Faster Genetic Programming GPquick via multicore and Advanced Vector Extensions

William B. Langdon (UCL) and Wolfgang Banzhaf (MSU)

Fax: +44 (0)20 7679 1397

Abstract

We evolve floating point Sextic polynomial populations of genetic programming binary trees for up to a million generations. Programs with almost 400,000,000 instructions are created by crossover. To support unbounded Long-Term Evolution Experiment LTEE GP we use both SIMD parallel AVX 512 bit instructions and 48 threads to yield performance of up to 139 billion GP operations per second, 139 giga GPops, on a single Intel Xeon Gold 6126 2.60 GHz server.
Figure 1: Evolution of mean absolute error in ten runs of Sextic polynomial [Koza (1992)] with population of 4000. (Runs aborted after first crossover to hit 15 million node limit.) End of run label gives number of generations when fitness got better (five shown at top right to avoid crowding).

**keywords:** genetic algorithms, genetic programming, GP, Convergence, Long-Term Evolution Experiment LTEE Extended unbounded evolution

1 Introduction

A couple of years we noted [Langdon (2017)] Rich Lenski’s experiments in long term evolution [Lenski et al. (2015)] in which the BEACON team evolved bacteria for more than 70,000 generations and found continued beneficial adaptive mutations. This prompted us to ask the same question in computation based evolution. We started to investigate what happens if we allow artificial evolution, specifically genetic programming (GP) with crossover [Koza (1992); Banzhaf et al. (1998); Poli et al. (2008)], to evolve for tens of thousands, even a hundred thousand generations. Since then new hardware has become available and we build a new GP engine based on Andy Singleton’s GPQUICK (Section 2). This allows us to switch from the Boolean to the continuous domain and run experiments of up to a million generations. Excluding some special applications or Boolean benchmarks based on graphics hardware (GPUs), at up to 139 billion GP operations per second (139 giga GPops), this appears to be the fastest single computer GP system [Langdon (2013), Tab. 3].

In the Boolean domain we found usually the population quickly found a best possible answer and then retained it exactly for thousands of generations. Nonetheless under subtree crossover we reported interesting population evolution with trees continuing to evolve. Indeed we were able to report the first signs of an eventual end of bloat due to fitness convergence of the whole population. We can now report in the continuous domain we do see (like the bacteria experiments) continual innovation and improvement in fitness. Figure 1 show that although the rate of innovation falls (as in Lenski’s E. Coli populations), typically better solutions are found even towards the end of the runs and, in these runs, there are several hundred or even a few thousand generations where sub-tree crossover between evolved parents gave a better child.

---

1 The E. Coli genome contains 4.6 million DNA base pairs.
We are going to run GP far longer than is normally done. Firstly in search of continual evolution but also noting that it is sometimes not safe to extrapolate from the first hundred or so generations. As an example, McPhee and Poli (2001, sect. 1.2) said that his earlier studies which had reported only the first 100 generations could not safely be extrapolated to 3000 generations.

It must be admitted that without size control we expect bloat, and so we need a GP system not only able to run for \(10^6\) generations but also able to process trees with well in excess of a 100 million nodes. The new system we use is based on Singleton’s GPQuick (Singleton, 1994; Keith and Martin, 1994; Langdon, 1998) but enhanced to take advantage of both multi-core computing using pthreads and Intel’s SIMD AVX parallel floating point operations (Section 2). Keith and Martin (1994) say GPQuick’s linearisation of the GP tree will be hard to parallelise. Nevertheless, GPQUICK was rewritten to use 16 fold Intel AVX-512 instructions to do all operations on each node in the GP tree immediately. Leading to a single eval pass and better cache locality but at the expense of keeping a \(T = 48\) wide stack of partial results per thread.

Although the populations never lose genetic diversity (Koza’s variety, Koza (1992)), with strong tournament selection (parent selection tournaments of size seven, see Table 1) even the larger populations do tend to converge to have identical fitness values. However 100% fitness convergence is only seen in long runs with smaller populations (500 or 48 trees). In contrast, in the Boolean domain Langdon (2017), even in bigger populations (500) there are many generation where the whole population has identical fitness (but again variety is 100%).

