12-1-2010

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From Program Synthesis to Optimal Program Synthesis

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Abstract—In many practical situations, we know the values of some quantities \( x_1, \ldots, x_n \), we know the relations between these quantities, the desired quantity \( y \), and maybe some auxiliary quantities, and we want to estimate \( y \). There exist automatic tools for such estimations—a called program synthesis tools.

A program synthesis tool usually generates a program for computing \( y \). In many cases, however, several such programs are possible, and it is desirable to generate the optimal (e.g., the fastest) program. In this paper, we describe algorithms aimed at such optimal program synthesis.

The problem can be interpreted in logical terms, as assigning fuzzy-style degrees to rules describing relations between variables.

I. NEED FOR DATA PROCESSING: A BRIEF REMINDER

One of the main objectives of science: an informal description. One of the main objectives of science is to describe the world.

One of the main objectives of science: formulation in precise terms. In precise terms, the above objective means that we would like to know the values of the numerical characteristics that characterize different objects.

Sometimes, this task is easy. Some of these values, we can simply measure.

In other cases, the task is more complex. There are many physical characteristics which are difficult or even impossible to measure directly, for example,

- the distance to a faraway star, or
- the amount of oil in a well.

Since we cannot measure these difficult-to-measure values \( y \) directly, we thus measure them indirectly: namely, we measure the values of related easier-to-measure characteristics \( x_1, \ldots, x_n \), and we find (and list) all possible relations between these characteristics \( x_i \) and the desired quantity \( y \) (and maybe some auxiliary difficult-to-measure characteristics).

Based on these relations, we design an algorithm \( f(x_1, \ldots, x_n) \) that, given the values \( x_1, \ldots, x_n \), returns the (estimate for the desired quantity \( y = f(x_1, \ldots, x_n) \).

First example: computations are simple and straightforward. To find the distance to a star, we can use the parallax method: we observe the position of this star on the celestial sphere in different seasons, when the Earth is at different sides of the Sun, and then use trigonometry to determine the desired distance.

In this example, the algorithm \( f(x_1, \ldots, x_n) \) can be described by explicit formulas—and indeed, these computations were successfully done “by hand” already many centuries ago, when no computers were available.

Second example: many more computations are needed. To estimate the amount of oil in a well, we perform a large number of seismic experiments, by

- making small explosions at different locations on the Earth surface, and
- measuring the time that the corresponding seismic waves took to travel to the sensors placed at different other surface locations.

Based on these times, we:

- solve the partial differential equations describing how seismic waves propagate, and
- thus find the velocity of sound at different locations and at different depths.

We then estimate the amount of oil by computing the volume at which the velocity of sound is consistent with oil; see, e.g., [1], [2], [5], [9], [18].

The corresponding algorithm \( f(x_1, \ldots, x_n) \) includes solving partial differential equations as an intermediate step. It requires millions and billions of computation steps, the amount that is very difficult (and often impossible) to do by hand. Thus, we need computer-based data processing.

The large number of computations is one of the main reasons why computers were invented in the first place—to perform data processing and thus, to determine the values of difficult-to-measure quantities.

Prediction: another problem for which data processing is needed. Another important goal of science is to predict the future state of the world, i.e., in more precise terms, to predict the future values of the desired physical quantities. We want to predict the trajectory of an asteroid approaching Earth, we want to predict the path of a hurricane, etc.

In some cases, we have explicit formulas \( y = f(x_1, \ldots, x_n) \) that describe the desired future values in terms of the current
values $x_i$ of different quantities. However, in many other cases, we only know some relations between $y$, $x_i$, and some additional auxiliary characteristics – such as the future values of other physical characteristics. Our objective is then to use these relations to design an algorithm that transforms the known values $x_1, \ldots, x_n$ into the desired prediction.

**Engineering design problems.** In contrast to science where our main objective is to understand the world and to predict how it will change, in engineering, we want to design a new object with given characteristics; for discussion of the difference between science and engineering, see, e.g., [6], [11] and references therein.

Sometimes, there are ready algorithms for such design. In many other cases, however, we only know

- the relations between different characteristics, and
- the desired values of some of these characteristics.

We need to find the values of the set characteristics for which we will achieve the desired goals.

