OPTIMAL ORDERED PROBLEM SOLVER

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Abstract

We present a novel, general, optimally fast, incremental way of searching for a universal algorithm that solves each task in a sequence of tasks. The Optimal Ordered Problem Solver (OOPS) continually organizes and exploits previously found solutions to earlier tasks, efficiently searching not only the space of domain-specific algorithms, but also the space of search algorithms. Essentially we extend the principles of optimal nonincremental universal search to build an incremental universal learner that is able to improve itself through experience.

The initial bias is embodied by a task-dependent probability distribution on possible program prefixes. Prefixes are self-delimiting and executed in online fashion while being generated. They compute the probabilities of their own possible continuations. Let $p^n$ denote a found prefix solving the first $n$ tasks. It may exploit previously stored solutions $p^i, i < n$, by calling them as subprograms, or by copying them and editing the copies before applying them. We provide equal resources for two searches that run in parallel until $p^{n+1}$ is discovered and stored. The first search is exhaustive; it systematically tests all possible prefixes on all tasks up to $n + 1$. The second search is much more focused; it only searches for prefixes that start with $p^n$, and only tests them on task $n + 1$, which is safe, because we already know that such prefixes solve all tasks up to $n$. Both searches are depth-first and bias-optimal: the branches of the search trees are program prefixes, and backtracking is triggered once the sum of the runtimes of the current prefix on all current tasks exceeds the prefix probability multiplied by the total search time so far.

In illustrative experiments, our self-improver becomes the first general system that learns to solve all $n$ disk Towers of Hanoi tasks (solution size $2^n - 1$) for $n$ up to 30, profiting from previously solved, simpler tasks involving samples of a simple context free language.

Keywords: oops, bias-optimality, incremental optimal universal search, metasearching & metalearning, self-improvement

Guide: The most important sections are 3.2 and 3.3 (and possibly 4, for those who are interested in experiments). All sections are illustrated by Figure 1 at the end of this paper. Frequently used symbols are collected in reference Table 3 (general OOPS-related symbols) and Table 5.1 (less important implementation-specific symbols, explained in the appendix, Section 5).
1 Basic Ideas and Concepts: Overview

Oops is a simple, general, theoretically sound, time-optimal way of searching for a universal behavior or program that solves each problem in a problem sequence, continually organizing and managing and reusing earlier acquired knowledge. For example, the $n$-th problem may be to compute the $n$-th event from previous events (prediction), or to find a faster way through a maze than found during the search for a solution to the $n-1$-th problem (optimization).

Primitives. We start with an initial set of user-defined primitive behaviors. Primitives may be assembler-like instructions or time-consuming software, such as, say, theorem provers, or matrix operators for neural network-like parallel architectures, or trajectory generators for robot simulations, or state update procedures for multiagent systems, etc. Each primitive is represented by a token.

Task-specific prefix codes. Complex behaviors are represented by token sequences or programs. To solve a given task, we try to sequentially compose an appropriate complex behavior from primitive ones, always obeying the rules of a given user-defined initial programming language. Programs are grown incrementally, token by token; their beginnings or prefixes are immediately executed while being created; this may modify some task-specific internal state or memory, and may transfer control back to previously selected tokens. To add a new token to some program beginning or prefix, we first have to wait until the execution of the prefix so far explicitly requests such a prolongation, by setting an appropriate signal in the internal state. That is, we obtain task-specific prefix codes on program space: with any given task, programs that halt because they have found a solution or encountered some error cannot request any more tokens. Given the current task, no halting program can be the prefix of another one \[27, 7\]. On a different task, however, the same program may continue to request additional tokens — this is important for our novel approach.

Access to previous solutions. Let $p^n$ denote a found prefix solving the first $n$ tasks. The search for $p^{n+1}$ may greatly profit from the information conveyed by $p^1, p^2, \ldots, p^n$ which are stored in special non-modifiable memory shared by all tasks, such that they are accessible to $p^{n+1}$. For example, $p^{n+1}$ might execute a token sequence that calls $p^{n-3}$ as a subprogram, or that copies $p^{n-17}$ into some internal task-specific memory, then modifies the copy a bit, then applies the slightly edited copy to the current task. In fact, much of the knowledge embodied by $p^i$ may be about how to access and edit and use older $p^j (i < j)$.

Bias. The searcher’s initial bias is embodied by initial, user-defined, task-dependent probability distributions on the finite or infinite search space of possible program prefixes. In the simplest case we start with a maximum entropy distribution on the tokens, and define prefix probabilities as the products of the probabilities of their tokens. But prefix continuation probabilities may also depend on previous tokens in context sensitive fashion.

Self-computed continuation probabilities. In fact, we permit that any executed prefix (halting programs are their own prefixes) computes a task-dependent probability distribution on its own possible continuations, to be stored and updated in task-specific internal memory. By, say, invoking previously frozen code that redefines the probability distribution on future prefix continuations, the currently tested prefix may completely reshape the most likely paths through the search space of its own continuations, based on experience ignored by Levin’s and Hutter’s nonincremental optimal search methods \[26, 17\]. This may introduce
significant problem class-specific knowledge derived from solutions to earlier tasks.

**TWO SEARCHES.** Novel oops provides equal resources for two near-bias-optimal searches (see below) that run in parallel until $p^{n+1}$ is discovered and stored. The first is exhaustive; it systematically tests all possible prefixes on all tasks up to $n + 1$. Alternative prefixes are tested on all current tasks in parallel while still growing; once a task is solved, we remove it from the current set; prefixes that fail on a single task are discarded. The second search is much more focused; it only searches for prefixes that start with $p^n$, and only tests them on task $n + 1$, which is safe, because we already know that such prefixes solve all tasks up to $n$.

**Bias-optimal backtracking.** In both searches of oops, alternative prefix continuations are evaluated by a novel, practical, token-oriented backtracking procedure that always ensures near-bias-optimality: no candidate behavior gets more time than it deserves, given the bias. This means that the total time spent on generating and testing any behavior will never significantly exceed its probability times the total search time so far. Essentially we conduct a depth-first search in program space, where the branches of the search tree are program prefixes, and backtracking is triggered once the sum of the runtimes of the current prefix on all current tasks exceeds the prefix probability multiplied by the total time so far.

We describe an efficient implementation of oops for a broad variety of initial programming languages. It features a miniature multitasking operating system which tracks the effects of tested prefixes (such as partially solved task sets and modifications of internal states and continuation probabilities) and is able to backtrack through or reset these effects in an optimally efficient fashion for subsequent tests of alternative prefix continuations.

In case of unknown, infinite task sequences we can typically never know whether we already have found an optimal solver for all tasks in the sequence. But once we unwittingly do find one, at most half of the total future run time will be wasted on searching for alternatives. Given the initial bias and subsequent bias shifts due to $p^1, p^2, \ldots$, no other bias-optimal searcher can expect to solve the $n + 1$-th task set substantially faster than oops. A by-product of this optimality property is that it gives us a natural and precise measure of bias and bias shifts, conceptually related to Solomonoff's conceptual jump size [50, 51].

**An example initial language.** For an illustrative application, we wrote an interpreter for a stack-based universal programming language inspired by Forth [31], with initial primitives for defining and calling recursive functions, iterative loops, arithmetic operations, and domain-specific behavior. Optimal metasearching for better search algorithms is enabled through the inclusion of bias-shifting instructions that can modify the conditional probabilities of future search options in currently running program prefixes.

**Experiments.** We first teach oops something about recursion, by training it to construct samples of the simple context free language $\{1^n2^n\}$ for $n$ up to 30. This takes roughly 0.3 days on a standard personal computer (PC). Thereafter, within a few additional days, by exploiting aspects of its previously discovered universal $1^n2^n$-solver, oops learns a universal solver for all $n$ disk Towers of Hanoi problems, solving all instances up to $n = 30$, where the shortest solution (not the search for it!) already costs more than $10^9$ moves. (Previous reinforcement learners and non-learning AI planners tend to fail for solution sizes exceeding $10^2$ and $10^5$, respectively.)

We conclude with an outlook on the physical limitations of oops.
2 Brief Introduction to Optimal Search

In this section we first briefly review general, asymptotically optimal, nonincremental search methods by Levin and Hutter, and introduce the concept of bias-optimal search, given a probabilistic bias. Then we point out why existing methods for incremental search either have very limited search spaces, or are not bias-optimal. The remainder of the paper offers a way of overcoming this.

2.1 Universal Search For Quickly Verifiable Solutions

Surprisingly, many books on search algorithms do not even mention a very simple asymptotically optimal algorithm for problems with quickly verifiable solutions:

Method 2.1 (Lsearch) Given a problem and a universal Turing machine \([53]\), every \(2^n\) steps on average execute one instruction of the \(n\)-th binary string (interpreted as a program) in an alphabetical list of all strings, until one of them finds a solution.

Given some problem class, if some unknown optimal program \(p\) requires \(f(k)\) steps to solve a problem instance of size \(k\), and \(p\) happens to be the \(m\)-th program in the alphabetical list, then LSEARCH (for Levin Search) \([26]\) will need at most \(O(2^m f(k)) = O(f(k))\) steps — the constant factor \(2^m\) may be huge but does not depend on \(k\). Compare \([28, p. 502-505]\) and \([17]\).

2.2 Asymptotically Fastest Nonincremental Problem Solver

Recently Hutter developed a more complex asymptotically optimal search algorithm for all well-defined problems \([17]\). HSEARCH (for Hutter Search) cleverly allocates part of the total search time to searching the space of proofs for provably correct candidate programs with provable upper runtime bounds; at any given time it focuses resources on those programs with the currently best proven time bounds. Unexpectedly, HSEARCH manages to reduce the constant slowdown factor to a value smaller than 5. In fact, it can be made smaller than \(1 + \epsilon\), where \(\epsilon\) is an arbitrary positive constant (M. Hutter, personal communication, 2002).

Unfortunately, however, the search in proof space introduces an unknown additive problem class-specific constant slowdown, which again may be huge. While additive constants generally are preferrable to multiplicative ones, both types may make universal search methods practically infeasible.

2.3 Nonuniversal Optimal Search and Bias-Optimality

In the real world, constants do matter. The last to cross the finish line in the Olympic 100 m dash may be only a constant factor slower than the winner, but this will not comfort him. And since constants beyond \(2^{500}\) do not even make sense within this universe, both LSEARCH and HSEARCH may be viewed as academic exercises demonstrating that the \(O()\) notation can sometimes be practically irrelevant despite its wide use in theoretical computer
science. LSEARCH & HSEARCH, however, do provide inspiration for nonuniversal but very practical methods which are optimal with respect to a limited search space, while suffering only from very small slowdown factors.

For example, designers of planning procedures often just face a binary choice between two options such as depth-first and breadth-first search. The latter is often preferrable, but its greater demand for storage may eventually require to move data from on-chip memory to disk. This can slow down the search by a factor of 10,000 or more. A straightforward solution in the spirit of LSEARCH is to start with a 50% bias towards either technique, and use both depth-first and breadth-first search in parallel — this will cause a slowdown factor of at most 2 with respect to the best of the two options (ignoring a bit of overhead for parallelization).

