C}} \\ &= \lambda^* f.\ ((\lambda x.\ f(x\ x))\ (\lambda x.\ f(x\ x)))_{\mathtt{C}} \\ &= \lambda^* f.\ ((\lambda x.\ f(x\ x))_{\mathtt{C}}\ (\lambda x.\ f(x\ x))_{\mathtt{C}}) \\ &= \lambda^* f.\ (\lambda^* x.\ (f(x\ x))_{\mathtt{C}})\ (\lambda^* x.\ (f(x\ x))_{\mathtt{C}}) \\ &= \lambda^* f.\ (\lambda^* x.\ f(x\ x))\ (\lambda^* x.\ f(x\ x)) \\ &= \mathbf{S}\ (\lambda^* f.\ \lambda^* x.\ f(x\ x))\ (\lambda^* f.\ \lambda^* x.\ f(x\ x)) \\ &= \mathbf{S}(\mathbf{S}(\mathbf{S}(\mathbf{KS})(\mathbf{S}(\mathbf{KK})\mathbf{I}))(\mathbf{K}(\mathbf{SII})))(\mathbf{S}(\mathbf{S}(\mathbf{KS})(\mathbf{S}(\mathbf{KK})\mathbf{I}))(\mathbf{K}(\mathbf{SII}))) \end{split}
```

8.3 Reduction machines

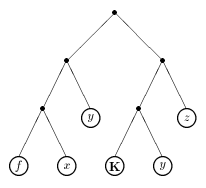
Until Dave Turner published his paper [68], combinators were regarded as a mathematical curiosity. In his paper Turner argued that translating functional languages, i.e. languages based on the λ -calculus, to combinators and then reducing the resulting expressions using the rewrites given on page 130 is a practical way of implementing these languages.

Turner's idea is to represent combinatory expressions by trees. For example, \mathbf{S} $(f\ x)$ $(\mathbf{K}\ y)$ z would be represented by:

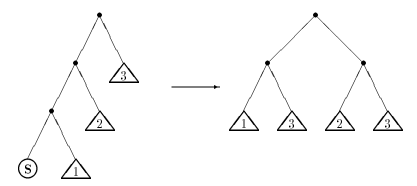


Such trees are represented as pointer structures in memory. Special hardware or firmware can then be implemented to transform such trees according to the rules of combinator reduction defining \xrightarrow{c} .

For example, the tree above could be transformed to:



using the transformation

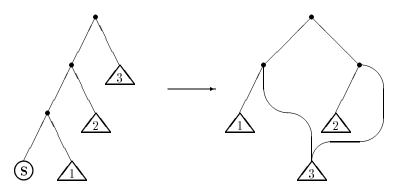


which corresponds to the reduction **S** E_1 E_2 $E_3 \xrightarrow[c]{} (E_1$ $E_2)$ $(E_2$ $E_3)$.

Exercise 90

What tree transformation corresponds to **K** E_1 E_2 \xrightarrow{c} E_1 ? How would this transformation change the tree above? \square

Notice that the tree transformation for ${\bf S}$ just given duplicates a subtree. This wastes space; a better transformation would be to generate one subtree with two pointers to it, i.e.



This generates a *graph* rather than a tree. For further details of such *graph reductions* see Turner's paper [68].

It is clear from the theorem above that a valid way of reducing λ -expressions is:

(i) Translating to combinators (i.e. $E \mapsto (E)_{c}$).

(ii) Applying the rewrites

K
$$E_1$$
 $E_2 \xrightarrow{c} E_1$
S E_1 E_2 $E_3 \xrightarrow{c} (E_1$ $E_3)$ $(E_2$ $E_3)$

until no more rewriting is possible.

An interesting question is whether this process will 'fully evaluate' expressions. If some expression E is translated to combinators, then reduced using \xrightarrow{c} , is the resulting expression as 'fully evaluated' as the result of λ -reducing E directly, or is it only partially evaluated? Surprisingly, there doesn't seem to be anything in the literature on this important question². However, combinator machines have been built and they appear to work [12]!

It is well known that if $E_1 \longrightarrow E_2$ in the λ -calculus, then it is *not* necessarily the case that $(E_1)_{\mathbb{C}} \longrightarrow (E_2)_{\mathbb{C}}$. For example, take

$$E_1 = \lambda y. \ (\lambda z. \ y) \ (x \ y)$$

$$E_2 = \lambda y. \ y$$

Exercise 91

With E_1 and E_2 as above show that $E_1 \longrightarrow E_2$ in the λ -calculus, but it is not the case that $(E_1)_{\mathbb{C}} \xrightarrow{\mathbb{C}} (E_2)_{\mathbb{C}}$. \square

A combinatory expression is defined to be in *combinatory normal form* if it contains no subexpressions of the form \mathbf{K} E_1 E_2 or \mathbf{S} E_1 E_2 E_3 . Then the normalization theorem holds for combinatory expressions, i.e. always reducing the leftmost combinatory redex will find a combinatory normal form if it exists.

Note that if E is in combinatory normal form, then it does not necessarily follow that it is a λ -expression in normal form.

Example: S K is in combinatory normal form, but it contains a β -redex, namely:

$$(\lambda f. (\lambda g \ x. (f \ x (g \ x))) (\lambda x \ y. \ x)$$

Exercise 92

Construct a combinatory expression E which is in combinatory normal form, but has no normal form. \square

 $^{^2}$ The most relevant paper I could find is one by Hindley [30]. This compares λ -reduction with combinatory reduction, but not in a way that is prima facie relevant to the termination of combinator machines.

8.4 Improved translation to combinators

The examples on page 134 show that simple λ -expressions can translate to quite complex combinatory expressions via the rules on page 133.

To make the 'code' executed by reduction machines more compact, various optimizations have been devised.

Examples

(i) Let E be a combinatory expression and x a variable not occurring in E. Then:

$$\mathbf{S} \ (\mathbf{K} \ E) \ \mathbf{I} \ x \longrightarrow_{\mathcal{C}} (\mathbf{K} \ E \ x) \ (\mathbf{I} \ x) \longrightarrow_{\mathcal{C}} E \ x$$

hence **S** (**K**E) **I** x = E x (because $E_1 \xrightarrow{c} E_2$ implies $E_1 \xrightarrow{} E_2$), so by extensionality (Section 4.7, see on page 71):

$$\mathbf{S} (\mathbf{K} E) \mathbf{I} = E$$

(ii) Let E_1 , E_2 be combinatory expressions and x a variable not occurring in either of them. Then:

$$\mathbf{S} \ (\mathbf{K} \ E_1) \ (\mathbf{K} \ E_2) \ x \xrightarrow[\mathbf{c}]{} \mathbf{K} \ E_1 \ x \ (\mathbf{K} \ E_2) \ x \xrightarrow[\mathbf{c}]{} E_1 \ E_2$$

Thus

$$S(K E_1)(K E_2) x = E_1 E_2$$

Now

$$\mathbf{K} (E_1 \ E_2) \ x \xrightarrow{c} E_1 \ E_2$$

hence **K** $(E_1 \ E_2) \ x = E_1 \ E_2$. Thus

$$S (K E_1) (K E_2) x = E_1 E_2 = K (E_1 E_2) x$$

It follows by extensionality that:

$$S (K E_1) (K E_2) = K (E_1 E_2)$$

Since **S** (**K** E) **I** = E for any E, whenever a combinatory expression of the form **S** (**K** E) **I** is generated, it can be 'peephole optimized' to just E. Similarly, whenever an expression of the form **S** (**K** E_1) (**K** E_2) is generated, it can be optimized to **K** (E_1 E_2).

Example: On page 134 it was shown that:

$$\lambda^* f. \ \lambda^* x. \ f(x \ x) = \mathbf{S} \ (\mathbf{S} \ (\mathbf{K} \ \mathbf{S}) \ (\mathbf{S} \ (\mathbf{K} \ \mathbf{K}) \ \mathbf{I})) \ (\mathbf{K} \ (\mathbf{S} \ \mathbf{I} \ \mathbf{I}))$$

Using the optimization **S** (**K** E) **I** = E this simplifies to:

$$\lambda^* f. \ \lambda^* x. \ f(x \ x) = \mathbf{S} \ (\mathbf{S} \ (\mathbf{K} \ \mathbf{S}) \ \mathbf{K}) \ (\mathbf{K} \ (\mathbf{S} \ \mathbf{I} \ \mathbf{I}))$$

8.5 More combinators

It is easier to recognize the applicability of the optimization \mathbf{S} (\mathbf{K} E) $\mathbf{I} = E$ if \mathbf{I} has not been expanded to \mathbf{S} \mathbf{K} \mathbf{K} , i.e. if \mathbf{I} is taken as a primitive combinator. Various other combinators are also useful in the same way; for example, \mathbf{B} and \mathbf{C} defined by:

LET
$$\mathbf{B} = \lambda f g x \cdot f (g x)$$

LET
$$C = \lambda f g x. f x g$$

These have the following reduction rules:

$$\mathbf{B} \ E_1 \ E_2 \ E_3 \xrightarrow[\mathbf{c}]{} E_1 \ (E_2 \ E_3)$$

$$\mathbf{C} \ E_1 \ E_2 \ E_3 \xrightarrow[\mathbf{c}]{} E_1 \ E_3 \ E_2$$

Exercise 93

Show that with **B**, **C** defined as above:

S (**K**
$$E_1$$
) E_2 = **B** E_1 E_2

$$\mathbf{S} \ E_1 \ (\mathbf{K} \ E_2) = \mathbf{C} \ E_1 \ E_2$$

(where E_1 , E_2 are any two combinatory expressions). \square

Using **B** and **C**, one can further optimize the translation of λ -expressions to combinators by replacing expressions of the form **S** (**K** E_1) E_2 and **S** E_1 (**K** E_2) by **B** E_1 E_2 and **C** E_1 E_2 .

8.6 Curry's algorithm

Combining the various optimizations described in the previous section leads to Curry's algorithm for translating λ -expressions to combinatory expressions. This algorithm consists in using the definition of $(E)_{\mathbb{C}}$ given on page 133, but whenever an expression of the form \mathbf{S} E_1 E_2 is generated one tries to apply the following rewrite rules:

- 1. **S** (**K** E_1) (**K** E_2) \longrightarrow (**K** E_1 E_2)
- 2. **S** (**K** E) **I** \longrightarrow E
- 3. S (K E_1) $E_2 \longrightarrow \mathbf{B} \ E_1 \ E_2$
- 4. $\mathbf{S}E_1(\mathbf{K}E_2) \longrightarrow \mathbf{C}E_1E_2$

If more than one rule is applicable, the *earlier* one is used. For example, \mathbf{S} (\mathbf{K} E_1) (\mathbf{K} E_2) is translated to \mathbf{K} (E_1 E_2), not to \mathbf{B} E_1 (\mathbf{K} E_2).

Exercise 94

Show that using Curry's algorithm, \mathbf{Y} is translated to the combinator:

Exercise 95

Show that:

$$S(S(KS)(S(KK)I))(K(SII)) = CB(SII)$$

8.7 Turner's algorithm

In a second paper, Turner proposed that Curry's algorithm be extended to use another new primitive combinator called S' [69]. This is defined by:

LET
$$S' = \lambda c f q x \cdot c (f x) (q x)$$

and has the reduction rule:

$$\mathbf{S}' \ C \ E_1 \ E_2 \ E_3 \longrightarrow C \ (E_1 \ E_3) \ (E_2 \ E_3)$$

where C, E_1 , E_2 , E_3 are arbitrary combinatory expressions. The reason why 'C' is used is that S' has the property that if C is a combinator (i.e. contains no variables), then for any E_1 and E_2 :

$$\lambda^* x \cdot C \ E_1 \ E_2 = \mathbf{S}' \ C \ (\lambda^* x \cdot E_1) \ (\lambda^* x \cdot E_2)$$

This can be shown using extensionality. Clearly x is a variable not occurring in $\lambda^* x$. C E_1 E_2 or \mathbf{S}' C $(\lambda^* x$. $E_1)$ $(\lambda^* x$. $E_2)$ (exercise: why?), so it is sufficient to show:

$$(\lambda^* x \cdot C \ E_1 \ E_2) \ x = (\mathbf{S}' \ C \ (\lambda^* x \cdot E_1) \ (\lambda^* x \cdot E_2)) \ x$$

From the definition of $\lambda^* x$ it easily follows that:

$$\lambda^* x \cdot C E_1 E_2 = \mathbf{S} \left(\mathbf{S} \left(\mathbf{K} C \right) \left(\lambda^* x \cdot E_1 \right) \right) \left(\lambda^* x \cdot E_2 \right)$$

hence

$$(\lambda^* x. C E_1 E_2) x = (\mathbf{S} (\mathbf{K} C) (\lambda^* x. E_1)) (\lambda^* x. E_2)) x$$

$$= \mathbf{S} (\mathbf{K} C) (\lambda^* x. E_1) x ((\lambda^* x. E_2)) x)$$

$$= \mathbf{K} C x ((\lambda^* x. E_1) x) ((\lambda^* x. E_2)) x)$$

$$= C ((\lambda^* x. E_1) x) ((\lambda^* x. E_2)) x)$$

But (S' C (λ^*x . E_1) (λ^*x . E_2)) x = C ((λ^*x . E_1) x) ((λ^*x . E_2)) x) also, and so:

$$(\lambda^* x. \ C \ E_1 \ E_2) \ x = (\mathbf{S}' \ C \ (\lambda^* x. \ E_1) \ (\lambda^* x. \ E_2)) \ x$$

Exercise 96

Where in the argument above did we use the assumption that C is a combinator? \square

Turner's combinator S' is useful when translating λ -expressions of the form $\lambda V_n \cdots V_2 V_1$. $E_1 E_2$ (it will be seen shortly why it is convenient to number the bound variables in descending order). To see this, following Turner [69], temporarily define

$$\begin{array}{lll} E' & \text{to mean} & \lambda^* V_1 . \ E \\ E'' & \text{to mean} & \lambda^* V_2 . \ (\lambda^* V_1 . \ E) \\ E''' & \text{to mean} & \lambda^* V_3 . \ (\lambda^* V_2 . \ (\lambda^* V_1 . \ E)) \\ & \vdots & \end{array}$$

Recall that:

$$(\lambda V_n \cdots V_2 \ V_1 \cdot E_1 \ E_2)_{\mathbf{c}} = \lambda^* V_n \cdot (\cdots (\lambda^* V_2 \cdot (\lambda^* V_1 \cdot (E_1 \ E_2)_{\mathbf{c}}))) \cdots)$$

The next exercise shows that:

$$\lambda^* V_n \dots \lambda^* V_2 \dots \lambda^* V_1 \dots (E_1 E_2)$$

gets very complicated as n increases.

Exercise 97

Show that:

- (i) $\lambda^* x_1$. $E_1 E_2 = \mathbf{S} E_1' E_2'$
- (ii) $\lambda^* x_2$. $(\lambda^* x_1$. $E_1 E_2) = \mathbf{S} (\mathbf{B} \mathbf{S} E_1'') E_2''$

(iii)
$$\lambda^* x_3$$
. $(\lambda^* x_2$. $(\lambda^* x_1$. $E_1 E_2)) = \mathbf{S} (\mathbf{B} \mathbf{S} (\mathbf{B} \mathbf{S}) E_1''')) E_2'''$

(iv)
$$\lambda^* x_4$$
. $(\lambda^* x_3$. $(\lambda^* x_2$. $(\lambda^* x_1$. $E_1 E_2))) =$ **S** (**B** S (**B** (**B** S) (**B** (**B** S))) E_1'''') E_2''''

The size of $\lambda^* V_n \dots \lambda^* V_2 \dots \lambda^* V_1 \dots (E_1 E_2)$ is proportional to the *square* of n. Using \mathbf{S}' , the size can be made to grow *linearly* with n:

$$\lambda^* x_2. (\lambda^* x_1. E_1 E_2) = \lambda^* x_2. \mathbf{S} E_1' E_2' = \mathbf{S}' \mathbf{S} (\lambda^* x_2. E_1') (\lambda^* x_2. E_2') = \mathbf{S}' \mathbf{S} E_1'' E_2''$$

$$\lambda^* x_3. (\lambda^* x_2. (\lambda^* x_1. E_1 E_2)) = \lambda^* x_3. \mathbf{S}' \mathbf{S} E_1'' E_2'' = \mathbf{S}' (\mathbf{S}' \mathbf{S}) (\lambda^* x_3. E_1') (\lambda^* x_3. E_2'') = \mathbf{S}' (\mathbf{S}' \mathbf{S}) E_1''' E_2'''$$

$$\begin{array}{lll} \lambda^* x_4. \; \lambda^* x_3. \; (\lambda^* x_2. \; (\lambda^* x_1. \; E_1 \; E_2))) & = \lambda^* x_4. \; \mathbf{S}' \; (\mathbf{S}' \; \mathbf{S}) \; E_1''' \; E_2''' \\ & = \mathbf{S}' \; (\mathbf{S}' \; (\mathbf{S}' \; \mathbf{S})) \; (\lambda^* x_4. \; E_1''') \; (\lambda^* x_4. \; E_2''') \\ & = \mathbf{S}' \; (\mathbf{S}' \; (\mathbf{S}' \; \mathbf{S})) \; E_1'''' \; E_2'''' \end{array}$$

Just as **B** and **C** were introduced to simplify combinatory expressions of the form **S** (**K** E_1) E_2 and **S** E_1 (**K** E_2) respectively, Turner also devised **B**' and **C**' with an analogous role for **S**'. The properties required are:

$$\mathbf{S}' \ C \ (\mathbf{K} \ E_1) \ E_2 = \mathbf{B}' \ C \ E_1 \ E_2$$

 $\mathbf{S}' \ C \ E_1 \ (\mathbf{K} \ E_2) = \mathbf{C}' \ C \ E_1 \ E_2$

(where C is any combinator, and E_1 , E_2 are arbitrary combinatory expressions). This is achieved if \mathbf{B}' and \mathbf{C}' are defined by:

LET
$$\mathbf{B}' = \lambda c \ f \ g \ x. \ c \ f \ (g \ x)$$

LET $\mathbf{C}' = \lambda c \ f \ g \ x. \ c \ (f \ x) \ g$

Clearly \mathbf{B}' and \mathbf{C}' will have the property that for arbitrary λ -expressions C, E_1 , E_2 and E_3 :

$$\mathbf{B'} \ C \ E_1 \ E_2 \ E_3 \xrightarrow[c]{} C \ E_1 \ (E_2 \ E_3)$$
$$\mathbf{C'} \ C \ E_1 \ E_2 \ E_3 \xrightarrow[c]{} C \ (E_1 \ E_3) \ E_2$$

Exercise 98

Show that for arbitrary λ -expressions C, E_1 , E_2 and E_3 :

(i)
$$\mathbf{S}' \ C \ (\mathbf{K} \ E_1) \ E_2 = \mathbf{B}' \ C \ E_1 \ E_2$$

(ii)
$$S' C E_1 (K E_2) = C' C E_1 E_2$$

(iii)
$$\mathbf{S}$$
 (\mathbf{B} \mathbf{K} E_1) $E_2 = \mathbf{S}'$ \mathbf{K} E_1 E_2

(iv) **B** (**K**
$$E_1$$
) E_2 = **B**' **K** E_1 E_2

(v) **C** (**B K**
$$E_1$$
) E_2 = **C**' **K** E_1 E_2

Turner's algorithm for translating λ -expressions to combinatory expressions is described by him [69] as follows:

Use the algorithm of Curry but whenever a term beginning in **S**, **B** or **C** is formed use one of the following transformations if it is possible to do so

$$\mathbf{S} (\mathbf{B} \ K \ A) \ B \longrightarrow \mathbf{S}' \ K \ A \ B,$$

$$\mathbf{B} (K \ A) \ B \longrightarrow \mathbf{B}' \ K \ A \ B,$$

$$\mathbf{C} (\mathbf{B} \ K \ A) \ B \longrightarrow \mathbf{C}' \ K \ A \ B.$$

Here A and B stand for arbitrary terms as usual and K is any term composed entirely of constants. The correctness of the new algorithm can be inferred from the correctness of the Curry algorithm by demonstrating that in each of the above transformations the left- and right-hand sides are extensionally equal. In each case this follows directly from the definitions of the combinators involved.

Since Turner's pioneering papers appeared, many people have worked on improving the basic idea. For example, John Hughes has devised a scheme for dynamically generating an 'optimal' set of primitive combinators (called *supercombinators*) for each program [36]. The idea is that the compiler will generate combinatory expressions built out of the supercombinators for the program being compiled. It will also dynamically produce 'microcode' to implement the reduction rules for these supercombinators. The result is that each program runs on a reduction machine tailored specially for it. Most current high-performance implementations of functional languages use supercombinators [2, 17]. Another avenue of research is to use combinators based on the De Bruijn notation briefly described on page 74. The 'Categorical Abstract Machine' [51] uses this approach.

Part III Implementing the Theories

A Quick Introduction to LISP

The LISP programming language is introduced. Enough detail is given to enable the reader to understand all the programs making up the theorem prover in Chapter 10, the verifier in Chapter 11 and the λ -calculus toolkit in Chapter 12.

In Part III of this book the ideas in the preceding two parts are illustrated by describing some LISP programs which embody the theories described there. These programs are small enough so that it is feasible to type them into a file and play with them; they are, however, often rather inefficient. The code has been written to be short and easy to follow; this has sometimes been at the expense of efficiency. The goal is that the programs be efficient enough for simple experiments; it is not expected that they will have anything more than educational value. Readers interested in implementing more efficient versions can use the prototypes provided here as a starting point and, with suitable intrumentation, can do experiments to determine those things that need optimizing. The purpose of this chapter is to provide a sufficient introduction to LISP so that the programs that follow will be comprehensible to readers who have not met the language before. For a more comprehensive introduction Wilensky's book LISP craft [73] is recommended. From now on this will be referred to as 'Wilensky'.

LISP is probably the second oldest programming language in widespread use (FORTRAN is the oldest). It is the first functional language to have been developed and its design was strongly based on the λ -calculus. Unfortunately, LISP functions differ in subtle and confusing ways from ordinary mathematical functions (and hence from the λ -calculus). This is partly for efficiency reasons and partly due to errors in the original design. For example, McCarthy, the inventor of the language, says in his paper on the history of LISP [52] that the dynamic binding of free variables (see Section 9.7) used by LISP's λ -expressions was a mistake.

It is possible to do imperative programming in LISP (see the description of the prog feature on page 159) and the resulting potential for side effects also complicates the relation between mathematical functions and LISP functions, so much so that some authors regard LISP as a bad influence. Here, for example, is a quotation by David Turner [70] from the discussion after his paper in the book Mathematical Logic and Programming Languages [35]:

It needs to be said very firmly that LISP, at least as represented by the dialects in common use, is not a functional language at all. LISP does have a functional subset, but that is a rather inconvenient programming language and there exists no significant body of programs written in it. Almost all serious programming in LISP makes heavy use of side effects and other referentially opaque features.

I think that the historical importance of LISP is that it was the first language to provide 'garbage-collected' heap storage. This was a very important step forward. For the development of functional programming, however, I feel that the contribution of LISP has been a negative one. My suspicion is that the success of LISP set back the development of a properly functional style of programming by at least ten years.

I have some sympathy for this view, but unfortunately there are no robust and efficient functional language implementations that I felt could reasonably be used instead of LISP. The best that are available are probably Standard ML [55] and Miranda [70]. Although current ML implementations¹ rival or exceed LISP in robustness and efficiency, they have not yet become widely distributed. Furthermore, ML, like LISP, has imperative features that complicate its semantics; however, unlike LISP, pure ML functions (i.e. ones without side effects) do correspond to mathematical functions. The Miranda system is very elegant, but it is currently not efficient enough for the kind of applications given here. Furthermore, Miranda is not available free and the expense of buying a system may unfortunately slow down its distribution.

If a suitable functional programming language becomes widely distributed, then it would probably be used in future editions of this book. To the extent that is practical, only the functional subset of LISP will be used here.

¹ Implementations of ML are available from Edinburgh University, Bell Laboratories, Imperial Software Technology and as part of Sussex University's Poplog system.

9.1 Features of LISP

According to John McCarthy, LISP is characterized by the ideas listed below (see his history of the language [52] and Mason's book [50]):

- Computing with symbolic expressions rather than numbers.
- Representation of symbolic expressions and other information by list structure in the memory of a computer.
- A small set of selector and constructor operations expressed as functions.
- Composition of functions as a tool for forming more complex functions.
- The use of conditional expressions for getting branching into function definitions.
- The recursive use of conditional expressions as a sufficient tool for building computable functions.
- The use of λ -expressions for naming functions.
- The representation of LISP programs as LISP data that can be manipulated by other programs.
- The conditional expression interpretation of Boolean connectives.
- The LISP function eval that serves both as a formal definition of the language and as an interpreter.
- Garbage collection.
- LISP statements as a command language for an interactive environment.

Some of these ideas were taken from other languages, but most were new (and due to McCarthy).

There are many versions of LISP in current use, for example:

- Maclisp. An influential LISP system produced at MIT.
- Zetalisp. An enhancement of Maclisp that runs on Symbolics LISP Machines.
- Franz LISP. A version of Maclisp that runs on Vaxes and Sun workstations.

- Interlisp. A sophisticated LISP programming environment developed at BBN and Xerox PARC. It runs on the Xerox workstations.
- Standard LISP. An early attempt to produce a standard lead by a group from the University of Utah.
- Cambridge LISP. This this was originally developed for IBM mainframes, but versions are now available for BBC microcomputers and Commodore Amigas. Cambridge LISP is quite similar to Standard LISP.
- Common LISP. A recent, fairly successful, attempt to produce a standard. Many people are now converting to Common LISP. Some of the horrors in the treatment of functions in other LISPs are corrected in Standard LISP.

In this book, the intention is to stay within a subset that is common to most LISPs. Where arbitrary notational choices are made, Franz LISP is used. This is the version of LISP used in Wilensky and public domain implementations are available for Vax and Sun computers running Unix. If no suitable purely functional languages become available, future editions of this book will probably use Common LISP because of its superior treatment of functions (lexical scoping is the default, see Section 9.7).

9.2 Some history

The earliest versions of LISP used a notation called M-expressions (described in [53]) that was 'intended to resemble FORTRAN as much as possible' [52]. The data manipulated by these M-expressions were things called S-expressions. The 'M' and 'S' in 'M-expression' and 'S-expression' abbreviate 'meta' and 'symbolic'.

To explain the semantics of LISP, McCarthy wrote an interpreter for LISP in LISP. This interpreter, called eval, used an encoding of M-expressions as S-expressions, so that programs (i.e. M-expressions) could be manipulated as data (i.e. S-expressions). This encoding provided an alternative syntax for LISP which rapidly replaced the M-expression syntax. Now only the S-expression notation is used and M-expressions are a historical relic. This little bit of history is worth knowing because it explains various features of the language. For example, McCarthy says [52]

The unexpected appearance of an interpreter tended to freeze the form of the language, and some of the decisions made rather lightheartedly . . . later proved unfortunate. These included the COND notation for conditional expressions which leads to unnecessary depth of parentheses.

Later, McCarthy adds:

Another reason for the initial acceptance of awkwardnesses in the internal form of LISP is that we still expected to switch to writing programs as M-expressions. The project of defining M-expressions precisely and compiling them or at least translating them into S-expressions was neither finalized nor explicitly abandoned. It just receded into the indefinite future, and a new generation of programmers appeared who preferred internal notation to any FORTRAN-like or ALGOL-like notation that could be devised.

9.3 S-expressions

LISP programs consist of functions for manipulating S-expressions. The functions themselves are also represented by S-expressions. The process of running a program consists in *evaluating* an S-expression representing the application of a function to some arguments.

An S-expression is either an atom or a dotted-pair $(S_1 cdot S_2)$ where S_1 and S_2 are S-expressions. Dotted pairs are sometimes called lists, but this term is used here in a different way (see Section 9.3.1 below). Small italics will be used for arbitrary S-expressions (e.g. S, S_1, S_2 etc.) and small typewriter font for particular S-expressions (e.g. foo, (foo . F00)).

Atoms are sequences of characters, for example:

$$x, y, foo, 22, +, -, *, =, <, >, >=, <=$$

The exact syntax of atoms is implementation dependent. Most of the atoms to be used will be the names of variables or functions. Such names will consist of sequences of letters or numbers starting with a letter. Other atoms represent numbers and mathematical operations. Some characters have a special meaning to the LISP reader. To include such characters in an atom, the atom name must be input with vertical bars (i.e. |'s) surrounding it. For example, |'''| is an atom whose name consists of three quote characters; note that the vertical bars are not part of the atom.

The atoms t and nil have a special role in LISP; as will be seen in Section 9.5.2 they represent 'true' and 'false' respectively. Some LISPs (e.g. Common LISP) do not distinguish upper and lower case; however Franz LISP does distinguish them: for example, for us (but not for some systems) the atoms F00 and foo are different.

Examples of S-expressions are:

