THE SAC-1 LIST PROCESSING SYSTEM®

by

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ABSTRACT

SAC-1 is a computer-independent system for Symbolic and Algebraic Calculation, programmed in Fortran IV (as defined by USASI). The system is especially directed towards efficient performance of large-scale calculations with multivariate polynomials and rational functions having exact coefficients. The SAC-1 List Processing System is the first of a series of SAC-1 subsystems (seven existing, three nearing completion, others planned). The present report is a new edition, with some corrections and minor changes, of the original report of July 1967.
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1. **Introduction**

SAC-1 is a computer-independent system for Symbolic and Algebraic Calculation (the suffix, -1, provides for subsequent versions of the system). SAC-1 is especially designed for efficiently performing a wide variety of operations on polynomials and rational functions in several variables, with the arithmetic operations on coefficients being performed exactly. These polynomials and rational functions, and other mathematical objects, are represented in the computer as lists, or "list structures". Therefore, the SAC-1 List Processing System is provided as the first, and most basic, of a series of SAC-1 subsystems. A second SAC-1 subsystem, [7], performs operations on infinite-precision integers. Other subsystems, [8,9,10,11,12], perform progressively more specialized and sophisticated operations on polynomials and rational functions. Three additional subsystems (for polynomial greatest common divisor and resultant calculation, for polynomial factorization and for exact solution of systems of linear equations) are presently near completion, and still others are being planned.

SAC-1 is computer-independent in the sense that it is programmed, with the exception of a small number of simple primitive subprograms, entirely in the ASA (now USASI) Fortran IV language, [3]. In order to implement SAC-1 on a particular type of computer, it suffices to compile the SAC-1 Fortran subprograms and to program in assembly language
according to specifications the primitive subprograms. A programer familiar
with the assembly language should be able to program and test the primitive
in no more than a week. The SAC-1 system has been implemented on CDC
1604, 3600, 6400, 6600 and 7600 computers, IBM 7094 and system 360 com-
puters (models 50, 65 and 67), GE 635 and 645, UNIVAC 1108, and others.

The SAC-1 List Processing System together with the SAC-1 integer arith-
metic and polynomial systems, [7] and [8], constitute a computer-independent
counterpart of the earlier PM system, [1], for the IBM 7094, in that similar
capabilities are provided using similar algorithms and data structures. In
particular, the SAC-1 List Processing System is a reference-count system
having origins in [1] and [4]. The SLIP system, [5], is another well-known
reference-count list processing system, which however differs in that it em-
loys a symmetric, or doubly-linked, representation of lists. For a thorough
discussion of list processing, including the relative advantages and disad-
vantages of the use of reference counts versus "garbage collection" methods,
see [2].

2. Concepts and Terminology

In [4] we introduced the notation of a list as an abstract mathematical
object, to be distinguished from its various possible symbolic and computer
memory representations. Since this distinction will be assumed in our des-
cription of the SAC-1 system, it is appropriate to repeat here some of the
relevant definitions and to introduce certain terminology.

Intuitively, a "list" is just a finite sequence, some of whose elements
may themselves be finite sequences, etc., to any desired depth. Those
elements which are not themselves finite sequences will be referred to as "atoms", and the "depth" to which one may descend before encountering an atom will be called the "order" of a list. This leads to the following more precise definitions. Let \( A \) be an arbitrary set. In contexts where \( A \) is fixed, the members of \( A \) will be called atoms. Now set \( \circ_{0}^{0} A = A \). Assuming \( B_{m} \) has been defined, define \( A_{m+1} \) to be the set of all finite sequences, each of whose elements is a member of \( B_{m} \), and then define \( B_{m+1} \) to be the union \( \bigcup_{i=0}^{m+1} A_{i} \). Clearly \( B_{0} \subseteq B_{1} \subseteq B_{2} \subseteq \ldots \). Set \( L_{m+1} = B_{m+1} - B_{m} \) (for \( m \geq 0 \)).

\( L_{m+1} \) is the set of all lists, over \( A \) of order \( m+1 \), and \( L = \bigcup_{i=1}^{\infty} L_{i} \) is the set of all lists over \( A \).

It is customary to denote a list by enclosing its elements in parentheses, separated by commas. For example, if we take \( A \) to be the set of natural numbers, then "(3, 1, 1, 2)" denotes a list of order 1, while "((3), 1, (1,2))" denotes a list of order 2. Note that the order of occurrence of elements is significant and that an element may have several occurrences. Thus, (3, 1, 1, 2), (3, 1, 2), and (1, 2, 3) are three distinct lists. The length of a list is its length considered as a finite sequence. For example, (3, 1, 1, 2), (3, 1, 2) and ((3), 1, (1,2)) are lists of lengths 4, 3 and 3 respectively. A list may have no elements at all; there is a unique such list, called the null list, which is denoted by "( )". Its length is zero, and its order is one. By contrast, (( )) is a list of length one and order two; its unique element is the null list.

The elements of ((3), 1, (1,2)) are (3), 1, and (1,2); 3 and 2 are not elements, but are constituents. The constituency relation is the transitive closure of the elementhood relation: every element of \( z \) is a constituent of \( z \), and if \( y \) is a constituent of \( z \) and \( x \) is a constituent of \( y \), then \( x \) is a constituent of \( z \).
Length and order are two measures (width and depth) of the complexity of a list. An area-like measure of a list is its extent. Let us say that the extent of an atom is zero. Then the extent of an arbitrary list \((x_1, \ldots, x_n)\) is defined, by induction on its order, to be \(n + \sum_{i=1}^{n} \text{extent}(x_i)\). We will see that in SAC-1 the extent of a list is the number of cells required for its computer memory representation (except when overlapping occurs).

The inverse of any list \((x_1, \ldots, x_n)\) is the list \((x_n, \ldots, x_1)\). Any \(x_i\) which are themselves lists are not inverted. For example, the inverse of \(((3), 1, (1,2))\) is \(((1,2), 1, (3))\). The concatenation of two lists \((x_1, \ldots, x_m)\) and \((y_1, \ldots, y_n)\) is the list \((x_1, \ldots, x_m, y_1, \ldots, y_n)\).

