# **Parsimony Pressure Made Easy**

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### **ABSTRACT**

The parsimony pressure method is perhaps the simplest and most frequently used method to control bloat in genetic programming. In this paper we first reconsider the size evolution equation for genetic programming developed in [26] and rewrite it in a form that shows its direct relationship to Price's theorem. We then use this new formulation to derive theoretical results that show how to practically and optimally set the parsimony coefficient dynamically during a run so as to achieve complete control over the growth of the programs in a population. Experimental results confirm the effectiveness of the method, as we are able to tightly control the average program size under a variety of conditions. These include such unusual cases as dynamically varying target sizes such that the mean program size is allowed to grow during some phases of a run, while being forced to shrink in others.

# Categories and Subject Descriptors

I.2.2 [Artificial Intelligence]: Automatic Programming

### **General Terms**

Algorithms, Performance

# Keywords

Genetic Programming, Bloat, Parsimony Pressure

# 1. INTRODUCTION

Starting in the early '90s researchers began to notice that in addition to progressively increasing their mean and best fitness, GP populations also exhibited certain other dynamics. In particular, it was often observed that, while the average size (number of nodes) of the programs in a population was initially fairly static (if noisy), at some point the average program size would start growing at a rapid pace. Typically this increase in program size was not accompanied by any corresponding increase in fitness.

This phenomena, which is known as *bloat*, has been the subject of intense study in GP, both because of its initially surprising

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nature, and because of its significant practical effects. (Large programs are computationally expensive to evolve and later use, can be hard to interpret, and may exhibit poor generalisation.) Over the years, many theories have been proposed to explain various aspects of bloat [25, Section 11.3], and while great strides have been made, we still lack a single unifying theory to explain the broad range of empirical observations. We review the key theoretical results on bloat in Section 2, with special emphasis on the size evolution equation [26] as it forms the basis for this new approach.

While the jury remains out on the causes of bloat, practitioners still have the practical problem of combating bloat in their runs. Consequently, a variety of practical techniques have been proposed to counteract bloat; we review these in Section 3. We will particularly focus on the parsimony pressure method [13, 35], which is perhaps the simplest and most frequently used method to control bloat in genetic programming. This method effectively treats the minimisation of size as a soft constraint and attempts to enforce this constraint using the penalty method, i.e., by decreasing the fitness of programs by an amount that depends on their size. The penalty is typically simply proportional to program size. The intensity with which bloat is controlled is, therefore, determined by one parameter called the parsimony coefficient. The value of this coefficient is very important: too small a value and runs will still bloat wildly; too large a value and GP will take the minimisation of size as its main target and will almost ignore fitness, thus converging towards extremely small but useless programs. Unfortunately, however, the "correct" values of this coefficient are highly dependent on particulars such as the problem being solved, the choice of functions and terminals, and various parameter settings. Very little theory, however, has been put forward to aid in setting the parsimony coefficient in a principled manner, forcing users to proceed by trial and error.

This paper presents a simple, effective, and theoretically grounded solution to this problem. In Section 4, we reconsider the size evolution equation for GP developed in [26], rewriting it in a form that shows its direct relationship to Price's theorem [27, 17, 25]. We then use this new formulation to derive theoretical results that tells us how to practically and optimally set the parsimony coefficient dynamically during a run so as to achieve very tight control over the average size of the programs in a population. We test our theory in Section 5 where we report extensive empirical results, showing how accurately the method controls program size in a variety of conditions. We then conclude in Section 6.

# 2. BLOAT IN THEORY

As mentioned above, there are several theories of bloat. For example, the *replication accuracy theory* [20] states that the success of a GP individual depends on its ability to have offspring

that are functionally similar to the parent. As a consequence, GP evolves towards (bloated) representations that increase replication accuracy. The removal bias theory [34] observes that inactive code in a GP tree (code that is not executed, or is executed but its output is then discarded) tends to be low in the tree, residing therefore in smaller-than-average-size subtrees. Crossover events excising inactive subtrees produce offspring with the same fitness as their parents. On average the inserted subtree is bigger than the excised one, so such offspring are bigger than average while retaining the fitness of their parent, leading ultimately to growth in the average program size. Another important theory, the nature of program search spaces theory [16, 18], predicts that above a certain size, the distribution of fitnesses does not vary with size. Since there are more long programs, the number of long programs of a given fitness is greater than the number of short programs of the same fitness. Over time GP samples longer and longer programs simply because there are more of them.

The explanations for bloat mentioned above are largely qualitative. There have, however, been some efforts to mathematically formalise and verify these theories. For example, Banzhaf and Langdon [3] defined a executable model where only the fitness, the size of active code and the size of inactive code were represented (i.e., there was no representation of program structures). Fitnesses of individuals were drawn from a bell-shaped distribution, while active and inactive code lengths were modified by a size-unbiased mutation operator. Various interesting effects were reported which are very similar to corresponding effects found in GP runs. Rosca proposed a similar, but slightly more sophisticated model which also included an analogue of crossover [29]. A strength of these types of models is their simplicity. A weakness is that they suppress or remove many details of the representation and operators typically used in GP. This makes it difficult to verify if all the phenomena observed in the model have analogues in GP runs, and if all important behaviours of GP in relation to bloat are captured by the model.