Generalizing this example, define a probability distribution $P$ on a finite or infinite set of programs. $P$ represents the searcher’s initial bias. A bias-optimal searcher will not spend more time on any solution candidate than it deserves, given the bias:

**Definition 2.1 (Bias-Optimal Searchers)** Given is a problem set $\mathcal{R}$, a search space $\mathcal{C}$ of solution candidates (where any problem $r \in \mathcal{R}$ should have a solution in $\mathcal{C}$), a task-dependent bias in form of conditional probability distributions $P(q | r)$ on the candidates $q \in \mathcal{C}$, and a predefined procedure that creates and tests any given $q$ on any $r \in \mathcal{R}$ within time $t(q, r)$ (typically unknown in advance). A searcher is $n$-bias-optimal ($n \geq 1$) if for any maximal total search time $T_{\text{max}} > 0$ it is guaranteed to solve any problem $r \in \mathcal{R}$ if it has a solution $p \in \mathcal{C}$ satisfying $t(p, r) \leq P(p | r) T_{\text{max}} / n$. It is bias-optimal if $n = 1$.

This definition makes intuitive sense: the most probable candidates should get the lion’s share of the total search time, in a way that precisely reflects the initial bias.

For example, given a set of problems with quickly verifiable solutions, and an arbitrary user-defined, quickly computable probability distribution $P$ on all programs of a universal Turing machine (e.g., $P$ could be based on program length, and conditioned on the problems), LSEARCH is $n$-bias-optimal, where $n$ may be huge: The precise slowdown factor depends on the arbitrary lexicographical order ignoring the bias $P$. The following method, however, is near-bias-optimal (compare Solomonoff’s paper [50] as well as [17, p. 502-505]). For simplicity, we notationally suppress conditional dependencies on the current problem:

**Method 2.2 (OSEARCH)** Set current time limit $T = 1$. While problem not solved do:

Test all programs $q$ such that $t(q)$, the maximal time spent on creating and running and testing $q$, satisfies $t(q) < P(q) T$. Set $T := 2T$.

The near-bias-optimality of OSEARCH is hardly affected by the fact that for each value of $T$ we repeat certain computations for the previous value. Roughly half the total search time is still spent on $T$’s maximal value (ignoring hardware-specific overhead for parallelization and nonessential speed-ups due to halting programs if there are any). Nonbinary, nonuniversal variants of OSEARCH were used to solve machine learning toy problems unsolvable by traditional methods [18, 17]. Probabilistic alternatives based on probabilistically chosen maximal program runtimes in Speed-Prior style [11, 12] also outperformed traditional methods on certain toy problems [13, 14].
2.4 Incremental Search?

Since Newell & Simon’s early attempts at building a “General Problem Solver” [32, 35], much work has been done to develop mostly heuristic machine learning algorithms that solve new problems based on experience with previous problems, by incrementally shifting the inductive bias [55]. Many pointers to learning by chunking, learning by macros, hierarchical learning, learning by analogy, etc. can be found in Mitchell’s book [30]. Relatively recent general attempts include program evolvers such as Olsson’s ADATE [33] and simpler heuristics such as Genetic Programming (GP) [8, 2]. Unlike logic-based program synthesizers [12, 57, 9], program evolvers use biology-inspired concepts of Evolutionary Computation [34, 48] and Genetic Algorithms [14] to evolve better and better computer programs. Most existing GP implementations, however, do not even allow for programs with loops and recursion, thus ignoring a main motivation for search in program space. They either have very limited search spaces (where solution candidate runtime is not even an issue), or are far from bias-optimal, or both. Similarly, traditional reinforcement learners [20] are neither general nor close to being bias-optimal.

Hsearch and Lsearch / Osearch (Sections 2.1, 2.2, 2.3) are nonincremental in the sense that they do not attempt to minimize their constant slowdowns by exploiting experience collected in previous searches. They simply ignore the constants — from an asymptotic point of view, incremental search does not buy anything. Practical applications, however, may not ignore the constants. A heuristic attempt to greatly reduce them through experience was called Adaptive Lsearch or Als [47] — compare Solomonoff’s related ideas [50, 51]. Essentially Als works as follows: whenever Osearch finds a program q that computes a solution for the current problem, q’s probability $P(q)$ is substantially increased using a “learning rate,” while probabilities of alternative programs decrease appropriately. Subsequent Osearches for new problems then use the adjusted $P$, etc. A nonuniversal variant of this approach was able to solve reinforcement learning (RL) tasks [20] in partially observable environments unsolvable by traditional RL algorithms [58, 47].

Each Osearch invoked by Als is bias-optimal with respect to the most recent adjustment of $P$. On the other hand, the rather arbitrary $P$-modifications themselves are not necessarily optimal. They may quickly distort the initial bias, and provoke a loss of near-bias-optimality with respect to the initial bias during exposure to subsequent tasks. One contribution of this paper is to overcome this drawback in a principled way.

Our method draws inspiration from several of our previous publications on learning to learn or metalearning [37], where the goal is to learn better learning algorithms through self-improvement without human intervention (compare Lenat’s human-assisted self-improver [25]). In particular, the concept of incremental search for improved, probabilistically generated code that modifies the probability distribution on the possible code continuations has been used before: our incremental self-improvers [46] use the success-story algorithm SSA [46] to undo those self-generated probability modifications that in the long run do not contribute to increasing the learner’s cumulative reward per time interval. Our earlier meta-GP algorithm [37] was designed to learn better GP-like strategies. We also combined Holland’s principles of reinforcement learning economies [15] with a “self-referential” metalearning approach [57]. Our gradient-based metalearning technique [38] for continuous program spaces
of differentiable recurrent neural networks (RNNs) was also designed to favor better learning algorithms; compare the remarkable recent success of Hochreiter’s related but technically improved RNN-based metalearner [13].

The algorithms above generally are not near-bias-optimal though. The method discussed in this paper, however, combines optimal search and incremental self-improvement (as informally proposed in the author’s previous SNF grant applications, e.g., [42]), and will be \( n\)-bias-optimal, where \( n \) is a small and practically acceptable number, such as 8.

3 Optimal Ordered Problem Solver (OOPS)

\( \text{oops} \) depends on self-delimiting behaviors or programs whose beginnings are executed as soon as they are generated. Such programs are unnecessary for the asymptotic optimality properties of \( L\text{search} \) and \( H\text{search} \). \( \text{Binary} \) self-delimiting programs were studied [27, 7] in the context of Turing machines [53] and the theory of Kolmogorov complexity and algorithmic probability [13, 22]. Here we will use a more practical, not necessarily binary framework that does not exclude long programs with high probability.

Subsection 3.1 will introduce notation. Subsection 3.2 will introduce a practical method for efficiently tracking and restoring effects due to online execution of program prefixes with self-generated continuation probabilities, written in any of many possible initial programming languages, and for efficiently searching for a prefix solving multiple tasks in parallel, given some code bias in the form of previously found code. Subsection 3.3 will then describe how solutions to new tasks may optimally exploit knowledge conveyed by solutions to previous tasks, and how to search for universal programs solving all tasks seen so far.

3.1 Basic Setup and Notation

Unless stated otherwise or obvious, to simplify notation, throughout the paper newly introduced variables are assumed to be integer-valued and to cover the range implicit in the context. Given some finite or countably infinite alphabet \( Q = \{Q_1, Q_2, \ldots \} \), let \( Q^* \) denote the set of finite sequences or strings over \( Q \), where \( \lambda \) is the empty string. Then let \( q, q^1, q^2, \ldots \in Q^* \) be (possibly variable) strings. \( l(q) \) denotes the number of symbols in string \( q \), where \( l(\lambda) = 0 \); \( q_n \) is the \( n \)-th symbol of string \( q \); \( q_m:n = \lambda \) if \( m > n \) and \( q_mq_{m+1} \ldots q_n \) otherwise (where \( q_0 := q_0:0 := \lambda \)). \( q^1q^2 \) is the concatenation of \( q^1 \) and \( q^2 \) (e.g., if \( q^1 = abc \) and \( q^2 = dac \) then \( q^1q^2 = abcdac \)).

Consider countable alphabets \( S \) and \( Q \). Strings \( s, s^1, s^2, \ldots \in S^* \) represent possible internal states of a computer; strings \( q, q^1, q^2, \ldots \in Q^* \) represent token sequences or code or programs for manipulating states. We focus on \( S \) being the set of integers and \( Q := \{1, 2, \ldots, n_Q\} \) representing a set of \( n_Q \) instructions of some programming language. (The first universal programming language due to G"odel [11] was based on integers as well, but ours will be more practical.) \( Q \) and \( n_Q \) may be variable: new tokens may be defined by combining previous tokens, just as traditional programming languages allow for the declaration of new tokens representing new procedures. Since \( Q^* \subset S^* \), substrings within states may also encode programs.
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Q$</td>
<td>variable set of instructions or tokens</td>
</tr>
<tr>
<td>$Q_i$</td>
<td>$i$-th possible token (an integer)</td>
</tr>
<tr>
<td>$n_Q$</td>
<td>current number of tokens</td>
</tr>
<tr>
<td>$Q^*$</td>
<td>set of strings over alphabet $Q$, containing the search space of programs</td>
</tr>
<tr>
<td>$q$</td>
<td>total current code $\in Q^*$</td>
</tr>
<tr>
<td>$q_n$</td>
<td>$n$-th token of code $q$</td>
</tr>
<tr>
<td>$q^{\pi}$</td>
<td>$n$-th frozen program $\in Q^*$, where total code $q$ starts with $q^1 q^2 \ldots$</td>
</tr>
<tr>
<td>$qp$</td>
<td>$q$-pointer to the highest address of code $q = q_{1:qp}$</td>
</tr>
<tr>
<td>$a_{last}$</td>
<td>start address of a program (prefix) solving all tasks so far</td>
</tr>
<tr>
<td>$a_{frozen}$</td>
<td>top frozen address, can only grow, $1 \leq a_{last} \leq a_{frozen} \leq qp$</td>
</tr>
<tr>
<td>$q_{1:a_{frozen}}$</td>
<td>current code bias</td>
</tr>
<tr>
<td>$R$</td>
<td>variable set of tasks, ordered in cyclic fashion; each task has a computation tape</td>
</tr>
<tr>
<td>$S$</td>
<td>set of possible tape symbols (here: integers)</td>
</tr>
<tr>
<td>$S^*$</td>
<td>set of strings over alphabet $S$, defining possible states stored on tapes</td>
</tr>
<tr>
<td>$s^i$</td>
<td>an element of $S^*$</td>
</tr>
<tr>
<td>$s(r)$</td>
<td>variable state of task $r \in R$, stored on tape $r$</td>
</tr>
<tr>
<td>$s_i(r)$</td>
<td>$i$-th component of $s(r)$</td>
</tr>
<tr>
<td>$l(s)$</td>
<td>length of any string $s$</td>
</tr>
<tr>
<td>$z(i)(r)$</td>
<td>equal to $q_i$ if $0 &lt; i \leq l(q)$ or equal to $s_{-i}(r)$ if $-l(s(r)) \leq i \leq 0$</td>
</tr>
<tr>
<td>$ip(r)$</td>
<td>current instruction pointer of task $r$, encoded on tape $r$ within state $s(r)$</td>
</tr>
<tr>
<td>$p(r)$</td>
<td>variable probability distribution on $Q$, encoded on tape $r$ as part of $s(r)$</td>
</tr>
<tr>
<td>$p_i(r)$</td>
<td>current history-dependent probability of selecting $Q_i$ if $ip(r) = qp + 1$</td>
</tr>
</tbody>
</table>

Table 1: Symbols used to explain the basic principles of OOPS (Section 3).

