Genetic algorithms in Forth

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Abstract

A method for automatically finding a program (bytecode) realizing the given algorithm is developed. The algorithm is specified as a set of tests \( \text{input} \rightarrow \text{output} \). Genetic methods made it possible to find the implementation of relatively complex algorithms: sorting, decimal digits, GCD, LCM, factorial, prime divisors, binomial coefficients, and others. The algorithms are implemented on a highly simplified version of Forth language.

Keywords: Genetic algorithm, Linear genetic programming, Evolutionary programming, Forth.

1 Introduction

We want to automatically receive a program in some programming language that implements an algorithm, defined as a set of tests. To construct such an algorithm, we will use methods that in some sense have analogies in biology, in genetics. This approach is called ”genetic programming” [2, 3, 10].

These ideas first appeared in the 60s of the last century [10], but remains quite popular until today. At least at the site scholar.google.com search for the words ”genetic algorithm” for 2017/18 year gives more than 46 000 links.

For example, the site [2] is devoted to genetic programming, it describes the results obtained.

There exist a special program for finding algorithms using the genetic methods [6], but examples of algorithms found with its help are known only to the simplest ones (function \( x \rightarrow x^2 + x + 1 \)).

The work [3] describes the results achieved on this way. Authors managed to find programs that implement the following functions:

- \( \text{nguen1}: x \rightarrow x^3 + x^2 + x \),
- \( \text{nguen2}: x \rightarrow x^4 + x^3 + x^2 + x \),
- \( \text{nguen3}: x \rightarrow x^5 + x^4 + x^3 + x^2 + x \),
- \( \text{nguen4}: x \rightarrow x^6 + x^5 + x^4 + x^3 + x^2 + x \),
- \( \text{keijzer4}: x \rightarrow x^3 e^{-x} \cos(x) \sin(x)(\sin^2(x) \cos(x) - 1) \),

and several other functions of this kind.

In the book [5] methods of genetic programming in Python are considered. Here, more than a dozen different problems are solved by genetic methods. However, the main difficulty in them is not the genetic algorithms themselves, but the suitable reformulation of the problem. After that, the problem can be solved by the genetic method, or directly.

The widely distributed machine learning package ”TensorFlow” [7] also contains a tools for genetic programming. You can read about the results achieved with it, for example, in [4].

In fact, in almost cases, the considered ”genetic” methods are reduced to the search for a minimum of some loss function from some, possibly, quite a considerable number of parameters (several dozen).

In any case, this approach takes us far enough from the initial idea of a genetic approach. In this paper we will try to stay within the framework of the originally formulated task.
2 Problem definition

We want to find a program in some programming language implementing given algorithm. The algorithm will be defined as a set of tests: \((\text{input}_\text{data}) \rightarrow (\text{output}_\text{data})\).

The unit of the test is the pair \(t = (x, y)\), where \((x \in X, y \in Y)\), and \(X, Y\) are the corresponding sets of input and output objects. In the future, we will consider as input and output objects arbitrary finite sequences of integers, more precisely, of type int, 32-bit signed numbers.

2.1 Test files

A test will be called an arbitrary finite set of test items \(T = (t_0, \ldots, t_{n-1})\).

Sometimes, when there are no confusion, we will call a single test element as a test.

In practice, the test is represented by a text file of the form:

```
#T SHIFTR_0 x,y -> x*2^y comment
<in> 9 3 </in><out> 1536 </out>
<in> 4 3 </in><out> 48 </out>
<in> 2 7 </in><out> 28 </out>
<in> 3 3 </in><out> 24 </out>
<in> 6 5 </in><out> 320 </out>
<in> 2 5 </in><out> 20 </out>
<in> 7 5 </in><out> 640 </out>
...
```

The first two characters \#T are a signature of the test file, the next word SHIFTR_0 is the recommended name of the program you are looking for, the rest of the line is a comment.

In each of the following lines, a set of input and output integers parameters of the desired algorithm is defined.

We want to find a program that from each set of input data generates the required output set.

**Definition 1.** Pair \((m, n)\), where \(m\) is a number of input integers and \(n\) is a number of output integers will be called the signature of the test element.

**Remark 1.** In general, it is allowed to have test elements with various signatures in one test. But this possibility was not used in current work.

The number of test items in each test is from several units to hundreds.

A large set of test files is prepared for testing the developed methods, they are available on the site [I]. Not for all of them was found implementing programs. Tests are collected by sections:

- Polynomial;
- Max/min and sorting;
- GCD, LCM, factorials;
- Number theory;
- Decimal and binary digits;
- Power;
- \(\text{sqrt, log, ...}\);
• Interval testing;
• Rest;

Some tests have multiple version: complete and simplified. For example, for GCD (GCD.tst) there is a test GCD.0.tst containing only positive input parameters. Such simplified tests are required in the case when the complete test can not be implemented immediately.