```
nil, FOO, (FOO . foo), ((A . nil) . ((B . nil) . nil))
```

Strings are a special kind of atom. They are input as a sequence of characters surrounded by quotes ("). For example,

```
"string1", "**?!", "This is a long string containing spaces."
```

Strings evaluate to themselves; they are useful for error messages and are more efficiently implemented then arbitrary atoms.

9.3.1 Lists

A list is the atom nil or a dotted-pair $(S_1 . S_2)$. Note that nil is the only S-expression that is both an atom and a list. It is called the *empty list*. Lists of the form:

$$(S_1 \cdot (S_2 \cdot (S_3 \cdot \cdots (S_n \cdot \text{nil}) \cdot \cdots)))$$

are called *linear lists* and are written as:

$$(S_1 \ S_2 \ S_3 \ \cdots \ S_n)$$

The notation () denotes the atom nil.

Warnings:

- (i) Sometimes the term 'list' is used for what we call 'linear list'.
- (ii) In some LISPs (e.g. Franz LISP) the expression (A.B) (where A and B are atoms) will be interpreted as the S-expression (A.B. nil), where A.B is a single atom. This is because some LISP readers do not recognize dotted pairs unless there are spaces around the dot. In such LISPs, (A.B) would be different from (A.B). A further complication is that in Franz LISP (1.2) is the linear list containing the floating point number 1.2; this atom is different from the atom |1.2| (see the descriptions of the predicates symbolp and number p in Section 9.12).

Each S-expression has a *value* which is obtained by evaluating it. The atoms t and nil evaluate to themselves, as do strings " ... " and the numbers 0, 1, 2 etc.

9.4 Variables and the environment

LISP atoms can be bound to S-expressions in the environment. Such atoms are called variables and when they are evaluated the result is the corresponding S-expression. Variables can either be part of the top-level interaction with LISP (global variables) or they can be local to the application of a function.

A variable x can be set to the value of an S-expression S by evaluating the S-expression (setq x S). For example, evaluating (setq one 1) sets the value of one to 1.

If S is an S-expression then (quote S) is an S-expression that evaluates to S. For example, (setq mike (quote (1 2 3))) binds the variable mike to (1 2 3). Note that evaluating (setq mike (1 2 3)) would result in an error, because setq would try to evaluate (1 2 3), but 1 is not a function; this is explained further in the next section.

The S-expression (quote S) can be abbreviated by 'S. Thus evaluating (setq mike '(1 2 3)) binds mike to (1 2 3).

9.5 Functions

If S is an S-expression representing a LISP function (see below) then

$$(S S_1 \ldots S_n)$$

is normally evaluated by

- (i) first evaluating S_1, \ldots, S_n (in that order)
- (ii) and then applying (see below) S to the resulting values.

For example, (add 2 3) evaluates to 5. If S is not an S-expression representing a function, then an error results. Thus, since 1 is not a function, evaluating (1 2 3) will cause an error. We will generally use f, g, h etc. to range over S-expressions representing functions.

Certains names are bound to special functions that do not evaluate their arguments. For example, setq does not evaluate its first argument. There is a function set that does evaluate its first argument: (setq x S) is equivalent to (set (quote x) S). Both setq and set have a side effect of binding a variable to a value. They also have a value, which is the value of their second argument. For example, (setq x 1) evaluates to 1.

Exercise 99

What are the values and side effects of:

- (i) (add 1 (setq x 2))
- (ii) (add (setq x 1) (setq x (add x 2)))
- (iii) (setq x (add (setq x 1) x))?

Functions that do not evaluate their arguments are normally built-in system functions (like setq). In most LISPs (but not Common LISP) one can define a kind of function called an fexpr which does not evaluate its arguments, however the use of fexprs is nowadays considered poor practice; see Chapter 12 of Wilensky for details. A facility that is very useful, however, is the definition of macros; these provide a way of preprocessing Sexpressions before they are evaluated. By defining macros that insert quotes around arguments, it is possible to control the evaluation of arguments in a clean and efficient way. This is described in detail in Section 9.10.

It is also possible to have functions that appear to take a variable number of arguments (e.g. cond, casesq, and and or in Section 9.5.2). Most such functions are built-in macros or fexprs.

User defined LISP functions are represented by S-expressions of the form (lambda $(x_1 \ldots x_n)$ S). Such functions are applied to arguments S_1, \ldots, S_n by binding them to x_1, \ldots, x_n respectively and then evaluating S in the resulting environment. For example, ((lambda x y z) (add (add x y) z)) 1 2 3) evaluates to 6. The binding of the arguments to the function variables x_1, \ldots, x_n is local to the evaluation of S.

A lambda-expression (lambda $(x_1 \ldots x_n)$ S) can be given a name f in the environment by evaluating:

$$(\text{def } f \text{ (lambda } (x_1 \ldots x_n) S))$$

Applying the name of a function is equivalent to applying the corresponding lambda-expression. Instead of defining functions with def as above, it is usual to use the equivalent simpler form:

$$(\text{defun } f (x_1 \ldots x_n) S)$$

defun is actually a macro; for its definition see page 171.

One of the awkward aspects of LISP is that function definitions are not held in the same environment in which variables are bound. Thus:

$$(setq f (quote (lambda (x_1 ... x_n) S)))$$

has a different effect from:

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```
(\text{def } f (\text{lambda} (x_1 \ldots x_n) S))
```

It is possible to apply functions that have been bound in the environment with setq. If f is such a function, then evaluating:

```
(funcall f S_1 \ldots S_n)
```

will apply the lambda-expression bound to f to the values of S_1, \ldots, S_n . Most LISPs (including Franz LISP) provide a special quoting mechanism for functions called function. Instead of

```
(setq f (quote (lambda (x_1 \ldots x_n) S)))
```

it is better style to write

```
(setq f (function (lambda (x_1 \ldots x_n) S)))
```

Although there is little semantic difference between quote and function in Franz LISP, other systems, especially Common LISP, try and make lambdas quoted with function behave more like true λ -expressions. There is more on the semantics of LISP lambda-expressions in Section 9.7.