The next section describes how GPQUICK was adapted to take advantage of Intel SIMD instructions able to process 16 floating point numbers in parallel and Posix threads to perform crossover and fitness evaluation on 48 cores simultaneously. Section 3 describes the floating point benchmark (Table 1 and Figure 4). Whilst Section 4 describes the evolution of fitness, size and depth in populations of 4000, 500 and 48 trees. It finds the earlier predictions of sub-quadratic bloat (Langdon, 1999a) and Flajolet limit (depth \(\approx \sqrt{2\pi |\text{size}|}\)) to essentially hold. In Section 5 there is a short discussion about the continuous evolution permitted by floating point benchmarks before we conclude in Section 6.

2 GPQUICK

First we describe how GPQUICK is used with the Sextic polynomial regression problem and then how GPQUICK has been modified to run in parallel, Sections 2.2 and 2.3.

2.1 Sextic and GPQuick

Andy Singleton’s GPQUICK (Singleton, 1994) is a well-established fast and memory efficient C++ GP framework. In steady state mode it stores GP trees in just one byte per tree node. (Originally it supported only steady state GAs. Ages ago a quick conversion to support generational GAs, with separate parent and child populations doubled this, although Koza (1992) shows doubling is not necessary.) The 8 bit opcode per tree node allows GPQUICK to support a number of different functions and inputs. Typically (as in these experiments) the remaining opcodes are used to support about 250 fixed ephemeral
Figure 2: 11 extended runs pop=48. Numbers on right indicate size of largest tree before the run stopped in millions of nodes. One run (*) converged so that more than 90% of the trees contain just five nodes. Three of the other four runs that reached 1 million generations (red) took between half a day and five days. In all but one run (*) we see repeated substantial bloat (> 64 million nodes) and subsequent tree size collapse. Seven runs, in black, terminated due to running out of memory (on server with 46GB).

Figure 3: Size and convergence in last 20% of a typical extended run pop=48. (First run “64” in Figure 2.) Solid blue curve shows almost all the time there are less than 2 trees with the maximum fitness (rescaled by $\times 5 \times 10^6$ and smoothed over 30 generations). In populations with huge trees, there are many generations where the whole population has identical fitness. Without a fitness differential, tree size may rise or fall.
random constants [Poli et al., 2008]. In the Sextic polynomial we have the traditional four binary floating point operations (+, −, × and protected division), an input (x) and 250 constants. The constants are chosen at random from the 2001 floating point numbers from -1.000 to +1.000. By chance neither end point nor 0.000 were chosen (see Table 1).

The continuous test cases (x) are selected at random from the interval -1 to +1. At the same time the target value y is calculated (Table 1). Since both x and y are stored in a text file, there may be slight floating point rounding errors from the standard float⇔string conversions.

Whereas the Sextic polynomial is usually solved with 50 test cases [Langdon et al., 1999], since the AVX hardware naturally supports multiples of 16, in our experiments we change this to 48 (i.e. $3 \times 16$) (Table 1). The multi-core servers we use each support 48 threads and for in the longest extended runs, we reduce the population to 48 (whereas in Langdon (2017) the smallest population considered contained 50 trees).

### 2.2 AVX GPQuick

GPQUICK stores the GP population by flattening each tree into a linear buffer, with the root node at the start. To avoid heap fragmentation the buffers are all of the same size. Traditionally the buffer is interpreted once per test case by multiple recursive calls to EVAL and the tree’s output is retrieved from the return value of the outer most EVAL. Each nested EVAL moves the instruction pointer on one position in the tree’s buffer, decodes the opcode there and calls the corresponding function. In the case of inputs x and constants a value is returned via EVAL immediately, whereas ADD, SUB, MUL and DIV will each call EVAL twice to obtain their arguments before operating on them and returning the result. For speed GPQUICK’s FASTEVAL, does an initial pass though the buffer and replaces all the opcodes by the address of the corresponding function that EVAL would have called. This expands the buffer 16 fold, but the expanded buffer is only used during evaluation and can be reused by every member of the population. Thus originally EVAL processed the tree $T + 1$ times (for $T$ test cases).