In [22, 26], a *size evolution equation* for genetic programming was developed, which is an exact formalisation of the dynamics of average program size:

$$E[\mu(t+1)] = \sum_{l} S(G_l) p(G_l, t),$$
 (1)

Here  $\mu(t+1)$  is the mean size of the programs in the population at generation t+1,  $E[\ ]$  is the expectation operator,  $G_l$  is the set of all programs having a particular shape l,  $S(G_l)$  is the size of programs in the set  $G_l$  (i.e., programs having the shape l), and  $p(G_l,t)$  is the probability of selecting programs from  $G_l$  (i.e., of shape l) from the population in generation t. This can be rewritten in terms of the expected change in average program size as:

$$E[\mu(t+1) - \mu(t)] = \sum_{l} S(G_l)(p(G_l, t) - \Phi(G_l, t)), \qquad (2)$$

where  $\Phi(G_l,t)$  is the proportion of programs of shape  $G_l$  in the current generation. Both equations apply to a GP system with selection and any form of symmetric subtree crossover.<sup>1</sup>

Note that Equations (1) and (2) do not directly explain bloat. They are, however, important because they constrain what can and cannot happen size-wise in GP populations. So, any explanation for bloat (including the theories summarised in this section) has to agree with these results. In particular, Equation (1) predicts that, for symmetric subtree-swapping crossover operators, the mean program size evolves as if selection only was acting on the population.

This means that if there is a variation in mean size (bloat, for example) it must be the result of some form of positive or negative selective pressure on some or all of the shapes  $G_l$ . Equation (2) shows that there can be bloat only if the selection probability  $p(G_l,t)$  is different from the proportion  $\Phi(G_l,t)$  for at least some l. In particular, for bloat to happen there will have to be some short  $G_l$ 's for which  $p(G_l,t) < \Phi(G_l,t)$  and also some longer  $G_l$ 's for which  $p(G_l,t) > \Phi(G_l,t)$  (at least on average). As we will see later, Equations (1) and (2) are the starting point for the work reported here.

We conclude this review with a recent explanation for bloat called the crossover bias theory [24, 5] which is based in significant part and is consistent with the size evolution equation above. On average, each application of subtree crossover removes as much genetic material as it inserts. So, crossover on its own does not produce growth or shrinkage. However, while the mean program size is unaffected, higher moments of the distribution are. In particular, crossover pushes the population towards a particular distribution of program sizes (a Lagrange distribution of the second kind), where small programs have a much higher frequency than longer ones. For example, crossover generates a very high proportion of singlenode individuals. In virtually all problems of practical interest, very small programs have no chance of solving the problem. As a result, programs of above average length have a selective advantage over programs of below average length. Consequently, the mean program size increases.

# 3. BLOAT CONTROL IN PRACTICE

Numerous empirical techniques have been proposed to control bloat [18, 33]. These include size fair crossover and size fair mutation [15, 4], which constrain the choices made during the execution of a genetic operation so as to actively prevent growth. In sizefair crossover, for example, the crossover point in the first parent is selected randomly, as in standard crossover. Then the size of the subtree to be excised is then used to constrain the choice of the second crossover point so as to guarantee that the subtree chosen from the second parent will not be "unfairly" big. Another recent technique, the Tarpeian method [23], controls bloat by acting directly on the selection probabilities in Equation 2 by setting the fitness of randomly chosen longer than average programs to 0. Recent methods also include the use of multi-objective optimisation (with two objectives: fitness and size) to control bloat. For example, [6] used a modified selection based on the Pareto criterion to reduce code growth without significant loss of solution accuracy, and [12] used a Pareto approach on regression problems in an industrial setting.

Older methods include several *mutation operators* that may help control the average tree size in the population while still introducing new genetic material. [10] proposes a mutation operator which prevents the offspring's depth being more then 15% larger than its parent. [14] proposes two mutation operators in which the new random subtree is on average the same size as the code it replaces. In *Hoist mutation* [11] the new subtree is selected from the subtree being removed from the parent, guaranteeing that the new program will be smaller than its parent. *Shrink mutation* [2] is a special case of subtree mutation where the randomly chosen subtree is replaced by a randomly chosen terminal. [21] provides theoretical analysis and empirical evidence that combinations of subtree crossover and subtree mutation operators can control bloat in linear GP systems.

None of the methods mentioned above, however, has gained as much widespread acceptance as the *parsimony pressure method* [13, 35]. The method works as follows. Let f(x) be the fitness of program x. When the parsimony pressure is applied we

<sup>&</sup>lt;sup>1</sup>In a symmetric operator the probability of selecting particular crossover points in the parents does not depend on the order in which the parents are drawn from the population.

define and use a new fitness function

$$f_p(x) = f(x) - c\ell(x) \tag{3}$$

where  $\ell(x)$  is the size of program x and c is a constant known as the parsimony coefficient.<sup>2</sup> [35] showed the benefits of adaptively adjusting the coefficient c at each generation in experiments on the evolution of Sigma-Pi neural networks with GP, but most implementations and results in the literature actually keep c constant. As we will see in Section 4, however, a dynamic c is in fact essential to obtain full control of bloat.

The parsimony pressure method can be seen as a way to address the generalisation-accuracy tradeoff common in machine learning [35, 31]. There are also connections between this method and the Minimum Description Length (MDL) principle used to control bloat in [8, 9, 7]. The MDL approach uses a fitness function which combines program complexity (expressed as the number of bits necessary to encode the program's tree) and classification error (expressed as the number of bits necessary to encode the errors on all fitness cases). Rosca also linked the parsimony pressure method to his approximate evolution equations for rooted-tree schemata [31, 28, 30, 32].