3 Programming language

Programs from our point of view are of two types: linear and structural. The advantages and disadvantages of each approach are quite obvious and there is no fundamental difference between them from our point of view. We have chosen a linear approach in our work, at least for the initial stage.

Usually, this is done by selecting an assembly-type instruction set (registers). But we decided that the stack approach, actually implemented in the language of Forth [8, 9], would be more effective.

3.1 Forth

To find the program with the genetic method, the most stripped-down version of the Forth language was chosen. Only those instructions are left, which are impossible to do without.

There is no interaction with the user, such as output to the screen, even variables are missing. The control structures are represented only by unconditional and conditional jump-instruction.

Language Forth is very compact, new words (functions) are introduced very easily. In Forth, the newly defined functions have exactly the same syntax as the built-in language elements. This is convenient for a genetic approach. Thanks to this, you do not need to go through programs with complex structure in the form of a tree, only the simplest ones.

An example of a program that calculates the sum of the squares of two numbers and the factorial:

: SUMSQ2 DUP * SWAP DUP * + ;
: FACTORIAL CONST 1 OVER -- -ROT * OVER IF -6 SWAP DROP ;

In this paper, we will call the language a Forth for short, although we must remember that this is a strongly truncated version of the language.

3.2 Structure of the program

All the programs and functions will work with the data stack. Only 4-byte integers are stored on the stack. The functions in the Forth have no arguments. The initial data they take from the stack and there they leave the results of their work.

We call the signature of a function a pair of nonnegative numbers \((p, q)\), which show the number of input and output arguments of the function in the stack.

The number of arguments can vary, depending on the state of the stack. Such a signature will be called ”floating”. For now, we will only consider functions with a fixed, static signature. At each point of such a program, the current stack depth is statically determined.

Later, functions will be called ”words”, or ”genes”. At each moment of time, the system has a certain set of built-in words (functions) and the current set of new genes, that is, new words built in the learning process (”evolution”). One step of evolution is the construction of one or more new words (functions) that solve one problem (test) in whole or in part.
3.3 Bytecode

The program in the Forth is a byte sequence, each byte is a separate command. There are two types of commands: built-in and implemented on the Forth itself. Thus, the total number of commands can not be more than 255.

In our version of Forth there are 33 built-in commands:

0: A sign of the end of a function or program. There should be no other zero bytes in the function (almost). Thus, the function can be considered as a NULL-terminated string.

1: Unconditional jump to the specified address. The command consists of 2 bytes: the command code (1) and the next byte indicating the relative address of the jump (−128...127). The functions considered in the system will be short, so the restriction on the amount of displacement is not essential. For example, in the code

\[ \text{OVER} + \text{GOTO} \ 2 \ \text{DUP} \ - \text{SWAP} \]

the command ”GOTO 2” means jump to the command ”-”, and in the code

\[ \text{ROT} \ \text{OVER} + \text{GOTO} \ -3 \ \text{DUP} \ - \text{SWAP} \]

command ”GOTO -3” means jump to the command ”OVER”.

The transition with 0 offset does not make sense, so the second byte of the command is always non-zero.

2: Conditional branch to the specified address. The command consists of 2 bytes: the command code (2) and the next byte indicating the relative jump address (−128...127). The second byte of the command is again always nonzero.

3: Numeric literal: put a number from interval −128...127 on the stack. The command consists of 2 bytes: code (3) and byte-value. If we do not want to use the command ”CONST 0” containing zero byte, we can add the word ”ZERO” to the dictionary, in this case the second byte of the command is always nonzero and the program will not contain zeroes, except for the final zero, treated as ”NULL-terminated string”.

The three commands described are two bytes, all others consist of one byte.

Stack manipulation commands.

| Command | Stack before | Stack after |
|---------|--------------|-------------|
| DUP     | a            | a a         |
| DROP    | a            |             |
| SWAP    | a b          | b a         |
| OVER    | a b          | a b a       |
| ROT     | a b c        | b c a       |
| −ROT    | a b c        | c a b       |

In addition, to manipulate the stack, there are two commands ”PICK” and ”ROLL”:

”PICK”: get the \(n\)-th number, where \(n\) is the top of the stack, for example: ”0 PICK” is equivalent to ”DUP”, ”1 PICK” is equivalent to ”OVER”.

”ROLL”: rotate \(n\) numbers, for example: ”1 ROLL” is equivalent to ”SWAP”, ”2 ROLL” is equivalent to ”ROT”.