The Intel AVX instructions process up to 16 floating point data simultaneously. The AVX version of EVAL was rewritten to take advantage of this. Indeed as we expect trees that are far bigger than the CPU cache (≈ 16 million bytes, depending on model), EVAL was rewritten to process each tree’s buffer only once. This is achieved by EVAL processing all of the test cases for each opcode, instead of processing the whole of the tree on one test case before moving on to the test case. Whereas before each recursive call to EVAL returned a single floating point value, now it has to return 48 floating point values. This was side stepped by requiring EVAL to maintain an external stack where each stack level contains 48 floating point values. The AVX instructions operate directly on the top of this stack and EVAL keeps track of which instruction is being interpreted, where the top of the stack is, and (with PTHREADS) which thread is running it. Small additional arrays are used to allow fast translation from opcode to address of eval function, and constant values. AVX instructions are used to speed loading each constant into the top stack frame. Similarly all 48 test cases (x) are rapidly loaded on to the top of the stack. However the true power comes from being able to use AVX to process the top of the stack and the adjacent stack frame (holding a total of 96 floats) in essentially three instructions to give 48 floating point results.

The depth of the evaluation stack is simply the depth of the GP tree. GPQUICK uses a fixed buffer length for everyone in the GP population. This is fixed by the user at the start of the GP run. Fixing the buffer size also sets the maximum tree size. Although in principle this only places a very weak limit on GP tree depth, it has been repeatedly observed [Langdon, 1999c, 2000b] that evolved trees are roughly shaped like random trees. The mathematics of trees is well studied [Sedgewick and Flajolet, 1996] in particular the depth of random binary trees tends to a limit $2\sqrt{\pi \frac{\text{treesize}}{2}} + O(\text{treesize}^{1/4+\epsilon})$ (Sedgewick and Flajolet, 1996, page 256). (See Flajolet limit in Figures 6, 7, 11 and 15.) Thus the user specified tree size limit can be readily converted into an expected maximum depth of evolved trees. The size of the AVX eval stack is set to this plus a suitable allowance for random fluctuations and $O(\text{treesize}^{1/4+\epsilon})$. Note, with very large trees, even allowing for the number of test cases and storing floats on the stack rather than byte sized opcodes, the evaluation stack is considerably smaller than the genome of the tree whose fitness it is calculating.
Although AVX allows reductions operations across a stack frame, these are not needed until the final conversion from output to fitness value. However although faster, the reduction operations manipulate the 48 numbers in a different order and so may (within floating point tolerances) produce different answers. Since the reduction is a tiny part of the whole fitness evaluation we decided instead to ensure the AVX version produces identical results to the original system and so the final fitness evaluation is done with a conventional for loop.

### 2.3 PTHREADS GPQuick

The second major change to GPQUICK was to delay fitness evaluation so that the whole new population can have its fitness evaluated in parallel. (This means PTHREADS can only be applied when GPQUICK is operating in generational mode.) If pThreads > 0, the population fitness evaluation is spread across pThreads. As trees are of different sizes, each fitness evaluation will take different times. Therefore which tree is evaluated by which thread is decided dynamically. Due to timing variations, in an identical run, which tree is evaluated by which thread may be different. However great care is taken so that this cannot affect the course of evolution. (Although, for example, where trees have identical fitness, it can affect which is found first and therefore which is reported to the user).

EVAL requires a few data arrays. These are all allocated near the start of the GP run. Those that are read only can be shared by the threads. Each thread requires its own instance of read-write data. To avoid “false sharing”, care is taken to align read-write data on cache line boundaries (64 bytes), e.g. with additional padding bytes and (aligned). So that each thread writes to its own cache lines and therefore these cached data are not shared with other threads.

Surprisingly an almost doubling of speed was obtained by also moving crossover operations to these parallel threads. Since crossover involves random choices of parents and subtrees these were unchanged but instead of performing the crossover immediately a small amount of additional information was retained and to be read later by the threads. This allows the crossover to be delayed and performed in one of 48 C++ pthreads. The results are identical but give an additional ≈two fold speed up.