Naturally, controlling bloat while at the same time maximising fitness turns the evolution of programs into either a multi-objective optimisation problem or, at least, into a constrained optimisation problem. Thus, as mentioned in Section 1, we should expect (and numerous results in the literature show this) that excessively aggressive methods to control bloat may lead to poor performance (in terms of ability to solve the problem at hand) of the evolved programs. The parsimony pressure method is not immune from this risk. So, although good control of bloat can be obtained with a careful choice of the parsimony coefficient, the choice of such a coefficient is an important but delicate matter. To date, however, trial and error remains the only general method for setting the parsimony coefficient. Furthermore, with a constant c the method can only achieve partial control over the dynamics of the average program size over time.

In this paper we aim to change all that, theoretically deriving and testing an easy and practical modification of the parsimony pressure technique which provides extremely tight control over the dynamics of the mean program size.

#### 4. **OPTIMAL PARSIMONY PRESSURE**

In this section we show the relationship between the size evolution equation and Price's Theorem [27]. We also show how to use this new form of the size evolution equation to solve for dynamic parsimony coefficients that will allow for various types of control of the average program size (e.g., Equations (14), (15), (19), and (20)). While some of these may look daunting at first, they are in fact straightforward to add to most GP systems and (as is shown in Section 5) can provide exceptionally tight control over the average population size.

Let us start by considering Equation (1) again. With trivial manipulations it can be rewritten in terms of length-classes, rather than tree shapes, obtaining

$$E[\mu(t+1)] = \sum_{\ell} \ell p(\ell, t) \tag{4}$$

where the index  $\ell$  ranges over all program sizes, and

$$p(\ell,t) = \sum_{l:S(G_l)=\ell} p(G_l,t)$$
 (5)

is the probability of selecting a program of length  $\ell$ . Similarly, we can then rewrite Equation (2) as

$$E[\Delta \mu] = E[\mu(t+1) - \mu(t)] = \sum_{\ell} \ell(p(\ell,t) - \Phi(\ell,t)),$$
 (6)

where  $\Phi(\ell,t) = \sum_{l:S(G_l)=\ell} \Phi(G_l,t)$ .

We now restrict our attention to fitness proportionate selection. In this case

$$p(\ell,t) = \Phi(\ell,t) \frac{f(\ell,t)}{\bar{f}(t)}, \tag{7}$$

where  $f(\ell,t)$  is the average fitness of the programs of size  $\ell$  and  $\bar{f}(t)$  is the average fitness of the programs in the population, both computed at generation t. Then from Equation (6) we obtain

$$\begin{split} E[\Delta \mu] &= \sum_{\ell} \ell \left( \Phi(\ell, t) \frac{f(\ell, t)}{\bar{f}(t)} - \Phi(\ell, t) \right) \\ &= \frac{1}{\bar{f}(t)} \sum_{\ell} \ell(f(\ell, t) - \bar{f}(t)) \Phi(\ell, t) \end{split}$$

$$= \underbrace{\frac{1}{\bar{f}(t)} \Big(\underbrace{\sum_{\ell} (\ell - \mu(t)) (f(\ell, t) - \bar{f}(t)) \Phi(\ell, t)}_{= \operatorname{Cov}(\ell, f) \text{ by definition}} \\ + \mu(t) \underbrace{\sum_{\ell} (f(\ell, t) - \bar{f}(t)) \Phi(\ell, t)}_{= \operatorname{0} \operatorname{by definition of } \bar{f}(t)},$$

where  $\mu(t) = \sum_{\ell} \ell \Phi(\ell, t)$  is the current average program size. So,

$$E[\Delta \mu] = \frac{\operatorname{Cov}(\ell, f)}{\bar{f}(t)}.$$
 (8)

This result is important because it shows that Equation (6), our coarse grained version of Equation (2), is in fact a form of Price's theorem (see [27, 17, 1] for a detailed review). While Price's theorem is generally applicable to "inheritable features" in an evolving system, only informal arguments have so far been made conjecturing that size might be one such feature [17]. Our result, instead, proves the conjecture.

We are now in a position to more clearly see the effects of parsimony pressure and, more generally, of any form of program-size control based on the following generalisation of Equation (3):

$$f_p(x,t) = f(x) - g(\ell(x),t) \tag{9}$$

where g is a function of program size,  $\ell(x)$ , and generation, t. To achieve this we consider Equation (8) when the fitness function f(x) is replaced by  $f_p(x,t)$ . We obtain

$$E[\Delta \mu] = \frac{\text{Cov}(\ell, f_p)}{\bar{f}_p} \tag{10}$$

$$= \frac{\operatorname{Cov}(\ell, f - g)}{\bar{f} - \bar{g}} \tag{11}$$

$$= \frac{\operatorname{Cov}(\ell, f - g)}{\bar{f} - \bar{g}}$$

$$= \frac{\operatorname{Cov}(\ell, f) - \operatorname{Cov}(\ell, g)}{\bar{f} - \bar{g}}$$
(11)

where we omitted t for brevity. So, absence of growth (and bloat),  $E[\Delta\mu] = 0$ , is obtained if

$$Cov(\ell, g) = Cov(\ell, f). \tag{13}$$

<sup>&</sup>lt;sup>2</sup>Naturally, while  $f_p$  is used to guide evolution, one needs to still use the original fitness function f to recognise solutions and stop

In many conditions, this equation makes it possible to determine penalty functions g that can be used to control the program size dynamics in GP runs.