$R$ is a set of currently unsolved tasks. Let the variable $s(r) \in S^*$ denote the current state of task $r \in R$, with $i$-th component $s_i(r)$ on a computation tape $r$ (a separate tape holding a separate state for each task). Since subsequences on tapes may also represent executable code, for convenience we combine current code $q$ and any given current state $s(r)$ in a single address space, introducing negative and positive addresses ranging from $-l(s(r))$ to $l(q) + 1$, defining the content of address $i$ as $z(i)(r) := q_i$ if $0 < i \leq l(q)$ and $z(i)(r) := s_{-i}(r)$ if $-l(s(r)) \leq i \leq 0$. All dynamic task-specific data will be represented at nonpositive addresses (one code, many tasks). In particular, the current instruction pointer $ip(r) := z(a_{ip}(r))(r)$ of task $r$ can be found at (possibly variable) address $a_{ip}(r) \leq 0$. Furthermore, $s(r)$ also encodes a modifiable probability distribution $p(r) = \{p_1(r), p_2(r), \ldots, p_{n_Q}(r)\}$ ($\sum_i p_i(r) = 1$) on $Q$. The current values of this variable distribution will be used to select a new instruction in case $ip(r)$ points to the first unused address right after the end of the current code $q$.

$a_{frozen} \geq 0$ is a variable address that can increase but not decrease. Once chosen, the code bias $q_{0:a_{frozen}}$ will remain unchangeable forever — it is a (possibly empty) sequence of programs $q^1 q^2 \ldots$, some of them prewired by the user, others frozen after previous successful searches for solutions to previous task sets (possibly completely unrelated to the current
task set $R$). Given $R$, the current goal is to solve all tasks $r \in R$, by a single program that either appropriately uses or extends the current code $q_{\text{frozen}}$ (no additional freezing will take place before all tasks in $R$ are solved). We will do this in a near-bias-optimal fashion (Def. 2.1), that is, no solution candidate will get much more search time than it deserves, given some initial probabilistic bias on program space $\subseteq Q^*$.

### 3.2 Prerequisites: Multitasking & Prefix Tracking By “Try”

The Turing machine-based setups for HSEARCH and LSEARCH assume potentially infinite storage. Hence they may largely ignore questions of storage management. In any practical system, however, we have to efficiently reuse limited storage. This, and multitasking, is what the present subsection is about. The recursive method Try below allocates time to program prefixes, each being tested on multiple tasks simultaneously, such that the sum of the runtimes of any given prefix, tested on all tasks, does not exceed the allocated total search time multiplied by the prefix probability (the product of the tape-dependent probabilities of the previously selected prefix components in $Q$).

#### 3.2.1 Overview of “Try”

Consider the left-hand side of Figure 1. All instruction pointers $ip(r)$ of all current tasks $r$ are initialized by some address, typically below the topmost code address, thus accessing the code bias common to all tasks, and/or using task-specific code fragments written into tapes. All tasks keep executing their instructions in parallel until interrupted or all tasks are solved, or until some task’s instruction pointer points to the yet unused address right after the topmost code address. The latter case is interpreted as a request for code prolongation through a new token, where each token has a probability according to the present task’s current state-encoded distribution on the possible next tokens. The deterministic method Try systematically examines all possible code extensions in a depth-first fashion (probabilities of prefixes are just used to order them for runtime allocation). Interrupts and backtracking to previously selected tokens (with yet untested alternatives) and the corresponding partial resets of states and task sets take place whenever one of the tasks encounters an error, or the product of the task-dependent probabilities of the currently selected tokens multiplied by the sum of the runtimes on all tasks exceeds a given total search time limit $T$.

To allow for efficient backtracking, Try tracks effects of tested program prefixes, such as task-specific state modifications (including probability distribution changes) and partially solved task sets, to reset conditions for subsequent tests of alternative, yet untested prefix continuations in an optimally efficient fashion (at most as expensive as the prefix tests themselves).

Since programs are created online while they are being executed, Try will never create impossible programs that halt before all their tokens are read. No program that halts on a given task can be the prefix of another program halting on the same task. It is important to see, however, that in our setup a given prefix that has solved one task (to be removed from the current task set) may continue to demand tokens as it tries to solve other tasks.
3.2.2 Details of "Try"

To allow us to efficiently undo state changes, we use global Boolean variables $mark_i(r)$ (initially FALSE) for all possible state components $s_i(r)$. We initialize time $t_0 := 0$; probability $P := 1$; $q$-pointer $qp := a_{\text{frozen}}$ and state $s(r)$ — including $ip(r)$ and $p(r)$ — with task-specific information for all task names $r$ in a so-called ring $R_0$ of tasks, where the expression "ring" indicates that the tasks are ordered in cyclic fashion; $| R |$ denotes the number of tasks in ring $R$. Given a global search time limit $T$, we Try to solve all tasks in $R_0$, by using existing code in $q = q_{1:qp}$ and / or by discovering an appropriate prolongation of $q$:

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**Method 3.1 (Boolean Try ($qp, r_0, R_0, t_0, P$))** ($r_0 \in R_0$; returns TRUE or FALSE; may have the side effect of increasing $a_{\text{frozen}}$ and thus prolonging the frozen code $q_{1:qp}$):

1. Make an empty stack $S$; set local variables $r := r_0$; $R := R_0$; $t := t_0$; $Done :=$ FALSE.

   **While** there are unsolved tasks ($| R | > 0$) **and** there is enough time left ($t \leq PT$) **and** instruction pointer valid ($\neg | s(r) | \leq ip(r) \leq qp$) **and** instruction valid ($1 \leq z(ip(r))(r) \leq n_Q$) **and** no halt condition is encountered (e.g., error such as division by 0, or robot bumps into obstacle; evaluate conditions in the above order until first satisfied, if any) **Do:**

   Interpret / execute token $z(ip(r))(r)$ according to the rules of the given programming language, continually increasing $t$ by the consumed time. This may modify $s(r)$ including instruction pointer $ip(r)$ and distribution $p(r)$, but not code $q$.

   Whenever the execution changes some state component $s_i(r)$ whose $mark_i(r) =$ FALSE, set $mark_i(r) :=$ TRUE and save the previous value $\hat{s}_i(r)$ by pushing the triple $(i, r, \hat{s}_i(r))$ onto $S$. Remove $r$ from $R$ if solved. If $| R | > 0$, set $r$ equal to the next task in ring $R$ (like in the round-robin method of standard operating systems). Else set $Done :=$ TRUE; $a_{\text{frozen}} := qp$ (all tasks solved; new code frozen, if any).

2. Use $S$ to efficiently reset only the modified $mark_i(k)$ to FALSE (the global mark variables will be needed again in step 3), but do not pop $S$ yet.

3. If $ip(r) = qp + 1$ (i.e., if there is an online request for prolongation of the current prefix through a new token): While $Done =$ FALSE and there is some yet untested token $Z \in Q$ (untried since $t_0$ as value for $q_{qp+1}$) **Do:**

   Set $q_{qp+1} := Z$ and $Done :=$ Try ($qp + 1, r, R, t, P * p(r)(Z)$), where $p(r)(Z)$ is $Z$’s probability according to current distribution $p(r)$.

4. Use $S$ to efficiently restore only those $s_i(k)$ changed since $t_0$, thus restoring all tapes to their states at the beginning of the current invocation of Try. This will also restore instruction pointer $ip(r_0)$ and original search distribution $p(r_0)$. Return the value of $Done$.  

---


In planning terminology [30], Try conducts a depth-first search in program space, where the branches of the search tree are program prefixes (each modifying a bunch of task-specific states), and backtracking is triggered once the sum of the runtimes of the current prefix on all current tasks exceeds the prefix probability multiplied by the total current time limit.

A successful Try will solve all tasks, possibly increasing $a_{frozen}$ and prolonging total code $q$. In any case Try will completely restore all states of all tasks. It never wastes time on recomputing previously computed results of prefixes, or on restoring unmodified state components and marks, or on currently irrelevant tasks — tracking / undoing effects of prefixes essentially does not cost more than their execution. So the $n$ in Def. 2.1 of $n$-bias-optimality is not greatly affected by the undoing procedure: we lose at most a factor 2, ignoring hardware-specific overhead such as the costs of single push and pop operations on a given computer, or the costs of measuring time, etc.

Since the distributions $p(r)$ are modifiable, we speak of self-generated continuation probabilities. As the variable suffix $q' := q_{a_{frozen} + 1:qp}$ of the total code $q = q_{1:qp}$ is growing, its probability can be readily updated:

$$P(q' \mid s^0) = \prod_{i=a_{frozen}+1}^{qp} P_i(q_i \mid s^i),$$

where $s^0$ is an initial state, and $P_i(q_i \mid s^i)$ is the probability of $q_i$, given the state $s^i$ of the task $r$ whose variable distribution $p(r)$ (as a part of $s^i$) was used to determine the probability of token $q_i$ at the moment it was selected. So we allow the probability of $q_{qp+1}$ to depend on $q_{0:qp}$ and intial state $s^0$ in a fairly arbitrary computable fashion. Note that unlike the traditional Turing machine-based setup [27, 7] (always yielding binary programs $q$ with probability $2^{-l(q)}$) this framework of self-generated continuation probabilities allows for token selection probabilities close to 1.0, that is, even long programs may have high probability.

Example. In many programming languages the probability of token “(“, given a previous token “WHILE”, equals 1. Having observed the “(” there is not a lot of new code to execute yet — in such cases the rules of the programming language will typically demand another increment of instruction pointer $ip(r)$, which will lead to the request of another token through subsequent increment of the topmost code address. However, once we have observed a complete expression of the form “WHILE (condition) DO (action),” it may take a long time until the conditional loop — interpreted via $ip(r)$ — is exited and the top address is incremented again, thus asking for a new token.

The round robin Try variant above keeps circling through all unsolved tasks, executing one instruction at a time. Alternative Try variants could also sequentially work on each task until it is solved, then try to prolong the resulting $q$ on the next task, and so on, appropriately restoring previous tasks once it turns out that the current task cannot be solved through prolongation of the prefix solving the earlier tasks. One potential advantage of round robin Try is that it will quickly discover whether the currently studied prefix causes an error for at least one task, in which case it can be discarded immediately.

Nonrecursive C-Code. An efficient iterative (nonrecursive) version of Try for a broad variety of initial programming languages was implemented in C. Instead of local stacks $S$, a single global stack is used to save and restore old contents of modified cells of all tapes / tasks.
Now suppose there is an ordered sequence of tasks \( r_1, r_2, \ldots \). Task \( r_j \) may or may not depend on solutions for \( r_i \) \((i, j = 1, 2, \ldots, j > i)\). For instance, in an optimization context, task \( r_j \) may be to find a faster way through a maze than the one found during the search for a solution to task \( r_{j-1} \). Or task \( r_j \) may just be to predict the \( j \)-th symbol in a sequence.