3. Computer Representation of Lists

In SAC-1, any list is represented in computer memory by a collection of cells. A cell consists of a fixed number, say \(k\), of memory locations. The number \(k\) is fixed for a particular implementation of SAC-1, but may be different for different implementations. For most computers \(k\) will be two, but an IBM 360 computer with a 32-bit word, for example, requires \(k = 3\). By convention, the address of a cell is the address of its first memory location.

Each cell consists of four fields: type field \((T)\), element field \((E)\), reference count field \((R)\), and successor field \((S)\). The computer representation of a list \((x_1, \ldots, x_n)\) consists of a set of \(n\) cells, say \(C_1, \ldots, C_n\), linked via their successor fields, together with representations, assumed already given, of any of the \(x_i\) which are themselves lists. I.e., the successor field of cell \(C_i\) contains the address of cell \(C_{i+1}\) \((1 \leq i < n)\); the successor field of \(C_n\) contains zero (it is assumed that zero is not the address of any cell, which is easily arranged). The type field of cell \(C_i\) contains the value zero or one according as \(x_i\) is an atom or a list. If \(x_i\) is an atom, then the
element field of \( C_i \) contains \( x_i \); if \( x_i \) is a list, then the element field of \( C_i \) contains the location of some representation of the list \( x_i \). In this connection, the location of the list \((x_1, \ldots, x_n)\) is taken to be the address of the first cell, \( C_i \), in its representation. Of course, a given list may have several representations, simultaneously.

Note that, according to this definition, whenever a representation of a list \((x_1, \ldots, x_n)\) is present, so is a representation of any final segment of it, \((x_k, \ldots, x_n)\), automatically. This turns out to be a very useful property for many applications, polynomial algebra in particular.

The above definition does not apply to the representation of the null list. By special convention, no cells are used in its representation, and the location of its representation is zero.

We draw diagrams of list representations in which each cell of the representation is depicted by a rectangle. Such a rectangle contains four subrectangles corresponding to the four fields of the cell. The geometry of the diagram is derived from the typical case in which there are two words per cell and the element field consists of exactly the second word. Here is a diagram of a cell:

```
<table>
<thead>
<tr>
<th>T</th>
<th>R</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>E</td>
</tr>
</tbody>
</table>
```

A diagram for our example list \(((3), 1, (1,2))\) would appear as follows:

```
1  | 0  | 1  
  |    |    
0  | 3  |    
  |    |    
```
The contents of successor fields are suppressed, since they are determined (to within an isomorphism) by the horizontal arrows; similarly for element fields containing the location of a list. The type fields could also be inferred from the diagram.

We have not yet specified the contents of reference count fields. The reference count field of any cell contains the number of arrows which terminate at it. However, this count must include a third type of arrow not shown in the above diagram, the program arrow. At any instant during the execution of a program, there exists a certain number of variables which are currently defined, and whose current values are locations of lists. Each such variable has associated with it a program arrow terminating at the first cell of the list which is its value.

It follows from this discussion, and from the following section on list erasure, that, in a correct program, all reference counts are greater than or equal to one.

With regard to field sizes, one bit obviously suffices for the type (but more could be used). The element field is assumed to be no longer than one word (but it could be shorter). The successor field must be long enough to contain any machine address (whether real or virtual). It is easy to see that the number of distinct computer memory locations (real or virtual) is an upper bound for any reference count. Hence if the reference count field is given the same length as the successor field, there will be no possibility of reference count overflow. In the present version of the system, no check is made for such overflow. However, an alternate version of the system is available in which a shorter field is allotted to the reference count, overflow is detected and, when it occurs, remedial action is taken, which consists in making a partial copy of the list in which the overflow would otherwise occur. This
alternate version involves a change in only a single system subprogram, namely BORROW.

4. Erasure and the Available Space List

As in other list processing systems, any cell is, at any instant, included in the representation of at least one list. This is provided for by the presence of a special list, the available space list, which is complementary to all other lists: a cell is in the available space list if and only if it is not in any other list.

In SAC-1, as in most other systems, the available space list, hereafter called AVAIL, is maintained in the form of a list of order one (SLIP, [5], is the known exception). For the sake of easy identification, each cell on AVAIL carries a reference count of zero.

So that it will be accessible to all the subprograms requiring it, the location of the available space list is kept in Common, specifically, in the first location of labelled common block TR1, with the symbolic designation AVAIL.

An initial available space list is created by the subroutine BEGIN (A, N), in which A is a one-dimensional integer array of N elements. N must be a multiple of the number, k, of words per cell. An available space list of N/k cells is created from the memory allocated to A, and its location is stored in AVAIL.

The only system subprograms which remove cells from AVAIL are PFA and PFL. This is a desirable feature to maintain, since it makes it easy to acquire statistics on the numbers of cells used in various applications.

Whenever a particular list representation is no longer needed in the continued execution of a program, its "release" must be programmed through a
call of the subroutine ERASE. ERASE returns to AVAIL all cells in the given list which are not shared with other lists, as determined by the reference counts, and decrements by one the reference counts of other cells as appropriate. This system has the advantage over the garbage collection of LISP, [6], that the processing time for reclaiming cells for re-use remains always in direct relation to the number of cells reclaimed (see [1] and [4]).

The reference count system makes possible a further gain in efficiency through the use of specialized subroutines for the erasure of lists of predetermined structure. One such specialized subroutine, ERLA, is included in the present system for erasing lists of atoms, i.e., lists of order 1. Other specialized subroutines for erasure will be available in other parts of SAC-1.

If a program fails to apply ERASE to a list where this is appropriate, the result is a permanent loss of a certain number of cells. If such an error occurs in a section of program which is executed repeatedly, the result may be a rapid, complete depletion of AVAIL. Hence, careful attention should be given to this matter.