These functions are not static: the required depth of the stack depends on the value of its vertex. In order to use only commands with a static signature, the following five static versions are added:

| Command | Stack before | Stack after |
|---------|--------------|-------------|
| 2PICK   | a b c        | a b c a     |
| 3PICK   | a b c d      | a b c d a   |
Arithmetic and bit commands take their arguments on the stack and leave the result there.

Arithmetic: \texttt{NEGATE} \ \ + \ \ - \ \ \ * \ \ / \ \ \ % \ \ \ /\% \ \ \ ++ \ \ --

Bit commands: \texttt{AND} \ \ \ \ OR \ \ \ \ XOR \ \ \ \ NOT

Logical (comparison): \texttt{>} \ \ < \ \ = \ \ 0 = \ \ > 0 \ \ > 0.

In addition to the built-in commands, the system may contain commands implemented on the Forth itself, for example, finding the sum of squares of two elements of the stack:

\begin{verbatim}
: SUMSQ2  DUP * SWAP DUP * + ;
\end{verbatim}

The colon at the beginning of the command is a sign of the beginning of a new word, after it the name of the word goes, which will be used in the future in the system. The semicolon at the end is a sign of the end of the word (function, program).

The word name can be an arbitrary string of characters that does not contain spaces, tabs to the end of the line. Individual words are separated by spaces or tabs or line ends.

An important advantage of Forth is that it is not required to develop a mechanism for "embedding genes", it already exists in the language. New words (procedures, functions, "genes") are used in exactly the same way as built-in ones.

Thus, the current list of words (the dictionary) under biological interpretation will be considered as a "genome".

4. The first results

Just try to go through all of programs are not too long.

When examining all programs, we immediately discard programs containing "forbidden" pairs of commands, such as \texttt{(DUP DROP)} or \texttt{(SWAP SWAP)}. It should be noted that if the second command of the forbidden pair has a jump, then such a pair must be resolved.

Even with such a simple approach, it is possible to find the implementation of some algorithms:

\begin{verbatim}
Squaring : SQUARE DUP * ;
Multiply by 2 : MUL2 DUP + ;
Test odd : ODD CONST 1 AND ;
Signum : SIGN DUP 0= IF 6 0> DUP IF 2 -- ;
Sum of squares : SUMSQ2 DUP * SWAP DUP * + ;
Absolute value : ABS DUP 0> IF 2 NEG ;
Descending sort of 2 numbers : SORT2R OVER OVER > IF 2 SWAP ;
Ascending sort of 2 numbers : SORT2 DUP 2PICK > IF 2 SWAP ;
Maximum of 2 numbers : MAX2 OVER OVER > IF 2 SWAP DROP ;
Minimum of 2 numbers : MIN2 DUP 0> IF 2 SWAP IF 2 ROT ;
Maximum of 3 numbers : MAX3 OVER 3ROLL > IF 2 SWAP DROP ;
Minimum of 3 numbers : MIN3 ROT 2PICK 0> IF -4 DROP DROP ;
\end{verbatim}

4.1 Polynomials

General polynomials of degrees 1 and 2 are easily:

\begin{verbatim}
: poly1 * + ; \quad (a,b,x) \rightarrow b\times x + a
: poly2 -ROT 2PICK * + * + ; \quad (a,b,c,x) \rightarrow c\times x^2 + b\times x + a
\end{verbatim}
Specific polynomials

: pol1_1 -- DUP + -- ; x -> 2x-3, unexpected decision
: pol1_2 -- CONST -3 * ++ ; x -> -3x+4, also surprised!
: pol1_3 -- ; x -> x-1
: pol1_4 DUP + -- ; x -> 2x-1
: pol1_5 CONST 10 * -- -- -- ; x -> 10x-3, Cost without "3"!
: pol1_6 CONST 11 * CONST 7 + ; x -> 11x + 7
: pol2_1 DUP OVER + * ; x -> 2x*x
: pol2_2 DUP CONST -3 * ++ * ; x -> -3x*x + x
: pol2_3 -- DUP * ; x -> x*x -2x + 1

Polynomials with division:

: pol2d_1 DUP ++ * CONST 3 -- / -- ; Quadratic: x->(x^2 + x -2)/2

Working time. Having 33 basic words, we get $\approx 10^9$ programs of length 6. If $\approx 10$ million programs per second are processed (one core, frequency $\approx 3$ GHz), then a full search will take about a minute and a half.

In an hour, you can go through all the programs of length 7.
The search of all programs of length 8 requires more than a day.
For an 8-core processor, it is realistic to check programs of length 9.
But this is not genetics and not biology. It is chemistry, and inorganic.
One can increase the length of the programs by reducing the list of basic words. However, at this stage it is not easy to do it automatically.