As an example, let us consider the case  $g(\ell(x),t) = c(t)\ell(x)$  where c(t) is a function of the generation number t. (This is essentially the traditional parsimony pressure in (3) but here the parsimony coefficient is allowed to change over time.) Then

$$Cov(\ell, g) = c(t) Cov(\ell, \ell) = c(t) Var(\ell).$$

Substituting this into Equation (13) and solving for c(t) one finds that, in order to completely remove growth (or shrinking) from a run, one needs to set

$$c(t) = \frac{\operatorname{Cov}(\ell, f)}{\operatorname{Var}(\ell)}.$$
 (14)

Note that c is a function of t because both numerator and denominator can change from generation to generation.

Let us now consider the more general case  $g(\ell(x),t) = c(t)\ell(x)^k$  where k is any real number (positive or negative). Here the no size-change condition requires

$$c(t) = \operatorname{Cov}(\ell, f) / \operatorname{Cov}(\ell, \ell^k). \tag{15}$$

As another example, let us consider the case  $g(\ell(x),t) = c(t)(\ell(x) - \mu(t))$ . Here

$$Cov(\ell,g) = c(t)Cov(\ell,\ell-\mu(t))$$
  
=  $c(t)Cov(\ell,\ell) - c(t)Cov(\ell,\mu(t)).$ 

But,  $Cov(\ell, \mu(t)) = 0$  and  $Cov(\ell, \ell) = Var(\ell)$ , so we end up with Equation (14) again (although the resulting penalty coefficient is then used in a different g).

What if we wanted  $\mu(t)$  to follow, in expectation, a particular function  $\gamma(t)$ , e.g., the ramp  $\gamma(t) = \mu(0) + b \times t$  or a sinusoidal function? The theory helps us in this case as well. Adding  $\mu(t)$  to both sides of Equation (10) we obtain:

$$\frac{\operatorname{Cov}(\ell, f) - \operatorname{Cov}(\ell, g)}{\bar{f} - \bar{g}} + \mu(t) = E[\mu(t+1)] = \gamma(t+1). \tag{16}$$

If g is a family of functions with a single degree of freedom (as is true of all the functions g considered above), then we can use this constraint to solve for the free variable. For example, if we want to control bloat with parsimony terms of the form  $g(\ell(x),t) = c(t)\ell(x)^k$  we can substitute this into Equation (16), obtaining

$$\frac{\operatorname{Cov}(\ell, f) - c(t)\operatorname{Cov}(\ell, \ell^k)}{\bar{f} - c(t)E[\ell^k]} + \mu(t) = \gamma(t+1). \tag{17}$$

Solving for c(t) gives:

$$c(t) = \frac{\operatorname{Cov}(\ell, f) - (\gamma(t+1) - \mu(t))\bar{f}}{\operatorname{Cov}(\ell, \ell^k) - (\gamma(t+1) - \mu(t))E[\ell^k]}$$
(18)

If k=1, i.e.,  $g=c(t)\ell(x)$  (as in the standard parsimony pressure), this simplifies to

$$c(t) = \frac{\operatorname{Cov}(\ell, f) - (\gamma(t+1) - \mu(t))\bar{f}}{\operatorname{Var}(\ell) - (\gamma(t+1) - \mu(t))\mu(t)}$$
(19)

Note that, in the absence of sampling noise (i.e., for an infinite population), requiring that  $E[\Delta\mu]=0$  at each generation causes Equation (13) to reduce to  $\mu(t)=\mu(0)$  for all t>0. However, in any finite population the parsimony pressure method can only achieve  $\Delta\mu=0$  in expectation, so there can be some random drift in  $\mu(t)$  w.r.t. its starting value of  $\mu(0)$ . Experimentally we have found that this tends to be significant only for very small populations and long

runs. If tighter control over the mean program size is desired, one can use Equation (18) with the choice  $\gamma(t) = \mu(0)$ , which leads to the following formula

$$c(t) = \frac{\text{Cov}(\ell, f) - (\mu(0) - \mu(t))\bar{f}}{\text{Cov}(\ell, \ell^k) - (\mu(0) - \mu(t))E[\ell^k]}.$$
 (20)

Note the similarities and differences between this and Equation (15). In the presence of any drift moving  $\mu(t)$  away from  $\mu(0)$ , this equation will actively strengthen the size control pressure to push the mean program size back to its initial value.<sup>3</sup>

As we will see in the following section, our technique gives users almost complete control over the dynamics of the mean program size, and control can be obtained in a single generation. It is thus possible to design interesting schemes where the covariance-based bloat control is switched on or off at different times, perhaps depending on the particular conditions of a run. In the next section we will, for example, test the idea of letting the GP system run undisturbed until the mean program size reaches a threshold, at which point we start applying bloat control to prevent further growth.

Finally, we note that while much of this theory assumes the use of fitness proportionate selection, Equation (6) is valid in general and one could imagine selection schemes that directly penalise the selection probabilities  $p(\ell,t)$  rather than fitnesses. As we will see in the experiments, however, the penalty coefficients estimated using the theory developed for fitness proportionate selection actually work very well without any modification in systems based on other forms of selection, such as tournament selection.

# 5. EXPERIMENTAL RESULTS

To verify the theory in a variety of different conditions, we conducted experiments using three different GP systems—two linear register-based GP systems and one tree-based GP system—and several problems. We briefly describe these systems and the problems in the next section, and then present some of our experimental results. Due to space limitations we will be able to report in detail on only a fraction of the tests we made, but the reported results generalize across a wide array of experiments.

# 5.1 GP Systems, Problems and Primitives

The first GP system we used is a linear generational GP system. It initialises the population by repeatedly creating random individuals with lengths uniformly distributed between 1 and 200 primitives. The primitives are drawn randomly and uniformly from a problem's primitive set. The system uses fitness proportionate selection and crossover applied with a rate of 100%. Crossover creates offspring by selecting two random crossover points, one in each parent, and taking the first part of the first parent and the second part of the second w.r.t. their crossover points. We used populations of size 100, 1,000 and 10,000. In each condition we performed 100 independent runs, each lasting 500 generations.