Of course, a list need not be immediately erased after its usefulness has terminated. The importance of immediate erasure depends on how extensive the list may be. A similar remark applies in garbage collection systems, where it may be desirable to program abandonment of lists, since automatic abandonment does not occur until a list variable is redefined or execution of a subprogram is completed.

5. The List Processing Primitives
Ten primitive subprgrms are required for the SAC-1 list processing system. Of these, two are associated with each of the four cell fields, for a total of eight; the other two are used only by the BEGIN subroutine. These primitives must, for the most part, be written in an assembly language, in the
correct form for linkage with the Fortran programs.

There are four primitive subroutines for storing information in cell fields. They are: \texttt{STYPE}(X,Y) - type field, \texttt{SCOUNT}(X,Y) - reference count field, \texttt{SSUCC}(X,Y) - successor field, and \texttt{ALTER}(X,Y) - element field. In each case the argument \( Y \) is the location of some cell and the argument \( X \) is a value which is to be stored in the appropriate field of the cell at \( Y \). In each case, the arguments \( X \) and \( Y \) are Fortran integers, with the possible exception of \texttt{ALTER}. Here, the correct interpretation of \( X \) may be something other than an integer (e.g. Hollerith), but with respect to Fortran, it is disguised as an integer; and \texttt{ALTER} merely transfers \( X \) from one memory location to another.

There are four corresponding field retrieval function subprograms: \( Z=\texttt{TYPE}(Y) \), \( Z=\texttt{COUNT}(Y) \), \( Z=\texttt{TAIL}(Y) \) - successor field, and \( Z=\texttt{FIRST}(Y) \) - element field. As above, each argument \( Y \) is the location of some cell. The appropriate field of cell \( Y \) is extracted and this value is returned as the Fortran integer \( Z \).

Two additional function subprograms are needed for \texttt{BEGIN}: \( Z=\texttt{NWPC}(X) \) and \( Z=\texttt{LOC}(X) \). In \texttt{NWPC}, the argument \( X \) is not referenced. The value \( Z \) returned is always the number of words per cell. \texttt{NWPC} could be programmed in Fortran, but it is classed as a primitive because it is computer-dependent. In \texttt{LOC}, the argument \( X \) is some variable (possibly subscripted). The value \( Z \) returned is the address of the memory location assigned to \( X \). \texttt{LOC} is unlike the other primitives in that there is no assurance implicit in the A.S.A. Fortran specifications that the subprogram \texttt{LOC} can be written, even in machine language. For most computers and their Fortran compilers, it can be; in those cases where it cannot, it may be necessary to rewrite \texttt{BEGIN} as a primitive.
6. **Description of the FORTRAN Subprograms**

In the following descriptions, function subprograms are indicated by a prototype of the form \( Z = F(X_1, \ldots, X_n) \), while subroutines are indicated by a prototype of the form \( S(X_1, \ldots, X_n) \). In SAC-1, the arguments of a function subprogram are never defined or redefined as a result of the execution of the subprogram.

**ADV(X,Y).** \( Y \) is a non-null list \((y_1, \ldots, y_n)\). After execution of **ADV**, the value of \( X \) is \( y_1 \) and the new value of \( Y \) is the list \((y_2, \ldots, y_n)\). The given list \((y_1, \ldots, y_n)\) remains intact. CALL **ADV** (\( X, Y \)) is equivalent to \( X = \text{FIRST}(Y), \; Y = \text{TAIL}(Y) \).

**BEGIN (A,N).** \( A \) is a one-dimensional integer array of \( N \) elements. \( N \) must be a multiple of \( K \), the number of words per cell in the SAC-1 implementation at hand. **BEGIN** creates an available space list from \( A \) consisting of \( N/K \) cells and initializes to null the system pushdown stack, \( \text{STAK} \). Any SAC-1 main program will therefore normally begin with a call of **BEGIN**.

**Z = BORROW (X).** \( X \) is the location of an arbitrary list. The reference count of this list is increased by one (unless \( X \) is null), and \( Z = X \), in the current implementation. However, in order to maintain compatibility with alternate implementations, it is recommended that users always reference **BORROW** in the form \( X = \text{BORROW}(X) \) rather than **CALL** **BORROW**(\( X \)).

The current implementation assumes the increased reference count will never overflow the reference count field. In implementations where such overflow is possible, an alternate version of **BORROW** will return a value which may be the location of a (partially) different representation of the given list when necessary to prevent overflow.
Z = CINV(X). X is the location of an arbitrary list \((x_1, \ldots, x_n)\). The inverse list \((x_n, \ldots, x_1)\) is constructed, and Z is its location. The list X remains intact. Elements \(x_i\) of X which are lists have their reference counts increased by one via an application of BORROW, and hence are not copied.

Z = CONC(X,Y). X and Y are lists, say \((x_1, \ldots, x_m)\) and \((y_1, \ldots, y_n)\). X and Y are concatenated to form the list \((x_1, \ldots, x_m, y_1, \ldots, y_n)\) by storing the location of Y in the successor field of the last cell of X. Hence X no longer exists (unless Y is null). None of the reference counts in either X or Y are modified. Application of CONC is usually inappropriate except when all main level reference counts of X and Y are one and X and Y are not needed after their concatenation.

DECAP(X,Y). Y is the location of a non-null list \((y_1, \ldots, y_n)\). After execution of DECAP, the value of X is \(y_1\) and the new value of Y is the list \((y_2, \ldots, y_n)\). The first cell of the given list is returned to the available space list, whatever its reference count was initially. No reference counts in \((y_2, \ldots, y_n)\) are changed. Application of DECAP is therefore ordinarily appropriate only when all main-level reference counts in the given list are known to be one.

ERASE(X). The arbitrary list X is "erased." I.e., those cells of X which are not parts of some other list representation, as determined by their reference counts, are returned to the available space list and the reference counts of other cells in X are decreased by one as appropriate.