The instruction set \( Q \) should contain instructions for invoking or calling code found below \( a_{\text{frozen}} \), for copying such code into \( s(r) \), and for editing the copies and executing the results. Examples of such instructions will be given in the appendix (Section 3).

We are searching for a single program that solves all tasks encountered so far. That is, the \( n \)-th problem is to solve the set of the first \( n \) tasks. For example, we might want to teach the system a program that computes \( \text{fac}(n) = 1 \times 2 \times \ldots n \) for any given \( n \). Naturally, the \( i \)-th task \( r_i \) in the “training sequence” will be to compute \( \text{fac}(i) \). A found program that solves all tasks up to \( r_5 \) may already be a universal solver for arbitrary \( r_i \).

Inductively suppose we have solved the first \( n \) tasks through programs stored below address \( a_{\text{frozen}} \), and that the most recently discovered program starting at address \( a_{\text{last}} \leq a_{\text{frozen}} \) actually solves all of them, possibly using information conveyed by earlier programs \( q^1, q^2, \ldots \). To find a program solving the first \( n+1 \) tasks, OOPS invokes \textbf{Try} as follows (using set notation for task rings, where the tasks are ordered in cyclic fashion):

\begin{verbatim}
Method 3.2 (oops (n+1)) Initialize current time limit \( T := 2 \) and q-pointer \( qp := a_{\text{frozen}} \) (top frozen address).
\begin{enumerate}
  \item Set instruction pointer \( ip(r_{n+1}) := a_{\text{last}} \) (start address of code solving all tasks up to \( n \)).
    \begin{enumerate}
      \item If \textbf{Try} \((qp, r_{n+1}, \{r_{n+1}\}, 0, \frac{1}{2})\) then exit.
    \end{enumerate}
    (This means that half the search time is assigned to the most recent \( q_{a_{\text{last}}}; a_{\text{frozen}} \) and all possible prolongations thereof).
  \item If it is possible to initialize all \( n+1 \) tasks within time \( T \):
    \begin{enumerate}
      \item Set local variable \( a := a_{\text{frozen}}+1 \) (first unused address); for all \( r \in \{r_1, r_2, \ldots, r_{n+1}\} \)
      \item set \( ip(r) := a \). If \textbf{Try} \((qp, r_{n+1}, \{r_1, r_2, \ldots, r_{n+1}\}, 0, \frac{1}{2})\) set \( a_{\text{last}} := a \) and exit.
    \end{enumerate}
    (This means that half the time is assigned to all new programs with fresh starts).
  \item Set \( T := 2T \), and go to 1.
\end{enumerate}
\end{verbatim}

That is, we spend roughly equal time on two simultaneous searches. The second (step 2) is exhaustive: it considers all tasks and all possible prefixes of programs starting after the topmost frozen address. The first (step 1), however, focuses only on task \( n+1 \) and the
most recent prefix starting at address $a_{\text{last}}$, and its possible continuations beyond $a_{\text{frozen}}$. In particular, $a_{\text{last}}$ does not increase as long as new tasks can be comparatively quickly solved by prolonging $q_{a_{\text{last}}a_{\text{frozen}}}$. Why is this justified? Because the code between $a_{\text{last}}$ and $a_{\text{frozen}}$, which cannot be altered, is already enough to solve all tasks up to $n$, and cannot request any additional tokens that could harm its performance on these previous tasks. That is, we already know by induction that all of its prolongations will solve all tasks up to $n$.

Therefore, given tasks $r_1, r_2, \ldots$, we first initialize $a_{\text{last}}$; then for $i := 1, 2, \ldots$ invoke oops$(i)$ to find programs starting at (possibly increasing) address $a_{\text{last}}$, each solving all tasks so far, possibly eventually discovering a universal solver for all tasks in the sequence.

As address $a_{\text{last}}$ increases for the $n$-th time, $q^n$ is defined as the program starting at $a_{\text{last}}$’s old value and ending right before its new value. Program $q^m$ ($m > i, j$) may exploit $q^i$ by calling it as a subprogram, or by copying $q^i$ into some state $s(r)$, then editing it there, e.g., by inserting parts of another $q^j$ somewhere, then executing the edited variant.

### 3.3.1 Near-Bias-Optimality of OOPS

oops is not only asymptotically optimal in Levin’s sense [26] (see Method 2.1), but also near bias-optimal (Def. 2.1). To see this, consider a program $p$ solving the current task set within $k$ steps, given current code bias $q_{\text{ba/frozen}}$ and $a_{\text{last}}$. Denote $p$’s probability by $P(p)$ (compare Eq. (1) and Method 3.2; for simplicity we omit the obvious conditions). A bias-optimal solver would find a solution within at most $k/P(p)$ steps. We observe that oops will find a solution within at most $2^3k/P(p)$ steps, ignoring a bit of hardware-specific overhead (for marking changed tape components, measuring time, switching between tasks, etc, compare Section 3.2): At most a factor 2 might be lost through allocating half the search time to prolongations of the most recent code, another factor 2 for the incremental doubling of $T$ (necessary because we do not know in advance the best value of $T$), and another factor 2 for Try’s resets of states and tasks. So the method is essentially $8$-bias-optimal (ignoring hardware issues) with respect to the current task. If we do not want to ignore hardware issues: on currently widely used computers we can realistically expect to suffer from slowdown factors smaller than acceptable values such as, say, 100.

The only changes in bias (or bias shifts) that occur in between searches are due to freezing code once it has solved all tasks in a task set. That is, unlike the learning rate-based bias shifts of Als [17] (section 2.4), those of oops do not reduce the probabilities of programs that were meaningful and executable before the addition of any new $q^i$. But consider the formerly meaningless, interrupted program prefixes trying to access code for earlier solutions when there weren’t any: such prefixes may suddenly become prolongable and successful, once some solutions to earlier tasks have been stored. That is, unlike with Als the acceleration potential of oops is not bought at the risk of an unknown slowdown due to nonoptimal changes of the underlying probability distribution through a heuristically chosen learning rate. As new tasks come along, oops remains near-bias-optimal with respect to the initial bias, while still being able to greatly profit from subsequent code bias shifts.

The advantages of oops materialize when $P(p) \gg P(p')$, where $p'$ is among the most probable fast solvers of the current task set that do not use previously found code. Ideally, $p$ is already identical to the most recently frozen code. Alternatively, $p$ may be rather short
and thus likely because it uses information conveyed by earlier found programs stored below $a_{frozen}$. For example, $p$ may call an earlier stored $q^i$ as a subprogram. Or maybe $p$ is a short and fast program that copies a large $q^i$ into state $s(r_j)$, then modifies the copy just a little bit to obtain $\bar{q}^i$, then successfully applies $\bar{q}^i$ to $r_j$. Clearly, if $p'$ is not many times faster than $p$, then OOPS will in general suffer from a much smaller constant slowdown factor than nonincremental asymptotically optimal search, precisely reflecting the extent to which solutions to successive tasks do share useful mutual information, given the set of primitives for copy-editing them.

Unlike nonincremental LSEARCH and HSEARCH, which do not require online-generated programs for their asymptotic optimality properties, OOPS does depend on such programs: by, say, invoking previously frozen code that redefines the probability distribution on future prefix continuations, the currently tested prefix may completely reshape the most likely directions in the search space of its own continuations, based on experience ignored by LSEARCH & HSEARCH. This may introduce significant problem class-specific knowledge derived from solutions to earlier tasks. Clearly, the advantages of such online search space modifications are not exploited by offline procedures that just alphabetically list all possible program candidates (possibly even including invalid ones with self-delimiting prefixes).

As we solve more and more tasks, thus collecting and freezing more and more $q^i$, it will generally become harder and harder to identify and address and copy-edit useful code segments within the earlier solutions. As a consequence we expect that much of the knowledge embodied by certain $q^j$ actually will be about how to access and edit and use programs $q^i$ ($i < j$) previously stored below $q^j$.

Given an optimal problem solver, problem $r$, current code bias $q_0,a_{frozen}$, the most recent start address $a_{last}$, and information about the starts and ends of previously frozen programs $q^1, q^2, \ldots, q^k$, the total search time $T(r, q^1, q^2, \ldots, q^k, a_{last}, a_{frozen})$ for solving $r$ can be used to define the degree of bias

$$B(r, q^1, q^2, \ldots, q^k, a_{last}, a_{frozen}) := 1/T(r, q^1, q^2, \ldots, q^k, a_{last}, a_{frozen}).$$

Compare Solomonoff’s concept of conceptual jump size $[50, 51]$.

### 3.3.2 How Often Can we Expect to Profit from Earlier Tasks?

How likely is it that any learner can indeed profit from earlier solutions? At first naive glance this seems unlikely, since most possible pairs of symbol strings (such as problem-solving programs) do not share any algorithmic information (e.g., $[28]$). Why not? Most possible combinations of strings $x, y$ are algorithmically incompressible, that is, the shortest algorithm computing $y$, given $x$, has the size of the shortest algorithm computing $y$, given nothing (typically a bit more than $l(y)$ symbols), which means that $x$ does not tell us anything about $y$. (Papers in evolutionary computation often mention no free lunch theorems $[59]$ which are variations of this ancient insight of theoretical computer science).

Typically, however, successive real world problems are not sampled from a uniform i.i.d. distribution on a large set of possible problems. Instead they tend to be closely related. In particular, teachers usually provide sequences of more and more complex tasks with very similar solutions. Problem sequences that humans consider to be interesting are atypical when
compared to arbitrary sequences of well-defined problems [11]. In fact, it is no exaggeration to claim that almost the entire field of computer science is focused on comparatively few atypical problem sets with exploitable regularities. For all interesting problems the consideration of previous work is justified, to the extent that interestingness implies relatedness to what’s already known [13]. Obviously, OOPS-like procedures are advantageous only where such relatedness does exist. In any case, however, they will at least not do much harm.

3.3.3 OOPS Variants

When we are not searching for a universal solver, but just intend to solve the most recent task \( r_{n+1} \), we should use a reduced variant of OOPS which replaces step 2 of Method 3.2 by:

2. Set \( a := a_{\text{frozen}} + 1 \); set \( ip(r_{n+1}) := a \). If \( \text{Try} (qp, r_{n+1}, \{r_{n+1}\}, 0, \frac{1}{2}) \), then set \( a_{\text{last}} := a \) and exit.

Similar OOPS variants will find a program that solves, say, just the \( m \) most recent tasks, etc. Other OOPS variants will assign more (or less) than half of the total time to the most recent code and prolongations thereof. Yet other OOPS variants will also assign fractions of the total time to the second most recent program and its prolongations, the third most recent program and its prolongations, etc.

We may also consider probabilistic OOPS variants in Speed-Prior style [11, 13].

One not necessarily useful idea: Suppose the number of tasks to be solved by a single program is known in advance. Now we might think of an OOPS variant that works on all tasks in parallel, again spending half the search time on programs starting at \( a_{\text{last}} \), half on programs starting at \( a_{\text{frozen}} + 1 \); whenever one of the tasks is solved by a prolongation of \( q_{a_{\text{last}}:a_{\text{frozen}}} \) (usually we cannot know in advance which task), we remove it from the current task ring and freeze the code generated so far, thus increasing \( a_{\text{frozen}} \) (in contrast to \( \text{Try} \) which does not freeze programs before the entire current task set is solved). If it turns out, however, that not all tasks can be solved by a program starting at \( a_{\text{last}} \), we have to start from scratch by searching only among programs starting at \( a_{\text{frozen}} + 1 \). Unfortunately, in general we cannot guarantee that this approach of early freezing will converge.