9.5.1 Primitive list-processing functions

The names car, cdr, cons, atom and eq are bound to primitive functions with the following semantics.

- 1. (car S) evaluates S and if the resulting value is a list $(v_1 \, ... \, v_2)$ then v_1 is returned.
- 2. (cdr S) evaluates S and if the resulting value is a list $(v_1 v_2)$ then v_2 is returned.
- 3. (cons S_1 S_2) evaluates S_1 to get a value v_1 and then S_2 to get a value v_2 and then returns the list $(v_1 \, . \, v_2)$ as result.
- 4. (atom S) evaluates to t if the value of S is an atom, otherwise it evaluates to nil.
- 5. (eq S_1 S_2) evaluates to t if
 - (a) S_1 and S_2 evaluate to the same atom, or
 - (b) S_1 and S_2 evaluate to pointers to the same list cell (this is explained in Section 9.8).

otherwise (eq S_1 S_2) evaluates to nil. The exact semantics of eq is quite subtle and messy as it depends on the representation of lists as pointer structures in memory (see Section 9.8).

The names car and cdr are a historical curiosity; apparently they abbreviate 'contents of address register' and 'contents of data register'. This reflects how the first LISP systems at M.I.T. were implemented.

9.5.2 Flow of control functions

The flow of control in LISP programs is specified using various functions that do not evaluate all their arguments. The most basic such function is cond; all the other ones can be defined in terms of it.

The conditional function cond

An S-expression:

should be read as:

```
if S_{11} then S_{12} followed by S_{13} ... followed by S_{1n_1} else if S_{21} then S_{22} followed by S_{23} ... followed by S_{2n_2} else \vdots if S_{m1} then S_{m2} followed by S_{m3} ... followed by S_{mn_m} else nil
```

The exact semantics is as follows: S_{11} is evaluated and if its value is not nil then S_{12} , S_{13} , ..., S_{1n_1} are evaluated in that order and the value of S_{1n_1} is returned. If the value of S_{11} is nil then S_{21} is evaluated and if its value is not nil then S_{22} , S_{23} , ..., S_{2n_2} are evaluated in that order and the value of S_{2n_2} is returned. If the value of S_{21} is nil then this process continues until the first S_{i1} is found that has a non-nil value and then S_{i2} , S_{i3} , ..., S_{in_i} are then evaluated in that order and the value of S_{in_i} is returned. If all of S_{21} , ..., S_{m1} evaluate to nil then the value of the cond-expression is nil.

Notice that nil represents 'false' and any S-expression not equal to nil represents 'true'.

The conditional function if

```
The S-expression
```

```
(if S S_1 S_2)
```

should be read as 'if S then S_1 else S_2 ' and is an abbreviation for

```
(cond (S S_1) (t S_2))
```

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The sequential conjunction and

The S-expression

(and
$$S_1 S_2 \ldots S_n$$
)

is evaluated by evaluating S_1, S_2 , ... in that order until the first S_i is found with value nil. The value of the and-expression is then nil. If all the S-expressions S_1, \ldots, S_n have non-nil values then the value of the and-expression is the value of S_n . The value of (and) is t.

Exercise 100

Show how to represent (and S_1 S_2 ... S_n) using cond. \square

As the name suggests, and can be used to compute the logical conjunction (i.e. \wedge) of a number of expressions. A common application is to do a sequence of tests like:

```
(and x (cdr x) (cdr(cdr x)))
```

which tests whether x has at least two elements. Evaluating the S-expression (cdr(cdr x)) would cause an error if x were an atom or one-element list (e.g. 1 or (1)); using (and x (cdr x) ...) prevents this.

The sequential disjunction or

The S-expression

(or
$$S_1$$
 S_2 ... S_n)

is evaluated by evaluating S_1, S_2, \ldots in that order until the first S_i is found whose value is not equal to nil. The value of the or-expression is then the value of S_i . If all of S_1, \ldots, S_n have nil values then the value of the or-expression is nil. The value of (or) is nil.

Exercise 101

Show how to represent (or S_1 S_2 ... S_n) using cond. \square

As the name suggests, or can be used to compute the logical disjunction (i.e. \vee) of a number of expressions.

An example of the use of and and or is the definition of the list reversing function reverse given on page 159.

The cases function casesq

A cases expression has the form:

A casesq-expression assumes that the value of S is an atom, A say. This value is compared with A_1, \ldots, A_n (in that order). If

- (i) A_i is t, or
- (ii) A_i is equal to A, or
- (iii) A_i is a list containing A,

then $(S_{i1} \ S_{i2} \ \dots \ S_{in_i})$ are evaluated in that order and the value of S_{in_i} is returned. If A is not equal to any A_i then nil is returned. Note that the A_1, \ldots, A_n are not evaluated.

A casesq-expression is roughly equivalent to (but more efficient than)

```
\begin{array}{c} (\text{cond} \\ \quad \  \, ((\text{or (eq $S$ t) (eq $S$ '$A_1) (memq $S$ '$A_1)) $S_{11} \dots S_{1n_1}) \\ \quad \  \, ((\text{or (eq $S$ t) (eq $S$ '$A_2) (memq $S$ '$A_2)) $S_{21} \dots S_{2n_2}) \\ \quad \  \, \vdots \\ \quad \  \, (\text{or (eq $S$ t) (eq $S$ '$A_m) (memq $S$ '$A_m)) ($S_{m1} \dots S_{mn_m}$)) \end{array}
```

The membership testing function memq is described on page 176.

Exercise 102

Why is the cond-expression above not exactly equivalent to a casesq-expression? *Hint*: Consider what happens if S is (setq x (add x 1)). \square

The local declaration function let

The expression:

```
(let ((x_1 \ S_1) \ \dots \ (x_n \ S_n)) \ S)
```

is evaluated as follows:

(i) S_1, \ldots, S_n are evaluated in that order and their values bound to the variables x_1, \ldots, x_n respectively;

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(ii) S is evaluated in the resulting environment.

A let-expression is equivalent to:

```
((lambda (x_1 \ldots x_n) S) S_1 \ldots S_n)
```

The program function prog

FORTRAN-like imperative programming can be done in LISP using the function prog. We only use this once, to format the output produced by our program verifier.

A prog-expression has the form shown below where each l_i is an atom acting as a label for the expression S_i . If a label for S_i is not needed then l_i can be omitted.

```
\begin{array}{ccc} (\texttt{prog} & (x_1 & \dots & x_n) \\ l_1 & S_1 \\ l_2 & S_2 \\ & & \vdots \\ l_n & S_n) \end{array}
```

A prog-expression is evaluated by evaluating the S-expressions S_1, \ldots, S_n in turn. The variables x_1, \ldots, x_n are local, so that a setq to them in an S_i will not be felt outside the prog-expression. They are initialized to nil on entry to the prog-expression. If an expression (return S) is evaluated inside an S_i , then the evaluation of the whole prog-expression is terminated and the value of S is returned. If an expression (go l_i) is evaluated then, control transfers to the corresponding expression S_i .

Here, for example, is the definition of a function reverse that reverses a list, i.e. (reverse ' $(S_1 \ldots S_n)$) evaluates to $(S_n \ldots S_1)$.

In Franz LISP, the local variable temp would not be needed because the function parameter x could be used instead. On exit from reverse, the value of x on entry is restored. Also, since local variables of prog-expressions are initialized to nil, the explicit initialization (setq result nil) is not needed. Thus a simpler definition of reverse would be:

This simpler definition of reverse is not really good style as it exploits specific features of Franz LISP and thus makes the definition less portable. It is good practice to always explicitly initialize local variables as this helps document the algorithm being implemented.

The functions catch and throw

Evaluating (catch S) causes S to be evaluated. If during this evaluation (throw S_1) is evaluated, then the evaluation of S is terminated and the value of S_1 returned as the value of (catch S). If a call to throw is not encountered, then the value of (catch S) is the value of S. Thus, to quote Wilensky, 'throw throws the value of its argument through any number of levels of intermediate function calls directly to the most recent catch'. The functions catch and throw are used in the pattern matcher described in Section 10.1.

9.5.3 Recursive functions

The description of function application given on page 154 was incomplete. If function f has been defined by

```
(\text{defun } f (x_1 \ldots x_n) S)
```

then when an S-expression ($f S_1 \ldots S_n$) is evaluated, S is evaluated in an environment in which:

- (i) the values of S_1, \ldots, S_n are bound to x_1, \ldots, x_n respectively, and
- (ii) the function name f is bound to (lambda $(x_1 \ldots x_n) S$).

Thus if f is applied within S, then this application will be interpreted as a recursive call because at the point of call f will be bound to (lambda $(x_1 \ldots x_n) S$).

Example: The following recursive definition defines append to be the function that concatenates two lists:

```
(defun append (x y)
  (if (atom x) y (cons (car x) (append (cdr x) y))))
```

Evaluating

```
(append '(X_1 \ldots X_n) '(Y_1 \ldots Y_n)) results in (X_1 \ldots X_n \ Y_1 \ldots Y_n)
```

9.6 The LISP top-level

LISP is interactive. The user inputs S-expressions and the LISP system evaluates them and prints out the result. Franz LISP prompts for input using ->. Here is a little session on the author's Sun workstation. The machine is called gwyndir and the Unix prompt is gwyndir%. The session starts with Franz LISP being run, then the values of the variables x and y are set to be 1 and 2, then the S-expression (add x y) is evaluated and finally the LISP session is terminated.

```
gwyndir% lisp
Franz Lisp, Opus 38.69
-> (setq x 1)
1
-> (setq y 2)
2
-> (add x y)
3
-> (exit)
gwyndir%
```

Functions are defined interactively too, for example:

```
-> (defun add3 (x y z) (add x (add y z)))
add3
-> (add3 1 2 3)
6
-> (add3 (add3 1 2 3) (add3 4 5 6) (add3 7 8 9))
45
->
```

During interactive sessions, LISP prints out values using the function print (see page 176). The printing of linear lists is as expected, but the printing of dotted pairs can look a bit funny. For example:

```
-> '(1 2 3)
(1 2 3)
-> '(1 . (2 . 3))
(1 2 . 3)
```

A sequence of S-expressions can be typed into a file called file and then read in as though they had been input interactively. This is done by evaluating:

```
(read 'file)
```

The function include is like read, but it doesn't evaluate its argument (see also Section 9.11.5).

9.7 Dynamic binding

In all common LISPs except Common LISP, free variables in functions are interpreted *dynamically*. This means that the value of the free variable is looked up in whatever environment happens to be in force when the variable is evaluated. Consider, for example, the following session with LISP:

```
-> (setq z 1)
1
-> (defun addz (x) (add x z))
addz
-> (setq z 2)
2
-> (addz 0)
2
-> ((lambda (z) (addz 0)) 3)
3
-> z
2
->
```

The first time (addz 0) is evaluated the value of z is 2 and thus the result of the evaluation is 2. Note, however, that when addz was defined the value of z was 1. The second time (addz 0) is evaluated it is inside (lambda (z) (addz 0)). By the time it is evaluated, z will have been locally bound to 3 by the enclosing ((lambda (z) ···) 3); the result is thus 3. Notice that the binding of z to 3 inside the lambda-expression is local. Thus when the evaluation is finished, z reverts to being bound to its previous value, namely 2.

Unfortunately, dynamic binding makes LISP lambda-expressions behave differently from λ -calculus λ -expressions. For example, the λ -expression

```
(\lambda x. ((\lambda f. (\lambda x. (f \mathbf{0})) \mathbf{1}) (\lambda y. x)) \mathbf{2}
```

reduces to 2, whereas the LISP S-expression

```
((lambda (x)
  ((lambda (f)
        ((lambda (x) (funcall f 0)) 1))
  (quote(lambda (y) x))))
2)
```

evaluates to 1.

Exercise 103

Verify the above assertion. □

The differences between LISP functions and conventional mathematical functions means that much of the theory developed for the λ -calculus does not apply to LISP. This is one of the reasons for the low opinion of LISP held by some functional programming enthusiasts (e.g. see the quote by Turner on page 148). According to McCarthy [52], dynamic binding was a bug in the first LISP implementation, but it soon became a feature.

There is a class of particulary horrible bugs, called context bugs, that result from LISP functions not behaving like mathematical functions. These can be very frustrating if one attempts to do any significant higher-order programming. Fortunately, Common LISP uses a binding scheme (called static binding or lexical scoping) that corresponds to the λ -calculus and thus, as people switch to Common LISP, context bugs may become a thing of the past. However, if one really wants to make serious use of higher-order functions, it would be much better to use a proper functional language like ML or Miranda. Even Common LISP is pretty awkward to use for higher-order programming.

Exercise 104

Reduce the following λ -expression to normal form:

```
\mathbf{Y} (\lambda f \ x. \ (y \to \mathbf{true} \mid (x \to \mathbf{false} \mid ((\lambda y. \ f \ y) \ \mathbf{true}))))
```

Compare your reduction to the evaluation of the S-expression

```
((lambda (y) (tricky nil)) nil)
```

where the function tricky is defined by:

Over the years there has been quite a bit of debate on the advantages and disadvantages of dynamic binding. The advantages are:

- (i) It is easy to implement as there is no need to remember old environments. Nothing special is needed to get recursion to work properly (for example, see the mutually recursive definitions of three functions making up the unparser on page 222).
- (ii) Functions can be invoked without all the functions they use being defined. This makes it easy to do a 'top-down' development of a program.
- (iii) Functions can be defined in any order; for example, in the programs in the next three chapters, it often happens that f_1 calls f_2 , but it is convenient for f_1 to be defined before f_2 in the source file.
- (iv) If a function is changed then this change will be instantly felt by all functions that call it. This is useful for debugging: functions can be modified to print out information when they are entered and exited and no recompilation of programs that call such traced functions is needed. Powerful debugging packages based on this idea are provided by most LISP systems.

The disadvantages are:

- (i) It makes the semantics more complex and precludes the use of standard theories like the λ-calculus. Formal theories of LISP do exist (e.g. see my own paper on the semantics of dynamic binding [22], or the specialized logics in Mason's book [50] and Manna and Waldinger's paper [48]); however, these theories are still rather experimental.
- (ii) It makes higher-order programming fraught with danger due to the possibility of subtle 'context bugs'.
- (iii) Existing programs can stop working if a function is changed. This makes large continuously evolving programs hard to maintain (modern LISP systems usually provide mechanisms to help alleviate this problem).

The LISP community seems to have at last decided that the disadvantages outweigh the advantages. This has always been the view of the funtional programming community.

9.8 List equality and identity

When an S-expression is read in, the LISP reader builds a structure in the computer's memory to represent it². The thing that is actually returned as a result of the read is the address of this structure. Atoms with the same name are always represented by the same structure. The first time an atom with a given name is encountered during a session, the system builds a structure to represent it. If the atom is typed in again, then the system will return the address of the structure that it has already built. When a dotted pair is read in, LISP always builds a new structure to represent it. Thus if two identical dotted pairs are input they will result in different addresses being returned. The primitive function eq works by testing the equality of addresses. Thus if the same atom is read in twice the resulting values will be eq, but if the same dotted pair is read in twice the resulting values will not be eq. Here is a session to illustrate this:

```
-> (setq x 'foo)
foo
-> (setq y 'foo)
foo
-> (eq x y)
t
-> (setq x '(1 2 3))
(1 2 3)
-> (setq y '(1 2 3))
(1 2 3)
-> (eq x y)
nil
->
```

Exercise 105

What are the values of the following two S-expressions?

```
(i) (eq 'mike 'mike)(ii) (eq '(mike) '(mike))
```

²The details in this section are somewhat simplified (e.g. we omit discussion of the representation of numbers and strings). The description given is, however, adequate for understanding the programs in this book.

If a variable x is bound to an S-expression S in the environment, then x will actually be associated with the machine address of the memory structure representing S. If (setq y x) is subsequently evaluated then y will also be associated with this address and thus x and y will be eq.

```
-> (setq x '(1 2 3))
(1 2 3)
-> (setq y x)
(1 2 3)
-> (eq x y)
t
->
```

The standard function equal tests the equality of S-expressions. If it was not already provided it could be defined recursively by the following definition (which is deliberately written in a rather obscure style to illustrate the sequential use of and and or).

9.9 Property lists

When the LISP reader inputs an atom, it builds a memory structure holding the characters making up the atom's name. It also builds a list called the property list of the atom. This list is initially nil. It is used to hold various S-expressions that are associated with the atom. In early LISP implementations, the values of variables and the definitions of functions were held on the property list of the corresponding name. In more recent implementations, like Franz LISP, property lists are no longer used for this purpose, but they are still very useful as a way of 'tagging' atoms with information. For example, the pattern matcher described in Section 10.1 looks at the property lists of atoms occuring in patterns to see if they represent constants or variables. In Chapter 7 of Wilensky there is an example of a library database system in which data about books (title, author, publisher etc.) are held on property lists.

The property list of an atom has the form

```
(property_1 \ value_1 \ property_2 \ value_2 \ \dots \ property_n \ value_n)
```

where $property_1, \ldots, property_n$ are S-expressions (normally atoms) naming 'properties' and $value_1, \ldots, value_n$ are the corresponding 'property values'. Arbitrary properties can be associated with arbitrary values using the following functions:

- 1. (putprop atom value property) modifies the property list of atom by associating value with the property property. If atom already has a property called property, then its value is modified; if not, (property value) is added to the front of the property list of atom. The value returned by (putprop atom value property) is the value of value; putprop evaluates all its arguments.
- 2. The function defprop is like putprop except that is does not evaluate any of its arguments.
- 3. (get atom property) returns the S-expression associated with the property property on the property list of atom. If there is no such property, nil is returned; get evaluates its arguments.
- 4. (plist atom) returns the property list of atom; put evaluates its argument.