5 Probabilistic and Markov approaches

5.1 Partial programs

In the simplest cases, as a result of a full search, we find a program that performs all the specified tests. In more complicated case to find solution only by a full search is impossible. However, we obtain a list of the programs that perform part of the tests, for example, the "factorial" of:

3 : DUP CONST 3 * + ;
4 : DUP * CONST 5 -- / -- ;
4 : NEG DUP * -- CONST 5 / ;
3 : DUP * DUP + ;
...

The number at the beginning of the line indicates the number of tests the program performs. Such programs will be called "partial programs", their list is the main result of a complete search of programs of small length.
All partial programs need not be memorized: a program that performs a single test is hardly of interest. In our implementation, we set the maximum number of partial programs ($N_p = 400$ by default) and remember only $N_p$ of the best, that is, those that perform the largest number of tests, are stored.
In addition, a histogram is calculated showing how many times each test was performed:
If after a full search in the list of partial programs some basic words are not found at all or very rare, you can restrict the list of basic words and repeat the full search with a limited set of valid words to programs of longer length.

5.2 Frequency table

Having a list of partial programs, we will construct a list of frequencies for the appearance of built-in words and the frequencies of the appearance of word pairs.

Remark 2. In our algorithm, words and pairs that have never met, still receive a small probability of $(1/1024)$.

Based on the frequency table, two methods of generating programs are implemented:

- Probabilistic, when the probability of occurrence of the next word in programs is taken from the frequency table;
- Markov, when the probability of the next word depends on the previous one.

The Markov approach should be more efficient, but usually the frequency table is not large enough to more or less accurately determine the frequencies of all pairs. Therefore, we propose a combination of probabilistic and Markov approaches.

The following is an obvious statement.

Proposition 1. Suppose you have a program $P$, consisting of the commands $P = c_1, \ldots, c_k$, the probability of the appearance of $c_i$ a is $p_i$. Denote by $\varepsilon$ the number $\varepsilon = p_1 \ldots p_k$, that is, the probability of the program $P$ and let $N = \text{round}(1/\varepsilon)$. Then the probability of the appearance of the program $P$ for $N$ tests is $\approx 63\%$, for $3N$ tests is $\approx 95\%$, for $5N$ tests is $\approx 99.3\%$.

5.3 Base step

Base step with parameters $(L_0, L_1, T)$ in a fixed dictionary will be called the following algorithm.

1. Performed a full search of programs up to length $\leq L_0$. Build a list of partial programs.
2. Using the partial programs, we build a list of word frequencies and a list frequencies of word pairs.
3. Generate within a specified interval of time $T$ seconds of a random program of length $L_0 + 1 \ldots L_1$, consistently, first a simple, then a Markov method.
4. We correct the frequency tables and repeat step 3, this cycle is performed 8 times.

The typical values of the parameters for working on one processor core is $(7, 14, 400)$, that is, first perform a full search of programs of length $\leq 7$, then 8 times for 50 seconds is performed alternately probabilistic/Markov search of programs of length from 8 to 14 with correction of the frequency table after each cycle.
5.4 Example: discriminant

The method of probabilistic search can find the implementation of many algorithms. Consider, for example, the discriminant: \((a, b, c) \rightarrow b^2 - 4ac\). Full search of the result was not given, but allowed to reduce the number of basic words (the size of the "genome")

\[
\begin{align*}
\text{CONST} & \ \text{DUP} \ \text{DROP} \ \text{SWAP} \ \text{OVER} \ \text{ROT} \ -\text{ROT} \ \text{NEG} \ + \ - \ * \ + \ + \ -- \\
\end{align*}
\]

On the basis of the constructed frequency table, a random search for programs of length 9 was started. Quite quickly the solution was found:

\[
: \text{discr} \ \text{OVER} \ \text{ROT} \ * \ -\text{ROT} \ * \ \text{CONST} \ 4 \ * \ - \\
\]

5.5 Arbitrary (random) functions

Let \(N\) be a natural number, denote by \(X\) the set of \(N\) elements \(X = \{0, \ldots, N - 1\}\). We consider all possible functions \(f: X \rightarrow X\). Each such function is fully described by a set of \(N\) tests:

\[
\begin{align*}
0 & \rightarrow f(0) \\
1 & \rightarrow f(1) \\
\vdots & \\
N - 1 & \rightarrow f(N - 1)
\end{align*}
\]

The number of different functions is \(N^N\).

\[
\begin{array}{c|c}
N & N^N \\
\hline
3 & 27 \\
4 & 256 \\
5 & 3125 \\
6 & 46656 \\
\end{array}
\]