3.3.4 Fundamental Limitations of OOPS

An appropriate task sequence may help OOPS to reduce the slowdown factor of plain OSEARCH through experience. Given a single task, however, OOPS does not by itself invent an appropriate series of easier subtasks whose solutions should be frozen first.

Of course, since both OSEARCH and OOPS may search in general algorithm space, some of the programs they execute may be viewed as self-generated subgoal-definers and subtask solvers. But with a single given task there is no incentive to freeze intermediate solutions before the original task is solved. The potential speed-up of OOPS does stem from exploiting external information encoded within an ordered task sequence. This motivates its very name.

Given some final task, a badly chosen training sequence of intermediate tasks may cost more search time than required for solving just the final task by itself, without any intermediate tasks.
OOPS is designed for resetable environments. In non-resetable environments it loses its theoretical foundation, and becomes a heuristic method. For example, it is possible to use OOPS for designing optimal trajectories of robot arms in virtual simulations. But once we are working with a real physical robot there may be no guarantee that we will be able to precisely reset it as required by backtracking procedure Try.

OOPS neglects one source of potential speed-up: it does not predict future tasks from previous ones, and currently does not spend a fraction of its time on solving predicted tasks. This is what an optimal universal reinforcement learner based on Hutter’s AIXI model [16, 18] would do. Future research should lead to a marriage of the asymptotically optimal AIXI and the near-bias-optimal OOPS.

3.4 Example Initial Programming Language

The efficient search and backtracking mechanism described in Section 3.2 is designed for a broad variety of possible programming languages, possibly list-oriented such as LISP, or based on matrix operations for recurrent neural network-like parallel architectures. Many other alternatives are possible.

A given language is represented by $Q$, the set of initial tokens. Each token corresponds to a primitive instruction. Primitive instructions are computer programs that manipulate tape contents, to be composed by OOPS such that more complex programs result. In principle, the “primitives” themselves could be large and time-consuming software, such as, say, traditional AI planners, or theorem provers, or multiagent update procedures, or learning algorithms for neural networks represented on tapes.

For each instruction there is a unique number between 1 and $n_Q$, such that all such numbers are associated with exactly one instruction. Initial knowledge or bias is introduced by writing appropriate primitives and adding them to $Q$. Step 1 of procedure Try (see Section 3.2) translates any instruction number back into the corresponding executable code (in our particular implementation: a pointer to a C-function). If the presently executed instruction does not directly affect instruction pointer $ip(r)$, e.g., through a conditional jump, or the call of a function, or the return from a function call, then $ip(r)$ is simply incremented.

Given some choice of programming language / initial primitives, we typically have to write a new interpreter from scratch, instead of using an existing one. Why? Because procedure Try (Section 3.2) needs total control over all (usually hidden and inaccessible) aspects of storage management, including garbage collection etc. Otherwise the storage clean-up in the wake of executed and tested prefixes could become suboptimal.

For the experiments (Section 4) we wrote (in C) an interpreter for an example, stack-based, universal programming language inspired by FORTH [31], whose disciples praise its beauty and the compactness of its programs.

The appendix (Section 5) describes the details. Data structures on tapes (Section 5.1) can be manipulated by primitive instructions listed in Sections 5.2.1, 5.2.2, 5.2.3. Section 5.3 shows how the user may compose complex programs from primitive ones, and insert them into total code $q$. Once the user has declared his programs, $n_Q$ will remain fixed.
4 Experiments

Experiments can tell us something about the usefulness of a particular initial bias such as the one incorporated by a particular programming language with particular initial instructions. In what follows we will describe illustrative problems and a few results obtained using the Forth-inspired language specified in the appendix (Section 5). The latter should be consulted for the details of the instructions appearing in programs found by oops.

While explaining the learning system’s setup, we will also try to identify several more or less hidden sources of initial bias.

4.1 Task-Specific Initialization

Besides the 61 initial primitive instructions from Sections 5.2.1, 5.2.2, 5.2.3 (appendix), the only user-defined (complex) tokens are those declared in Section 5.3 (except for the last one, tailrec). That is, we have a total of 61 + 7 = 68 initial non-task-specific primitives.

Given any task, we add task-specific instructions. In the present experiments, we do not provide a probabilistic syntax diagram defining conditional probabilities of certain tokens, given previous tokens. Instead we simply start with a maximum entropy distribution on the \( n_Q > 68 \) tokens \( Q \), initializing all probabilities \( p_i = \frac{1}{n_Q} \), setting all \( p[\text{curp}][i] := 1 \) and \( \text{sum[curp]} := n_Q \) (compare Section 5.1).

Note that the instruction numbers themselves significantly affect the initial bias. Some instruction numbers, in particular the small ones, are computable by very short programs, others are not. In general, programs consisting of many instructions that are not so easily computable, given the initial arithmetic instructions (Section 5.2.1), tend to be less probable. Similarly, as the number of frozen programs grows, those with higher addresses in general become harder to access, that is, the address computation may require longer subprograms.

For the experiments we insert substantial prior bias by assigning the lowest (easily computable) instruction numbers to the task-specific instructions, and by boosting (see instruction boostq in Section 5.2.3) the appropriate “small number pushers” (such as \( c1, c2, c3 \); compare Section 5.2.1) that push onto data stack \( ds \) the numbers of the task-specific instructions, such that they become executable as part of code on \( ds \). We also boost the simple arithmetic instructions \( by2 \) (multiply top stack element by 2) and \( dec \) (decrement top stack element), such that the system can easily create other integers from the probable ones. For example, without these boosts the code sequence \( (c3 \ by2 \ by2 \ dec) \) (which returns integer 11) would be much less likely. Finally we express our initial belief in the occasional future usefulness of previously useful instructions, by also boosting boostq itself.

The following numbers represent maximal values enforced in the experiments: state size: \( l(s) = 3000 \); absolute tape cell contents \( s_i(r) \): \( 10^9 \); number of self-made functions: 100, of self-made search patterns or probability distributions per tape: 20; callstack pointer: \( \text{maxcp} = 100 \); data stack pointers: \( \text{maxdp} = \text{maxDp} = 200 \).
4.2 Towers of Hanoi: the Problem

Given are \( n \) disks of \( n \) different sizes, stacked in decreasing size on the first of three pegs. One may move some peg’s top disk to the top of another peg, one disk at a time, but never a larger disk onto a smaller. The goal is to transfer all disks to the third peg. Remarkably, the fastest way of solving this famous problem requires \( 2^n - 1 \) moves \((n \geq 0)\).

The problem is of the reward-only-at-goal type — given some instance of size \( n \), there is no intermediate reward for achieving instance-specific subgoals.

The exponential growth of minimal solution size is what makes the problem interesting: Brute force methods searching in raw solution space will quickly fail as \( n \) increases. But the rapidly growing solutions do have something in common, namely, the short algorithm that generates them. Smart searchers will exploit such algorithmic regularities. Once we are searching in general algorithm space, however, it is essential to efficiently allocate time to algorithm tests. This is what oops does, in near-bias-optimal incremental fashion.

Untrained humans find it hard to solve instances \( n > 6 \). Anderson [1] applied traditional reinforcement learning methods and was able to solve instances up to \( n = 3 \), solvable within at most 7 moves. Langley [24] used learning production systems and was able to solve instances up to \( n = 5 \), solvable within at most 31 moves. (Side note: Baum and Durdanovic also applied an alternative reinforcement learner based on Holland’s artificial economy [15] to a simpler 3 peg blocks world problem where any disk may be placed on any other [3]; thus the required number of moves grows only linearly with the number of disks, not exponentially; we were able to replicate their results for \( n \) up to 5 [23].) Traditional AI planning procedures (e.g., chapter V of [36], [21]) do not learn but systematically explore all possible move combinations, using only absolutely necessary task-specific primitives (while oops will later use more than 70 general instructions, most of them unnecessary). On current personal computers AI planners tend to fail to solve Hanoi problem instances with \( n > 15 \) due to the exploding search space (Jana Koehler, IBM Research, personal communication, 2002). oops, however, searches program space instead of raw solution space. Therefore, in principle it should be able to solve arbitrary instances by discovering the problem’s elegant recursive solution: given \( n \) and three pegs \( S, A, D \) (source peg, auxiliary peg, destination peg), define procedure

\[
\text{hanoi}(S, A, D, n) : \text{If } n = 0 \text{ exit; Else Do:}
\]

\[
\text{call hanoi}(S, D, A, n-1); \text{ move top disk from } S \text{ to } D; \text{ call hanoi}(A, S, D, n-1).
\]

4.2.1 Task Representation and Domain-Specific Primitives

The \( n \)-th problem is to solve all Hanoi instances up to instance \( n \). Following our general rule, we represent the dynamic environment for task \( n \) on the \( n \)-th task tape, allocating \( n + 1 \) addresses for each peg, to store the order and the sizes of its current disks, and a pointer to its top disk (0 if there isn’t one).

We represent pegs \( S, A, D \) by numbers 1, 2, 3, respectively. That is, given an instance of size \( n \), we push onto data stack \( ds \) the values 1, 2, 3, \( n \). By doing so we insert substantial, nontrivial prior knowledge about the fact that it is useful to represent each peg by a symbol, and to know the problem size in advance. The task is completely defined by \( n \); the other
3 values are just useful for the following primitive instructions added to the programming language of Section 5.1: Instruction \textit{mvdsk()} assumes that \( S, A, D \) are represented by the first three elements on data stack \( ds \) above the current base pointer \( \text{cs[cp].base} \) (Section 5.1). It operates in the obvious fashion by moving a disk from peg \( S \) to peg \( D \). Instruction \textit{xSA()} exchanges the representations of \( S \) and \( A \), \textit{xAD()} those of \( A \) and \( D \) (combinations may create arbitrary peg patterns). Illegal moves cause the current program prefix to halt. Overall success is easily verifiable since our objective is achieved once the first two pegs are empty.

4.3 Solve Simpler Tasks First: Context Free Language \( 1^n2^n \)

Despite the near-bias-optimality of oops, within reasonable time (a week) on a personal computer the system was not able to solve instances involving more than 3 disks. What does this mean? \textit{Since search time of an optimal searcher is a natural measure of initial bias}, it just means that the already non-negligible bias towards our task set was still too weak.

This actually gives us an opportunity to demonstrate that oops can indeed profit from its incremental learning abilities. In what follows, we first train it on additional, easier tasks, to teach it something about recursion, hoping that the resulting code bias shifts will help to solve the Hanoi tasks as well.

For this purpose we use a seemingly unrelated problem class based on the context free language \( \{1^n2^n\} \): given input \( n \) on the data stack \( ds \), the goal is to place symbols on the auxiliary stack \( Ds \) such that the \( 2n \) topmost elements are \( n \) 1’s followed by \( n \) 2’s. Again there is no intermediate reward for achieving instance-specific subgoals.