These functions are illustrated below:

```
-> (plist 'mike)
nil
-> (putprop 'mike t 'nice)
t
-> (plist 'mike)
(nice t)
-> (defprop mike nil lazy)
nil
-> (plist 'mike)
(lazy nil nice t)
-> (get 'mike 'nice)
t
-> (get 'mike 'hungry)
nil
->
```

Warning: The functions putprop, defprop and get use eq not equal to compare properties. For example:

```
-> (defprop sid t (likes gas))
t
-> (plist 'sid)
((likes gas) t)
-> (setq prop '(likes gas))
(likes gas)
-> (putprop 'sid 'maybe prop)
maybe
-> (plist 'sid)
((likes gas) maybe (likes gas) t)
-> (putprop 'sid 'yes prop)
yes
-> (plist 'sid)
((likes gas) yes (likes gas) t)
->
```

The association between atoms and their property lists is global; there is no notion of local properties analogous to local variables.

Warning: Strings cannot have properties. For example:

```
-> (plist "mike")
Error: Only Atoms and disembodied property lists allowed for
        plist
<1>: (reset)

[Return to top level]
-> (putprop "mike" t 'nice)
Error: putprop: Bad first argument: mike
<1>:
```

The function reset resets LISP to top level from the error state (being in an error state is indicated by a prompt of the form $\langle n \rangle$:). See Wilensky for details of Franz LISP's error handling features.

9.10 Macros

Macros provide a way of preprocessing S-expressions before they are evaluated. A typical macro definition looks like:

```
(defmacro M (x_1 \ldots x_n) S)
```

If M is so defined then an S-expression $(M S_1 \ldots S_n)$ is evaluated as follows:

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(i) First S is evaluated in an environment in which S_1, \ldots, S_n are bound unevaluated to x_1, \ldots, x_n respectively. This is called the macro expansion phase.

(ii) Second, the value resulting from evaluating S during the expansion phase is evaluated. This is called the *evaluation phase*.

Thus, evaluating $(M S_1 \ldots S_n)$ results in two evaluations.

Example: The following macro definition makes hd an abbreviation for car

```
(defmacro hd (x) (list 'car x))
```

This works as follows:

- (i) If (hd S) is evaluated, then during the expansion phase (list 'car x) is evaluated in an environment in which x is bound to S. This results in (car S).
- (ii) The S-expression (car S) is then evaluated and this returns the car of the value of S.

An important fact about macros is that the expansion phase is done at compile time (see Section 9.11 below). Thus, with compiled programs, there is no run-time overhead in using a macro. If hd had been defined as a function, then every time an expression of the form (hd S) is evaluated there would be an extra function call; thus using hd instead of car would reduce efficiency.

Example: Suppose only set had been defined, but not setq; then setq could be defined as a macro by:

```
(defmacro setq (x y)
  (list 'set (list 'quote x) y))
```

During the expansion phase of evaluating (setq x S) the S-expression (list 'set (list 'quote x) y) is evaluated in an environment in which x is bound to x and y is bound to S. This results in (set (quote x) S). This is then evaluated to bind x to the value of S. \square

Notice that since setq does not evaluate its first argument there is no way it could have been defined as a function using defun.

9.10.1 The backquote macro

A very useful built-in macro is the backquote macro. A backquoted expression 'S is like a quoted expression 'S except that any S-expressions in S that is preceded by a comma is evaluated. Here is a LISP session illustrating the difference between ' and ':

```
-> (setq y '(1 2 3))
(1 2 3)
-> '(x y z)
(x y z)
-> '(x ,y z)
(x (1 2 3) z)
-> '(x ,(car y) z)
(x 1 z)
-> '(x (car ,y) z)
(x (car (1 2 3)) z)
->
```

Inside a backquote, an expression can also be preceded by ,0 (comma followed by an 'at' sign); the expression is then evaluated and the result 'spliced in', i.e. inserted without its outermost brackets. This is best illustrated with an example. Continuing the session above:

```
-> '(x ,@y z)
(x 1 2 3 z)
-> '(a b c ,(list 1 2 3) d e f)
(a b c (1 2 3) d e f)
-> '(a b c ,@(list 1 2 3) d e f)
(a b c 1 2 3 d e f)
->
```

Wilensky describes the semantics of backquote rather nicely as follows:

The idea of the backquote macro is the following. Usually, in LISP, we evaluate expressions, and prevent evaluation by preceding an expression by quote. However, an expression that is backquoted works just the opposite way: all its elements are not evaluated *unless* they are preceded by something that explicitly indicates evaluation.

Backquote is an example of a read-macro. These are single characters that have macros associated with them and are invoked whenever this character is encountered by the reader (with the following complete

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S-expression as argument). When '(x ,y z) is read in, it is converted to (cons 'a (cons y '(z))) and when '(x ,@y z) is read in, it is converted to (append 'a (append y '(z))). We will not describe how read-macros are defined in Franz LISP; Chapter 14 of Wilensky is devoted to this. Another read-macro is the ordinary quote character '. When 'S is read in it is converted to (quote S).

Example: The macros hd and setq could be defined more lucidly using backquote as follows.

```
(defmacro hd (x) '(car ,x))
(defmacro setq (x y) '(set (quote ,x) ,y))
```

Example: Suppose only def had been defined, but not defun; then defun could be defined as a macro by:

A useful collection of macros are first, second, third, ..., ninth, which get the first, second, third, ..., ninth elements of lists. These are just sequences of cars and cdrs and can be defined using the built-in LISP functions with names of the form $c\cdots d$, where ' \cdots ' is any sequence of a's or d's. The idea of these functions is best conveyed by a few examples: (cadr S) abbreviates (car(cdr S)), (cadar S) abbreviates (car(cdr(car S))) and (cddaar S) abbreviates (cdr(cdr(car(car S)))). In general, the sequence of a's and d's determines the corresponding sequence of car's and cdr's. Using these functions one can define the much more mnemonically named first, second etc. by:

```
(defmacro first (x) '(car ,x))
(defmacro second (x) '(cadr ,x))
(defmacro third (x) '(caddr ,x))
(defmacro fourth (x) '(cadddr ,x))
(defmacro fifth (x) '(caddddr ,x))
(defmacro sixth (x) '(cadddddr ,x))
(defmacro seventh(x) '(caddddddr ,x))
(defmacro eighth (x) '(cadddddddr ,x))
(defmacro ninth (x) '(caddddddddr ,x))
(defmacro tenth (x) '(cadddddddddr ,x))
```

Also cdr is given a more mnemonic name:

```
(defmacro rest (x) '(cdr ,x))
```

Exercise 106

Define a macro backquote such that (backquote S) is like (quote S) except that any expression inside S of the form (comma S_1) is replaced by the value of S_1 , and any expression inside S of the form (comma@ S_1) results in the value of S_1 being spliced into S.

Thus backquote is a macro analogous to the read-macro 'with comma and comma@ being analogous to, and, @ respectively. \Box

When debugging macros it is useful to be able to see the result of expansion. Evaluating (macroexpand S) returns the result of macro expanding any macros occurring in the value of S. For example, suppose hd, setq and defun are defined to be macros as above, then:

```
-> (macroexpand '(hd '(1 2 3)))
(car '(1 2 3))
-> (macroexpand '(setq x (hd '(1 2 3))))
(set 'x (car '(1 2 3)))
-> (macroexpand '(defun foo (x y z) (list x y z)))
(def foo (lambda (x y z) (list x y z)))
```

9.11 Compilation

The LISP compiler is a program that translates function definitions into machine code. Compiling a function usually makes it execute several times faster. In many LISP systems there is a system function compile which when applied to function names has the side effect of replacing their S-expression definitions by equivalent machine code. Unfortunately, the Franz LISP compiler does not work this way. Instead, the functions to be compiled must be put into a file and then the compiler, called liszt, must be invoked on this file from outside LISP. For example, suppose the file rev.1 contains the definition of reverse.

Then all the functions in the file rev.l (actually there is only one, reverse) can be compiled by executing list rev as shown below.

```
gwyndir% ls rev.*
rev.l
gwyndir% liszt rev
Compilation begins with Liszt 68000 version 8.36
source: rev.l, result: rev.o
reverse
%Note: rev.l: Compilation complete
%Note: rev.l: Time: Real: 0:1,CPU: 0:0.13,GC: 0:0.00 for 0 gcs
%Note: rev.l: Assembly begins
%Note: rev.l: Assembly completed successfully
gwyndir% ls rev.*
rev.l rev.o
gwyndir%
```

The compiler has produced a new file rev.o, called the *object file*, which contains the compiled code. To use the compiled reverse the file rev.o must be loaded into LISP.

```
gwyndir% lisp
Franz Lisp, Opus 38.69
-> (load 'rev)
[fasl rev.o]
t
-> (reverse '(1 2 3 4 5 6 7 8 9))
(9 8 7 6 5 4 3 2 1)
->
```

The message [fasl rev.o] from LISP indicates that the object file is being loaded (fasl is a function, 'fast load', for loading files produced by liszt). Both the Unix command liszt and the LISP function load do not require the file extension '.1'; they will look for a file called file.1 if they cannot find file. In addition, as illustrated above, load will look first for a file with extension '.o' and load that if it finds it; the source file file.1 will only be loaded if there is no object file file.o.

9.11.1 Special variables

To work properly, the compiler needs to be told the names of *special* variables. These are variables that occur free in a function, i.e. occur in a function but are not local to it. For example, z is free in the function addz defined by

```
(defun addz (x) (add x z))
```

and so z must be declared special. This is done by placing

```
(declare (special z))
```

before the definition of addz. The compiler liszt will often automatically declare a variable to be special, but there are cases when it doesn't and this can lead to very obscure bugs. It is good practice to explicitly declare all special variables.

9.11.2 Compiling macros

The Franz LISP compiler liszt will expand any macro definitions that are in force when it compiles a function. This means that the compiled code has no reference to the macros and so their definitions are not needed at run time. The object file produced by liszt thus does not contain the macro definitions. Sometimes it is useful to keep the macros around at run time. This would be the case if it was planned to input new S-expressions containing the macro after the file containing the macro definitions was loaded. The compiler can be instructed to include in the object file all macro definitions in the source file with the declaration

```
(declare (macros t))
```

9.11.3 Local functions

Functions f_1, f_2, \ldots, f_n , can be declared local to a file by executing

```
(declare (localf f_1 \ldots f_n))
```

Declaring functions to be local has two advantages:

- (i) The compiler liszt can produce much faster code for local functions.
- (ii) Local functions are only known to the other functions in the file in which they are declared local. This provides a way of using the same name in two files without them getting confused. This is useful when several people are writing programs; it goes some way to overcome disadvantage (iii) of dynamic binding (see page 164).

9.11.4 Transfer tables

Evaluating the S-expression (sstatus translink on) will result in subsequently loaded compiled code running much faster. It also makes certain debugging tools (like baktrace) useless. For an explanation of what this magic actually does see Wilensky, Section 20.6, together with the documentation on the particular version of Franz LISP that you use. This arcane feature is mentioned here as it is useful for speeding up the painfully slow λ -calculus reducer described in Chapter 12.

Warning: After executing (sstatus translink on) you may find that loading redefinitions of previously loaded functions has no effect. To prevent this you should execute (sstatus translink nil) before doing the redefinition. When you have got them working, do (sstatus translink on) again.

9.11.5 Including files

If (include file) occurs at top level in a file, then the compiler will treat the contents of file (or file.1) as though they were textually inserted in place of the S-expression (include file). This enables the compiler to generate an object file in which there are fast links between functions in file and functions in the file in which file is included. Furthermore, any local functions in file will be available in both files.

9.12 Some standard functions and macros

Many of the functions and macros listed in this section can be defined in terms of the primitive functions, but they are provided for convenience. LISP systems differ in the predefined functions that they provide. For the examples in this book only a minute subset of the functions available in Franz LISP are used. The functions listed below are in alphabetical order; they all evaluate all their arguments.

- 1. (add S_1 S_2) evaluates to the sum of the values of S_1 and S_2 . An error results if either of these values is not a number.
- 2. (add1 S) is equivalent to (add S 1).
- 3. (append S_1 S_2) appends the values of S_1 and S_2 (see the example in Section 9.5.3 on page 161).
- 4. (assoc S, $((S_{11}, S_{12}), ..., (S_{n1}, S_{n2}))$) returns (S_{i1}, S_{i2}) where S_{i1} is the first value in the sequence S_{11} , S_{21} etc. that is equal to the value of S; assq is like assoc but uses eq instead of equal.

- 5. (concat A_1 A_2) concatenates atoms and/or strings; A_1 and A_2 can evaluate to either atoms or strings, but the result is always an atom.
- 6. (equal S_1 S_2) evaluates to t if the values of S_1 and S_2 are equal, otherwise it returns nil. Note that (eq '(1 2) '(1 2)) evaluates to to nil, but (equal '(1 2) '(1 2)) evaluates to t. There is a definition of equal on page 166.
- 7. (list S_1 S_2 ... S_n) evaluates to a list of the values of S_1 , S_2 , ..., S_n ; it is equivalent to

```
(cons S_1 (cons S_2 ( ... (cons S_n nil) ... )))
```

- 8. (listp S) evaluates to t if the value of S is a list (i.e. nil or a dotted pair) and to nil otherwise.
- 9. (mapcar f '(S_1 ..., S_n)) applies f successively to the S-expressions S_1 , ..., S_n and returns a list of the resulting values. For example, (mapcar (function add1) '(1 2 3)) evaluates to (2 3 4).
- 10. (member S ' $(S_1 \ldots S_n)$) evaluates to t if the value of S is equal to one of S_1, \ldots, S_n ; it evaluates to nil otherwise.
- 11. (memq S ' $(S_1 \ldots S_n)$) evaluates to t if the value of S is eq to one of S_1, \ldots, S_n ; it evaluates to nil otherwise. The functions member and memq differ in that they test for membership using equal and eq, respectively.
- 12. (not S) evaluates to t if S evaluates to nil, otherwise it evaluates to nil.
- 13. (null S) is equivalent to (eq S nil).
- 14. (number S) evaluates to t if the value of S is a number and to nil otherwise (see the description of symbol p below).
- 15. (pp-form S) pretty-prints the value of S, i.e. prints S in a nicely indented format with line-breaks at sensible places. The value of S is returned.
- 16. (princ S) prints the value of S and then returns t. If the value of S contains atoms or strings then no quotes (") or vertical bars (|) are printed.
- 17. (print S) prints the value of S followed by a carriage return and then returns nil. Atoms and strings are printed with vertical bars (if necessary) and quotes surrounding them.

- 18. (prog1 $S_1 \ldots S_n$) evaluates S_1, \ldots, S_n in that order and then returns the value of S_1 as result.
- 19. (prog2 $S_1 \ldots S_n$) evaluates S_1, \ldots, S_n in that order and then returns the value of S_2 as result.
- 20. (progn $S_1 \ldots S_n$) evaluates S_1, \ldots, S_n in that order and then returns the value of S_n as result.
- 21. (reset) resets LISP back to the top-level. It is the thing to do after detecting an error.
- 22. (sublis '($(S_{11} . S_{12}) ... (S_{n1} . S_{n2})$) S) returns the result of substituting S_{i2} for S_{i1} (for all i such that $1 \le i \le n$) in the value of S
- 23. (subst S_1 S_2 S_3) substitutes the value of S_1 for all occurrences of the value of S_2 in the value of S_3 .
- 24. (symbolp S) evaluates to t if the value of S is a symbol and to nil otherwise (e.g. if the value is a number). What this means is described in Chapter 18 of Wilensky; for our puposes it is sufficient to think of symbols as names (i.e. sequences of letters and numbers starting with a letter), the only other data type we use is numbers (see the description of number above).
- 25. (terpri) outputs a carriage return.

A Simple Theorem Prover

A simple theorem prover is described. It is based on higher-order rewriting operators and is implemented in LISP.

In this chapter, a very simple theorem proving program is described. The aim is to provide an example system that can automatically prove many of the verification conditions arising from the examples in Chapter 3. The reader should be aware that there are two respects in which the techniques described here are inadequate:

- (i) The proof strategy used is very ad hoc and is not representative of the current state of research. An example of a powerful general-purpose theorem prover is the Boyer-Moore system [7].
- (ii) The programming style used is insecure and not suitable for building theorem provers for safety-critical applications. This is discussed further in Section 10.3.

Readers interested in learning more about automatic theorem proving should consult the texts Symbolic Logic and Mechanical Theorem Proving by Chang and Lee [10], Logical Foundations of Artificial Intelligence by Genesereth and Nilsson [19] and Automated Reasoning: Introduction and Applications by Wos et al. [74].

The theorem prover described below is based on rewriting: a statement is proved by repeatedly replacing (i.e. rewriting) subexpressions by equivalent subexpressions until truth, represented by the atom T, is obtained, or the prover gets stuck. The prover rewrites E_1 to E_2 if:

- 1. $(E_1 = E_2)$ is an instance of an equation provided to the system, or
- 2. E_1 represents $S_1 \Rightarrow S_2$ and E_2 represents $S_1 \Rightarrow (S_2[T/S_1])$, or
- 3. E_1 represents $(x = E) \Rightarrow S$ and E_2 represents $(x = E) \Rightarrow (S[E/x])$

where, as usual, $S[S_1/S_2]$ denotes the result of substituting S_1 for all occurrences of S_2 in S.