ERLA(X). X is the location of an arbitrary list of atoms, i.e., a list of order one. X is erased; i.e., the effect of ERLA is the same as that of ERASE. However, knowledge of the structure of X is used by ERLA,
resulting in appreciably faster execution.

\[ \text{Z=} \text{INV(X). } \text{X is the location of a list } (x_1, \ldots, x_n), \text{ possibly null.} \]

The given list is modified into the inverse list \((x_n, \ldots, x_1)\), whose location, \(Z\), is returned as value. The given list \(X\) no longer exists. The inversion is carried out by modifying the successor field only of each of the \(n\) cells in the main level of the given list. Hence INV is appropriately applied only to a list all of whose main level reference counts are one (typically a list which has just been constructed).

\[ \text{Z=} \text{LENGTH(X). } \text{X is an arbitrary list } (x_1, \ldots, x_n). \text{ Z is the length, } n, \text{ of } X, \text{ expressed as a Fortran integer.} \]

**NOAVLS.** A system subroutine called by PFA or PFL whenever the available space list is empty. It prints the diagnostic "out of available space" and terminates job execution.

\[ \text{Z=} \text{PFA(X,Y). } \text{X is an atom, Y is the location of an arbitrary list } (y_1, \ldots, y_n). \text{ X is prefixed to the list } Y, \text{ resulting in the list } (X, y_1, \ldots, y_n). \]

The given representation of \(Y\) remains intact, and its reference count remains unchanged. The reference count of the resulting list is one and the value \(Z\) returned is its location. A single cell is removed from the available space list, into which is stored the element \(X\).

\[ \text{Z=} \text{PFL(X,Y). PFL is exactly like PFA except that here the argument } X \text{ is a list. The reference count of } X \text{ is not modified, since typically the list } X \text{ is not independently needed, or else its reference count has previously been augmented by an application of BORROW. Note that, if } X = (x_1, \ldots, x_m) \text{ and } Y = (y_1, \ldots, y_n), \text{ then the resulting list is } (X, y_1, \ldots, y_n), \text{ a list of length } n+1 \text{ whose first element is the list } X, \text{ not the list } (x_1, \ldots, x_m, y_1, \ldots, y_n) \text{ as in concatenation.} \]
STACK2 (X,Y). X and Y are placed on the system pushdown stack - first X, then Y - by two applications of PFA. X and/or Y may, however, be the locations of lists; this causes no difficulty since the pushdown stack is never processed as a general list in which the types of the elements must be checked.

STACK3(X,Y,Z). Like STACK2 except for the third argument Z, which is stacked last.

UNSTK2(X,Y). The top element on the pushdown stack is removed (via DECAP) and assigned to Y; then the next is removed and assigned to X. Note that a CALL STACK2(X,Y) will normally be paired with a CALL UNSTK2(X,Y) - as opposed to CALL UNSTK2(Y,X).

UNSTK3(X,Y,Z). The extension of UNSTK2 to three arguments. Z is unstacked first.
7. **Recursion in SAC-1**

One of the main deficiencies in A.S.A. Fortran as a language for list processing and for symbolic and algebraic manipulation is its failure to provide for recursive subprograms (see [3], p. 615 and p. 618). For use in SAC-1 a method has been developed for circumventing the prohibition against, and lack of provision for, recursion. The method is somewhat inconvenient to use, but completely general. The SLIP system, [5], provides a mechanism for recursion in Fortran, but it requires the use of two additional primitives; moreover these primitives violate certain A.S.A restrictions and, indeed, there is no assurance that these primitives can be realized in a way that is consistent with an arbitrary Fortran compiler.

A subprogram is **recursive** in case the completion of a particular execution of that subprogram may require the prior initiation of a new execution. This may come about either **directly**, as a result of the subprogram referencing itself, or **indirectly**, as when the given subprogram references another subprogram which in turn references the former (either directly or indirectly). Recursion in Fortran poses three distinct problems, which we shall now discuss, one by one.

The first problem is the syntactic prohibition, stated in [3], that a subprogram shall not reference itself, directly or indirectly. Its effect is that any subprogram which references itself directly will likely result in a diagnostic, and will fail to be compiled. The syntactic prohibition against indirect recursive reference, on the other hand, is unenforceable since its detection would require that the several subprograms in the circular chain of references be compiled simultaneously. Since a direct self-reference is easily translatable into an indirect self-reference, by the introduction of a new subprogram which does nothing more than to call the original subprogram with the
same argument list, this first problem is easily circumventable.

The other two problems pertain to possible inadequacies in the compiled programs. When an interrupted execution is later resumed, it is essential that all its variables have the same definitional status and values as prior to the interruption, although these may have been altered by intervening executions. The A.S.A. specifications do not require this restoration and it must be explicitly programmed. The solution to this problem is quite simple with the use of a pushdown stack.

The SAC-1 system has a single pushdown stack, which is in the form of a list of order one, in which all reference counts are one. Values are placed on the stack by prefixing them to the list, and removed by the list decapitation process. The location of the pushdown stack is kept in the second memory location of the labelled common block TR1, bearing the symbolic name STAK. Special system subroutines, STACK2, STACK3, UNSTK2 and UNSTK3, described in Section 6, are provided to facilitate stacking and unstacking. Just prior to a recursive reference, current values of program variables are stacked and, immediately following the recursive reference, these values are unstacked and reassigned to the same variables. Not every program variable needs to be so preserved and restored – only those whose current value will be used in completing the interrupted execution. This will be illustrated in the example below.

The third problem connected with recursion pertains to the flow of control, and the Fortran Return statement. When a subprogram may have several executions in progress simultaneously, there is associated with each execution a return location which must be saved along with the current values of program variables. Since, however, these return locations are not, in Fortran, associated with any program variable, it is impossible to use the stack mechanism. We believe, in fact, that, for this reason, it is impossible to program in A.S.A. Fortran a
recursive subprogram in the strict sense. We can, however, devise an effective substitute for a recursive subprogram, which we call a recursive procedure.