After every executed instruction we test whether the objective has been achieved. By definition, the time cost per test (measured in unit time steps; Section 5.2) equals the number of considered elements of \( Ds \). Here we have an example of a test that may become more expensive with instance size.

We add two more instructions to the initial programming language: instruction \textit{1toD()} pushes 1 onto \( Ds \), instruction \textit{2toD()} pushes 2. Now we have a total of five task-specific instructions (including those for Hanoi), with instruction numbers 1, 2, 3, 4, 5, for \textit{1toD}, \textit{2toD}, \textit{mvdsk}, \textit{xSA}, \textit{xAD}, respectively, which gives a total of 73 initial instructions.

So we first boost (Section 5.2.3) the “small number pushers” \( c1, c2 \) (Section 5.2.1) for the first training phase where the \( n \)-th task (\( n = 1, \ldots , 30 \)) is to solve all \( 1^n2^n \) problem instances up to \( n \). Then we undo the \( 1^n2^n \)-specific boosts of \( c1, c2 \) and boost instead the Hanoi-specific instruction number pushers \( c3, c4, c5 \) for the subsequent training phase where the \( n \)-th task (again \( n = 1, \ldots , 30 \)) is to solve all Hanoi instances up to \( n \).

4.4 C-Code

All of the above was implemented by a dozen pages of code written in C, mostly comments and documentation: Multitasking and storage management through an iterative variant of \textit{round robin Try} (Section 5.2); interpreter and 62 basic instructions (Section 5); simple
user interface for complex declarations (Section 5.3); applications to 1^n2^n-problems (Section 4.3) and Hanoi problems (Section 4.2). The current nonoptimized implementation considers between one and two million discrete unit time steps per second on an off-the-shelf PC (1.5 GHz).

4.5 EXPERIMENTAL RESULTS FOR BOTH TASK SETS

Within roughly 0.3 days, oops found and froze code solving all thirty 1^n2^n-tasks. Thereafter, within 2-3 additional days, it also found a universal Hanoi solver. It is instructive to study the sequence of intermediate solutions. In what follows we will transform integer sequences discovered by oops back into readable programs (compare instruction details in Section 5).

1. For the 1^n2^n-problem, within less than a second, after 480,000 time steps, oops found non-general but working code for \( n = 1 \): \( (\text{defnp 2toD}) \).

2. At time \( 10^7 \) it had solved the 2nd instance by simply prolonging the previous code, using the old, unchanged start address \( a_{last} \): \( (\text{defnp 2toD grt c2 c2 endnp}) \). So this code solves the first two instances.

3. At time \( 10^8 \) it had solved the 3rd instance, again through prolongation:
   \( (\text{defnp 2toD grt c2 c2 endnp boostq delD delD bsf 2toD}) \).
   Here instruction \( \text{boostq} \) greatly boosted the probabilities of the subsequent instructions.

4. At time \( 2.85 \times 10^9 \) (less than 1 hour) it had solved the 4th instance through prolongation:
   \( (\text{defnp 2toD grt c2 c2 endnp boostq delD delD bsf 2toD fromD delD delD delD delD fromD bsf by2 bsf}) \).

5. At time \( 3 \times 10^9 \) it had solved the 5th instance through prolongation:
   \( (\text{defnp 2toD grt c2 c2 endnp boostq delD delD bsf 2toD fromD delD delD delD delD fromD by2 bsf}) \).
   The code found so far was lengthy and unelegant. But it does solve the first 5 instances.

6. Finally, at time \( 30,665,044,953 \) (roughly 0.3 days), oops had created and tested a new, elegant, recursive program (no prolongation of the previous one) with a new increased start address \( a_{last} \), solving all instances up to 6: \( (\text{defnp c1 calltp c2 endnp}) \).
   That is, it was cheaper to solve all instances up to 6 by discovering and applying this new program to all instances so far, than just prolonging the old code on instance 6 only.

7. The program above turns out to be a near-optimal universal 1^n2^n-problem solver. On the stack, it constructs a 1-argument procedure that returns nothing if its input argument is 0, otherwise calls the instruction \( \text{1toD} \) whose code is 1, then calls itself with a decremented input argument, then calls \( \text{2toD} \) whose code is 2, then returns.
That is, all remaining 1ⁿ2ⁿ-tasks can profit from the solver of instance 6. Reusing this current program qₐₜₐₜₐₐₜₐₜₐₐₜₐₚₐₚₐₚₐₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜäßig
1^n2^n-solver. This online bias shift greatly increases the probability that `defnp, calltp, endnp, will appear in the remainder of the online-generated program. These instructions in turn are helpful for building (on the data stack `ds) the double-recursive procedure generated by the suffix `(defnp c4 calltp c3 c5 calltp endnp), which essentially constructs (on data stack `ds) a 4-argument procedure that returns nothing if its input argument is 0, otherwise decrements the top input argument, calls the instruction `xAD whose code is 4, then calls itself on a copy of the top 4 arguments, then calls `mvdsk whose code is 5, then calls `xSA whose code is 3, then calls itself on another copy of the top 4 arguments, then makes yet another (unnecessary) argument copy, then returns (compare the standard Hanoi solution).

The total probability of the final solution, given the previous codes, is calculated as follows: since `nQ = 73, given the boosts of c3, c4, c5, by2, dec, boostq, we have probability `(1+73)^3/(7+73) for the prefix `(c3 dec boostq); since this prefix further boosts `defnp, c1, calltp, c2, endnp, we have probability `(1+73)^7/(12+73) for the suffix `(defnp c4 calltp c3 c5 calltp endnp). That is, the probability of the complete 10-symbol code is 9.3 * 10^{-11}. On the other hand, the probability of the essential Hanoi-specific suffix `(defnp c4 calltp c3 c5 calltp endnp), given just the initial boosts, is only `(1+73)^3/(7+73)^4 = 4.5 * 10^{-14}, which explains why it was not quickly found without the help of the solution to an easier problem set. (Without any initial boosts its probability would actually have been similar: `(4/73)^7 = 9 * 10^{-14}.) So in this particular setup the simple recursion for the 1^n2^n-problem indeed provided useful training for the more complex Hanoi recursion.

On the other hand, the search for the universal solver for all 1^n2^n-problems (first found with instance `n = 6) did not at all profit from solutions to earlier solved tasks. Of course, instances `n > 6 did profit.

Note that the time spent by the final 10-token Hanoi solver on increasing the probabilities of certain instructions and on constructing executable code on the data stack (less than 50 time steps) quickly becomes negligible as the Hanoi instance size grows. In this particular application, most time is spent on executing the code, not on constructing it.

Note also that we could continue to solve Hanoi tasks up to `n > 40. The execution time required to solve such instances with an optimal solver greatly exceeds the search time required for finding the solver itself. There it does not matter much whether OOPS already starts with a prewired Hanoi solver, or first has to discover one, since the initial search time for the solver becomes negligible anyway.

Different initial bias will yield different results. E.g., we could set to zero the initial probabilities of most of the 73 initial instructions (most are unnecessary for our two problem classes), and then solve all 2 × 30 tasks more quickly (possibly at the expense of obtaining a nonuniversal initial programming language). The point of this experimental section, however, is not to find the most reasonable initial bias for particular problems, but to illustrate the general functionality of the first general, near-bias-optimal, incremental learner.

Future research may focus on devising particularly compact, particularly reasonable sets of initial codes with particularly broad practical applicability. It may turn out that the most useful initial languages are not traditional programming languages similar to the FORTH-like one from Section 6, but instead based on a handful of primitive instructions for massively parallel cellular automata [54, 56, 60], or on a few nonlinear operations on matrix-
like data structures such as those used in recurrent neural network research \cite{5}. For example, we could use the principles of OOPS to create a non-gradient-based, near-bias-optimal variant of Hochreiter’s successful recurrent network metalearn\cite{13}. It should also be of interest to study probabilistic Speed Prior \cite{45}-based OOPS variants, and to devise applications of OOPS-like methods as components of universal reinforcement learners, based on Hutter’s asymptotically optimal AI\textsuperscript{U}\textsuperscript{L} model \cite{16,18} for non-resetable environments whose reactions to control signals are sampled from computable probability distributions.

### 4.7 Physical Limitations of OOPS

Due to its generality and its optimality properties, OOPS should scale to large problems in an essentially unbeatable fashion, thus raising the question: Which are its physical limitations? To give a very preliminary answer, we first observe that with each decade computers become roughly 1000 times faster by cost, reflecting Moore’s empirical law first formulated in 1965. Within a few decades nonreversible computation will encounter fundamental heating problems associated with high density computing \cite{1}. Remarkably, however, OOPS can be naturally implemented using reversible computing strategies \cite{10}, since it completely resets all state modifications due to the programs it tests. But even when we naively extrapolate Moore’s law, within the next century OOPS will hit the Bremermann limit \cite{6}: approximately \(10^{51}\) operations per second on \(10^{32}\) bits for the “ultimate laptop” \cite{29} with 1 kg of mass and 1 liter of volume. Clearly, the Bremermann limit constrains the maximal “conceptual jump size” \cite{50,51} from one problem to the next. For example, given some prior code bias derived from solutions to previous problems, within 1 minute, a sun-sized OOPS (roughly \(2 \times 10^{30} kg\)) might be able to solve an additional problem that requires finding an additional 200 bit program with, say, \(10^{20}\) steps runtime. But within the next centuries, OOPS will fail on new problems that require additional 300 bit programs of this type, since the speed of light greatly limits the acquisition of additional mass, through a function quadratic in time.

Still, even the comparatively modest hardware speed-up factor \(10^9\) expected for the next 30 years appears quite promising for OOPS-like systems. For example, with the 73 token language used in the experiments (Section \ref{sec:experiments}), we could learn from scratch (within a day or so) to solve the 20 disk Hanoi problem (> \(10^6\) moves), without any need for boosting task-specific instructions, or for incremental search through instances < 20, or for additional training sequences of easier tasks. Comparable speed-ups should be achievable much earlier, by distributing OOPS across large computer networks.

### Acknowledgments

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References


<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ds</td>
<td>data stack holding arguments of functions, possibly also edited code</td>
</tr>
<tr>
<td>dp</td>
<td>stack pointer of ds</td>
</tr>
<tr>
<td>Ds</td>
<td>auxiliary data stack</td>
</tr>
<tr>
<td>Dp</td>
<td>stack pointer of Ds</td>
</tr>
<tr>
<td>cs</td>
<td>call stack or runtime stack to handle function calls</td>
</tr>
<tr>
<td>cp</td>
<td>stack pointer of cs</td>
</tr>
<tr>
<td>cs[cp].ip</td>
<td>current function call’s instruction pointer ( ip(r) := cs<a href="r">cp</a>.ip )</td>
</tr>
<tr>
<td>cs[cp].base</td>
<td>current base pointer into ds right below the current input arguments</td>
</tr>
<tr>
<td>cs[cp].out</td>
<td>number of return values expected on top of ds above cs[cp].base</td>
</tr>
<tr>
<td>fns</td>
<td>stack of currently available self-made functions</td>
</tr>
<tr>
<td>fnp</td>
<td>stack pointer of fns</td>
</tr>
<tr>
<td>fns[fnp].code</td>
<td>start address of code of most recent self-made function</td>
</tr>
<tr>
<td>fns[fnp].in</td>
<td>number of input arguments of most recent self-made function</td>
</tr>
<tr>
<td>fns[fnp].out</td>
<td>number of return values of most recent self-made function</td>
</tr>
<tr>
<td>pats</td>
<td>stack of search patterns (probability distributions on ( Q ))</td>
</tr>
<tr>
<td>patp</td>
<td>stack pointer of pats</td>
</tr>
<tr>
<td>curp</td>
<td>pointer to current search pattern ( 0 \leq curp \leq patp )</td>
</tr>
<tr>
<td>p[curp][i]</td>
<td>( i )-th numerator of current search pattern</td>
</tr>
<tr>
<td>sum[curp]</td>
<td>denominator; the current probability of ( Q_i ) is ( p[curp][i]/sum[curp] )</td>
</tr>
</tbody>
</table>

Table 2: Frequently used implementation-specific symbols, relating to the data structures used by a particular FORTH-inspired programming language (Section 5). Not necessary for understanding the basic principles of OOPS.