Exercise 107

Show that

- (i) $\vdash S_1 \Rightarrow S_2$ if and only if $\vdash S_1 \Rightarrow (S_2[T/S_1])$.
- (ii) $\vdash (x = E) \Rightarrow S$ if and only if $\vdash (x = E) \Rightarrow (S[E/x])$.

Part of the theorem prover is a fairly general rewriting engine. This will also be used to implement a (very inefficient) combinator reducer in Chapter 12. The rewriter takes a list of equations ($(E_{11}=E_{12})$... $(E_{n1}=E_{n2})$) and an expression E to be rewritten, and then repeatedly replaces instances of E_{1i} by the corresponding instances of E_{2i} until no more changes occur.

To avoid having to write a parser, a LISP-like syntax for terms and statements will be used. This is illustrated in the table below:

Mathematical notation	Representation in LISP
$(X \times Y) + Z$	((X * Y) + Z)
(-X) - Y	((-X) - Y)
$\sin(X) + \cos(Y)$	$((\sin X) + (\cos Y))$
$P(E_1, \ldots, E_n)$	$(P E_1 \ldots E_n)$
$S_1 \wedge S_2$	$(S_1 \ { m and} \ S_2)$
$S_1 \vee S_2$	$(S_1 \text{ or } S_2)$
$\neg S$	$(\verb"not" S")$
$S_1 \Rightarrow S_2$	$(S_1$ implies $S_2)$

To implement the rewriting engine two main programs are needed:

- A pattern matcher: this determines if the left-hand side of a supplied equation matches an expression.
- (ii) A scanner: this repeatedly goes through expressions matching lefthand sides of equations with subexpressions and replacing them with the corresponding instances of the right-hand side, if a match is found.

Here a straightforward recursive pattern matcher is used. The approach to scanning is based on ideas from a paper by Paulson [59].

The code for our theorem prover is presented in a sequence of boxes. To experiment with this code, you should create a file called prover.1 which contains the S-expressions in these boxes in the order in which they occur in this chapter. At the beginning of this file are some declarations of special variables (see Section 9.11.1), an instruction to the compiler to include macros in the object file (see Section 9.11.2) and the magic transfer table trick to speed up compiled code (see Section 9.11.4).

```
(declare (special rewrite-flag culprit eqns facts))
(declare (macros t))
(sstatus translink on)
```

The macros first, second, third etc. and rest are also defined. They are described on page 171.

```
(defmacro first (x) '(car ,x))
  (defmacro second (x) '(cadr ,x))
  (defmacro third (x) '(caddr ,x))
  (defmacro fourth (x) '(cadddr ,x))
  (defmacro fifth (x) '(caddddr ,x))
  (defmacro sixth (x) '(cadddddr ,x))

  (defmacro rest (x) '(cdr ,x))
```

10.1 A pattern matcher

A function match will be defined such that evaluating (match pat exp):

(i) returns an S-expression $((X_1 . E_1) ... (X_n . E_n))$ such that

(sublis '((
$$X_1 . E_1$$
) ... ($X_n . E_n$)) pat)

evaluates to exp where X_1, \ldots, X_n are variables (see below) occurring in pat; and

(ii) returns the atom fail if no such substitution exists.

A variable in a pattern is any atom that is not a number and that does not have a property with name constant and value t. Thus a constant is explicitly marked on its property list and any other atom (that isn't a number) is a variable. The non-number constants to be used are:

```
+, -, <, <=, >, >=, *, =, T, F, DIV, not, and, or, implies
```

To set their constant property to t simply evaluate:

To test whether something is a variable define the macro

```
(defmacro is-variable (x)
  '(not (or (null ,x) (numberp ,x) (get ,x 'constant))))
```

Now define a function matchfn that matches a pattern against an expression in the context of a substitution. Suppose sub is a substitution

```
((X_1 . E_1) ... (X_n . E_n))
```

then (matchfn pat exp sub) matches pat against exp, but treats any occurrence of a variable X_i (where $1 \le i \le n$) in pat as though it were an occurrence of the constant E_i . Here are some examples to illustrate what matchfn does:

```
-> (matchfn '(x 2 z) '(1 2 3) nil)
((z . 3) (x . 1))
-> (matchfn '(x 2 z) '(1 2 3) '((x . 1)))
((z . 3) (x . 1))
-> (matchfn
    '((x + y) * (z * y)) '((3 + 1) * (3 * 1)) '((z . 3)))
((y . 1) (x . 3) (z . 3))
-> (matchfn
    '((x + y) * (z * y)) '((x + 1) * (x * 1)) '((z . x)))
((y . 1) (x . x) (z . x))
```

If pat does not match exp then matchfn exits by doing (throw 'fail); this will be caught by the function match that calls matchfn. Here is the definition of matchfn:

Now match can simply be defined to call matchfn with an empty initial substitution; it must also catch any throws from matchfn:

```
(defun match (pat exp) (catch (matchfn pat exp nil)))
```

Here are some examples of match in use:

```
-> (match '(x + y) '(1 + 2))
((y . 2) (x . 1))
-> (match '(x + x) '(1 + 2))
fail
->
```

10.2 Some rewriting tools

Evaluating (rewrite1 ' $(E_1 = E_2)$ E), where the function rewrite1 is defined below, treats E_1 as a pattern and matches it against the value of E. If the match succeeds then the corresponding instance of E_2 is returned, otherwise E is returned unchanged. For example:

```
-> (rewrite1 '((x + 0) = x) '((1 * 2) + 0))
(1 * 2)
-> (rewrite1 '((x + 0) = x) '((1 * 2) * 0))
((1 * 2) * 0)
->
```

The obvious definition of rewrite1, but not the one we will use, is:

```
(defun rewrite1 (eqn exp)
  (let ((l (first eqn)) (r (third eqn)))
    (let ((sub (match l exp)))
      (if (eq sub 'fail) exp (sublis sub r)))))
```

This is not used because it is convenient to provide a facility for tracing those rewrites that are done. A global variable rewrite-flag is used to control tracing: if it has value t then tracing is activated, otherwise it isn't. The default is no tracing (i.e. rewrite-flag set to nil). To implement tracing the definition of rewrite1 above is modified so that if rewrite-flag is t, then it prints out the instance of any equation that it succeeds in rewriting with. Thus the definition of rewrite1 is:

Here is the new rewrite1 in action:

```
-> (setq rewrite-flag t)
t
-> (rewrite1 '((x + 0) = x) '((1 * 2) + 0))
(((1 * 2) + 0) = (1 * 2))
(1 * 2)
->
```

Next define a function rewrite that takes a list of equations rather than a single one and rewrites with each equation in turn:

```
(defun rewrite (eqns exp)
(if (null eqns)
exp
(rewrite (rest eqns) (rewrite1 (first eqns) exp))))
```

Note that rewrite rewrites using the equations in the order they are given. When designing rewrite lists one has to take this order into account.

```
-> (setq rewrite-flag nil)
nil
-> (rewrite '(((x + 0) = x) ((x * 1) = x)) '((X * 1) + 0))
X
-> (rewrite '(((x * 1) = x) ((x + 0) = x)) '((X * 1) + 0))
(X * 1)
->
```

Shortly it will be shown how to repeatedly apply rewrite to all subexpressions of an expression, but first the two special rewrites described at the begining of the chapter (2 and 3 on page 179) must be implemented. Both these rewrites transform expressions of the form (P implies Q). To simplify their definition, four macros are useful: the macro is-imp tests whether its argument has the form (P implies Q); the macro mk-imp builds such an implication given P and Q; the macro antecedent extracts P from such an implication and the macro consequent extracts Q.

The function imp-subst-simp defined below transforms any S-expression of the form

```
(P \text{ implies } Q)
```

to one of the form

```
(P \text{ implies } Q[T/P]).
```

The function ${\tt imp-and-simp}$ defined below transforms any S-expression of the form

```
((X = E) \text{ implies } Q)
```

to one of the form

```
((X = E) \text{ implies } Q[E/X])
```

First define a macro is-eqn that tests whether its argument has the form $(E_1 = E_2)$:

The function imp-and-simp is defined by:

The function imp-simp first applies imp-subst-simp and then applies imp-and-simp to the result:

```
(defun imp-simp (exp) (imp-and-simp (imp-subst-simp exp)))
```

10.2.1 Higher-order rewriting functions

The higher-order functions repeat, depth-conv, top-depth-conv and re-depth-conv described in this section are based on rewriting operators from Paulson's elegant paper [59]. These operators specify the order in which subexpressions are rewritten.

Suppose f is a function that when applied to an expression will either change it or leave it alone, then (repeat f E) repeatedly applies f until no more change occurs.

```
(defun repeat (f exp)
  (let ((exp1 (funcall f exp)))
   (if (equal exp exp1)
       exp
       (repeat f exp1))))
```

The function depth-conv repeatedly applies a function f to all subexpressions of an expression exp in a 'bottom-up' order:

- (i) if exp is an atom then f is repeatedly applied to exp;
- (ii) if exp has the form $(E_1 ext{ . } E_2)$ then
 - (a) E_1 is recursively rewritten to $\hat{E_1}$,
 - (b) E_2 is recursively rewritten to $\hat{E_2}$,
 - (c) f is repeatedly applied to $(\hat{E_1} \cdot \hat{E_2})$ and the result returned.

Now use depth-conv to define a function depth-imp-simp that applies imp-simp to all subexpressions of an expression:

```
(defun depth-imp-simp (exp)
  (depth-conv
  (function (lambda (x) (imp-simp x)))
  exp))
```

Here is an example illustrating depth-imp-simp in action:

```
-> (depth-imp-simp '((x = 1) implies ((y = x) implies (y = z)))) ((x = 1) implies ((y = 1) implies (1 = z))) ->
```

Next define a function called top-depth-conv that is like depth-conv except that it rewrites 'top-down' before rewriting 'bottom-up': if exp is a dotted pair $(E_1 ext{ } E_2)$, then (top-depth-conv $f ext{ } exp$) repeatedly applies f to E_1 and E_2 before descending inside them.

Using top-depth-conv define a function for repeatedly rewriting all subexpressions of an expression in top-down order using a supplied list of equations. The top-down order is important.

```
(defun top-depth-rewrite (eqns exp)
  (top-depth-conv
  (function (lambda (x) (rewrite eqns x)))
  exp))
```

Notice that the variable eqns is free in the lambda-expression in which it occurs and must thus be declared special (see Section 9.11.1). To see that the top-down order is important consider the following example:

If a rewriter depth-rewrite is defined that uses depth-conv instead of top-depth-conv, i.e.

```
(defun depth-rewrite (eqns exp)
  (depth-conv
   (function (lambda (x) (rewrite eqns x)))
  exp))
```

then a different answer results:

It is a difficult problem to decide which rewriting order to use: some problems might be solved with depth-rewrite but not solved with top-depth-rewrite. However, for simple 'brute force' rewriting, it seems that top-depth-rewrite is usually satisfactory. Interactive theorem provers like LCF [60] provide the user with control over the order of rewriting by letting him or her decide whether to use a top-down or bottom-up order, or maybe even some other order specially tailored to the problem at hand. This flexibility is the theme of Paulson's paper [59]; he provides a powerful kit for building both specialized and general purpose rewriting tools.

Yet another 'depth conversion' is re-depth-conv; this will be useful in Chapter 12 for rewriting combinatory expressions (see Section 12.3):

The corresponding rewriting function is:

```
(defun re-depth-rewrite (eqns exp)
  (re-depth-conv
   (function (lambda (x) (rewrite eqns x)))
   exp))
```

Exercise 108

What is the difference between the functions re-depth-conv, depth-conv and top-depth-conv? Think up a list of equations eqns and an expression exp such that (depth-rewrite eqns exp), (top-depth-rewrite eqns exp) and (re-depth-rewrite eqns exp) all give different answers. \Box

The simple theorem prover can now be assembled by defining a function prove that repeatedly applies depth-imp-simp followed by rewriting using top-depth-rewrite:

```
(defun prove (eqns exp)
  (repeat
    (function
        (lambda (x) (top-depth-rewrite eqns (depth-imp-simp x))))
    exp))
```

The equations used for rewriting will be structured into two lists:

- (i) A list called logic containing various properties of implies and and.
- (ii) A list called arithmetic containing various arithmetical facts.

The verifier in Chapter 11 uses prove with (append logic arithmetic) as its list of equations. Both logic and arithmetic have been somewhat 'tuned' (in their content and order) so that they solve many of the verification conditions for the examples discussed in Chapters 2 and 3. The need that the equations in the lists themselves be mechanically verified is discussed in Section 10.3. Here is the definition of logic:

```
(setq
  logic
  '(
        ((T implies X) = X)
        ((F implies X) = T)
        ((X implies T) = T)
        ((X implies X) = T)
        ((X implies X) = X)
        ((X and X) = X)
        ((X and T) = X)
        ((F and X) = F)
        ((X and F) = F)
        ((X and F) = T)
        (((X and Y) implies Z) = (X implies (Y implies Z)))
        ))
        ))
        ((X and Y) implies Z) = (X implies (Y implies Z)))
        ))
        ((X and Y) implies Z) = (X implies (Y implies Z)))
        ((X implies Z))
        ((X implies Z)))
        ((X implies Z))))
        ((X implies Z))))
        ((X implies Z))))
```

Here is the definition of arithmetic. The particular collection of arithmetical facts included in it is very ad hoc. They have been chosen just so that various examples go through. The order in which they are listed has also been carefully 'tuned'. An interesting exercise is to switch tracing on (by (setq rewrite-flag t)) and then experiment both with different collections of facts and also with the same facts but in a different order. This will give a good feel for how rewriting works.

```
(setq
 arithmetic
 '(
   ((X + 0) = X)
   ((0 + X) = X)
   ((X * 0) = 0)
   ((0 * X) = 0)
   ((X * 1) = X)
   ((1 * X) = X)
   ((not(X \le Y)) = (Y < X))
   ((not(X >= Y)) = (Y > X))
   ((not(X < Y)) = (X >= Y))
   (((-X) >= (-Y)) = (X <= Y))
   (((-X) >= Y) = (X <= (-Y)))
   ((-0) = 0)
   (((X < Y) \text{ implies } (X <= Y)) = T)
   ((X - X) = 0)
   (((X + Y) - Z) = (X + (Y - Z)))
   (((X - Y) * Z) = ((X * Z) - (Y * Z)))
   ((X * (Y + Z)) = ((X * Y) + (X * Z)))
   (((X + Y) * Z) = ((X * Z) + (Y * Z)))
   (((X \ge Y) \text{ implies } ((X < Y) \text{ implies } Z)) = T)
   (((X \le Y) \text{ implies } ((Y \le X) \text{ implies } Z)) = T)
   ((0 DIA X) = 0)
   (((X DIV Y) + Z) = ((X + (Y * Z)) DIV Y))
   (((X - Y) + Z) = (X + (Z - Y)))
   ((2 * X) = (X + X))
```

The list facts is then defined by:

```
(setq facts (append logic arithmetic))
```

An example of something that can be proved is:

```
-> (prove facts
'(((X = (((N - 1) * N) DIV 2)) and ((1 <= N) and (N <= M)))
implies
((X + N) = ((((N + 1) - 1) * (N + 1)) DIV 2))))
T
->
```

An example of something that cannot be proved with the theorem prover is:

```
-> (prove facts
'(((T and (X >= Y)) implies (X = (max X Y)))
and
((T and (not (X >= Y))) implies (Y = (max X Y)))))
(((X >= Y) implies (X = (max X Y)))
and
((Y > X) implies (Y = (max X Y))))
```

These could be proved by adding some arithmetical facts about max (see page 216). Here is a trace of the instances of equations used during a proof, obtained by setting rewrite-flag to t:

```
-> (setq rewrite-flag t)
t
-> (prove
    facts
    '(((R = X) and (Q = 0)) implies (X = (R + (Y * Q)))))

((((R = X) and (Q = 0)) implies (X = (R + (Y * Q)))))
=
((R = X) implies ((Q = 0) implies (X = (R + (Y * Q))))))

((Y * 0) = 0)

((X + 0) = X)

((X = X) = T)

(((Q = 0) implies T) = T)

T
```

Exercise 109

Consider the following definition of prove1:

```
(defun prove1 (eqns exp)
  (re-depth-conv
   (function (lambda (x) (rewrite eqns (imp-simp x))))
   exp))
```

- (i) Is prove1 equivalent to prove?
- (ii) If it is equivalent, is it more or less efficient? If it is not equivalent, devise an example to show the difference.

10.3 Validity of the theorem prover

How can one be sure that there are not bugs in our theorem prover which enable it to prove things that are not true? These bugs could be of two kinds:

- (i) bugs in the underlying algorithm,
- (ii) bugs in the equations in the list facts.

An example of (ii) would be:

```
(((X DIV Z) + (Y DIV Z)) = ((X + Y) DIV Z))
```

which was originally included in the list arithmetic until it was noticed that it was not true (take X, Y and Z to be 3, 5 and 2, respectively). So this was replaced with

```
(((X DIV Y) + Z) = ((X + (Y * Z)) DIV Y))
```

However, how can one be sure this is true? A well known verifier developed by a U.S. company has a large number of rewrite rules in its knowledge base and every so often someone spots that one of them is wrong, usually as a result of a simple typing error when the equations were typed in.