A **recursive procedure** is a sequence of executable statements together with a list of variables occurring in these statements, which function as procedure parameters. Among the procedure parameters, one acts as the return location of the procedure. In the case of a function procedure, one also serves to carry the function value. All other procedure parameters play the roles of the dummy arguments of a subprogram.

A recursive procedure has all the essential features of a subprogram and, in addition, a variable associated with its return location. The actual value of the return location variable is not a location, but a statement number, or a code number for use in a computed-go-to statement. This value can be stacked along with the values of other variables. An inessential, although very convenient, feature of subprograms, not shared by recursive procedures, is the automatic association of actual arguments with dummy arguments, as well as the possession of a symbolic name.

We now illustrate the use of recursive procedures by an example. Suppose we wish to write a function subprogram $Z=$ORDER($X$) which, when applied to an arbitrary list $X$, returns as value the order, $Z$, of $X$, as a Fortran integer. This function is naturally defined by recursion since the order of $X$ is just one greater than the maximum order of any element of $X$ which is a list, or one if all elements of $X$ are atoms. Thus, if recursion were permissible, we might define ORDER as follows.
INTEGER FUNCTION ORDER(X)
INTEGER FIRST, TAIL, TYPE
INTEGER X,T
T = X
N = 0
IF (T.EQ.0) GO TO 3
1 IF (TYPE(T).EQ.0) GO TO 2
   M = ORDER (FIRST(T))
   IF (M.GT.N) N = M
2 T = TAIL(T)
   IF (T.NE.0) GO TO 1
3 ORDER = N + 1
RETURN
END

We now rewrite the ORDER subprogram, so that it contains within it a recursive procedure for the order function and is not, itself, recursive. We first choose the procedure parameters: Y for the function argument, N for the function value, and R for the return location. The procedure will be "called" from two places: from outside the procedure, but within the ORDER subprogram and from just one place, in this case, within the procedure itself. We assign code numbers, 1 and 2, to these two places. The following program is obtained, in which comments have been inserted to delimit the structural parts.

INTEGER FUNCTION ORDER(X)
INTEGER FIRST, TAIL, TYPE
INTEGER X,T,Y,R,E

C PROCEDURE CALL
Y = X
R = 1
GO TO 4
5
ORDER = N
C
END PROCEDURE CALL
RETURN
C
RECURSIVE PROCEDURE N = ORDER(Y), RETURN TO R
4
T = Y
N = 0
IF (T.EQ.0) GO TO 3
1
IF (TYPE(T).EQ.0) GO TO 2
E = FIRST(T)
C
RECURSIVE PROCEDURE CALL
CALL STACK3 (N,T,R)
Y = E
R = 2
GO TO 4
6
M = N
CALL UNSTK3(N,T,R)
C
END RECURSIVE PROCEDURE CALL
IF (M.GT.N) N = M
2
T=TAIL(T)
IF (T.NE.0) GO TO 1
3
N = N + 1
C
PROCEDURE RETURN
GO TO (5,6), R
C
END RECURSIVE PROCEDURE N=ORDER(Y), RETURN TO R
END
This program is by no means optimal because, by intent, it illustrates a more or less algorithmic translation of the former program. Also, this simple function could be quite easily programmed without the aid of recursion. But there are much more complex situations where recursion is practically indispensable.

The example above illustrates the case of direct recursion. In the general case of indirect recursion, one has subprograms $A_1, \ldots, A_n$ such that $A_i$ calls $A_{i+1}$, for $1 \leq i < n$, and $A_n$ calls $A_1$. It is then necessary to set up procedures for each $A_i$ within a single subprogram, say $B$. In addition, there is a subprogram for each $A_i$, the sole function of which is to accept computational requests from the system and relay them to subprogram $B$ for the actual computation, together with a code number $i$, which tells subprogram $B$ which procedure to invoke.

Suppose, for simplicity, that $n = 2$ and that each $A_i$ is a function of one argument. Subprograms $A_1$, $A_2$ and $B$ (in outline) would then appear as follows.

```
INTEGER FUNCTION A1(X)
INTEGER X,B
A1 = B(1,X)
RETURN
END

INTEGER FUNCTION A2(X)
INTEGER X,B
A2 = B(2,X)
RETURN
END
```
INTEGER FUNCTION B(I,X)
INTEGER X,U,R,W,V,S,Z,...
IF (I.EQ.2) GO TO 2
1   U = X
   R = 1
   GO TO 3
4   B = W
   RETURN
2   V = X
   S = 2
   GO TO 5
6   B = Z
   RETURN
3   (procedure A1)
5   (procedure A2)
END

Here U, R and W (V,S and Z) are the argument, return location, and value variables for procedure A1 (A2).

The above programs serve to illustrate another programming device not related to recursion, but to another inconvenience of Fortran. In a system such as SAC-1, we generally want to eliminate the distinction of the various types of variables because it may, for example, result in unwanted conversions from fixed to floating-point, or vice versa. Since we do, however, wish some data to be interpreted as integers, the solution is to disguise all data to Fortran as integers. It is unnatural and anti-mnemonic to begin every variable and function name with I, J, K, L, M or N. So we choose identifiers freely and, after writing any program or subprogram, list in an integer declaration all identifiers which
occur. This is a simple mechanical process which has been found to be non-
error-inducing, and is recommended to users of SAC-1 as well. It also results
in the useful byproduct that all function subprograms referenced by a given sub-
program are listed at the beginning of the subprogram; we usually list them in
a separate declaration.

8. **The SAC-1 Character Set and Internal Code**

Since different computers have different character sets and, also, different
internal codes for the same characters, some scheme is required for maintaining
computer independence in the SAC-1 input-output system. The obvious solution
to the problem of differing character sets is the choice of a small set of characters
common to all computers. Such a choice is provided by the A.S.A. Fortran
character set ([3], p. 593), consisting of all the characters used in writing an
A.S.A. Fortran program. The adoption of this set as the SAC-1 character set
implies that all input and output data processed by SAC-1 will consist of these
characters only.