With intermediate sized trees, there may be some efficiency gain by evaluating the newly created chromosome immediately (i.e. whilst still in the hardware cache).

One thing that was less successful was to implement a strategy for minimising memory consumption by deleting parents immediately all their children have been created. (Koza’s generational scheme with crossover means at most $M + 2$ individuals need to be stored (where $M$ is the population size). In principle this grows to $M + 2t$ (where $t$ is the number of threads). This of course gives no saving where the population is the same size or smaller than the number of threads. Although getting the threads to free memory immediately works, C++ free is surprisingly slow in a multi-threaded environment (within threads it has to wait on locks to avoid corrupting its heap). This seriously impacted performance (unless the time taken by EVAL dominates). Therefore the calls to delete parents were moved back out of the threads to the surrounding sequential code.

In principle, even with multiple threads $t > 1$, the efficient reduction of memory from $2M$ to $M + 2t$ chromosomes should be possible. The number and size of all buffers is known in advance and it would be possible to manipulate the allocation and freeing explicitly in GPQUICK’s chrome.cxx C++ source directly (rather than using new and delete). Thus effectively building a specialised heap for the population but this has not been attempted.

### 3 Experiments

We use the well known Sextic polynomial benchmark (Koza[1994], Tab. 5.1). Briefly the task given to GP is to find an approximation to a sixth order polynomial, $x^6 - 2x^4 + x^2$, given only a fixed set of samples. I.e. a fixed number of test cases. For each test input $x$ we know the anticipated output $f(x)$, see
Figure 4: 48 test cases for Sextic Polynomial Benchmark.

Table 1: Long term evolution of Sextic polynomial symbolic regression binary trees

| Terminal set: X, 250 constants -0.995 to 0.997 |
| Function set: MUL, ADD, DIV, SUB |
| Fitness cases: 48 fixed input -0.97789 to 0.979541 (randomly selected from -1.0 to +1.0 input). |
| Selection: tournament size 7 with fitness = $\frac{1}{48} \sum_{i=1}^{48} |GP(x_i) - y_i|$ |
| Population: Panmictic, non-elitist, generational. |
| Parameters: Initial population (4000) ramped half and half Koza (1992) depth between 2 and 6. 100% unbiased subtree crossover. 10,000 generations (stop run if any tree reaches limit $15 \times 10^6$). |
| DIV is protected division $(y! = 0)$? $x/y : 1.0f$ |

Figure 4 and Table 1. Of course the real point is to investigate how GP works. How do GP populations evolve over time? In particular, even for such a simple continuous problem, it is possible for GP to continue to find improvements (as Lenski’s E.Coli are doing) or, like the Boolean case Langdon (2017), will the GP population get stuck early on and from then on never make further progress?

We ran three sets of experiments. In the first the new GP systems was set up as like the original Sextic polynomial runs which reported phenotypic convergence [Langdon et al. 1999, Fig. 8.5]. The first group use a population of 5000, the next 500 and the last 48. (Section 2.1 above has already described a few technical differences between these and our earlier experiments [Langdon et al. 1999].)

### 3.1 Crossover

Each generation is created entirely using Koza’s two parent subtree crossover [Koza (1992)]. (GPQuick creates one offspring per crossover.) For simplicity and in the hope that this would make GP populations easier to analyse, both subtrees (i.e. to be removed and to be inserted) are chosen uniformly at random. That is, we do not use Koza’s bias in favour of internal nodes (functions) at the expense of external nodes (leafs or inputs). Instead the root node of the subtree (to be deleted or to be copied) is chosen uniformly at random from the whole of the parent tree. This means there is more chance of subtree crossover simply moving leaf nodes and so many children will differ from the root node donating parent (the mum) by just one leaf.
As mentioned above (Section 2.3) once fitness evaluation has been sped up by parallel processing, for very long trees, producing the child is surprisingly large part of the remaining run time and so it too can be done in parallel. However the choice of crossover points is done in the usual way (i.e. in sequential code, not done in parallel) and so remains unaffected by multithreading. This ensures the variability introduced by multiple parallel threads does not change the course of evolution.