With this linear GP system we used two artificial test problems. The first was the Hole problem, which simply allocates a fitness of 0.001 to programs of size smaller than 10 nodes, and a fitness of 1.0 to all other programs. This problem was used because it presents the minimal conditions for bloat to occur (according to the crossover-bias theory described in Section 2). The second problem, which we will call Square Root, was one where the fitness of programs was simply the square root of their size, i.e.,  $f(x) = \sqrt{\ell(x)}$ . This problem also satisfies the conditions for bloat, but, unlike the previous one, here the entire fitness landscape is expected to favour

<sup>&</sup>lt;sup>3</sup>We talk about size control pressure rather than parsimony pressure because  $\mu(t)$  can drift both above and below  $\mu(0)$ .

Table 1: Primitive sets used in our experiments.

	Poly	6-MUX	
ı	R1 =	RIN	AND
ı	R2 =	RIN	OR
ı	R1 =	R1 + R2	NAND
ı	R2 =	R1 + R2	NOR
ı	R1 =	R1 * R2	
ı	R2 =	R1 * R2	
ı	Swap	R1 R2	

bloat (not just sections containing the very short programs) because the correlation between length and fitness is very high for all sizes. Because of its very strong tendency to bloat, we consider this problem a good stress-test of our method. The fitness for both Hole and Square Root is determined completely by the size of the program, so any choice of primitive set produces the same results.

The second GP system we used is also linear and generational. It uses the same crossover (with the same rate) and the same form of initialisation as the first system, but initial program lengths are in the range 1 to 50. Runs lasted 100 generations. The system uses *tournament selection* (with tournament size 2) instead of fitness proportional selection. This allows us to test the generality of our method for controlling program size.

With this system we solved two classical symbolic regression problems. The objective was to evolve a function which fits a polynomial of the form  $x+x^2+\cdots+x^d$ , where d is the degree of the polynomial, for x in the range [-1,1]. In particular we considered degrees d=6 and d=8 and we sampled the polynomials at the 21 equally spaced points  $x \in \{-1,-0.9,\ldots,0.9,1.0\}$ . We call the resulting symbolic regression problems Poly-6 and Poly-8. Polynomials of this type have been widely used as benchmark problems in the GP literature.

Fitness (to be maximised) was 1/(1 + error) where error is the sum of the absolute differences between the target polynomial and the output produced by the program under evaluation over these 21 fitness cases. The primitive set used to solve these problems is shown in the first column of Table 1. The instructions refer to three registers: the input register RIN which is loaded with the value of x before a fitness case is evaluated and the two registers R1 and R2 which can be used for numerical calculations. R1 and R2 are initialised to x and 0, respectively. The output of the program is read from R1 at the end of its execution.

The third GP system was a classical generational tree-based GP system using binary tournament selection, with subtree crossover applied with 100% probability. 35 independent runs were done for each of the five different targets for the average program size. The population size in each case was 2,000, and each run went for 500 generations. The populations were initialized using the PTC2 tree creation algorithm [19] with the initial trees having size 150.

With the tree-based GP system we used the 6-Multiplexer problem. This is a classical Boolean function indiction problem where the objective is to evolve a Boolean function with 6 inputs designed as A0, A1, D0, D1, D2, D3 which produces as output a copy of one of the inputs D0-D3. These are known as the data lines of the multiplexer. The particular input copied over is determined by the inputs A0 and A1 (known as the address lines of the multiplexer), as follows: if A0 = 0 and A1 = 0 then Out = D0, if A0 = 1 and A1 = 0 then Out = D1, if A0 = 0 and A1 = 1 then Out = D2, if A0 = 1 and A1 = 1 then Out = D3. The function has 64 possible combinations of inputs, so we have 64 fitness cases. Fitness is the number of fitness cases a program correctly computes. The primitive set used is shown in the second column of Table 1.

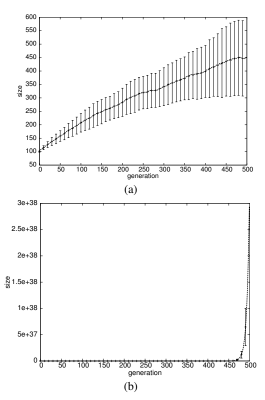


Figure 1: Behaviour of the mean program size in a linear GP system when solving the Hole problem (a) and the Square Root problem in the absence of bloat control for populations of size 1,000. Results are averages over 100 independent runs. The error bars indicate the standard deviation across the runs. Note the log scale on plot (b).

# 5.2 Results

We start by looking at the Hole and Square Root problems. As Figures 1(a)–(b) show for populations of size 1,000, bloat is present in both cases, with the  $\sqrt{\ell}$  fitness function bloating fiercely. Results for populations of size 100 and 10,000 are qualitatively similar.

To give a sense of the degree of control that can be achieved with our technique, Figure 2 illustrates the behaviour of mean program size for the Hole and Square Root problems when five different flavours of our size control scheme are used. Results are for populations of size 1,000, but other population sizes provide similar behaviours. Note the small amount of drift present when Equation (14) is used (first column). This is completely removed when instead we use Equation (20) (column 5, note the different scale, and also column 3 after transient). As columns 2 and 4 show, the user is free to impose any desired mean program size dynamics thanks to the use of Equation (19).