At \(N = 3\), all 27 functions have an implementation of the length \(\leq 5\), which is found by a full search, for example (the numbers at the beginning of the string mean the values \(f(0), f(1), f(2)\), respectively):

\[
\begin{align*}
0 \ 2 \ 1 : & \ 
\text{DUP} \ 0> \ + \ + \ \text{OR} \ -- \ -- \\
1 \ 0 \ 2 : & \ 
\text{DUP} \ 0= \ + \ + \ \text{OR} \ -- \ -- \\
2 \ 0 \ 1 : & \ 
-- \ \text{DUP} \ 0< \ + \ + \ \text{AND} \ -- \\
2 \ 2 \ 0 : & \ 
\text{CONST} \ -3 \ \text{OR} \ \text{NEG} \ -- \ -- \\
\end{align*}
\]

At \(N = 4\) of 256 functions, about 200 have an implementation of the length \(\leq 6\). For all the checked functions of the remaining, using probabilistic methods, an implementation of lengths from 7 to 12 was found, for example:

\[
\begin{align*}
0 \ 3 \ 1 \ 0 : & \ 
\text{DUP} \ -- \ -- \ 0= \ 0= \ \text{SWAP} \ -- \ 0= \ + \ + \ \text{XOR} \ ; \quad (\text{len}=10) \\
0 \ 3 \ 1 \ 1 : & \ 
\text{DUP} \ -- \ 0= \ \text{DUP} \ + \ + \ \text{ROT} \ 0= \ -- \ + \ ; \\
3 \ 2 \ 0 \ 3 : & \ 
-- \ \text{DUP} \ -- \ 0= \ 0= \ + \ + \ \text{SWAP} \ 0= \ 0= \ \text{XOR} \ ; \\
3 \ 2 \ 3 \ 1 : & \ 
\text{DUP} \ \text{DUP} \ + \ + \ \text{OR} \ + \ + \ \text{XOR} \ + \ + \ \text{DUP} \ -- \ -- \ -- \ / ; (\text{len}=11) \\
3 \ 3 \ 2 \ 0 : & \ 
\text{DUP} \ 0= \ 0= \ + \ + \ / \ + \ + \ + \ + \ \text{XOR} \ ; (\text{len}=9) \\
3 \ 3 \ 3 \ 0 : & \ 
\text{DUP} \ -- \ \text{AND} \ \text{DUP} \ + \ + \ \text{DUP} \ \text{OR} \ -- \ 0= \ + \ + \ \text{XOR} \ ; (\text{len}=12) \\
\end{align*}
\]

However, here there is no guarantee that the length is minimal. For example, the last program can be replaced with

\[
: \ -- \ -- \ -- \ 0< \ \text{CONST} \ 3 \ * \ ; \quad (\text{len}=7)
\]
5.6 Right shift

Consider the function \((n, x) \rightarrow 2^n \cdot x\). A test file containing 13 test items was taken. Full search was unsuccessful, but allowed to reduce the list of basic words ("genome") to:

```
IF CONST DUP DROP SWAP OVER ROT -ROT NEG + - -- ++ *
```

After that, probabilistic search received as many as 5(!) programs that perform 9 tests (of 13). They were very similar:

9: `DUP + SWAP -- -ROT IF -7 + ;`
9: `DUP ROT -- -ROT + OVER IF -7 + ;`
9: `SWAP -- OVER ROT + OVER IF -7 + ;`
9: `SWAP -- SWAP DUP + OVER IF -7 + ;`
9: `DUP + SWAP -- SWAP OVER IF -7 + ;`

And what tests are not performed? It turns out that the simplest, when \(n = 0\):

```
<in> 0 3 </in><out> 3 </out>
<in> 0 5 </in><out> 5 </out>
<in> 0 5 </in><out> 5 </out>
<in> 0 7 </in><out> 7 </out>
```

Here you have a ready gene!

6 Genetics

In the biological interpretation of our results, the probabilistic and Markov approach can be considered as "organic chemistry". Let's take one more step and move on to "genetics".

As already mentioned, the list of words in the dictionary plays the role of "genome". Each gene added can be either "good", "useful", or "unsuccessful", "useless". The quality of the function-gene itself cannot be determined, only as part of the genome, aimed at solving a specific problem. Therefore, we need to determine the quality of the genome (dictionary) in relation to this set of tests. If the new gene improves the quality, we will consider it a good one.

Since our goal is to find a program that passes all the tests, the quality it would naturally to consider the largest number of tests that one of the programs passed. When this value is equal to the total number of tests, the answer is found!

However, in practice this definition of quality is insufficient. Often even a obviously successful gene does not increase the maximum number of tests passed: it was 3 and there remained 3.

The "quality" of the genome will be the average number of tests performed for one million randomly generated programs.