A common internal code for these characters is also required. Since the
internal hardware codes of different computers are inconsistent, a conversion
from the hardware code to the SAC-1 code upon input, and the inverse conversion
just prior to output, is, in general, required. Hence, instead of choosing the
hardware code of one particular computer as the SAC-1 code, a code has been
chosen which facilitates the kind of character testing required in the SAC-1 input
conversion subprograms. In this code, functionally similar characters (digits,
letters, operators and punctuators) are assigned contiguous codes. The coding
system is given in the following table.

<table>
<thead>
<tr>
<th>Characters</th>
<th>Decimal Codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 - 9</td>
<td>0 - 9</td>
</tr>
<tr>
<td>A - Z</td>
<td>10 - 35</td>
</tr>
<tr>
<td>+ - * /</td>
<td>36 - 39</td>
</tr>
<tr>
<td>( ) , . blank</td>
<td>40 - 44</td>
</tr>
<tr>
<td>=</td>
<td>45</td>
</tr>
<tr>
<td>$</td>
<td>46</td>
</tr>
</tbody>
</table>

9. The Primitive Subroutines READ and WRITE.

The READ subroutine reads a standard size record from a specified unit U, performs the conversion from hardware code to SAC-1 code, and delivers the input characters as the elements of an array A. Correspondingly, the WRITE subroutine converts the elements of an array A from SAC-1 code to hardware code, and writes the result on unit U as a standard size record. U and A are the subroutine arguments: READ(U,A), WRITE(U,A).

The standard record length is 72. Thus, A is a one-dimensional integer array of 72 elements. Input records may be longer (e.g. 80 in the case of card-generated input), but only the first 72 characters will read and stored in A. Output records generated by WRITE will be of either length 72 or 73, depending on the unit number U. Certain unit numbers, according to local option (as reflected in the implementation of WRITE), are regarded as designating printer output units (either printers or tapes destined for printing). When such a unit is designated, a record of length 73 is written of which the first character
is blank, resulting in single line printer spacing, and \( A(I) \) corresponds to the
\((I + 1)\)-th character of the record for \( 1 \leq I \leq 72 \). Otherwise, \( A(I) \) always

\[ \text{corresponds to the } I \text{-th character of the record (input or output). A 73-character } \]

\[ \text{record so written cannot, of course, be later read as meaningful input.} \]

If, when the READ subroutine is called, unit \( U \) is positioned at an end-of-

\[ \text{file, } A(I) \text{ is given a value of } -1 \text{ as a means of relaying this information to the} \]

\[ \text{calling program. Execution of WRITE may result in redefining the array } A \text{ (e.g.,} \]

\[ A \text{ may then contain the hardware code for the record written).} \]

Subprograms provided as parts of the SAC-1 system will process only records

\[ \text{containing only characters from the Fortran character set. However, individual} \]

\[ \text{users of SAC-1 may wish to make extensions of the system to process additional} \]

\[ \text{characters. This can be done by assigning codes to these characters in the} \]

\[ \text{range 47-63 and implementing READ and WRITE so as to perform the required} \]

\[ \text{code conversions for the expanded character set.} \]

10. **Reading and Writing Lists**

SAC-1 contains two subprograms, LREAD and LWRITE, for reading and writing

\[ \text{arbitrary lists. This requires that an arbitrary list have an "external representation"} \]

\[ \text{as a sequence of characters. This external representation has been illustrated} \]

\[ \text{in Section 2 for the case that atoms are integers represented in standard decimal} \]

\[ \text{notation. In SAC-1, atoms are always sequences of } m \text{ binary digits, where } m \]

\[ \text{is the word length of a particular computer.} \]

(We assume throughout this section that SAC-1 is implemented on a binary

\[ \text{computer. All parts of SAC-1 other than those described in this section are} \]
independent of this assumption, including the parts to be described in subsequent papers. The subprograms here described are useful, but inessential, adjuncts to the system.)

These $m$ binary digits are clustered into groups of three, thereby determining a sequence of octal digits. (If the word length is not a multiple of 3, one or two leading binary zeros are considered implicit.) The external representation is then a sequence of left and right parentheses, commas, and octal digits. The main list and its constituent lists are delimited by parentheses; list elements are separated by commas. Since a sequence of octal digits representing an atom will, in common practice, frequently have several leading zeros, special provision is made for their suppression. In input to LREAD, any number of leading zeros may optionally be suppressed. In output produced by LWRITE all leading zeros will be suppressed. An exception to this rule is that if all digits are zero, one zero must be retained (and is retained by LWRITE). The number of characters in an external list representation may be arbitrarily large, and hence span an arbitrary number of records. The first character always occurs in the first position of the first record, the 73rd character, if it exists, occurs in the first position of record two, etc. The last record is filled out with blanks as required; intervening blanks are not permitted.

$Z=LREAD(U)$ is a function subprogram which reads a list from unit $U$ and generates an internal representation of the list read, returning the location of the internal representation as the value $Z$. If, when LREAD was called, unit $U$ was positioned at an end-of-file, then the value $Z$ returned is instead $-1$. 
LREAD makes various checks on validity of the input list. If the input is invalid, then the value \( Z = -2 \) is returned.

LWRITE(U, X) is a subroutine, which converts the list X to external representation and writes it on unit U. The internal representation remains intact and unerased. Since LWRITE uses WRITE, the record length may be either 72 or 73 depending on U.