We turn to the Poly-6 and Poly-8 problems. As Figures 4(a)–(b) show, bloat is present in both problems. The behaviour of mean program size is brought under complete control with our technique as shown in Figure 3. Here, we used the same targets as in Figure 2 (although with slightly different parameters), but, to illustrate a further alternative, we used a parsimony term of the form  $g(t)=c(t)/\ell$ . This effectively promotes the shorter programs rather than penalising the longer ones.

Excellent size control was also obtained in tree-based GP when solving the 6-MUX problem, as shown in Figure 5. We used the same targets as in Figure 2, but again with slightly different parameters. Here the drift that's possible when using Equation (14) (the

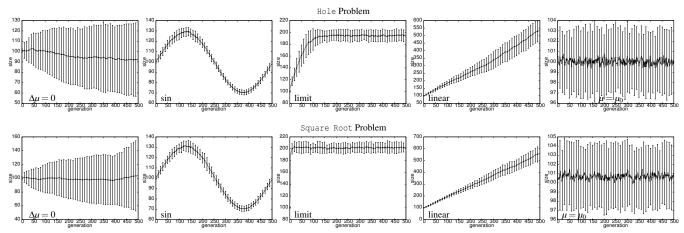


Figure 2: Size control obtained in the Hole and Square Root problems with populations of size 1,000 using the penalty function  $g(\ell(x),t)=c(t)(\ell(x)-\mu(t))$ . c(t) is computed via Equation (14) so that  $E[\Delta\mu]=0$  for the plots in the first column. Equation (19) was used for the other plots so that  $E[\mu(t)]=\gamma(t)$ .  $\gamma(t)=30\sin(t/80)+\mu(0)$  in the second column,  $\gamma(t)=t+\mu(0)$  in the fourth column and  $\gamma(t)=\mu(0)$  in the fifth column. In the plots in the third column bloat control with  $\gamma(t)=200$  was activated when mean program size reached 200. Results are the average mean size over 100 independent runs. Note: the range of the y-axes vary across the plots.

"Local" case in the figure) is quite apparent when compared to the very tight control obtained in the other configurations. When one considers, however, that there is no size limit or other form of bloat control being used, having the mean sizes in that case remain below 300 for 500 generations is still a significant achievement.

Performance comparisons are not the focus of this paper. However, since virtually all bloat control methods need to balance parsimony and solution accuracy, it is reasonable to ask what sort of performance implications the use of our covariance-based bloat-control technique implies. As shown in Table 2 for the two polynomial regression problems, there is generally very little performance loss associated with the use of our technique, and several of the configurations in fact increase the success rates.

### 6. CONCLUSIONS

For many years scientists, engineers and practitioners in the GP community have used the parsimony pressure method to control bloat in genetic programming. Although more recent and sophisticated techniques exist, parsimony pressure remains the most widely known and used technique.

The method suffers from two problems. Firstly, although good control of bloat can be obtained with a careful choice of the parsimony coefficient, such a choice is difficult and is often simply done by trial and error. Secondly, while it is clear that a constant parsimony coefficient can only achieve partial control over the dynamics of the average program size over time, no practical method to choose the parsimony coefficient dynamically and efficiently is available. The work presented in this paper changes all of this.

Starting from the size evolution equation developed in [26], we have developed a theory that tells us how to practically and optimally set the parsimony coefficient dynamically during a run so as to achieve complete control over the growth of the programs in a population. The method is extremely general, applying to a large class of control strategies of which the classical parsimony pressure method is an instance. Experimental results with three different GP systems, using different selection strategies and 5 different problems all strongly confirm the effectiveness of the method.

Dolo	A mti	Standard		
Poly	Anti-bloat		Success	
Degree	target	penalty	Rate	Deviation
6	none		0.77	0.04
6	$\Delta\mu=0$	$c\ell$	0.83	0.04
6	sin	$c\ell$	0.77	0.04
6	limit	$c\ell$	0.86	0.03
6	linear	$c\ell$	0.83	0.04
6	$\mu = \mu_0$	$c\ell$	0.83	0.04
6	$\Delta \mu = 0$		0.70	0.05
6	sin	$c\ell^{-1}$	0.77	0.04
6	limit	$c\ell^{-1}$	0.80	0.04
6	linear	$c\ell^{-1}$	0.79	0.04
6	$\mu = \mu_0$	$c\ell^{-1}$	0.71	0.05
8	none		0.24	0.04
8	$\Delta\mu=0$	$c\ell$	0.37	0.05
8	sin	$c\ell$	0.47	0.05
8	limit	$c\ell$	0.41	0.05
8	linear	$c\ell$	0.36	0.05
8	$\mu = \mu_0$	$c\ell$	0.35	0.05
8	$\Delta\mu=0$	$c\ell^{-1}$	0.26	0.04
8	sin	$c\ell^{-1}$	0.32	0.05
8	limit	$c\ell^{-1}$	0.26	0.04
8	linear	$c\ell^{-1}$	0.23	0.04
8	$\mu = \mu_0$	$c\ell^{-1}$	0.20	0.04

Table 2: Success rate comparison for Poly-6 and Poly-8 runs with different bloat control settings.