The new gene will be called "successful" if the maximum number of tests performed has grown or remained the same, but the quality of the genome has increased by at least 1 percent.

As the "candidates for genes" we will take the most frequent chains of bytes of length 2 or 3 among partial programs.

6.1 Selecting a list of admissible words

The same idea can be used to select a list of admissible words.

1. First, we take the minimum possible list of basic words ((11 words) as admissible.

2. In turn, we add one system word to this list.

3. Perform the basic step, find the quality of the genome (dictionary).

4. If the gene is successful, leave it in the list, otherwise delete it.
6.2 General algorithm

During the process we will have two lists of programs (genes): "candidates for genes" and "unsuccessful genes". There should be no duplicate elements in them.

1. Find a list of valid words. Upon completion of the search, we get, besides this list, also a list of partial programs.
2. Clear the list "candidates for the genes" and "bad genes".
3. In the lists of "candidates for genes" add the most frequent chains of length 2, 3, and only those that are not in the list of "unsuccessful genes".
4. If there are no candidates for genes, we finish the job.
5. Add one of the candidates to the dictionary.
6. Perform a basic step with this genome, find its quality.
7. If the gene was unsuccessful, remove it from the dictionary and add it to the list of "unsuccessful genes".
8. Goto item (3).

7 Results

Let’s describe the results obtained with the help of the developed program.

7.1 Program description

The program gene.exe along with the detailed description is available on the site [1]. Its main features:

- setting a list of basic and additional words;
- full search of programs up to length \( L \) inclusive;
- full search of programs within no more than \( T \) seconds;
- search probabilistic for \( T \) seconds;
- Markov search for \( T \) seconds;
- full search to the length \( L \) and the probabilistic/Markov search to the length \( L_1 \) within \( T \) seconds;
- search for valid words.

7.2 Digits

The lowest digits were successfully found by direct search:

```
: digit0   CONST 10 % ;
: digit1   CONST 10 / CONST 10 % ;
: digit2   CONST 10 DUP * / CONST 10 % ;
```

To find the remaining algorithms, a list of basic words was found in which the constant 10 was added:
The 3rd digit was found by probabilistic search. Here is the result (of length 8):

```
: digit3 C10 / C10 / C10 / C10 % ;
```

Similarly, the highest digit was found (program of length 9):

```
: digitH DUP C10 / DUP -ROT IF -5 SWAP DROP ;
```

But the k-th figure in this way could not be found. To find the program, a genetic approach has already been needed. During the search, the genes were checked:

```
: F_036 DROP C10 ;
: F_036 SWAP DROP ;
: F_036 % SWAP ;
: F_037 C10 % ;
: F_038 DROP C10 % ;
: F_039 SWAP DROP C10 ;
: F_039 % SWAP DROP ;
: F_039 C10 % SWAP ;
: F_039 C10 F_036 ;
: F_039 F_036 DROP ;
: F_039 C10 F_036 DROP ;
: F_039 C10 F_036 0= ;
```

"Useful" of them were recognized:

```
: F_036 % SWAP ;
: F_037 C10 % ;
: F_038 DROP C10 % ;
: F_039 C10 F_036 0= ;
```

Finally, the answer of length 12 was found:

```
: digitK DUP IF 4 SWAP C10 / OVER -- ROT IF -7 F_038 ;
```

As you can see, from the "useful" genes it was used only one F_038. If "open" the gene, we get the program of length 14:

```
: digitK DUP IF 4 SWAP C10 / OVER -- ROT IF -7 DROP C10 % ;
```

7.3 Factorial

For the function "factorial", \( n \rightarrow n! \), take a test file consisting of 13 tests for \( n \leq 13 \). We start the search for a set of admissible words and get it:

```
GOTO IF CONST DUP DROP SWAP OVER ROT -ROT + - * = --
```

After that we try to build the required program.

Unfortunately, it is not possible to do this directly. For better understanding, let’s throw out the test line \( 1! = 1 \). On the remaining set, start the search for the program by genetic method.

Because the process is probabilistic, different launches will produce different results.

At the first start, the word appeared as a gene

```
: F_035 DUP DUP ;
```
and already at the next step was found the program that performs all the tests:

```plaintext
: FACTORIAL_1 F_035 IF 5 DUP ROT * SWAP -- DUP IF -7 IF -12 ;
```

At the second start, the word appeared as a gene \( F_035 \), but the finished program could not be found at once. Two additional genes were first constructed:

```plaintext
: F_036 DUP -- ; : F_037 -- DUP ;
```

and only after this the result is obtained:

```plaintext
: FACTORIAL_1 F_036 DUP -ROT * SWAP F_037 IF -6 + ;
```

If you substitute intermediate genes, insert them in the final program, we get:

```plaintext
: FACTORIAL_1 DUP DUP IF 5 DUP ROT * SWAP -- DUP IF -7 IF -13 ;
```

in the first case (length 14)

```plaintext
: FACTORIAL_1 DUP -- DUP -ROT * SWAP -- DUP IF -7 + ;
```

in the second (length 11).