**Comment.** From the computational viewpoint, the difference between science and engineering is not as clear as it may seem. For example, when we talk about trajectories, then

- in science, a typical problem would be to predict a trajectory – e.g., to decide whether a given asteroid will hit the Moon,
- while in engineering, a typical problem may be to find the starting velocity and the starting direction for which the spaceship reaches the Moon.

However, from the computational viewpoint, we may encounter the exact same problem in science as well: e.g., when we know that a meteorite found in Antarctica originated on the Mars surface, it is then desirable to find the trajectory that led it to the Earth. Substance-wise, this is a typical scientific problem, because we are not talking about designing a new object or a new trajectory, but computationally, it is exactly the same problem as computing the trajectory of a spaceship that will go from Mars to Earth.

II. **Program Synthesis: Successes and Open Problems**

**A general problem.** In all the above situations:

- we know the values of some of these quantities $x_1, \ldots, x_n$, and
- we know the relations between these quantities and the desired quantity $y$ (and maybe some auxiliary quantities).

Our objective is to use the known values and the known relations to find the values of the desired quantities.

**How this problem is traditionally solved.** Traditionally, in science and engineering, researchers and practitioners use their own creativity to come up with an algorithm $f(x_1, \ldots, x_n)$ that

- inputs the values of the known quantities, and
- outcomes an estimate for the desired quantity $y$.

Designing such an algorithm is a very difficult task.

**The main idea of program synthesis.** It turns out that in many cases, we can have a system that automatically analyzes the relations and synthesizes the desired program. Such automatic systems are called systems for *program synthesis*; see, e.g., [19].

**Toy example.** Let us use a simple example of a triangle to illustrate how program synthesis works.

A triangle is described by its angles $A, B, C$ and side lengths $a, b, c$. We know the following relations between them:

- $A + B + C = \pi$ (the sum of the angles is $180^\circ$, or $\pi$ radians),
- $a^2 + b^2 - 2ab\cos(C) = c^2$ and similar expressions for $a$ and $b$ (cosine theorem), and
- $\frac{a}{\sin(A)} = \frac{b}{\sin(B)} = \frac{c}{\sin(C)}$ (sine theorem).

Now we can ask all kinds of questions:

- If we know $a, b$ and $c$, can we determine $A$? and if yes, how?
- If we know $a, b$ and $A$, how to compute $b$? and if yes, how?

**Preparing for the program synthesis.** First, we analyze which quantities are directly computable from which.

Suppose that we have a relation $F(A, B, C) = 0$. If we know all of these values but one (for example, $A$ and $B$), then we have an equation with one unknown, from which in general we can compute $C$. So, if we already know $A$ and $B$, then we are able to compute $C$. We will describe this implication, for short, as $A, B \rightarrow C$.

Similarly, if we know $A$ and $C$, then we can compute $B$, and from $B$ and $C$ we can compute $A$.

So each equation leads to as many computability relations as there are unknowns in it. In our case we get three computability relations: $A, B \rightarrow C$; $A, C \rightarrow B$; and $B, C \rightarrow A$.

**Example.** In the triangle case, the relations turn into the following formulas:

- $A, B \rightarrow C$; $B, C \rightarrow A$; $A, C \rightarrow B$; (these three come from the equation $A + B + C = \pi$)
- $A, a, b \rightarrow B$; $A, a, B \rightarrow B$; . . . (from sine theorem), and
- $a, b, C \rightarrow c$; $a, b, c \rightarrow C$; $a, c, C \rightarrow b$; $b, c, C \rightarrow a$; . . . (from cosine theorem).

**Wave algorithm for program synthesis.** Let us describe a natural algorithm for deciding whether the desired quantity $y$ can be computed based on the known quantities $x_i$.

According to this algorithm,

- We first mark the variables that we know.
- Then, when we look at all the rules, find those, for which all the conditions are marked and the conclusion is not, and mark the conclusion.
- Then we repeat the same procedure again and again.

After each iteration,

- either we did not add anything – which means that we are done (nothing else can be computed),
or we add at least one marked variable.

Since there are finitely many variables, this process will eventually stop:

- If the desired \( y \) is marked, then we can compute it,
- else we cannot compute \( y \).