Various methods have been developed to decrease the frequency of errors in mechanical theorem provers. One approach, invented by Robin Milner, is to employ a programming language type discipline to ensure that only 'safe' operations can be used to deduce new theorems. It is not possible to go into this here; interested readers are referred to the original LCF book

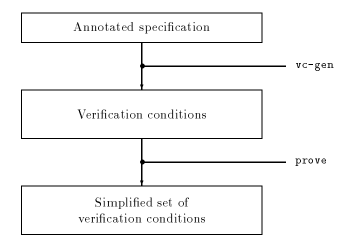
[25], my tutorial paper on the use of ML [24] or Paulson's recent monograph on Cambridge LCF [60]. Another approach is to have the theorem prover generate a representation of a formal proof rather than just magically announce 'T'. This proof can then be separately checked by a special proof-checking program. Such a program can be quite simple, and can be subject to especially rigorous analysis so that it has a very low probability of being wrong. Thus the correctness of an arbitrary program can be reduced to the correctness of a single proof checking program. This idea was originally proposed by Malcolm Newey in the early 1970s. More recently it has been discussed by Bob Boyer and J Moore and also advocated, in the context of hardware verification, by Keith Hanna.

As far as I know no one has implemented such a system. One can never attain complete certainty of the correctness of a theorem prover, but by methods like the ones just discussed, one can gain a very high degree of confidence.

A Simple Program Verifier

A simple program verifier implemented in LISP is described. This consists of a verification condition generator and the theorem prover described in Chapter 10.

The verifier described in this chapter is based on the principles explained in Chapter 3 and uses the theorem prover described in Chapter 10. It consists of two main LISP functions, vc-gen and prove, whose role is shown in the diagram below.



The input to the verifier is an annotated partial correctness specification. This is then checked to ensure that it is syntactically well formed and that annotations have been properly inserted according to the rules on page 44. The verification conditions are then generated and passed to the theorem prover. The final result is either an announcement 'all proved' indicating

that the specification has been proved or a list of simplified verification conditions that the prover could not prove.

The bulk of the LISP code is in the well-formedness checker. The verification condition generator is a simple one-pass recursive function.

To avoid having to write a parser, a LISP-like syntax will be used for partial correctness specifications. The table below illustrates this.

Standard notation	Representation in LISP
$\{P\}$ C $\{Q\}$	(SPEC P C Q)
V:=E	(ASSIGN V E)
$C_1;\{R_2\} \ldots ;\{R_n\}C_n$	$(\mathtt{SEQ}\ C_1\ R_2\ \dots\ R_n\ C_n)$
BEGIN $ \begin{array}{ccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$
BEGIN C_1 ; $\{R_2\}$; $\{R_n\}C_n$ END	$(\mathtt{BLOCK}\ C_1\ R_2\ \dots\ R_n\ C_n)$
IF S THEN C	(IF1 S C)
IF S THEN C_1 ELSE C_2	$(\text{IF2 } S \ C_1 \ C_2)$
WHILE S DO $\{R\}$ C	(WHILE S R C)
FOR $V\!:=\!E_1$ UNTIL E_2 DO $\{R\}$ C	(FOR V E_1 E_2 R C)

Terms and statements will be represented as illustrated in the table on page 180. Annotations are represented by (ASSERT S) where S is the representation of the annotating statement. For example:

```
{T}
       BEGIN
         R := X :
         Q:=0; {R=X \land Q=0}
         WHILE Y<R DO \{X = R+Y\times Q\}
            BEGIN R:=R-Y; Q:=Q+1 END
       END
     \{X = R + Y \times Q \land R < Y\}
would be represented by:
    (SPEC
      T
      (BLOCK
       (ASSIGN R X)
       (ASSIGN Q 0)
       (ASSERT ((R = X) and (Q = 0)))
       (WHILE
          (<=YR)
          (ASSERT (X = (R + (Y * Q))))
          (BLOCK (ASSIGN R (R - Y)) (ASSIGN Q (Q + 1)))
       ((R < Y) \text{ and } (X = (R + (Y * Q)))))
```

As with the theorem prover in Chapter 10, the code for our program verifier is presented in a sequence of boxes. To experiment with the verifier, you should create a file called verifier.1 containing the S-expressions in these boxes in the order in which they occur in this chapter. Since the verifier will use the theorem prover, the code for the prover is included with the LISP function include (see page 175). The effect of this is as though the file prover.1 was textually inserted at the beginning of the file verifier.1; this enables the compiler to compile the prover and verifier into a single object file:

```
(include prover)
```

The implementation begins with some functions and macros for manipulating LISP representations of terms, statements, commands and specifications.

11.1 Selectors, constructors and predicates

The verifier uses many macros to make the code more readable. These can be grouped into various kinds.

- (i) Selectors: these extract components of data structures. Their names correspond the type of component they are selecting. For example, the macroprecondition selects the the precondition part P of a partial correctnes specification (SPEC P C Q).
- (ii) Constructors: these construct a data-structure from its components. Their names have the form mk-thing, where thing is the kind of thing being constructed. For example, the macro mk-spec constructs a specification (SPEC P C Q) from the three components P, C and Q.
- (iii) Predicates: these are predicates that test whether an S-expression is a representation of a particular kind of object. They return t if the test succeeds and nil otherwise. Their names have the form is-thing, where thing is the kind of thing being tested. For example, the macro is-assign tests whether an S-expression represents an assignment command, i.e. whether it has the form (ASSIGN S₁ S₂).

11.1.1 Selector macros

The following three macros extract the components of a specification (SPEC P C Q):

```
(defmacro precondition (x) '(second ,x))
(defmacro command (x) '(third ,x))
(defmacro postcondition (x) '(fourth ,x))
```

The macro command-type gets the constructor from a command; it is used in case-switches in the definition of chk-and-cmd (page 203), assigned-vars (page 205) and vc-gen (page 206):

```
(defmacro command-type (c) '(first ,c))
```

Next, two macros to get the left-hand side (lhs) and right-hand side (rhs) of an assignment command:

```
(defmacro lhs (c) '(second ,c))
(defmacro rhs (c) '(third ,c))
```

The next three macros get the list of commands in a sequence, the components (i.e. local variable declarations and commands) of a block and the name declared in a local variable declaration:

```
(defmacro seq-commands (c) '(rest ,c))
(defmacro block-body (c) '(rest ,c))
(defmacro var-name (v) '(second ,v))
```

The following three macros get the components of conditionals:

```
(defmacro if-test (c) '(second ,c))
(defmacro then-part (c) '(third ,c))
(defmacro else-part (c) '(fourth ,c))
```

and the next three macros get the components of an annotated WHILE-commands:

```
(defmacro while-test (c) '(second ,c))
(defmacro while-annotation (c) '(third ,c))
(defmacro while-body (c) '(fourth ,c))
```

There are five macros to get the components of an annotated FOR-command:

```
(defmacro for-var (c) '(second ,c))
(defmacro lower (c) '(third ,c))
(defmacro upper (c) '(fourth ,c))
(defmacro for-annotation (c) '(fifth ,c))
(defmacro for-body (c) '(sixth ,c))
```

Finally, a macro to get the statement contained in an annotation (ASSERT S):

```
(defmacro statement (a) '(second ,a))
```

11.1.2 Constructor macros

Now define macros for constructing partial correctness specifications, negated statements, implications and conjunctions (\land s):

```
(defmacro mk-spec (p c q) '(list (quote SPEC) ,p ,c ,q))
(defmacro mk-not (s) '(list (quote not) ,s))
(defmacro mk-imp (s1 s2) '(list ,s1 (quote implies) ,s2))
(defmacro mk-and (s1 s2) '(list ,s1 (quote and) ,s2))
```

Three macros are defined to construct arithmetic expressions of the form m+n, m < n and m < n:

```
(defmacro mk-add (m n) '(list ,m (quote +) ,n))
(defmacro mk-less (m n) '(list ,m (quote <) ,n))
(defmacro mk-less-eq (m n) '(list ,m (quote <=) ,n))
```

11.1.3 Test macros

Only two predicates are used; usually it is better to abort if a test fails, rather than return nil. In such cases an error check (see Section 11.2 below) is more appropriate.

The macro is-var is a predicate that tests whether something represents a local variable declaration, i.e. has the form (VAR x), where x is a symbol:

The function is-assign is a predicate to test whether a command is an assignment, i.e. has the form (ASSIGN S_1 S_2):

11.2 Error checking functions and macros

To check that various things are well formed and that side conditions to the block and FOR rules are met, a number of functions and macros are defined that are like predicates, except that if the check fails, instead of just returning nil they print out an error message and reset LISP back to top-level. The names of such checks have the form chk-thing, where thing is the kind of thing being checked. Sometimes a check has a second argument which is the message to be printed if the test fails; for example, evaluating (chk-assert S message) returns t if S has the form (ASSERT S₁), but if it

does not then message is printed out and LISP reset. Sometimes a check generates its own error message; for example (chk-ann-cmd S) checks whether the value of S is a well-formed command (with annotations correctly embedded within it); if not an appropriate error message (e.g. Bad lhs of ASSIGN) is generated.

Checks are implemented with the function error. This prints out an error message, puts the offending thing in the global variable culprit (so that the user can look at what caused the error, if the message isn't sufficient) and then resets LISP to top level:

```
(defun error (message thing)
(progn (princ message)
(terpri)
(setq culprit thing)
(reset)))
```

11.2.1 Checking wellformedness

The function chk-typ checks whether the first element of something has a given name; t is returned if it does, an error is generated if it doesn't:

```
(defun chk-typ (name thing msg)
  (if (or (null thing) (not (eq (first thing) name)))
        (error msg thing)
      t))
```

Evaluating (chk-parts (constructor $x_1 \ldots x_m$) n) checks that m=n and if not raises the error "Parts of constructor missing" with culprit (constructor $x_1 \ldots x_n$):

The macro chk-sym checks whether something is a LISP symbol:

```
(\texttt{defmacro chk-sym (v msg) '(or (symbolp ,v) (error ,msg ,v)))}
```

The function chk-rev-ann-seq checks whether a sequence is correctly annotated (see page 44). It is given its argument in reverse order by chk-ann-cmd; this makes the recursion easier. See also the discusion on page 208 preceding the definition of rev-seq-vc-gen.

The function chk-ann-block checks whether the sequence of commands in a block is properly annotated. It uses the function block-commands that gets the sequence of commands in a block; it also uses chk-rev-ann-seq:

```
(defun chk-ann-block (c)
        (chk-rev-ann-seq (reverse (block-commands c))))

(defun block-commands (c)
        (strip-locals (block-body c)))

(defun strip-locals (c-list)
        (if (is-var (first c-list))
              (strip-locals (rest c-list))
              c-list))
```

The next two macros are intended to check whether statements and expressions (terms) are well-formed. These checks have not been implemented in this version of the verifier, so the macros just expand to t.

```
(defun chk-stat (s msg) t)
(defun chk-exp (e msg) t)
```

The function chk-assert checks whether something is an annotation and raises an error otherwise:

```
(defun chk-assert (s msg)
(and (chk-typ 'ASSERT s msg)
(chk-parts s 1)))
```

Here is the definition of the function chk-ann-cmd that checks whether a command is properly annotated. This function is just a cases-switch on the command type:

```
(defun chk-ann-cmd (c)
(caseq (command-type c)
        (ASSIGN (and (chk-parts c 2)
                     (chk-sym (lhs c) "Bad lhs of ASSIGN")
                     (chk-exp (rhs c) "Bad rhs of ASSIGN")))
        (IF1
                (and (chk-parts c 2)
                     (chk-stat (if-test c) "Bad test in IF1")
                     (chk-ann-cmd (then-part c))))
        (IF2
                (and (chk-parts c 3)
                     (chk-stat (if-test c) "Bad test in IF2")
                     (chk-ann-cmd (then-part c))
                     (chk-ann-cmd (else-part c))))
        (WHILE (and (chk-parts c 3)
                     (chk-stat
                      (while-test c)
                      "Bad test in WHILE")
                     (chk-assert
                      (while-annotation c)
                      "Bad annotation in WHILE")
                     (chk-ann-cmd (while-body c))))
        (FOR
                (and (chk-parts c 5)
                     (chk-sym (for-var c) "Bad FOR variable")
                     (chk-exp (lower c) "Bad lower bound")
                     (chk-exp (upper c) "Bad upper bound")
                     (chk-assert
                      (for-annotation c)
                      "Bad annotation in FOR")
                     (chk-ann-cmd (for-body c))))
        (SEQ
                (chk-rev-ann-seq (reverse (seq-commands c))))
        (BLOCK (chk-ann-block c))
                  "Unknown command type"
                  (command-type c))))
```

The function chk-ann-spec checks whether something is a correctly annotated partial correctness specification and raises an error if it isn't:

```
(defun chk-ann-spec (a)
  (and (chk-typ 'SPEC a "Bad specification constructor")
        (chk-parts a 3)
        (chk-stat (precondition a) "Bad precondition")
        (chk-ann-cmd (command a))
        (chk-stat (postcondition a) "Bad postcondition")))
```

11.2.2 Checking side conditions

The functions chk-block-side-condition and chk-for-side-condition, which are defined at the end of this section, are used during verification condition generation to check that the syntactic side conditions for blocks and FOR-commands are met (see pages 48 and 50 respectively). First it is necessary to define some auxiliary functions.

Evaluating (get-locals ((VAR x_1) ... (VAR x_n) C_1 ... C_n)) returns $(x_1 \ldots x_n)$:

The function vars-in computes the variables in a term; it is fairly crude in that it just 'flattens' the list:

The function append-lists appends together the members of a list of lists:

The function assigned-vars computes a list of the variables assigned to in a command. It is used by the function chk-for-side-condition:

```
(defun assigned-vars (c)
(caseq (command-type c)
        (ASSIGN (list (lhs c)))
        (IF1 (assigned-vars(then-part c)))
        (IF2 (append (assigned-vars(then-part c))
                     (assigned-vars(else-part c))))
        (WHILE (assigned-vars(while-body c)))
        (FOR (assigned-vars(for-body c)))
        (SEQ
         (append-lists
          (mapcar (function assigned-vars)
                  (seq-commands c))))
        (BLOCK
         (append-lists
          (mapcar (function assigned-vars)
                  (block-commands c))))
        (t nil)))
```

The definition of the function disjoint below depends on the arguments of and and or being evaluated in left-to-right order:

Evaluating (chk-block-side-condition P C Q) checks that if C represents a block, i.e. has the form

```
(BLOCK (VAR V_1) ... (VAR V_m) C_1 ... C_n)
```

then none of V_1, \ldots, V_m occur in P or Q. This check is done during verification condition generation by the function vc-gen of Section 11.3 below.

Evaluating (chk-for-side-condition (FOR V E_1 E_2 R C)) checks that neither V, nor any variable occurring in E_1 or E_2 is assigned to inside C:

11.3 The verification condition generator

Evaluating (vc-gen P C Q) returns the verification conditions from an annotated specification (SPEC P C Q). It calls the functions assign-vc-gen, if1-vc-gen, if2-vc-gen, while-vc-gen, for-vc-gen, seq-vc-gen and block-vc-gen, which are defined below.

The function assign-vc-gen generates a list of verification conditions according to the rule in the box on page 45:

The function if1-vc-gen generates a list of verification conditions according to the rule in the box on page 45:

The function if2-vc-gen generates a list of verification conditions according to the rule in the box on page 46:

The function while-vc-gen generates a list of verification conditions according to the rule in the box on page 49:

The function for-vc-gen generates a list of verification conditions according to the rule in the box on page 50. The syntactic side condition is checked by chk-for-side-condition defined on page 206.

```
(defun for-vc-gen (p c q)
(let ((v (for-var c))
       (e1 (lower c))
       (e2 (upper c))
       (r (statement(for-annotation c)))
       (c1 (for-body c)))
  (and (chk-for-side-condition c)
       (append
        (list
         (mk-imp p (subst e1 v r))
         (mk-imp (subst (mk-add e2 1) v r) q)
         (mk-imp (mk-and p (mk-less e2 e1)) q))
        (vc-gen
         (mk-and
         r
          (mk-and
           (mk-less-eq e1 v)
           (mk-less-eq v e2)))
         (subst (mk-add v 1) v r))))))
```

The function seq-vc-gen generates a list of verification conditions according to the rule in the box on page 46. The process of generating these conditions is recursive, with the recursion being done at the end of the sequence. This is most simply implemented by having the recursion in a subsidiary function rev-seq-vc-gen which is passed the reversed list of commands by seq-vc-gen and can thus recurse on the cdr of its argument.

```
(defun seq-vc-gen (p c q)
(rev-seq-vc-gen p (reverse (seq-commands c)) q))
```

The function rev-seq-vc-gen is a bit messy; this reflects the messiness of the algorithm for generating verification conditions from sequences. From the definition of seq-vc-gen, the verification conditions are generated from (SPEC P (SEQ $C_1 \ldots C_n$) Q) by evaluating

```
(rev-seq-vc-gen P (C_n \ldots C_1) Q)
```

There are two cases:

1. If C_n is an assignment (ASSIGN V E), then

```
(rev-seq-vc-gen P (C_{n-1} ... C_1) Q[E/V])
```

is evaluated. This corresponds to case 2 on page 46.

2. If C_n is not an assignment, then, by the rules of proper annotation (see page 44 and the definition of the function chk-rev-ann-seq on page 202), the penultimate element in the list $(C_1 \ldots C_n)$ passed to seq-vc-gen must represent an annotation, R say. This will be the second element of the reversed list $(C_n \ldots C_1)$ passed to rev-seq-vc-gen. Thus the verification conditions corresponding to case 1 on page 46 are computed by evaluating

```
(rev-seq-vc-gen P (C_{n-1} ... C_1) R)
```

and then appending to the result of this the verification conditions generated from (SPEC R C_n Q).