3.2 Fitness Function

The fitness of every member of every generation is calculated using the same fitness function as That is, baring rounding errors (page 4), fitness is given by the mean of the absolute difference between the value returned by the GP tree on each test case and the Sextic polynomial’s value for the same test input (see Table 1). However unlike Koza, we use tournament selection to chose both parents.

Like (Koza [1994] Tab. 5.1), we also keep track of the number of test case where each tree is close to the target (i.e. within 0.01, known as a “hit”). The number of hits is used for reporting the success of a GP run. It is not used internally during a GP run. Also our GP runs do not stop when a solution is found (48 hits) but continue until either the user specified number of generations is reached or bloat means the GP runs out of memory.

Where needed, floating point calculations are done in a fixed order, to avoid parallelism creating minor changes in calculated fitness, which could quickly cause otherwise identical runs to diverge because of implementation differences in parallel calculations. (Also mentioned above above in Section 2.2.)

4 Results

4.1 Results Population 4000 trees

In the first set of experiments, we use the standard population of 4000 trees. Table 2 summarises the results of 10 runs. In all cases GP found a reasonable approximation to the target (the Sextic polynomial). Indeed in all but one run (47 hits) the best trees score 48 out of 48 possible hits. I.e. they are within 0.01 on all 48 test cases. Indeed in most cases the average error was less than $10^{-4}$. Figure 1 shows that GP tends to creep up on the best match to the training data. Typically after several thousand generations, GP has progressively improved by more than a thousand increasingly small steps. (See Table 2 column 3 and Figure 1).

In all cases we do see enormous increases in size. In all ten runs with a population of 4000, the runs are stopped as they hit the size limit (15 000 000) before reaching 10 000 generations. Column 5 in Table 2 gives the size (in millions) of the largest evolved tree in each run. The log-log plot in Figure 5 shows a typical pattern of sub quadratic Langdon (2000a) increase in tree size. The straight line, shows a power law fit. In this run the best fit has an exponent of 1.2. Column 6 of Table 2 shows that the best fit between generations ten and a thousand for all 10 runs varies between 1.1 and 1.9.

As expected not only do programs evolve to be bigger but also they increase in depth. As described above (in Section 2.2) highly evolved trees tend to be randomly shaped and so as expected tend to lie near the Flajolet limit, depth $\approx \sqrt{\frac{2\pi}{\text{size}}}$ (see Figures 6 and 7).

In all ten runs we see some phenotypic convergence. The last column in Table 2 shows the peak fitness convergence. That is, out of 4000, the number of trees having exactly the same fitness as the best in the population. Typically at the start of the run (see Figure 8), the population contains mostly poorer trees, but later in the run the population begins to converge and towards the end of the run we may see hundreds of generations where more than 90% of the population have identical fitness. Under these circumstances, even with a tournament size as high as 7, many tournaments include potential parents with identical fitness. These, and hence the parents of the next generation, are decided entirely randomly. However, even in the most converged population there are at least two individuals with worse fitness. (In Figure 8 it is at least 19.) As we saw with the Boolean populations Langdon (2017), even this small number can be enough to drive bloat (albeit at a lower rate).
Table 2: 10 Sextic polynomial runs with population 4000

| Gens | smallest error | impr. | hits | size10^6 | power law | conv |
|------|----------------|-------|------|----------|-----------|------|
| 6370 | 0.0000064487   | 2139  | 48   | 14.329   | 1.200     | 3981 |
| 8298 | 0.000145796    | 2040  | 48   | 14.102   | 1.916     | 3982 |
| 2323 | 0.000642006    | 389   | 47   | 13.441   | 1.387     | 3995 |
| 7119 | 0.000507600    | 608   | 48   | 13.668   | 1.589     | 3997 |
| 11750| 0.0000000001   | 3583  | 48   | 13.854   | 1.364     | 3989 |
| 3412 | 0.000065562    | 1277  | 48   | 14.348   | 1.625     | 3986 |
| 5106 | 0.000071289    | 1615  | 48   | 14.233   | 1.146     | 3988 |
| 6112 | 0.000728757    | 1871  | 48   | 14.500   | 1.254     | 3983 |
| 6679 | 0.000028853    | 1741  | 48   | 14.022   | 1.396     | 3998 |
| 4454 | 0.000067817    | 790   | 48   | 14.900   | 1.227     | 3997 |

Figure 1 gives number of generations which improve on their parents, whereas here we give strictly better than anything previously evolved. Hence slight differences.