Once we have a sequence of rules that lead to computing \( y \), we can combine the corresponding algorithms and come up with a program for computing \( y \) based on \( x_i \).

**Example.** Suppose that in the triangle, we know \( A \) and \( B \), and we want to compute \( C \) and \( a \).

Then, according to the algorithm, we first mark \( A \) and \( B \).

There is only one rule whose conditions are marked: the rule \( A, B \rightarrow C \). So, we mark \( C \).

On the second iteration, we find three rules whose conditions are marked: \( A, B \rightarrow C \); \( B, C \rightarrow A \); and \( B, C \rightarrow A \), but their conclusion have already been marked. So, we stop.

As a result, \( C \) is marked, which means that we can compute \( C \). Moreover, we know how to compute \( C \); \( C \) was obtained from a rule \( A, B \rightarrow C \) that stems from \( A + B + C = \pi \), so we must solve an equation \( A + B + C = \pi \), in which \( A \) and \( B \) are known, and \( C \) is the only unknown.

As for \( a \), it is not marked, and therefore, cannot be computed.

**Comment.** It should be mentioned that while the above description captures the main idea of program synthesis, the actual program synthesis algorithms implemented, e.g., [19], are more complex.

**Boolean logic interpretation of the wave algorithm.** The program synthesis problem can be reformulated in logical terms. Namely, we can interpret each rule \( A, B \rightarrow C \) that stems from the relations as a propositional formula \( A \& B \rightarrow C \) with variables \( A, B, \ldots \) that can take the values “true” or “false”:

- “true” means that we can compute the corresponding variable, and
- “false” means that we cannot.

So our knowledge can be represented as a set of propositional formulas that include all the rules and all the atoms \( A \) that represent the known variables \( x_i \).

We want to know whether the value \( y \) is computable, or, in the propositional terms, whether the variable that corresponds to \( y \) is true. So, in logical terms, we want to know whether these variables are deducible from the knowledge base.

We can therefore use logical deduction tools to check whether \( y \) is deducible from \( x_i \); see, e.g., [12], [13], [14], [15], [16], [17], [19], [20].

**Triangle example.** In the triangle example, we have a knowledge base \( A \& B \rightarrow C \); \( B \& A \rightarrow C \); \( A \& B \); and we want to know whether \( C \) and \( a \) follow from these formulas.

**An example of a practical application.** The logical deduction tools have been used to automate program synthesis in space missions such as the NASA Cassini mission to Saturn; see, e.g., [12], [13], [14], [15], [16], [17], [19], [20].

**Limitations of the Boolean logic approach.** There exist cases in which this logical approach does not work. Let us consider the case when we want to know the values of two unknowns \( y_1 \) and \( y_2 \), and we know two relations between them: \( y_1 + y_2 - 1 = 0 \) and \( y_1 - y_2 - 2 = 0 \). In this case, we can determine both \( y_1 \) and \( y_2 \), because we have a system of two linear equations with two unknowns.

However, in this example, the above logical approach will not work; indeed:

- The first equation will translate into two rules \( Y_1 \rightarrow Y_2 \), \( Y_2 \rightarrow Y_1 \), where propositional variables \( Y_i \) correspond to \( y_i \).
- The second equation will lead to these same rules.

From these two formulas we cannot logically conclude that \( Y_1 \) is true (because if \( Y_1 \) are both false, still both rules are true), and therefore, we cannot conclude that \( y_1 \) are computable.

**Possibility to use fuzzy-type logic.** In [3], it was shown that such examples can be handled if we replace the original Boolean logic with a more complex fuzzy-type logic.

**Remaining problem.** The existing program synthesis algorithms produce a program for computing \( y \). In many cases, there are several possible ways of computing \( y \).

In the triangle example, if we know \( A, B, a, b, \) and \( c \), and we want to compute \( C \), then we have several alternatives:

- we can use the rule \( A, B \rightarrow C \) corresponding to the relation \( A + B + C = \pi \), and find \( C \) as \( \pi - A - B \);
- alternatively, we can use the sine rule \( A, a, c \rightarrow C \) corresponding to the relation \( c \cdot \sin(A) = \frac{a \cdot \sin(C)}{\sin(C)} \), and compute \( C \) as \( C = \arcsin \left( \frac{c \cdot \sin(A)}{a} \right) \);
- we can use even more complex formulas related to the cosine rule.