The definition of rev-seq-vc-gen is thus:

The function block-vc-gen generates a list of verification conditions according to the rule in the box on page 48. It needs to check the syntactic side condition, using chk-block-side-condition defined on page 206; it then generates the verification from the sequence of commands in its body. This is done by directly applying rev-seq-vc-gen to the reversed sequence:

```
(defun block-vc-gen (p c q)
(and (chk-block-side-condition p c q)
(rev-seq-vc-gen p (reverse (block-commands c)) q)))
```

11.4 The complete verifier

Using the verification condition generator, vc-gen, and the theorem prover, prove, described in Chapter 10, the verification system can now be completed. The function verify takes an annotated specification and:

- (i) checks that it is annotatated correctly using chk-ann-spec;
- (ii) generates the verification conditions using vc-gen,
- (iii) prints them out;
- (iv) attempts to prove them with the theorem prover prove described in Chapter 10 (using the facts in the global variable facts);
- (v) prints out "all proved" if it succeeds in proving all the verification conditions, otherwise it prints out the things it cannot prove;
- (vi) returns t if all the verifications are proved and nil otherwise.

Before defining verify a couple of auxiliary functions are needed. The first of these, print-list, prints out the elements of a list preceded by a blank line:

```
(defun print-list (list)
  (mapcar
   (function(lambda (x) (terpri) (pp-form x)))
   list))
```

The verifier prints (using print-list) the verification conditions that it cannot prove. This list is constructed by the function unproved-vcs, which removes occurrences of T (representing proved verification conditions) from the list returned by the theorem prover prove.

Here, at last, is the definition of the verifier verify:

```
(defun verify (a)
(prog (vcs vcs1)
      (terpri)
       (princ "Checking syntax of annotated program ... ")
       (chk-ann-spec a)
       (setq vcs (vc-gen (precondition a)
                         (command a)
                         (postcondition a)))
       (princ "OK.")
       (terpri)
       (princ "The verification conditions are:")
       (print-list vcs)
       (terpri) (terpri)
       (princ "Trying to prove verification conditions ...")
       (setq vcs1 (unproved-vcs (prove facts vcs)))
       (cond ((null vcs1)
              (princ " all proved."))
             (t (princ " can't prove:")
                (print-list vcs1)))
       (terpri) (terpri)
       (return (eq vcs1 nil))))
```

11.5 Examples using the verifier

Here are some examples showing the verifier in action. First, an example where the verifier (correctly) fails:

-> (verify '(SPEC

Here is the debugged program for swapping the values of two variables (see page 47):

```
((X = x) \text{ and } (Y = y))
                (SEQ (ASSIGN R X) (ASSIGN X Y) (ASSIGN Y R))
                ((X = y) \text{ and } (Y = x)))
Checking syntax of annotated program ... OK.
The verification conditions are:
 (((X = x) \text{ and } (Y = y)) \text{ implies } ((Y = y) \text{ and } (X = x)))
Trying to prove verification conditions ... all proved.
Here is the division program described on page 42:
-> (verify '(SPEC
                (BLOCK
                 (ASSIGN R X)
                 (ASSIGN Q 0)
                 (ASSERT ((R = X) and (Q = 0)))
                 (WHILE (Y \le R)
                   (ASSERT (X = (R + (Y * Q))))
                   (BLOCK (ASSIGN R (R - Y))
                            (ASSIGN Q (Q + 1))))
                ((R < Y) \text{ and } (X = (R + (Y * Q))))))
Checking syntax of annotated program ... OK.
The verification conditions are:
 (T implies ((X = X) \text{ and } (0 = 0)))
 (((R = X) \text{ and } (Q = 0)) \text{ implies } (X = (R + (Y * Q))))
 (((X = (R + (Y * Q))) \text{ and } (\text{not } (Y \le R)))
implies
 ((R < Y) \text{ and } (X = (R + (Y * Q)))))
 (((X = (R + (Y * Q))) \text{ and } (Y \le R))
```

```
implies
 (X = ((R - Y) + (Y * (Q + 1)))))
Trying to prove verification conditions ... all proved.
t
The next example is from Exercise 15 on page 31:
-> (verify
     '(SPEC
       T
       (SEQ (ASSIGN X 0)
             (ASSERT (X = 0))
             (FOR N 1 M (ASSERT (X = (((N - 1) * N) DIV 2)))
              (ASSIGN X (X + N)))
       (X = ((M * (M + 1)) DIV 2))))
Checking syntax of annotated program ... OK.
The verification conditions are:
 (T \text{ implies } (0 = 0))
 ((X = 0) \text{ implies } (X = (((1 - 1) * 1) DIV 2)))
 ((X = ((((M + 1) - 1) * (M + 1)) DIV 2))
 implies
 (X = ((M * (M + 1)) DIV 2)))
 (((X = 0) \text{ and } (M < 1)) \text{ implies } (X = ((M * (M + 1)) \text{ DIV } 2)))
 (((X = (((N - 1) * N) DIV 2)) and ((1 <= N) and (N <= M)))
 implies
 ((X + N) = ((((N + 1) - 1) * (N + 1)) DIV 2)))
Trying to prove verification conditions ... can't prove:
((X = 0) \text{ implies } ((M < 1) \text{ implies } (0 = (((M * M) + M) \text{ DIV } 2)))))
nil
```

The unproved verification condition is not true unless M >= 1. This suggests changing the precondition from T to M >= 1:

```
-> (verify
    '(SPEC
       (M >= 1)
       (SEQ (ASSIGN X 0)
            (ASSERT (X = 0))
            (FOR N 1 M (ASSERT (X = (((N - 1) * N) DIV 2)))
             (ASSIGN X (X + N)))
       (X = ((M * (M + 1)) DIV 2))))
Checking syntax of annotated program ... OK.
The verification conditions are:
((M >= 1) \text{ implies } (0 = 0))
((X = 0) \text{ implies } (X = (((1 - 1) * 1) DIV 2)))
((X = ((((M + 1) - 1) * (M + 1)) DIV 2))
 implies
 (X = ((M * (M + 1)) DIV 2)))
(((X = 0) \text{ and } (M < 1)) \text{ implies } (X = ((M * (M + 1)) \text{ DIV } 2)))
(((X = (((N - 1) * N) DIV 2)) and ((1 <= N) and (N <= M)))
implies
((X + N) = ((((N + 1) - 1) * (N + 1)) DIV 2)))
Trying to prove verification conditions ... can't prove:
((X = 0) \text{ implies } ((M < 1) \text{ implies } (0 = (((M * M) + M) DIV 2))))
nil
```

This did not quite work. It is necessary to add the annotation that M >= 1 just before the FOR-command. A smarter verifier could have done this automatically by propagating annotations forwards if variables in them are not assigned to.

Exercise 110

Implement an improved verification condition generator that will propagate annotations in such a way that the example above is proved. \Box

The following program demonstrates that, with the extra annotation, the example goes through:

```
-> (verify
        '(SPEC (M >= 1)
           (SEQ (ASSIGN X 0)
                 (ASSERT ((X = 0) and (M >= 1)))
                 (FOR N 1 M (ASSERT (X = (((N - 1) * N) DIV 2)))
                 (ASSIGN X (X + N)))
           (X = ((M * (M + 1)) DIV 2))))
    Checking syntax of annotated program ... OK.
    The verification conditions are:
    ((M >= 1) \text{ implies } ((0 = 0) \text{ and } (M >= 1)))
    (((X = 0) \text{ and } (M >= 1)) \text{ implies } (X = (((1 - 1) * 1) DIV 2)))
    ((X = ((((M + 1) - 1) * (M + 1)) DIV 2))
     implies
     (X = ((M * (M + 1)) DIV 2)))
    ((((X = 0) \text{ and } (M >= 1)) \text{ and } (M < 1))
     implies
     (X = ((M * (M + 1)) DIV 2)))
    (((X = (((N - 1) * N) DIV 2)) and ((1 <= N) and (N <= M)))
    implies
    ((X + N) = ((((N + 1) - 1) * (N + 1)) DIV 2)))
    Trying to prove verification conditions ... all proved.
    t
Here is another example that fails:
    -> (verify
        '(SPEC
           (IF2 (X \ge Y) (ASSIGN MAX X) (ASSIGN MAX Y))
           (MAX = (max X Y)))
    Checking syntax of annotated program ... OK.
    The verification conditions are:
    ((T \text{ and } (X \ge Y)) \text{ implies } (X = (max X Y)))
```

```
((T and (not (X >= Y))) implies (Y = (max X Y)))
Trying to prove verification conditions ... can't prove:
((X >= Y) implies (X = (max X Y)))
((Y > X) implies (Y = (max X Y)))
nil
```

Some facts about ${\tt max}$ must be added to the knowledge base of the theorem prover:

```
-> (let ((facts (append
facts
'(((X >= Y) implies (X = (max X Y)))
((Y > X) implies (Y = (max X Y)))))))

(verify
'(SPEC
T
(IF2 (X >= Y) (ASSIGN MAX X) (ASSIGN MAX Y))
(MAX = (max X Y)))))

Checking syntax of annotated program ... OK.
The verification conditions are:

((T and (X >= Y)) implies (X = (max X Y)))

((T and (not (X >= Y))) implies (Y = (max X Y)))

Trying to prove verification conditions ... all proved.
```

A λ -calculus Toolkit

Programs for experimenting with the λ -calculus and combinators are described. These include a normal-order reduction engine, a program for compiling λ -expressions to combinators and a program for doing combinator reduction.

The programs described in this chapter provide tools, implemented in LISP, for:

- (i) translating λ -expressions into and out of an internal representation, (Section 12.1);
- (ii) reducing λ -expressions by β -conversion (Section 12.2);
- (iii) translating λ -expressions to combinators (Section 12.3);
- (iv) reducing combinatory expressions by combinator reduction (Section 12.4).

These tools provide a workbench for playing and experimenting with the theories described in Part II. In particular, they provide tools for exploring the inefficiencies of β -reduction and combinator reduction. Recent research has led to compiling techniques that yield implementations that are orders of magnitude faster than direct implementations of these reduction methods. These techniques are still very experimental; for an introduction to some of them see Peyton Jones's book [61].

12.1 Parsing and printing λ -expressions

To enable λ -expressions to be input and output in a readable form, a simple parser and pretty printer are provided for a 'user-friendly' external syntax. However, to make things simple this external syntax will still be LISP-like; it is designed to minimize brackets according to the conventions described in

Section 4.2. All the parser and printer really do is insert or remove brackets according to these conventions. The table below shows the external syntax and internal representation for λ -expressions. In this table, \hat{E} denotes the internal representations of E.

λ -expression	External syntax	Internal representation
x	x	x
$\lambda x. E$	(L x E)	(L x \hat{E})
$\lambda x_1 \ x_2 \ \dots \ x_n . E$		(L x_1 (L x_2 (L x_n \hat{E})))
E_1 E_2	$(E_1 \ E_2)$	$(\hat{E}_1 \ \hat{E}_2)$
$E_1 \ E_2 \ \dots \ E_n$	$(E_1 \ E_2 \ \dots \ E_n)$	$((\ldots (\hat{E_1} \ \hat{E_2}) \ldots) \ \hat{E_n})$

As in the previous two chapters, programs are presented in a sequence of boxes. To experiment with these, you should create a file called lambda.l containing the S-expressions in these boxes in the order in which they occur in this chapter. Since the programs for translating to combinators and reducing them will use the rewriting engine, the code for prover is included with the LISP function include (see page 175). The effect of this is as though the file prover.l was textually inserted at the beginning of the file lambda.l:

(include prover)

In the next section some macros are defined for constructing S-expressions that represent λ -expressions. These macros consist of selectors, constructors and tests, the general idea of which is described in Section 11.1.

12.1.1 Selectors, constructors and predicates

The following selector macros extract components of the representations of λ -expressions. The macros for names are redundant, but are defined for consistency and to make it easy to change representations later.

• name gets the name of a variable.

```
(name 'x) = x
```

• by and body get the bound variable and body of abstractions.

$$(bv '(L x E)) = x$$

 $(body '(L x E)) = E$

• rator and rand get the operator and operand of an application.

$$(rator '(E_1 E_2)) = E_1$$

 $(rand '(E_1 E_2)) = E_2$

These macros are defined as follows:

```
(defmacro name (e) e)

(defmacro bv (e) '(second ,e))
(defmacro body (e) '(third ,e))

(defmacro rator (e) '(first ,e))
(defmacro rand (e) '(second ,e))
```

The following constructor macros build internal representations of λ -expressions:

• mk-var builds the internal representation of a variable.

$$(mk-var, x) = x$$

• mk-abs builds the internal representation of a λ -abstraction.

$$(mk-abs 'x 'E) = (L x E)$$

• mk-comb builds the internal representation of a combination.

$$(mk-comb 'E_1 E_2) = (E_1 E_2)$$

Here are the definitions in LISP:

```
      (defmacro mk-var (x) x)

      (defmacro mk-abs (x e) '(list 'L ,x ,e))

      (defmacro mk-comb (e1 e2) '(list ,e1 ,e2))
```

The macros defined below are predicates that test whether S-expressions represent variables, abstractions or combinations:

```
(defmacro is-var (e)
    '(and (symbolp ,e) (not (get ,e (quote DEFN)))))

(defmacro is-abs (e)
    '(and ,e (listp ,e) (eq (first ,e) 'L)))

(defmacro is-comb (e)
    '(and ,e (listp ,e) (not (eq (first ,e) 'L)) (rest ,e)))
```

12.1.2 Definitions

A λ -expression E can be given a name name by evaluating

```
(LET name E)
```

The expression E is parsed (see Section 12.1.3) and then stored on the property list of name under the property DEFN. Names defined with LET can occur both in the external syntax and the internal representation of λ -expressions. The LISP definition of LET is given in Section 12.1.5.

The macro defn gets the expression associated with a name and the macro is-defn is a predicate that tests whether an atom is the name of an expression. When reading the definition of is-defn remember that any non-null S-expression counts as 'true' for and and that get returns nil if there is no appropriate property:

```
(defmacro defn (e) '(get ,e (quote DEFN)))

(defmacro is-defn (e)
  '(and (symbolp ,e) (get ,e (quote DEFN))))
```

To simplify the writing of functions that process expressions, we define selectors and tests that expand out names (if necessary). The names of these macros all end with *.

If the value of S is a name, then (defn* S) evaluates to the expression associated with the name; if S is not a name then (defn* S) evaluates to the value of S. It is possible that one name could be defined to be another, for example

```
(LET K (L x y x))
(LET TRUE K)
```

We want (defn* TRUE) to evaluate to $(L \times y \times)$ and thus defn* may need to be called recursively. This recursion is done by the auxiliary function defn-fun below:

```
(defun defn-fun (e)
  (if (symbolp e) (defn-fun (defn e)) e))

(defmacro defn* (e)
  '(if (is-defn ,e) (defn-fun (defn ,e)) ,e))
```

The selectors and predicates defined below expand definitions using defn*:

```
(defmacro name* (e) '(name (defn* ,e)))

(defmacro bv* (e) '(bv (defn* ,e)))

(defmacro body* (e) '(body (defn* ,e)))

(defmacro rator* (e) '(rator (defn* ,e)))

(defmacro rand* (e) '(rand (defn* ,e)))

(defmacro is-var* (e) '(is-var (defn* ,e)))

(defmacro is-abs* (e) '(is-abs (defn* ,e)))
(defmacro is-comb* (e) '(is-comb (defn* ,e)))
```

12.1.3 A parser

The parser is now defined; this translates from external syntax to internal representation. The function parse is a straightforward recursive function; it calls an auxiliary function parse-seq to parse sequences:

```
(parse-seq E (E_1 E_2 ... E_n)) = (( ... ((E \hat{E}_1) \hat{E}_2) ... ) \hat{E}_n)
```

where, as before, \hat{E}_i is the result of parsing E_i .

The main parsing function parse can then be defined by:

It is useful to define a macro parseq that is like parse, but quotes its argument:

```
(defmacro parseq (x) '(parse (quote ,x)))
```

Here is an example session with LISP illustrating the parser:

```
-> (parseq (L x y x))
(L x (L y x))
-> (parseq (L f ((L x (f x x))(L x (f x x)))))
(L f ((L x ((f x) x)) (L x ((f x) x))))
->
```

It is convenient to have an unparser to convert back to the external syntax.

12.1.4 An unparser

The function unparse converts the internal representation of a λ -expression to the external syntax. It is defined by mutual recursion with two other functions, unparse-abs and unparse-comb for unparsing abstractions and combinations. This sort of mutual recursion can be very convenient and illustrates one of the benefits of dynamic binding (see Section 9.7).

```
(unparse-abs '(L x_1 (L x_2 ... (L x_n E) ... )))

= (L x_1 x_2 ... x_n E)

(unparse-comb '(( ... ((E E_1) E_2) ... ) E_n))

= (E E_1 E_2 ... E_n)
```

The definition of unparse-abs uses an auxiliary function strip-abs that strips off the variables of an abstraction:

```
(strip-abs '(abs (var x_1) (abs (var x_2) ... (abs (var x_n) E) ... ))) = ((x_1 \ x_2 \ \dots \ x_n) \ . E)
```

The definition of unparse-comb is just a simple recursion:

The following session illustrates unparse:

```
-> (setq Y (parseq (L f ((L x (f x x)) (L x (f x x))))))
(L f ((L x ((f x) x)) (L x ((f x) x))))
-> (unparse Y)
(L f ((L x (f x x)) (L x (f x x))))
->
```

Exercise 111

Could the parser and unparser be implemented by rewriting? If so devise a suitable set of equations. If not, why not? \Box

12.1.5 LET and LETREC

The following macro LET is used for making definitions. It first parses the expression that is being given a name and then puts the result on the property list of the name under the property DEFN. It also binds the name to the expression using setq (this is just for the user's convenience).