5 APPENDIX: EXAMPLE PROGRAMMING LANGUAGE

OOPS can be seeded with a wide variety of programming languages. For the experiments, we wrote an interpreter for a stack-based universal programming language inspired by FORTH [31]. We provide initial instructions for defining and calling recursive functions, iterative loops, arithmetic operations, and domain-specific behavior. Optimal metasearching for better search algorithms is enabled through bias-shifting instructions that can modify the conditional probabilities of future search options in currently running self-delimiting programs. Sections 5.1 explains the basic data structures; Sections 5.2.1, 5.2.2, and 5.2.3 define basic primitive instructions; Section 5.3 shows how to compose complex programs from primitive ones, and explains how the user may insert them into total code \( q \).

5.1 DATA STRUCTURES ON TAPES

Each tape \( r \) contains various stack-like data structures represented as sequences of integers. For any stack \( Xs(r) \) introduced below (here \( X \) stands for a character string reminiscent of the stack type) there is a (frequently not even mentioned) stack pointer \( Xp(r); 0 \leq Xp(r) \leq \)
maxXp, located at address aXp, and initialized by 0. The n-th element of Xs(r) is denoted Xs[n](r). For simplicity we will often omit tape indices r. Each tape has:

1. A data stack ds(r) (or ds for short, omitting the task index) for storing function arguments. (The corresponding stack pointer is dp : 0 ≤ dp ≤ maxdp).

2. An auxiliary data stack Ds.

3. A runtime stack or callstack cs for handling (possibly recursive) functions. Callstack pointer cp is initialized by 0 for the “main” program. The k-th callstack entry (k = 0, . . . , cp) contains 3 variables: an instruction pointer cs[k](r).ip (or simply cs[k].ip, omitting task index r) initialized by the start address of the code of some procedure f, a pointer cs[k].base pointing into ds right below the values which are considered input arguments of f, and the number cs[k].out of return values ds[cs[k].base + 1], . . . , ds[dp] expected on top of ds once f has returned. cs[cp] refers to the topmost entry containing the current instruction pointer ip(r) := cs[cp](r).ip.

4. A stack fns of entries describing self-made functions. The entry for function fn contains 3 integer variables: the start address of fn’s code, the number of input arguments expected by fn on top of ds, and the number of output values to be returned.

5. A stack pats of search patterns. pats[i] stands for a probability distribution on search options (next instruction candidates). It is represented by nQ + 1 integers p[i][n] (1 ≤ n ≤ nQ) and sum[i] (for efficiency reasons). Once ip(r) hits the current search address l(q) + 1, the history-dependent probability of the n-th possible next instruction Qn (a candidate value for qip(r)) is given by p[curp][n]/sum[curp], where curp is another tape-represented variable (0 ≤ curp ≤ patp) indicating the current search pattern.

6. A binary quoteflag determining whether the instructions pointed to by ip will get executed or just quoted, that is, pushed onto ds.

7. A variable holding the index r of this tape’s task.

8. A stack of integer arrays, each having a name, an address, and a size (not used in this paper, but implemented and mentioned for the sake of completeness).

9. Additional problem-specific dynamic data structures for problem-specific data, e.g., to represent changes of the environment. An example environment for the Towers of Hanoi problem is described in Section 4.

5.2 PRIMITIVE INSTRUCTIONS

Most of the 61 tokens below do not appear in the solutions found by OOPS in the experiments (Section 4). Still, we list all of them for completeness’ sake, and to provide at least one example way of seeding OOPS with an initial set of behaviors.
In the following subsections, any instruction of the form $\text{inst}(x_1, \ldots, x_n)$ expects its $n$ arguments on top of data stack $ds$, and replaces them by its return values, adjusting $dp$ accordingly — the form $\text{inst}()$ is used for instructions without arguments.

Illegal use of any instruction will cause the currently considered program prefix to halt. In particular, it is illegal to set variables (such as stack pointers or instruction pointers) to values outside their prewired given ranges, or to pop empty stacks, or to divide by zero, or to call a nonexistent function, etc.

Since CPU time measurements on our PCs turned out to be unreliable, we defined our own, rather realistic time scales. By definition, most instructions listed below cost exactly 1 unit time step. Some, however, consume more time: Instructions making copies of strings with length $n$ (such as $\text{cpn}(n)$) cost $n$ time steps; so do instructions (such as $\text{find}(x)$) accessing an $a \text{ priori}$ unknown number $n$ of tape cells; so do instructions (such as $\text{boostq}(k)$) modifying the probabilities of an $a \text{ priori}$ unknown number $n$ of instructions.

5.2.1 Basic Data Stack-Related Instructions

1. **Arithmetic.** $c0()$, $c1()$, $c2()$, ..., $c5()$ return constants 0, 1, 2, 3, 4, 5, respectively; $\text{inc}(x)$ returns $x + 1$; $\text{dec}(x)$ returns $x - 1$; $\text{by2}(x)$ returns $2x$; $\text{add}(x,y)$ returns $x + y$; $\text{sub}(x,y)$ returns $x - y$; $\text{mul}(x,y)$ returns $x \cdot y$; $\text{div}(x,y)$ returns the smallest integer $\leq x/y$; $\text{powr}(x,y)$ returns $x^y$ (and costs $y$ unit time steps).

2. **Boolean.** Operand $\text{eq}(x,y)$ returns 1 if $x = y$, otherwise 0. Analogously for $\text{geq}(x,y)$ (greater or equal), $\text{grt}(x,y)$ (greater). Operand $\text{and}(x,y)$ returns 1 if $x > 0$ and $y > 0$, otherwise 0. Analogously for $\text{or}(x,y)$. Operand $\text{not}(x)$ returns 1 if $x \leq 0$, otherwise 0.

3. **Simple Stack Manipulators.** $\text{del}()$ decrements $dp$; $\text{clear}()$ sets $dp := 0$; $\text{dp2ds}()$ returns $dp$; $\text{setdp}(x)$ sets $dp := x$; $\text{ip2ds}()$ returns $\text{cs}[\text{ip}].\text{ip}$; $\text{base}()$ returns $\text{cs}[\text{cp}].\text{base}$; $\text{fromD}()$ returns $Ds[\text{Dp}]$; $\text{toD}()$ pushes $ds[dp]$ onto $Ds$; $\text{delD}()$ decrements $Dp$; $\text{topf}()$ returns the integer name of the most recent self-made function; $\text{intopf}()$ and $\text{outopf}()$ return its number of requested inputs and outputs, respectively; $\text{popf}()$ decrements $\text{fnp}$, returning its old value; $\text{xmn}(m,n)$ exchanges the $m$-th and the $n$-th elements of $ds$, measured from the stack’s top; $\text{ex}()$ works like $\text{xmn}(1,2)$; $\text{xnmb}(m,n)$ exchanges the $m$-th and the $n$-th elements above the current base $ds[\text{cs}[\text{cp}].\text{base}]$; $\text{outn}(n)$ returns $ds[dp+n]$; $\text{outb}(n)$ returns $ds[\text{cs}[\text{cp}].\text{base} + n]$ (the $n$-th element above the base pointer); $\text{inn}(n)$ copies $ds[dp]$ onto $ds[dp+n]$; $\text{innb}(n)$ copies $ds[dp]$ onto $ds[\text{cs}[\text{cp}].\text{base} + n]$.

4. **Pushing Code.** Instruction $\text{getq}(n)$ pushes onto $ds$ the sequence beginning at the start address of the $n$-th frozen program (either user-defined or frozen by OOPS) and ending with the program’s final token. $\text{insq}(n,a)$ inserts the $n$-th frozen program above $ds[\text{cs}[\text{cp}].\text{base} + a]$, then increases $dp$ by the program size. Useful for copying and later editing previously frozen code.

5. **Editing Strings on Stack.** Instruction $\text{cpn}(n)$ copies the $n$ topmost $ds$ entries onto the top of $ds$, increasing $dp$ by $n$; $\text{up}()$ works like $\text{cpn}(1)$; $\text{cpnb}(n)$ copies $n$ $ds$ entries above $ds[\text{cs}[\text{cp}].\text{base}]$ onto the top of $ds$, increasing $dp$ by $n$; $\text{mmv}(a,b,n)$ copies the $n$ $ds$ entries starting with $ds[\text{cs}[\text{cp}].\text{base} + a]$ to $ds[\text{cs}[\text{cp}].\text{base} + b]$ and following
cells, appropriately increasing \( dp \) if necessary; \( ins(a,b,n) \) inserts the \( n \) \( ds \) entries above \( ds[cs[cp].base+a] \) after \( ds[cs[cp].base+b] \), appropriately increasing \( dp \); \( deln(a,n) \) deletes the \( n \) \( ds \) entries above \( ds[cs[cp].base + a] \), appropriately decreasing \( dp \); \( find(x) \) returns the stack index of the topmost entry in \( ds \) matching \( x \); \( findb(x) \) the index of the first \( ds \) entry above base \( ds[cs[cp].base] \) matching \( x \). Many of the above instructions can be used to edit stack contents that may later be interpreted as executable code.

### 5.2.2 Control-Related Instructions

Each call of callable code \( f \) increments \( cp \) and results in a new topmost callstack entry. Functions to make and execute functions include:

1. Instruction \( def(m,n) \) defines a new integer function name (1 if it is the first, otherwise the most recent name plus 1) and increments \( fnp \). In the new \( fns \) entry we associate with the name: \( m \) and \( n \), the function’s expected numbers of input arguments and return values, and the function’s start address \( cs[cp].ip + 1 \) (right after the address of the currently interpreted token \( def \)).

2. Instruction \( dof(f) \) calls \( f \): it views \( f \)’s address and input number \( m \) and output number \( n \), increments \( cp \), lets \( cs[cp].base \) point right below the \( m \) topmost elements (arguments) in \( ds \) (if \( m < 0 \) then \( cs[cp].base = cs[cp−1].base \)), that is, all \( ds \) contents corresponding to the previous instance are viewed as arguments), sets \( cs[cp].out := n \), and sets \( cs[cp].ip \) equal to \( f \)’s address, thus calling \( f \).