In such situations, it is desirable to come up with the fastest program – or, more generally, a program that spends the smallest amount of resources.

This may include the resources needed to measure the input characteristics \( x_i \).

In other words, instead of program synthesis, we need optimal program synthesis.

**What we do in this paper.** In this paper, we interpret the problem in logical terms, as assigning fuzzy-style degrees to rules describing relations between variables. As a result, we develop algorithms aimed at optimal program synthesis.

**III. TOWARDS ALGORITHMS FOR OPTIMAL PROGRAM SYNTHESIS**

**Formulation of the problem: what is known.** We assume that for each known variable \( a \), we know the corresponding weight \( w(a) \) – describing the amount of resources that are needed to measure the value \( a \).

For example, if the main resource in which we are interested is time, then \( w(a) \) is the time needed to perform the corresponding measurement. This amount of time is known from the description of the corresponding measuring instrument.
If the main resource of interest is energy (as in the space mission to distant planets), then \( w(a) \) is the amount of energy that is needed to perform this measurement. This amount can also be extracted from the description of the measuring instrument.

We also assume that for each rule \( r \) of the type \( a, b \rightarrow c \), we know the corresponding weight \( w(r) \) — describing the amount of resources that are needed to compute \( c \) based on the known values \( a \) and \( b \).

We assume that for each rule of the type \( a, b \rightarrow c \), we already have a well pre-tested program that, given \( a \) and \( b \), computes \( c \). If our main resource is time, then \( w(r) \) is the average computation time needs for this program to work; this average amount can be easily computed based on the known results of pre-testing the program. Similarly, when our main resource is energy, we can determine the average energy \( w(r) \) needed to run this program based on the known results of pre-testing this program.

We assume that the amount of resources needed to perform a sequence of measurements and computations is equal to the sum of the corresponding weights.

**Formulation of the problem: what we want.** Our objective is to select, among all paths that lead to the desired quantity \( y \), the path with the shortest overall weight.

**We can simplify the description by only assigning weights to edges.** In the standard program synthesis, a variable that can be measured is assumed to be known.

Actually, in practice, we can only measure those variables that we need for deriving \( y \), and we do not have to measure the values of all other variables. However, in the standard program synthesis, we do not take into account the resources that go into measurements, so we do not have to distinguish between measurements that we actually do and measurements that we can potentially perform.

However, in our (more realistic) formulation, we do take measurement efforts into account. To properly describe these efforts, we:

- denote by \( 0 \) the starting point, where no resources have been spent and thus, no measurements have been made, and
- describe each potentially measurable variable \( a \) by a rule \( 0 \rightarrow a \) with the weight \( w(a) \) (describing how many resources we need to spend to measure its value).

In this new description, weights \( w(r) \) are only assigned to rules \( r \).

**Logical interpretation of weights.** We can interpret these weights as *degrees* in the style of fuzzy logic (see, e.g., [8], [10]): if the derivation takes a long time, this means that we should not be using this implication unless it is absolutely necessary.

To make a description similar to the standard fuzzy logic, where the degrees are from the interval \([0, 1] \), we can *normalize* the weights, i.e., select a value \( W \) which is larger than or equal to all the weights \( w(r) \), and, instead of the original weights, consider the degrees \( \mu(r) = w(r)/W \).

So, we can interpret the difference \( 1 - \mu(r) \) as the “degree of confidence” in using this rule: the smaller the weight, the more confident we are in using this rule.

**A possible solution: exhaustive search.** In the toy example of a triangle, we can simply enumerate all possible paths and select the one with the smallest weight.

However, the number of possible paths grows exponentially with the number of variables. In the practical applications (like the space navigation problem mentioned earlier), we have dozens and hundreds of variables, so exhaustive search will take too long. For example, for \( n \approx 300 \), \( 2^n \) computations require longer time that the lifetime of the Universe :-(!

**Simplest case: rules of the type \( a \rightarrow b \).** To look for faster algorithms, let us start with the simplest case when all the rules have the form \( a \rightarrow b \), i.e., when all the rules only have one input. This simplest case can be described by a directed graph in which nodes are variables, and variables \( a \) and \( b \) are connected by an edge if there is a rule \( a \rightarrow b \). Non-negative weights are now assigned to edges.