Figure 5: Evolution of tree size in first Sextic run (population 4000). (This run aborted after 6370 generations by first crossover to hit 15 million node limit.) Straight line shows best RMS error power law fit between generation 10 and 1000, \( y = 8.65 x^{1.2001} \)
Figure 6: Evolution of depth and size in population of 4000 trees in typical Sextic polynomial run. In this run, as predicted [Langdon 1999b], average trees lie within one or two standard deviations of random binary trees (Flajolet limit, depth \( \approx \sqrt{2\pi|\text{size}|} \), Sedgewick and Flajolet 1996, dotted parabola). See also Figure 7.

Figure 7: Plot of size and depth of the best individual in each generation for 10 Sextic polynomial runs with population of 4000. Binary trees must lie between short fat trees (lower curve “Full”) and “Tall” stringy trees. Most trees are randomly shaped and lie near the Flajolet limit (depth \( \approx \sqrt{2\pi|\text{size}|} \), solid line, note log-log scales). Figure 6 shows the first run in more detail.
4.2 Results Population 500 trees

We repeated the GP runs but allowed still larger trees to evolve by splitting the available memory between fewer trees by reducing the population from 4000 to 500. Figure 9 shows the evolution of the best fitness with the reduced population size. Notice two runs do not really solve the problem getting less than half the test cases (see “hits” column in Table 3). Nonetheless in all cases evolution continues to make progress and each GP runs finds several hundred or more small improvements (third column in Table 3).

Since we have deliberately extended the space available to the GP trees, it is no surprise that the trees grow even bigger than before (column 5 in Table 3). Again the bloat is approximately a power law. Although in one unsuccessful run we do see a power law exponent greater than 2, mostly growth is at a similar (sub-quadratic) rate as with the bigger population runs (1.4–2.2 vs 1.1–1.9, column 6 in Table 2 (pop 4000)). Figure 10 shows as expected we again see sub-quadratic growth in tree size between generations 10 and 1000. In fact the power law fit (<2.0), in the first GP run with a population of 500, seems to extrapolate well, even though the population starts to converge in later generations (see Figure 12).

Again randomly shaped trees evolve. Figure 11 shows the relationships between depth and size of the best trees in the population. As expected in all six runs it lies near the Flajolet limit, depth ≈ $\sqrt{2\pi |\text{size}|}$, for large binary trees.

Unlike with the large populations, all the runs with populations of 500 trees showed some cases of complete fitness convergence (last column in Table 3 is 500). Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run. Figure 12 shows fitness convergence in first Sextic polynomial pop=500 run.
Figure 9: Evolution of mean absolute error in first six runs of Sextic polynomial \cite{Koza1992} with population of 500. (Runs aborted running out of memory, 400 million node limit.) End of run label gives number of generations when fitness got better.

Table 3: 6 Sextic polynomial runs with population 500

| Gens    | smallest error | imp[p] hits | size10^6 | power law | conv |
|---------|----------------|-------------|-----------|-----------|------|
| 111582  | 0.000538248    | 3545        | 399.594   | 1.558     | 500  |
| 23937   | 0.034313100    | 757         | 202.439   | 1.736     | 500  |
| 35783   | 0.000307189    | 3484        | 227.488   | 1.436     | 500  |
| 43356   | 0.018373600    | 929         | 267.416   | 2.181     | 500  |
| 27713   | 0.000137976    | 5852        | 327.253   | 1.928     | 500  |
| 103953  | 0.001765590    | 664         | 230.106   | 1.408     | 500  |

Figure 9 gives number of generations which improve on their parents, whereas here we give strictly better than anything previously evolved. Hence slight differences.
Figure 10: Evolution of tree size in first Sextic run (population 500). (This run aborted after 111582 generations by first crossover to hit 400 million node limit.) Straight line shows best RMS error power law fit between generation 10 and 1000, $y = 1.3x^{1.56}$.