And the hand-made program:

```plaintext
: FACTORIAL_1 DUP -- SWAP OVER * SWAP -- DUP IF -7 DROP ; (len=11)
```

Add a deleted item to the tests:

```plaintext
<in> 1</in><out> 1 </out>
```

and an additional gene (FACTORIAL_0). After this, the desired program is already found:

```plaintext
: FACTORIAL_0 DUP -- 0= IF 2 FACTORIAL_1 ;
```

If we substitute genes, we get a result of length 16:

```plaintext
: FACTORIAL_0 DUP -- 0= IF 12 DUP -- SWAP OVER * SWAP -- DUP IF -7 DROP ;
```

### 8 Genetic-2

The results obtained above can be considered as bottom-up genetics, that is, the construction of small genes, from which the final program is built. You can consider another approach, when the genes taken as a whole program that performs some task. Unfortunately, this process has not been fully automated yet. Therefore, we will look at it in examples, when genes are constructed in a "semi-manual" mode.
8.1 GCD, LCM

Consider a test file GCD.tst of 180 elements:

```
#T GCD  x,y-> GCD(x,y)
<in>  4  4 </in><out> 4 </out>
<in>  5  5 </in><out> 5 </out>
<in>  0  9 </in><out> 9 </out>
<in>  7  0 </in><out> 9 </out>
<in>  2 10 </in><out> 2 </out>
<in>  4 10 </in><out> 2 </out>
<in>  4 12 </in><out> 4 </out>
<in> 12  5 </in><out> 1 </out>
<in> 12  3 </in><out> 3 </out>
...
<in>-33964 90856 </in><out> 4 </out>
<in> 38817 -90856 </in><out> 1 </out>
```

Brute force can not find a solution. The probabilistic method give a partial program:

```
: GCD_0 DUP -ROT % DUP IF -5 - ;
```

which gives the correct answer for all positive \((x,y)\). Adding it as a gene, we find a partial program:

```
: GCD_1 OVER ROT IF 3 GOTO -5 GCD_0 ;
```

which gives the correct answer for all non-negative \((x,y)\). Adding it as a gene and a abs-program:

```
: ABS DUP 0> IF 2 NEG ;
```

we fine the correct result:

```
: GCD ABS SWAP ABS GCD_1 ;
```

On the basis of the found genes can easily be able to find an algorithm for LCM:

```
: LCM ABS OVER OVER GCD / * ABS ;
```

In fact, this algorithm will give an error for the pair \((0.0)\). We will not pay attention to this.

8.2 What we want to get (Fibonacci)

On the example of Fibonacci numbers.

We will look for a program that calculates Fibonacci numbers. Now, even with the help of a genetic approach, this does not succeed. Let’s try to manipulate the genes ”manually” to understand how this should look in a more general situation.

We introduce an additional gene from the side, ”from space”. (see p. 9 in next section):

```
: fib1 -- -ROT SWAP OVER + ROT ; (x,y,k) -> (y, x+y, k-1)
```

With its help we find the program (length 12):

```
: Fibonacci0 ONE ONE ROT -- -- fib1 DUP IF -3 DROP SWAP DROP ;
```

It only works at \(n \geq 3\). The general program will be as follows:

```
: Fibonacci3 DUP CONST 2 > IF 4 0> GOTO 2 Fibonacci0 ;
```

After expanding get (length 27):

```
: Fibonacci DUP CONST 2 > IF 4 0> GOTO 19 CONST 1 DUP ROT
    -- -- -- -ROT SWAP OVER + ROT DUP IF -8 DROP SWAP DROP ;
```

Thus, we can implement very complex algorithms, although the automatic implementation of some steps looks rather problematic.
9 Conclusion and future work

Unfortunately, the developed methods do not allow us to find the implementation of many algorithms. Some areas for further research can be proposed now. Under this proposals, we can hope to make significant progress in our direction.

1. Preparation of program templates.
2. Ranking of test items.
3. Frequency of appearance of triples.
4. Initial filling of the frequency table.
5. Saturation check for probabilistic search of a given length.
6. Initial genes ”from space”.
7. Setting the algorithm for selecting candidates for genes.
8. Transition to multi-core processor.
9. Merge multiple test files into one and pre-process it.
10. Efficient generation of structural programs.
11. Genetic-2.