In this case, estimating \( y \) simply means that we have a sequence of rules \( 0 \rightarrow a \rightarrow \ldots \rightarrow b \rightarrow y \), i.e., that we have a path from the starting node \( 0 \) to the desired node \( y \). The optimal program corresponds to the shortest path from \( 0 \) to \( y \).

**Efficient algorithms for the simplest case.** Efficient algorithms are known for computing shortest path in a graph; see, e.g., [4]. Thus, we can use these algorithms to solve the above simplest case of the optimal program synthesis problem.

Our objective is to generalize these algorithm to the more general case. In view of this objective, let us describe these algorithms in some detail.

Most of the efficient shortest path algorithms are based on the following *dynamic programming* idea. We want to find the length of the shortest path from the fixed node \( 0 \) to different nodes \( y \). The shortest path cannot visit a node twice; otherwise, we can shorten it by cutting out the part between the two visits. Thus, on a graph with \( n \) nodes, we only need to consider paths with \( \leq n - 1 \) edges. For each node \( x \), let \( d_k(x) \) denote the shortest of all the paths from \( 0 \) to \( x \) which have \( \leq k \) edges (\( \infty \) in no such path exists).

Then, for \( k = 0 \), we have \( d_0(0) = 0 \) and \( d_0(x) = \infty \) for all \( x \neq 0 \).

For \( k > 0 \), the shortest path of length \( \leq k \):

- either has the length \( \leq k - 1 \), in which case its length is equal to \( d_{k-1}(x) \);
- or has length exactly \( k \), in which case it spends \( k - 1 \) edges to get to some node \( y \) and then the last edge to get from \( y \) to \( x \); in this case, its length is

\[
\begin{align*}
d_{k-1}(y) + w(y \rightarrow x). 
\end{align*}
\]

Thus, the length \( d_k(x) \) of the shortest path with \( \leq k \) edges is equal to the smallest of these values:

\[
\begin{align*}
d_k(x) = \min \left( d_{k-1}(x), \min_y (d_{k-1}(y) + w(y \rightarrow x)) \right). 
\end{align*}
\]
For each $k$ and for each $x$, we need a linear time to compute this expression (by counting all $y$s). For each $k$, we need to repeat this computation for each of $n$ nodes $x$—which requires $n \cdot n = O(n^2)$ time.

Finally, as we have mentioned, the shortest path is equal to $d_{n-1}(x)$, we need to repeat all computations for all $k = 1, \ldots, n-1$—so the overall time is $(n-1) \cdot O(n^2) = O(n^3)$. This is thus indeed a polynomial-time algorithm.

Comment. Once we computed the values $d_{n-1}(x)$, we can also find the shortest paths: when the minimum of $d_k(x)$ is attained for $d_{k-1}(y) + w(y \rightarrow x)$, this means that the previous step before last should be the rule $y \rightarrow x$. Similarly, we can find the best rule leading to $y$, etc.

A seemingly natural extension of this idea to a more general case. Since the above dynamic programming idea works for the simplest case, it seems reasonable to try it for the general case as well. Specifically, for each variable $x$, let $d_k(x)$ be the smallest amount of resources that we need to compute $x$ by using $\leq k$ rules.

Similarly to the above, for $k = 0$, we have $d_0(0) = 0$ and $d_0(x) = \infty$ for all $x \neq 0$. For $k > 0$, we have

$$d_k(x) = \min (d_{k-1}(x), d_k'(x)),$$

where

$$d_k'(x) \overset{\text{def}}{=} \min \limits_{a, \ldots, b \rightarrow x} (d_{k-1}(a) + \ldots + d_{k-1}(b) + w(a, \ldots, b \rightarrow x)),$$

and the minimum is taken over all rules that result in $x$.