3. \( ret() \) causes the current function call to return; the sequence of the \( n = cs[cp].out \) topmost values on \( ds \) is copied down such that it starts in \( ds \) right above \( ds[cs[cp].base] \), thus replacing the former input arguments; \( dp \) is adjusted accordingly, and \( cp \) decremented, thus transferring control to the \( ip \) of the previous callstack entry (no copying or \( dp \) change takes place if \( n < 0 \) — then we effectively return the entire stack contents above \( ds[cs[cp].base] \)). Instruction \( rtl(x) \) calls \( ret() \) if \( x \leq 0 \) (conditional return).

4. \( oldq(n) \) calls the \( n \)-th frozen program (either user-defined or frozen by OOPS) stored in \( q \) below \( a_{frozen} \), assuming (somewhat arbitrarily) zero inputs and outputs.

5. Instruction \( jmp1(val, n) \) sets \( cs[cp].ip \) equal to \( n \) provided that \( val \) exceeds zero (conditional jump, useful for iterative loops); \( pip(x) \) sets \( cs[cp].ip := x \) (also useful for defining iterative loops by manipulating the instruction pointer); \( bsjmp(n) \) sets current instruction pointer \( cs[cp].ip \) equal to the \( address \) of \( ds[cs[cp].base+n] \), thus interpreting stack contents above \( ds[cs[cp].base + n] \) as code to be executed.

6. \( bsf(n) \) uses \( cs \) in the usual way to call the code starting at \( ds[cs[cp].base + n] \) (as usual, once the code is executed, we will return to the address of the next instruction right after \( bsf \)); \( exec(n) \) interprets \( n \) as the number of an instruction and executes it.

7. \( qot() \) flips a binary flag \( quoteflag \) stored at address \( a_{quoteflag} \) on tape as \( z(a_{quoteflag}) \). The semantics are: code in between two \( qot \)’s is quoted, not executed. More precisely,
instructions appearing between the $m$-th ($m$ odd) and the $m + 1$st qot are not executed; instead their instruction numbers are sequentially pushed onto data stack $ds$. Instruction $\text{nop()}$ does nothing and may be used to structure programs.

In the context of instructions such as $\text{getq}$ and $\text{bsf}$, let us quote from reference [19] (reprinted with friendly permission by Philip J. Koopman Jr., 2002):

Another interesting proposal for stack machine program execution was put forth by Tsukamoto (1977). He examined the conflicting virtues and pitfalls of self-modifying code. While self-modifying code can be very efficient, it is almost universally shunned by software professionals as being too risky. Self-modifying code corrupts the contents of a program, so that the programmer cannot count on an instruction generated by the compiler or assembler being correct during the full course of a program run. Tsukamoto’s idea allows the use of self-modifying code without the pitfalls. He simply suggests using the run-time stack to store modified program segments for execution. Code can be generated by the application program and executed at run-time, yet does not corrupt the program memory. When the code has been executed, it can be thrown away by simply popping the stack. Neither of these techniques is in common use today, but either one or both of them may eventually find an important application.

Some of the instructions introduced above are almost exactly doing what has been suggested by Tsukamoto [52]. Remarkably, they turn out to be useful in the experiments (Section 4).

5.2.3 Bias-Shifting Instructions to Modify Search Probabilities

The concept of online-generated probabilistic programs with “self-referential” instructions that modify the probabilities of instructions to be executed later was already implemented earlier [47]. Here we use the following primitives:

1. $\text{incQ}(i)$ increases the current probability of $Q_i$ by incrementing $p[curp][i]$ and $\text{sum}[curp]$. Analogously for $\text{decQ}(i)$ (decrement). It is illegal to set all $Q$ probabilities (or all but one) to zero; to keep at least two search options. $\text{incQ}(i)$ and $\text{decQ}(i)$ do not delete argument $i$ from $ds$, by not decreasing $dp$.

2. $\text{boostq}(n)$ sequentially goes through all instructions of the $n$-th self-discovered frozen program; each time an instruction is recognized as some $Q_i$, it gets boosted: its numerator $p[curp][i]$ and the denominator $\text{sum}[curp]$ are increased by $nQ$. (This is less specific than $\text{incQ}(i)$, but can be useful, as seen in the experiments, Section 4.)

3. $\text{pushpat()}$ stores the current search pattern $\text{pat}[curp]$ by incrementing $\text{patp}$ and copying the sequence $\text{pat}[\text{patp}] := \text{pat}[curp]$; $\text{poppat()}$ decrements $\text{patp}$, returning its old value. $\text{setpat}(x)$ sets $\text{curp} := x$, thus defining the distribution for the next search, given the current task.

The idea is to give the system the opportunity to define several fairly arbitrary distributions on the possible search options, and switch on useful ones when needed in a given computational context, to implement conditional probabilities of tokens, given a computational history.
Of course, we could also explicitly implement tape-represented conditional probabilities of tokens, given previous tokens or token sequences, using a tape-encoded, modifiable probabilistic syntax diagram for defining modifiable n-grams. This may facilitate the act of ignoring certain meaningless program prefixes during search. In the present implementation, however, the system has to create / represent such conditional dependencies by invoking appropriate subprograms including sequences of instructions such as incQ(), pushpat() and setpat().

5.3 Initial User-Defined Programs: Examples

“If it isn’t 100 times smaller than 'C' it isn’t Forth.” (Charles Moore)

The user can declare initial, possibly recursive programs by composing the tokens described above. Programs are sequentially written into q, starting with q₁ at address 1. To declare a new token (program) we write decl(m, n, NAME, body), where NAME is the textual name of the code. Textual names are of interest only for the user, since the system immediately translates any new name into the smallest integer > nQ which gets associated with the topmost unused code address; then nQ is incremented. Argument m denotes the code’s number of expected arguments on top of the data stack ds; n denotes the number of return values; body is a string of names of previously defined instructions, and possibly one new name to allow for cross-recursion. Once the interpreter comes across a user-defined token, it simply calls the code in q starting with that body’s first token; once the code is executed, the interpreter returns to the address of the next token, using the callstack cs. All of this is quite similar to the case of self-made functions defined by the system itself — compare instruction def in section 5.2.2.

Here are some samples of user-defined tokens or programs composed from the primitive instructions defined above. Declarations omit parantheses for argument lists of instructions.

1. decl(0, 1, c999, c5 c5 mul c5 c4 c2 mul mul mul dec ret) declares c999(), a program without arguments, computing the constant 999 and returning it on top of data stack ds.

2. decl(2, 1, testexp, by2 by2 dec c3 by2 mul mul up mul ret) declares testexp (x,y), which pops two values x, y from ds and returns \((6x(4y - 1))^2\).

3. decl(1, 1, fac, up c1 ex rt0 del up dec fac mul ret) declares a recursive function fac(n) which returns 1 if n = 0, otherwise returns \(n \times \text{fac}(n-1)\).

4. decl(1, 1, fac2, c1 c1 def up c1 ex rt0 del up dec topf def mul ret) declares fac2(n), which defines self-made recursive code functionally equivalent to fac(n), which calls itself by calling the most recent self-made function even before it is completely defined. That is, fac2(n) not only computes fac(n) but also makes a new fac-like function.

5. The following declarations are useful for defining and executing recursive procedures (without return values) that expect as many inputs as currently found on stack ds, and call themselves on decreasing problem sizes. defnp first pushes onto auxiliary
stack $Ds$ the number of return values (namely, zero), then measures the number $m$ of inputs on $ds$ and pushes it onto $Ds$, then quotes (that is, pushes onto $ds$) the begin of the definition of a procedure that returns if its topmost input $n$ is 0 and otherwise decrements $n$. $callp$ quotes a call of the most recently defined function / procedure. Both $defnp$ and $callp$ also quote code for making a fresh copy of the inputs of the most recently defined code, expected on top of $ds$. $endnp$ quotes the code for returning, grabs from $Ds$ the numbers of in and out values, and uses $bsf$ to call the code generated so far on stack $ds$ above the input parameters, applying this code (possibly located deep in $ds$) to a copy of the inputs pushed onto the top of $ds$.

dcl(-1,-1,defnp, c0 toD pushdp dec toD qot def up rt0 dec intpf cpn qot ret)
dcl(-1,-1,calltp, qot topf dof intpf cpn qot ret)
dcl(-1,-1,endnp,qot ret qot fromD cpnb fromD up delD fromDex bsf ret)

6. Since our entire language is based on integer sequences, there is no obvious distinction between data and programs. The following illustrative example demonstrates that this makes functional programming very easy:

dcl(-1, -1, TAILREC, qot c1 c1 def up qot c2 outb qot ex rt0 del up dec topf dof qot c3 outb qot ret qot c1 outb c3 bsjmp) declares a tail recursion scheme TAILREC with a functional argument. Suppose the data stack $ds$ holds three values $n$, $val$, and $codenum$ above the current base pointer. Then TAILREC will create a function that returns $val$ if $n = 0$, else applies the 2-argument function represented by $codenum$, where the arguments are $n$ and the result of calling the 2-argument function itself on the value $n - 1$.

For example, the following code fragment uses TAILREC to implement yet another version of $fac(n)$: $(qot c1 mul qot tailrec ret)$. Assuming $n$ on $ds$, it first quotes the constant $c1$ (the return value for the terminal case $n = 0$) and the function $mul$, then applies TAILREC.

The primitives of Section 3 collectively embody a universal programming language, computationally as powerful as Gödel’s [11] or FORTH or ADA or C. In fact, a small fraction of the primitives would already be sufficient to achieve this universality. Higher-level programming languages can be incrementally built based on the initial low-level FORTH-like language.

To fully understand a given program, one may need to know which instruction has got which number. For the sake of completeness, and to permit precise re-implementation, we include the full list here:

Figure 1: Storage snapshot during an OOPS application. **Left:** general picture (Section 3). **Right:** language-specific details for a particular FORTH-like programming language (Sections 5 and 6).

**Left:** current code \( q \) ranges from addresses 1 to \( qp \) and includes previously frozen programs \( q^1, q^2, q^3 \). Three unsolved tasks require three tapes (lower left) with addresses \(-3000\) to 0. Instruction pointers \( ip(1) \) and \( ip(3) \) point to code in \( q \), \( ip(2) \) to code on the 2nd tape. Once, say, \( ip(3) \) points right above the topmost address \( qp \), the probability of the next instruction (at \( qp + 1 \)) is determined by the current probability distribution \( p(3) \) on the possible tokens. OOPS spends equal time on programs starting with prefix \( q_{\text{last}}a_{\text{frozen}} \) (tested only on the most recent task, since such programs solve all previous tasks, by induction), and on all programs starting at \( a_{\text{frozen}} + 1 \) (tested on all tasks).

**Right:** Details of a single tape. There is space for several alternative self-made probability distributions on \( Q \), each represented by \( n_Q \) numerators and a common denominator. Pointer \( curp \) determines which distribution to use for the next token request. There is a stack \( fns \) of self-made function definitions, each with a pointer to its code’s start address, and its numbers of input arguments and return values expected on data stack \( ds \) (with stack pointer \( dp \)). The dynamic runtime stack \( cs \) handles all function calls. Its top entry holds the current instruction pointer \( ip \) and the current base pointer into \( ds \) below the arguments of the most recent call. There is also space for an auxiliary stack \( Ds \), and for representing modifiable aspects of the environment. See text for details.