The table below shows how some of the definitions at the beginning of Chapter 5 are represented using our S-expression syntax.

```
LET \mathbf{true} = \lambda x \cdot \lambda y \cdot x (LET TRUE (L x y x))

LET \mathbf{false} = \lambda x \cdot \lambda y \cdot y (LET FALSE (L x y y))

LET \mathbf{not} = \lambda t \cdot t \mathbf{false} \mathbf{true} (LET NOT (L t (t FALSE TRUE)))

LET (E \to E_1 \mid E_2) = (E \mid E_1 \mid E_2) (LET IF (L t x y (t x y)))

LET \mathbf{fst} = \lambda p \cdot p \mathbf{true} (LET FST (L p (p TRUE)))

LET \mathbf{snd} = \lambda p \cdot p \mathbf{false} (LET SND (L p (p FALSE)))

LET (E_1, E_2) = \lambda f \cdot f \mid E_1 \mid E_2 (LET PAIR (L x y f (f x y)))
```

The definition of the fixed-point operator Y is:

```
(LET\ Y\ (L\ f\ ((L\ x\ (f\ (x\ x)))\ (L\ x\ (f\ (x\ x))))))
```

The macro LETREC provides a convenient way of defining functions recursively with Y. Evaluating

```
(LETREC f (x_1 \ldots x_n) E) is equivalent to (\text{LET } f \ (Y \ (\text{L } f' \ x_1 \ldots x_n \ E[f'/f])))
```

Note that this definition is not completely robust: it only works if f' does not occur in E. One could use f instead of f' since, logically, there should be no confusion between names of abbreviations and bound variables. Unfortunately such a confusion is possible with our simple implementation (see Exercise 112 below). To implement LETREC we need to define a macro that adds a prime to an atom:

```
(defmacro prime (x) '(concat ,x '|'))
```

The macro prime is illustrated by:

```
-> (prime 'a)
|a'|
-> (prime (prime 'a))
|a''|
```

Here is the definition of LETREC:

Suppose ZERO, IS-ZERO, ADD and PRE have been defined (see page 232), then the multiplication function MULT can be defined by:

Exercise 112

Devise an example to show that the following definition of the macro LETREC does not work.

Modify the treatment of variables and/or definitions so that this will work.

12.2 A λ -calculus reducer

In this section some programs for doing normal order λ -reduction are described. The tricky part of this is making sure that the substitutions done during β -reduction are valid. First a function frees is defined to compute the free variables in a λ -expression. This uses the function union for computing the union of two lists; (union ' $(x_1 \ldots x_m)$ ' ' $(y_1 \ldots y_n)$) evaluates to the result of adding to the front of $(y_1 \ldots y_n)$ those members of $(x_1 \ldots x_m)$ that are not in it. For example:

```
(union '(1 2 3) '(2 4 6)) = (1 3 2 4 6)
```

The definition of frees is now straightforward. First define an auxiliary function frees-fun such that (frees-fun E vars) computes the free variables in E that are not in vars. The *-ed macros are used to ensure that free variables in definitions are noticed.

12.2.1 Substitution

The substitution algorithm of this section automatically renames variables to avoid capture. To implement this a method of generating new names is needed. Evaluating

```
(variant 'x '(x_1 \ldots x_n))
```

returns x'^{\cdots} which denotes x with sufficient primes added so that the resulting variable is different from x_i for $1 \le i \le n$.

```
(defun variant (v vlist)
  (cond ((and (not(member v vlist)) (not(is-defn v))) v)
          (t (let ((v1 (mk-var (prime (name* v)))))
                (variant v1 vlist)))))
```

This is illustrated by:

```
-> (variant 'a '(a |a'| |a''|)) |a'''|
->
```

The table below is based on the one on page 72, but with different variable names and the rows reordered. With these changes it corresponds to the definition of the function substitute below.

E	$E[E_1/x]$
x	E_1
$y \qquad \text{(where } x \neq y\text{)}$	y
$\lambda x. E_2 \text{ (where } x = y)$	λx . E_2
$\lambda y. E_2$ (where $x \neq y$ and y is not free in E_1)	$\lambda y. E_2[E_1/x]$
$\lambda y. E_2$ (where $x \neq y$ and y is free in E_1)	λy_1 . $E_2[y_1/y][E_1/x]$ where y_1 is a variable not free in E_1 or E_2
E' E''	$E'[E_1/x]$ $E''[E_1/x]$

The LISP function substitute, defined below, implements this substitution algorithm. The idea is that:

(substitute
$$E$$
 E_1 x) = $E[E_1/x]$

Note that from the table it follows that:

- (i) If x is not free in E then substitute just returns E.
- (ii) If x is free in E, then substitute does a recursive descent through E according to the table above.

Although the definition of the LISP function substitute given below looks complicated, it is actually quite straightforward. The way to understand it is to compare it line by line with the table.

```
(defun substitute (e e1 x)
(if (not (member x (frees e)))
     (cond ((is-var* e)
            (if (eq (name* e) (name x)) e1 e))
           ((is-abs*e)
            (let ((y (bv*e))
                  (e2 (body* e)))
             (cond ((equal y x) e)
                   ((not(member y (frees e1)))
                    (mk-abs y (substitute e2 e1 x)))
                    (let ((y1 (variant
                                (append
                                 (frees e1)
                                 (frees e2)))))
                     (mk-abs y1
                             (substitute
                                (substitute e2 y1 y)
                               е1
                               x)))))))
           ((is-comb* e)
            (mk-comb
             (substitute (rator* e) e1 x)
             (substitute (rand* e) e1 x))))))
```

The two sessions below illustrate substitute on examples (i) and (ii) in Exercise 50 on page 73:

```
-> (setq e (parseq (L y (x (L x x)))))
(L y (x (L x x)))
-> (setq e1 (parseq (L y (y x))))
(L y (y x))
-> (setq x (parseq x))
x
-> (setq e2 (substitute e e1 x))
(L y ((L y (y x)) (L x x)))
->
-> (setq e (parseq (y (L z (x z)))))
(y (L z (x z)))
```

```
-> (setq e1 (parseq (L y (z y))))
(L y (z y))
-> (substitute e e1 x)
(y (L |z'| ((L y (z y)) |z'|)))
->
```

12.2.2 β -reduction

The function beta-conv, defined below, reduces a β -redex; it returns nil if applied to non-redexes. The global variable beta-count will be used to hold a cumulative count of the number of β -reductions done. It is convenient to define a macro inc for incrementing variables: (inc x) expands to (setq x (add1 x)).

The function reduce1 finds the leftmost β -redex in an expression and returns the expression that results when the redex is reduced; beta-count is incremented. If the expression contains no β -redexes then nil is returned (and beta-count is not incremented). The algorithm for finding the leftmost redex in E is as follows:

- 1. If E is λx . E_1 then reduce1 is applied to E_1 to get E_{11} . If E_{11} in not nil (i.e. a redex was reduced) then λx . E_{11} is returned. If E_{11} is nil then there are no β -redexes in E and so nil is returned.
- 2. If E is (E₁ E₂) then if E is a β-redex it is reduced and the result returned, otherwise reduce1 is applied to E₁ to get E₁₁. If this is not nil then (E₁₁ E₂) is returned. If E₁₁ is nil then reduce1 is applied to E₂ to get E₂₂. If this is not nil then (E₁ E₂₂) is returned. If E₂₂ is nil then there are no β-redexes in E and so nil is returned.

The definition of reduce1 in LISP may be clearer than this explanation in English:

The function reduce keeps reducing an expression until there is no more change. If the global flag reduce-flag is t then intermediate redexes are printed out:

The macros trace-on and trace-off switch the printing of reductions on and off respectively:

```
(setq reduce-flag nil)
  (defun trace-on () (setq reduce-flag t))
  (defun trace-off () (setq reduce-flag nil))
```

The macro REDUCE sets beta-count to 0 and then reduces the result of parsing its argument (which is not evaluated). It returns the unparse of the result of the reduction:

```
(defmacro REDUCE (e)
'(prog2 (setq beta-count 0)
(unparse (reduce (parseq ,e)))))
```

Here is an example illustrating the reducer:

```
-> (LET TRUE (L x y x))

TRUE

-> (LET FALSE (L x y y))

FALSE

-> (LET IF (L t x y (t x y)))

IF

-> (REDUCE (IF TRUE E1 E2))

E1

-> (REDUCE (IF FALSE E1 E2))

E2

->
```

The reduction with tracing switched on is as follows:

```
-> (trace-on)
t
-> (REDUCE (IF TRUE E1 E2))
((L x y (TRUE x y)) E1 E2)
((L y (TRUE E1 y)) E2)
(TRUE E1 E2)
((L y E1) E2)
E1
E1
->
```

The reducer can be used to experiment with Church's representation of numbers. The definitions of ZERO, SUC and IS-ZERO are explained in Section 5.3.

```
-> (LET ZERO (L f x x))
ZERO
-> (LET SUC (L n f x (n f (f x))))
SUC
-> (LET IS-ZERO (L n (n (L x FALSE) TRUE)))
IS-ZERO
-> (trace-off)
```

```
nil
    -> (REDUCE (IS-ZERO ZERO))
    -> (REDUCE (IS-ZERO (SUC ZERO)))
    FALSE
    -> (REDUCE (IS-ZERO (SUC X)))
    (X (L x FALSE) FALSE)
   Here is a session illustrating the addition function ADD:
    \rightarrow (LET ADD (L m n f x (m f (n f x))))
    ADD
    -> (REDUCE (ADD (SUC ZERO) (SUC (SUC ZERO))))
    (L f x (f (f (x))))
   The pairing functions FST, SND and PAIR are defined by:
    -> (LET FST (L p (p TRUE)))
    FST
    -> (LET SND (L p (p FALSE)))
    SND
    -> (LET PAIR (L x y f (f x y)))
    PAIR
    -> (REDUCE (FST (PAIR E1 E2)))
    -> (REDUCE (SND (PAIR E1 E2)))
    E2
    ->
   The reducer can be used to check that the almost incomprehensible
predecessor function described on page 84 works:
    -> (LET
         PREFN
         (L f p (PAIR FALSE (IF (FST p) (SND p) (f (SND p))))))
    -> (LET PRE (L n f x (SND (n (PREFN f) (PAIR TRUE x)))))
    -> (REDUCE (PRE (SUC (SUC ZERO))))
    (L f x (f x))
    -> beta-count
```

The multiplication function ${\tt MULT}$ was defined above to illustrate LETREC. Here is an example:

The function tricky in Exercise 104 on page 163 is defined in the session below:

The corresponding definition in the λ -calculus is:

Notice that the LISP tricky and the λ -calculus TRICKY give different results.

12.3 Translating to combinators

A modification of the rewriting engine of Chapter 10 can be used to translate λ -expressions to combinators. The idea (see page 131) is simply to rewrite using the equations

```
\lambda x. x = \mathbf{I} 

\lambda x. y = \mathbf{K} y \text{ (if } x \neq y) 

\lambda x. C = \mathbf{K} C \text{ (if } C \text{ is a combinator)} 

\lambda x. E_1 E_2 = \mathbf{S} (\lambda x. E_1) (\lambda x. E_2)
```

This suggests defining

```
(LET I (L x x))
(LET K (L x y x))
(LET S (L f g x ((f x) (g x))))
```

and then using the rewriter with the list of equations remove-lambdas defined below:

```
(setq
  remove-lambdas
'(
    ((L v v) = I)
    ((L v v1) = (K v1))
    ((L v c) = (K c))
    ((L v (e1 e2)) = ((S (L v e1)) (L v e2)))
))
```

Unfortunately this won't work because, with the matcher defined in Chapter 10, there is no way of constraining v1 only to match variables and c only to match combinators. To enforce these necessary constraints the matcher must be modified. This is done by having it look at the property lists of variables occurring in patterns. If such a variable has a predicate P as its TYPE, then the variable will only match expressions E such that (P E) evaluates to t. Here, for example, is a predicate is-combinator that only matches combinators:

```
(defun is-combinator (e) (null(frees(parse e))))
```

To declare that v, v1 and v2 only match variables and c, c1 and c2 only match combinators, the following function is used:

```
(defun map-putprop (list val prop)

(mapcar (function (lambda (x) (putprop x val prop))) list))

(map-putprop '(v v1 v2) (function symbolp) 'TYPE)

(map-putprop '(c c1 c2) (function is-combinator) 'TYPE)
```

Here is the modified matcher; it should be compared with the one on page 183:

```
(defun matchfn (pat exp alist)
 (if (atom pat)
     (if (is-variable pat)
         (if (assoc pat alist)
             (if (equal (cdr(assoc pat alist)) exp)
                 alist
                 (throw 'fail))
             (if (get pat 'TYPE)
                 (if (funcall (get pat 'TYPE) exp)
                      (cons (cons pat exp) alist)
                      (throw 'fail))
                 (cons (cons pat exp) alist)))
         (if (eq pat exp) alist (throw 'fail)))
     (if (atom exp)
         (throw 'fail)
         (matchfn
          (rest pat)
          (rest exp)
          (matchfn (first pat) (first exp) alist)))))
```

Note that if this definition is loaded after the definition on page 183 then, by dynamic binding, it is the later definition that will be used. Thus, since the file prover.1 has been included, the functions match, depth-conv etc. can be used and the new version of matchfn will be invoked by them.

It is necessary to make L into a constant for matching; later I, K, S, B, C, S1, B1 and C1 will also need to be constants.

```
(map-putprop '(L I K S B C S1 B1 C1) t 'constant)
```

To translate a λ -expression to a combinatory expression, the function re-depth-rewrite defined on page 189 can be used. A function trans is defined to take a list of equations and an expression and then to:

- (i) parse the expression;
- (ii) expand any names defined with LET to their definitions;
- (iii) rewrite the result using re-depth-conv and the supplied list of equations; and
- (iv) return the unparsed result of the rewriting.

For (ii) the function expand-defns is required that recursively goes through an expression replacing names by their definitions:

Here is an example illustrating expand-defns:

```
-> (expand-defns TRUE)
(L x (L y x))
-> (expand-defns IF)
(L t (L x (L y ((t x) y))))
-> (expand-defns (parseq (IF TRUE E1 E2)))
((((L t (L x (L y ((t x) y)))) (L x (L y x))) E1) E2)
```

The function trans is defined as follows:

```
(defun trans (eqns e)
(unparse (re-depth-rewrite eqns (expand-defns (parse e)))))
```

Here are the translations of TRUE and FALSE using the equations in remove-lambdas (we assume these have been defined using LET as above):

```
-> (trans remove-lambdas TRUE)
(S (K K) I)
-> (trans remove-lambdas FALSE)
(K I)
->
```

Here is the translation of \mathbf{Y} (we assume it has been defined with LET as above):

```
-> (trans remove-lambdas Y)
(S (S (K S) (S (K K) I)) (K (S I I)))
(S (S (K S) (S (K K) I)) (K (S I I))))
```

To implement Curry's algorithm (Section 8.6) an appropriate set of equations called curry is defined:

```
(setq
curry
'(((L v v) = I)
    ((L v v1) = (K v1))
    ((L v c) = (K c))
    (((S (K e1)) (K e2)) = (K (e1 e2)))
    (((S (K e)) I) = e)
    (((S (K e1)) e2) = ((B e1) e2))
    (((S e1) (K e2)) = ((C e1) e2))
    ((L v (e1 e2)) = ((S (L v e1)) (L v e2)))
))
```

These equations can be used to give the translation of \mathbf{Y} in Exercise 94 on page 140:

```
-> (trans curry Y)
(S (C B (S I I)) (C B (S I I)))
```

Exercise 113

Write a set of equations corresponding to Turner's algorithm (Section 8.7). \Box

12.4 A combinator reducer

Combinators can be reduced using our rewriting engine. An appropriate list of equations for Curry's algorithm is:

```
(setq
  curry-reductions
'(
        ((I x) = x)
        (((K x) y) = x)
        ((((S f) g) x) = ((f x) (g x)))
        ((((B x) y) z) = (x (y z)))
        ((((C x) y) z) = ((x z) y))
        ))
```

The function curry-reduce translates an expression to combinators and then reduces it:

```
(defun curry-reduce (e)
(unparse
(re-depth-rewrite
curry-reductions
(re-depth-rewrite curry (expand-defns(parse e))))))
```

This 'combinator machine' is incredibly slow, but you can use it for experiments. Here is an example:

```
-> (curry-reduce '(IF TRUE E1 E2))
E1
-> (curry-reduce '(IF FALSE E1 E2))
E2
-> (curry-reduce '(ADD (SUC ZERO) ZERO))
(S (B B (S (B B (K I)) I)) (K I))
-> (curry-reduce '(ADD (SUC ZERO) ZERO f x))
(f x)
-> (curry-reduce '(ADD (SUC ZERO) (SUC (SUC ZERO)) f x))
(f (f (f x)))
->
```

Exercise 114

Devise a set of equations corresponding to Turner's algorithm, as described by him in the quotation on page 143. \Box

Exercise 115

Implement as efficient a combinator reducer as you can. Compare its performance with the one described here. \Box

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