The above seemingly natural approach does not always lead to an optimal program synthesis. Let us show that this approach does not always work. Let us assume that we have rules $0 \rightarrow a$, $a \rightarrow b$, $a \rightarrow c$, and $b, c \rightarrow d$, all with weight 1. Then,

- for $k = 0$, we have $d_0(0) = 0$, $d_0(a) = d_0(b) = d_0(c) = \infty$;
- for $k = 1$, we have $d_1(0) = 0$, $d_1(a) = 1$, $d_1(b) = d_1(c) = d_1(d) = \infty$;
- for $k = 2$, we have $d_2(0) = 0$, $d_2(a) = 1$, $d_2(b) = d_2(c) = 2$, $d_2(d) = \infty$;
- for $k = 3$, we have $d_3(0) = 0$, $d_3(a) = 1$, $d_3(b) = d_3(c) = 2$, $d_3(d) = 5$;
- after that, the values $d_k(x)$ do not change.

We are thus tended to conclude that the length of the shortest path to $d$ is 5. However, we can compute $d$ by using only 4 resource units:

- first, we spend one unit to measure $a$, i.e., to use the rule $0 \rightarrow a$;
- then, we spend two units to apply the rules $a \rightarrow b$ and $a \rightarrow c$;
- finally, we spend the last (fourth) unit to apply the rule $b, c \rightarrow d$.

Analysis of the situation. The reason why the above algorithm over-estimated is that when we had inputs $b$ and $c$, we added their costs — without taking into account that computing $b$ and $c$ has a common part—the part of measuring $a$. As a result, in our computations, we counted the resources needed to measure $a$ twice:

- once as part of estimating resources needed to compute $b$,
- second time as part of estimating resources needed to compute $c$.

Solution to the problem: a more efficient algorithm. Let us first consider the case when all the rules use either a single symbol or a pair in the left-hand side—i.e., when all the rules have either a form $a \rightarrow b$ or a form $a, b \rightarrow c$.

In this case, instead of only iteratively estimating the cost of computing each individual quantity $a$ (as in the shortest path algorithm), we also iteratively estimate the cost of computing all possible pairs. In other words, for each $k$, in addition to the values $w_k(a)$, we also estimate the smallest cost $d_k(a, b)$ of computing both $a$ and $b$ in $k$ rules. In general, we are computing values $d_k(A)$, where $A$ is either a single variable or a pair of variables.

To be able to perform computations, we need to expand the original set of rules covering variables to rules covering pairs. Specifically: we generate a new rule $A \rightarrow x$ if

- either $x$ is already an element of $C$ (in this case, the weight is 0),
- or there is an original rule $r$ of the type $C \rightarrow x$ with $C \subseteq A \cup \ldots \cup B$ (i.e., in which available inputs $A, \ldots, B$ include inputs needed for this rule); the new rule gets the weight $w(r)$.

Similarly, we generate a new rule $A, \ldots, B \rightarrow X$ if every element $x \in X$ is:

- either already in the inputs $x \in A \cup \ldots \cup B$,
- or is covered by a rule $r_x$ of the type $C_x \rightarrow x$ with $C_x \subseteq A \cup \ldots \cup B$.

The weight of the new rule is then defined as the sum of the weights of these original rules.

Then, we apply the same algorithm to the new nodes $A$.

Comment. While we increase the number of rules, this increase is still polynomial, so we still get a polynomial-time algorithm.

Example. Let us show that in the above example, the new algorithm lead to the correct estimate $d(d) = 4$. Indeed, with pairs, we now have rules

- $0 \rightarrow a$ (of weight 1),
• $a \rightarrow \{b, c\}$ (of weight 2), and
• $\{b, c\} \rightarrow d$ (of weight 1).

Applying these rules one after another we get the desired shortest path of length 4.

**General case.** In general, we may have a rule that has three inputs like $a, b, c \rightarrow d$, or even $k > 3$ inputs. In this case, we have to consider sets $A$ consisting of 3 (or, correspondingly, $k$) variables.

**Comment.** For each $k$, the resulting algorithm is still polynomial. Since the size is usually limited, we thus have still a polynomial algorithm.

However, it is worth mentioning that the computation time grows as $n^k$ – i.e., exponentially with $k$. This exponential growth is inevitable, since we are facing a general problem of finding a shortest path in a hyper-graph (= graph with “edges” of the type $a, \ldots, b \rightarrow c$), and this problem is known to be NP-hard [7].

**ACKNOWLEDGMENTS**

The author is thankful to Vladik Kreinovich and Steve Roach for valuable discussions, and to the anonymous referees for useful suggestions.

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