Three additional primitive subprograms, LSHIFT, RSHIFT and NBPW, are required by LREAD and LWRITE. These primitives are not used elsewhere in SAC-1 and are therefore dispensable if LREAD and LWRITE are omitted from the system. LREAD and LWRITE serve two main purposes: debugging and the conservation of addressable memory. LREAD may be used to input simple test lists. LWRITE may be used to output intermediate results during program testing. Main memory may be conserved during a calculation by writing lists temporarily on tape, erasing them, and later reading them with LREAD. Later increments to SAC-1 will contain input-output subprograms for lists representing particular types of objects, e.g., large integers, polynomials, rational functions, etc., but the I/O conversions employed are more complex and time consuming, and are not needed for saving intermediate results on tape.

\[ Z = \text{LSHIFT}(X, Y, N) \] is a function subprogram in which \( X = X_1 \ldots X_m \) and \( Y = Y_1 \ldots Y_m \) are atoms, regarded as sequences of bits, and \( N = n \) is a shift amount, expressed as a Fortran integer. \( Z \) is the atom resulting from left-shifting \( X \) \( n \) places and inserting the last \( n \) bits of \( Y \) into the vacated bit positions. I.e., \( Z = X_{n+1} \ldots X_m Y_{m-n+1} \ldots Y_m \). \( X \) and \( Y \) are not redefined.
RSHIFT(X, Y, N) is a subroutine in which \( X = X_1 \ldots X_m \) and \( Y \) are atoms and \( N = n \) is a shift amount. \( X \) is right-shifted \( n \) places so that the redefined value of \( X \) is \( 0 \ldots 0 X_1 \ldots X_{m-n} \). \( Y \) need not be initially defined; its resulting value is \( 0 \ldots 0 X_{m-n+1} \ldots X_m \).

\( Z = \text{NBPW}(X) \) is a function subprogram whose argument \( X \) is ignored and whose value \( Z \) is the number of bits per word in the computer of implementation, expressed as a Fortran integer.

LREAD and LWRITE depend on one additional assumption, which holds for all known computers, but which should probably be made explicit, nevertheless. This assumption is that non-negative Fortran integers in the range \( 0 \leq N \leq 7 \) are represented in standard base two notation; i.e., 0 by 0 \ldots 0, 1 by 0 \ldots 0 1, 2 by 0 \ldots 010, etc.
References


SUBROUTINE ADV(EL, LOC)
COMM /TR1/AVAIL, STAK, RECORD(72)
INTEGER AVAIL, STAK, RECORD
INTEGER EL, LOC, L, TAIL, FIRST
L = LOC
EL = FIRST(L)
LOC = TAIL(L)
RETURN
END

SUBROUTINE BEGIN(ARRAY, N)
COMM /TR1/AVAIL, STAK, RECORD(72)
INTEGER AVAIL, STAK, RECORD
INTEGER ARRAY
DIMENSION ARRAY(N)
J = NWPC(L)
L = LOC(ARRAY(1))
M = N - J
AVAIL = L
DO 1 K = 1, M, J
LN = LOC(ARRAY(K + J))
CALL SSUCC(LN, L)
CALL SCOUNT(0, L)
CALL STYPE(0, L)
CALL ALTE(0, L)
1 L = LN
CALL SSUCC(0, L)
CALL SCOUNT(0, L)
CALL STYPE(0, L)
CALL ALTE(0, L)
STAK = 0
RETURN
END

INTEGER FUNCTION BORROW(LIST)
COMM /TR1/AVAIL, STAK, RECORD(72)
INTEGER AVAIL, STAK, RECORD
INTEGER COUNT
BORROW = LIST
IF (BORROW .EQ. 0) RETURN
CALL SCOUNT(COUNT(BORROW) + 1, BORROW)
RETURN
END

INTEGER FUNCTION CINV(X)
COMM /TR1/AVAIL, STAK, RECORD(72)
INTEGER AVAIL, STAK, RECORD
INTEGER TYPE, PFA, PFL, BORROW
INTEGER X, Z, R, E, T
Z = 0
R = X
1 IF (R .EQ. 0) GO TO 3
T = TYPE(R)
CALL ADV(E, R)
IF (T.EQ.1) GO TO 2
Z=PFA(E,Z)
GO TO 1

2 Z=PFL(BORROW(E),Z)
GO TO 1

3 CINV=Z
RETURN
END

INTEGER FUNCTION CONC(X,Y)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER X,Y,XAD,XNEXT,XSUC,TAIL
CONC=Y
XSUC=CONC
XNEXT=X
IF (XNEXT.EQ.0) RETURN
CONC=XNEXT
IF (XSUC.EQ.0) RETURN
XAD=XNEXT
XNEXT=TAIL(XNEXT)
IF (XNEXT.NE.0) GO TO 1
CALL SSUCEXSUC,XAD)
RETURN
END

SUBROUTINE DECAP(EL,LOC)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER B,EL,TAIL,FIRST
B=AVAIL
AVAIL=LOC
EL=FIRST(AVAIL)
LOC=TAIL(AVAIL)
CALL SSUCXEL,AVAIL)
CALL SCOUNTO,B,AVAIL)
RETURN
END

SUBROUTINE ERASE(XX)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER COUNT,FIRST,TAIL,TYPE
INTEGER X,XX,T,RIN
X=XX
RIN=0
1 IF (X.EQ.0) GO TO 3
2 K=COUNT(X)-1
CALL SCOUNT(K,X)
IF (K.GT.0) GO TO 3
IF (TYPE(X).EQ.1) GO TO 4
T=TAIL(X)
CALL SSUCXAVAIL,X)
AVAIL=X
X=T

3 GO TO 2
4 RETURN
GO TO 1
3 IF(BIN.EQ.0)RETURN
X=FIRST(BIN)
T=TAIL(BIN)
CALL SSUC(AVAIL,BIN)
AVAIL=BIN
BIN=T
GO TO 1
4 T=TAIL(X)
CALL SSUC(BIN,X)
BIN=X
X=T
GO TO 1
END

SUBROUTINE ERLA(X)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER X,U,V,TAIL,COUNT
IF(X.EQ.0)GO TO 3
U=0
V=X
1 N=COUNT(V)-1
CALL SCOUNT(N,V)
IF(N.NE.0)GO TO 2
U=V
V=TAIL(V)
IF(V.NE.0)GO TO 1
2 IF(U.EQ.0)GO TO 3
CALL SSUC(AVAIL,U)
AVAIL=X
3 RETURN
END