### 7. REFERENCES

- [1] L. Altenberg. The Schema Theorem and Price's Theorem. In L. D. Whitley and M. D. Vose, editors, *Foundations of Genetic Algorithms 3*, Estes Park, Colorado, USA, 31 July–2 Aug. 1994. Morgan Kaufmann. Published 1995.
- [2] P. J. Angeline. An investigation into the sensitivity of genetic programming to the frequency of leaf selection during subtree crossover. In J. R. Koza, et al., editors, *Genetic Programming 1996: Proceedings of the First Annual Conference*, pages 21–29, Stanford University, CA, USA, 28–31 July 1996. MIT Press.
- [3] W. Banzhaf and W. B. Langdon. Some considerations on the

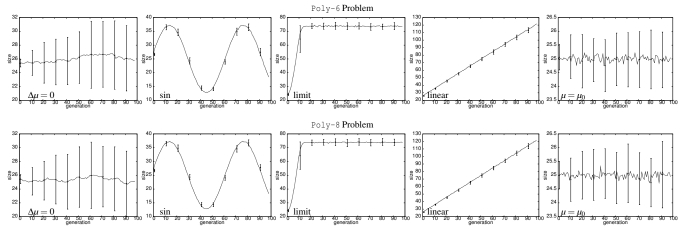


Figure 3: Size control obtained in the Poly-6 and Poly-8 problems with populations of size 1,000 using the penalty function  $g(\ell(x),t)=\frac{c(t)}{\ell(x)}$ . c(t) is computed via Equation (15) with k=-1 so that  $E[\Delta\mu]=0$  for the plots in the first column. It was computed via Equation (18) (with the same k) so that  $E[\mu(t)]=\gamma(t)$  for the remaining plots.  $\gamma(t)=12.5\sin(t/10)+\mu(0)$  in the second column,  $\gamma(t)=t+\mu(0)$  in the fourth column and  $\gamma(t)=\mu(0)$  in the fifth column. In the plots in the third column bloat control with  $\gamma(t)=75$  was activated only when mean program size reached 50. Results are the average mean size over 100 independent runs. Population sizes 100 and 10,000 provided qualitatively similar behaviours. Note: the range of the y-axes vary across the plots.

- reason for bloat. *Genetic Programming and Evolvable Machines*, 3(1):81–91, Mar. 2002.
- [4] R. Crawford-Marks and L. Spector. Size control via size fair genetic operators in the PushGP genetic programming system. In W. B. Langdon, et al., editors, GECCO 2002: Proceedings of the Genetic and Evolutionary Computation Conference, pages 733–739, New York, 9-13 July 2002. Morgan Kaufmann Publishers.
- [5] S. Dignum and R. Poli. Generalisation of the limiting distribution of program sizes in tree-based genetic programming and analysis of its effects on bloat. In D. Thierens, et al., editors, GECCO '07: Proceedings of the 9th annual conference on Genetic and evolutionary computation, volume 2, pages 1588–1595, London, 7-11 July 2007. ACM Press.
- [6] A. Ekart and S. Z. Nemeth. Selection based on the pareto nondomination criterion for controlling code growth in genetic programming. *Genetic Programming and Evolvable Machines*, 2(1):61–73, Mar. 2001.
- [7] H. Iba. Complexity-based fitness evaluation for variable length representation. Position paper at the Workshop on Evolutionary Computation with Variable Size Representation at ICGA-97, 20 July 1997.
- [8] H. Iba, H. de Garis, and T. Sato. Genetic programming using a minimum description length principle. In K. E. Kinnear, Jr., editor, *Advances in Genetic Programming*, chapter 12, pages 265–284. MIT Press, 1994.
- [9] H. Iba, H. de Garis, and T. Sato. Temporal data processing using genetic programming. In L. Eshelman, editor, *Genetic Algorithms: Proceedings of the Sixth International Conference (ICGA95)*, pages 279–286, Pittsburgh, PA, USA, 15-19 July 1995. Morgan Kaufmann.
- [10] K. E. Kinnear, Jr. Evolving a sort: Lessons in genetic programming. In *Proceedings of the 1993 International Conference on Neural Networks*, volume 2, pages 881–888, San Francisco, USA, 28 Mar.-1 Apr. 1993. IEEE Press.
- [11] K. E. Kinnear, Jr. Fitness landscapes and difficulty in genetic programming. In *Proceedings of the 1994 IEEE World*

- Conference on Computational Intelligence, volume 1, pages 142–147, Orlando, Florida, USA, 27-29 June 1994. IEEE Press.
- [12] M. Kotanchek, G. Smits, and E. Vladislavleva. Pursuing the pareto paradigm tournaments, algorithm variations & ordinal optimization. In R. L. Riolo, et al., editors, *Genetic Programming Theory and Practice IV*, volume 5 of *Genetic and Evolutionary Computation*, chapter 3. Springer, Ann Arbor, 11-13 May 2006.
- [13] J. R. Koza. Genetic Programming: On the Programming of Computers by Means of Natural Selection. MIT Press, Cambridge, MA, USA, 1992.
- [14] W. B. Langdon. The evolution of size in variable length representations. In 1998 IEEE International Conference on Evolutionary Computation, pages 633–638, Anchorage, Alaska, USA, 5-9 May 1998. IEEE Press.
- [15] W. B. Langdon. Size fair and homologous tree genetic programming crossovers. *Genetic Programming and Evolvable Machines*, 1(1/2):95–119, Apr. 2000.
- [16] W. B. Langdon and R. Poli. Fitness causes bloat. In P. K. Chawdhry, et al., editors, *Soft Computing in Engineering Design and Manufacturing*, pages 13–22. Springer-Verlag London, 23-27 June 1997.
- [17] W. B. Langdon and R. Poli. *Foundations of Genetic Programming*. Springer-Verlag, 2002.
- [18] W. B. Langdon, T. Soule, R. Poli, and J. A. Foster. The evolution of size and shape. In L. Spector, et al., editors, *Advances in Genetic Programming 3*, chapter 8, pages 163–190. MIT Press, Cambridge, MA, USA, June 1999.
- [19] S. Luke. Two fast tree-creation algorithms for genetic programming. *IEEE Transactions on Evolutionary Computation*, 4(3):274–283, Sept. 2000.
- [20] N. F. McPhee and J. D. Miller. Accurate replication in genetic programming. In L. Eshelman, editor, Genetic Algorithms: Proceedings of the Sixth International Conference (ICGA95), pages 303–309, Pittsburgh, PA, USA, 15-19 July 1995. Morgan Kaufmann.
- [21] N. F. McPhee and R. Poli. Using schema theory to explore