More detailed.

**Preparation of program templates.** During the search, most of the algorithms are discarded due to incorrect syntax, invalid signature, etc. It is suggested to compile a complete list of templates of syntactically correct programs with this signature in advance and then select only from this list.

**Ranking of test items.** Now, when building a list of partial programs, all the test items are equivalent. This is not entirely correct. One test is performed by very many programs, and the other by a few units. Therefore, for each item, we consider how many generated programs it is performed and in evaluating the quality of the following programs we give preference to those that perform more complex (rare) test items.

**Frequency of appearance of triples.** At present, probabilistic generation of programs takes into account only the probabilities of occurrence of a single word and the probability of the appearance of pairs. Practice shows that some triples have a rather high frequency of appearance, noticeably greater than many pairs. Therefore, for a probabilistic approach, such triples should be taken into account separately.

**Initial filling of the frequency table.** Now the frequency table is built entirely on the basis of a partial programs. List of these programs can be relatively small, so the frequencies of individual words and especially pairs are too rough. It makes sense to construct an initial filling of the frequency table of built-in words and their pairs on the basis of an analysis of a large number of real examples.

**Saturation check for probabilistic search of a given length.** At random/Markov program search of a short length (8, 9, 10) after a while the generated programs begin to repeat. If such a phenomenon is found, it should be assumed that programs of a given length have already been exhausted and this length is not further considered.

**Initial genes ”from space”.** When discussing the problem of the appearance of life on Earth, one of the hypotheses is the introduction of life from space. In our case, this means the appearance of genes from an external source.
More precisely, to the original 33-th built-in words add a few dozens or even hundreds of additional words. If you use the algorithm of finding admissible words (paragraph 6.1), the initial dictionary will not grow too much. Further work with such a dictionary can be much more effective.

In this case, most likely, it is worth combining the basic words in some groups with a fixed frequency ratio within the group. Each group of words should be included in the basic dictionary only in its entirety and determined only by the general probability of the occurrence of a word from the group. The ratio of the frequencies within the group should remain unchanged.

**Setting the algorithm for selecting candidates for genes.**

Now candidates for genes are several of the most frequent substrings of length 2 and 3 in the list of partial programs. In fact, it is necessary to select substrings of different length, up to 5 – 6, but taking into account the quality of each partial program.

**Transition to multi-core processor.** This is an extensive approach, but now (2018-07-01) there are 28 and 32-core processors. If not even consider supercomputers, such an increase in processing power can lead to a noticeable increase in results.

**Merge multiple test files into one and pre-process it.** Sometimes it turns out to be impossible to find a program for this test, genetic algorithms do not provide a sufficiently powerful genome. In this case, a more complex test can help, which includes several separate tests, but is related to a common theme. Then genes built with the implementation of one test will help to advance and implement another.

**Efficient generation of structural programs.** Now with the random generation of programs up to the actual execution of the program on tests reaches about one program out of thousands, the rest are discarded before that on various checks. Based on this, you can try to immediately generate only the correct program. This will greatly increase the complexity of the generation algorithm, but the potential gain in speed is worth it.

**Genetic-2**

Now we’re adding to the genome some of the frequently occurring sub-strings in partial programs, this approach can be considered as genetics “bottom-up”, we start with simple programs and build more complex ones from them. You can try to consider another approach, ”up-bottom”.

**Definition 2.** Let the partial program $P$ be performed on a certain subset $S'$ of all test items $S$. Let us denote a new test in $S''$, in which the input values coincide with the input values of the elements $S$, and the return value is only one and is equal to 0 or 1, and 1 on a subset of $S'$ and 0 on its addition.

**Definition 3.** A partial program $P$ is said to be good if:

- It performs a significant part of all tests;
- It is possible to find the program $Q_P$ realizing the test $S_P$.

Let $P$ be some good partial program for the $S$ test. We denote by $S''$ a test consisting of items on which $P$ NOT holds. Suppose that we managed to find the program $R$ realizing the test $S''$. Then the program that implements the original $S$ test looks like this:

: P2 Q IF 4 P GOTO 2 R ;

**9.1 Conclusion**

The proposed method of genetic construction of programs allows us to move quite far in this direction. Namely, we can implement programs for algorithms sorting of 2 and 3 elements, finding the maximum and minimum, many general and individual polynomials of one, two and three variables, binary shifts, decimal and binary digits, GCM, LCM, find a minimal prime divisor of a number and many others. However, not all algorithms can be implemented. Further development of the proposed methods should lead to a significant increase in the capabilities.
In addition to the theoretical interest, you can already see its practical application. For example, taking the assembly language of a certain processor instead of the Forth language, you can optimize individual sections of the program.

References

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