FUNCTION INV(LIST)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER TAIL
INV=0
NEXT=LIST
IF(NEXT.EQ.0)RETURN
1 ISTORE=NEXT
NEXT=TAIL(ISTORE)
CALL SSUC(INV,ISTORE)
INV=ISTORE
IF(NEXT.NE.0)GO TO 1
RETURN
END

FUNCTION LENGTH (LIST)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER TAIL
LENGTH=0
K=LIST
1 IF(K.EQ.0)RETURN
K=TAIL(K)
LENGTH=LENGTH+1
GO TO 1
END

INTEGER FUNCTION LREAD(U)
COMMON /TR1/AVAL,STAK,RECORD(72)
INTEGER AVAL,STAK,RECORD
INTEGER PFA,PFL,INV,LSHIFT,NBPW
INTEGER B,C,D,E,P,T,I,J,X,Y,Z
T=0
X=0
Z=0
I=1
B=NBPW(X)
J=B
CALL READ(U,RECORD)
LREAD=-1
IF (RECORD(1).EQ.-1) RETURN
IF (RECORD(1).NE.40) GO TO 90
10 P=RECORD(I)
I=I+1
IF (I.LE.72) GO TO 11
I=1
CALL READ(U,RECORD)
IF (RECORD(1).LT.0) GO TO 90
11 C=RECORD(I)
IF (P.EQ.40) GO TO 20
IF (P.EQ.41) GO TO 21
IF (P.EQ.42) GO TO 22
GO TO 23
20 IF (C.EQ.40) GO TO 30
IF (C.EQ.41) GO TO 31
IF (C.LT.8) GO TO 32
GO TO 90
21 IF (C.EQ.41) GO TO 31
IF (C.EQ.42) GO TO 30
GO TO 90
22 IF (C.EQ.40) GO TO 30
IF (C.LT.8) GO TO 32
GO TO 90
23 IF (C.EQ.41) GO TO 33
IF (C.EQ.42) GO TO 33
IF (C.LT.8) GO TO 32
GO TO 90
30 Z=PFL(X,Z)
X=0
GO TO 10
31 X=INV(X)
IF (Z.EQ.0) GO TO 80
CALL DECAP(Y,Z)
X=PFL(X,Y)
GO TO 10
T=PFA(C,T)
J=J-3
GO TO 10
33 IF (J.LT.-2) GO TO 90
T=INV(T)
35 IF (J.LE.0) GO TO 36
T=PFA(0,T)
J=J-3
GO TO 35
36 CALL DECAP(D,T)
E=LSHIFT(E,D,3)
IF (T.NE.0) GO TO 36
J=B
X=PFA(E,X)
40 IF (C.EQ.41) GO TO 31
GO TO 10
80 LREAD=X
RETURN
90 CALL ERLA(T)
CALL ERASE(X)
CALL ERASE(Z)
LREAD=-2
RETURN
END

SUBROUTINE LWRITE(U,Y)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER NBPW,FIRST,TAIL,TYPE,PFA
INTEGER B,C,D,S,T,U,X,Y
S=0
X=Y
I=1
B=NBPW(T)
10 C=40
GO TO 50
11 IF (X.NE.0) GO TO 12
20 C=41
GO TO 50
13 IF (S.EQ.0) GO TO 60
CALL DECAP(X,S)
14 X=TAIL(X)
15 IF (X.EQ.0) GO TO 20
40 C=42
GO TO 50
12 T=FIRST(X)
15 IF (TYPE(X).EQ.0) GO TO 30
S=PFA(X,S)
X=T
GO TO 10
30 L=0
DO 31 J=1,B+3
CALL RSHIFT(T,D,3)
L=PFA(D*L)
CALL DECAP(C*L)
IF (C*NE.0) D=C
IF (L*EQ.0) GO TO 50
IF (D*NE.0) GO TO 50
GO TO 32
IF (L*NE.0) GO TO 32
GO TO 14
IF (I*LE.72) GO TO 51
CALL WRITE(U,RECORD)
I=1
RECORD(I)=C
I=I+1
IF (C*LT.8) GO TO 33
IF (C*EQ.42) GO TO 12
IF (C*EQ.41) GO TO 13
GO TO 11
IF (I*EQ.73) GO TO 62
DO 61 J=I,72
RECORD(J)=44
61 CALL WRITE(U,RECORD)
RETURN
END

SUBROUTINE NOAVLS
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
PRINT 1
STOP
1 FORMAT (23H OUT OF AVAILABLE SPACE)
END

INTEGER FUNCTION PFA(ATOM,LIST)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER ATOM,TAIL
PFA=AVAIL
IF (PFA*EQ.0) CALL NOAVLS
AVAIL=TAIL(AVAIL)
CALL SSUCCLIST,PFA)
CALL STYPE(0,PFA)
CALL SCOUNT(1,PFA)
CALL ALTER(ATOM,PFA)
RETURN
END

INTEGER FUNCTION PFL(LIST1,LIST)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER TAIL
PFL=AVAIL
IF (PFL*EQ.0) CALL NOAVLS
AVAIL=TAIL(AVAIL)
CALL SSUCCLIST,PFL)
CALL STYPE(1,PFL)
CALL SCOUNT(1,PFL)
CALL ALTER(LIST1,PFL)
RETURN
END

SUBROUTINE STACK2(X,Y)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER X,Y,PFA
STAK=PFA(Y,PFA(X,STAK))
RETURN
END

SUBROUTINE STACK3(X,Y,Z)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER X,Y,Z,PFA
STAK=PFA(Z,PFA(Y,PFA(X,STAK)))
RETURN
END

SUBROUTINE UNSTK2(X,Y)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER X,Y
CALL DECAP(Y,STAK)
CALL DECAP(X,STAK)
RETURN
END

SUBROUTINE UNSTK3(X,Y,Z)
COMMON /TR1/AVAIL,STAK,RECORD(72)
INTEGER AVAIL,STAK,RECORD
INTEGER X,Y,Z
CALL DECAP(Z,STAK)
CALL DECAP(Y,STAK)
CALL DECAP(X,STAK)
RETURN
END