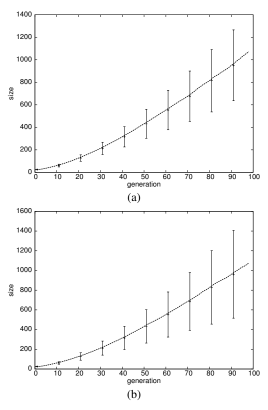


Figure 4: Behaviour of the mean program size in a linear GP system when solving the Poly-6 problem (a) and the Poly-8 problem in the absence of bloat control for populations of size 1,000. Results are averages over 100 independent runs.

- interactions of multiple operators. In W. B. Langdon, et al., editors, *GECCO 2002: Proceedings of the Genetic and Evolutionary Computation Conference*, pages 853–860, New York, 9-13 July 2002. Morgan Kaufmann Publishers.
- [22] R. Poli. General schema theory for genetic programming with subtree-swapping crossover. In *Genetic Programming*, *Proceedings of EuroGP 2001*, LNCS, Milan, 18-20 Apr. 2001. Springer-Verlag.
- [23] R. Poli. A simple but theoretically-motivated method to control bloat in genetic programming. In C. Ryan, et al., editors, Genetic Programming, Proceedings of the 6th European Conference, EuroGP 2003, LNCS, pages 211–223, Essex, UK, 14-16 Apr. 2003. Springer-Verlag.
- [24] R. Poli, W. B. Langdon, and S. Dignum. On the limiting distribution of program sizes in tree-based genetic programming. In M. Ebner, et al., editors, *Proceedings of the* 10th European Conference on Genetic Programming, volume 4445 of LNCS, pages 193–204, Valencia, Spain, 11-13 Apr. 2007. Springer.
- [25] R. Poli, W. B. Langdon, and N. F. McPhee. *A field guide to genetic programming*. Published via http://lulu.com and freely available at http://www.gp-field-guide.org.uk, 2008. (With contributions by J. R. Koza).
- [26] R. Poli and N. F. McPhee. General schema theory for genetic programming with subtree-swapping crossover: Part II. Evolutionary Computation, 11(2):169–206, June 2003.
- [27] G. R. Price. Selection and covariance. *Nature*, 227, August 1:520–521, 1970.
- [28] J. Rosca. Generality versus size in genetic programming. In

### Avg size vs. time, different target size functions

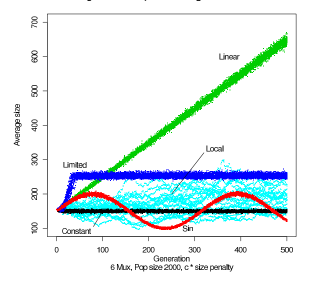


Figure 5: Scatterplot of the average size over multiple runs of the 6-MUX problem with various size target functions. The population size was 2,000 and we used the penalty function  $f-c\ell$ . The "Constant" case had a constant target size of 150. "Sin" had the target size function  $\sin((t+1)/50.0)\times 50.0+150$ . "Linear" had the target function 150+t. "Limited" used no size control until the size exceeded the limit 250, after which a constant target of 250 was used. "Local" used a target of  $\Delta\mu=0$ .

- J. R. Koza, et al., editors, *Genetic Programming 1996: Proceedings of the First Annual Conference*, pages 381–387,
  Stanford University, CA, USA, 28–31 July 1996. MIT Press.
- [29] J. Rosca. A probabilistic model of size drift. In R. L. Riolo and B. Worzel, editors, *Genetic Programming Theory and Practice*, chapter 8, pages 119–136. Kluwer, 2003.
- [30] J. P. Rosca. Analysis of complexity drift in genetic programming. In J. R. Koza, et al., editors, *Genetic Programming 1997: Proceedings of the Second Annual Conference*, pages 286–294, Stanford University, CA, USA, 13-16 July 1997. Morgan Kaufmann.
- [31] J. P. Rosca and D. H. Ballard. Complexity drift in evolutionary computation with tree representations. Technical Report NRL5, University of Rochester, Computer Science Department, Rochester, NY, USA, Dec. 1996.
- [32] J. P. Rosca and D. H. Ballard. Rooted-tree schemata in genetic programming. In L. Spector, et al., editors, *Advances* in *Genetic Programming 3*, chapter 11, pages 243–271. MIT Press, Cambridge, MA, USA, June 1999.
- [33] T. Soule and J. A. Foster. Effects of code growth and parsimony pressure on populations in genetic programming. *Evolutionary Computation*, 6(4):293–309, Winter 1998.
- [34] T. Soule and J. A. Foster. Removal bias: a new cause of code growth in tree based evolutionary programming. In 1998 IEEE International Conference on Evolutionary Computation, pages 781–186, Anchorage, Alaska, USA, 5-9 May 1998. IEEE Press.
- [35] B.-T. Zhang and H. Mühlenbein. Balancing accuracy and parsimony in genetic programming. *Evolutionary Computation*, 3(1):17–38, 1995.