Figure 11: Plot of size and depth of the best individual in each generation for six Sextic polynomial runs with population of 500. Binary trees must lie between short fat trees (lower curve “Full”) and “Tall” stringy trees. Most trees are randomly shaped and lie near the Flajolet limit (depth $\approx \sqrt{2\pi|\text{size}|}$, solid line, note log-log scales).
Figure 12: Fitness convergence in first Sextic polynomial pop=500 run. In 30% of this run, the whole population has identical fitness (y=0). Mean prog size (linear scale, black) and best fitness (log, blue) plotted in the background.

4.3 Results Population 48 trees

In the final experiments the population was reduced still further to allow still larger trees to be evolved. Whereas in the Boolean experiments Langdon (2017) we reduced the population to 50, this was before we had access to servers with 48 cores. Therefore these smallest population runs were run with a population of 48, since this should readily map well to the available Intel mult-core servers.

With the small population, none of the runs solve the problems. Indeed only three runs got close on 40 or more test cases (see Figure 13 and Table 4). Of the remaining eight, only one finds a large number of fitness improvements. Seven runs contain only between 3 and 30 generations containing fitness improvement, column 3 in Table 4. In three of these, the population gets trapped at trees with just three nodes which evaluate to constants 0.0626506, 0.069169 and 0.0830508. Although eventually the population does eventually escape and large trees evolve by the end of the run. Except for these three runs, all the other runs contain populations where every member of the population has identical fitness. Therefore their maximum convergence is 48 (see last column in Table 4).

As expected, as with larger populations, the evolved highly binary trees are again approximately the same shape as random trees. See Figure 15.

Figure 16 shows for almost the whole run the best fitness in the population is fixed but once trees get big enough further size changes are essentially random. Notice fitness depends only on the sum of the absolute difference between the value returned by the GP tree and the target value. In particular the “hits” is only used for reporting. The best fitness found in this run is given by robust trees which always return a midpoint value (cf. Figure 4) which only passes close to four test points. Trees which closely matched more test points were discovered in the first nineteen generation of this run. However, in terms of fitness, they scored worse than a constant and so went extinct.
Figure 13: Evolution of mean absolute error in 11 runs of Sextic polynomial [Koza, 1992] with population of 48. (Up to a million generation or aborted on running out of memory, 500 million node limit.) End of run label gives number of generations when fitness got better. (Seven shown at top right to avoid crowding.)

Table 4: 11 Sextic polynomial runs with population of 48

| Gens    | smallest error | hits | size10^6 | power law | conv |
|---------|----------------|------|----------|-----------|------|
| 1000000 | 0.046215700    | 11   | 63.920   | 1.633     | 48   |
| 491618  | 0.002748230    | 745  | 396.576  | 2.060     | 48   |
| 1000000 | 0.046215700    | 7    | 190.654  | 1.448     | 48   |
| 689414  | 0.004857990    | 448  | 159.949  | 1.260     | 48   |
| 1000000 | 0.046215700    | 8    | 50.365   | 1.701     | 48   |
| 143251  | 0.046215700    | 11   | 99.541   | 1.672     | 48   |
| 212528  | 0.046650600    | 30   | 257.766  | na        | 42   |
| 1000000 | 0.046730800    | 3    | 0.000    | na        | 42   |
| 958147  | 0.023259300    | 1683 | 308.958  | 1.791     | 48   |
| 294098  | 0.047174400    | 3    | 308.121  | na        | 43   |
| 757830  | 0.002985980    | 2921 | 294.821  | 1.320     | 48   |

Figure 13 gives number of generations which improve on their parents, whereas here we give strictly better than anything previously evolved. Hence slight differences.
Figure 14: Evolution of tree size in first Sextic run with a population of 48. This run ran for a million generations. Straight line shows best RMS error power law fit between generation 10 and 1000, $y = 0.75x^{1.63}$.

Figure 15: Plot of size and depth of the best individual in each generation for eleven Sextic polynomial runs with population of 48.
Figure 16: Fitness convergence in first Sextic polynomial run with population of 48 trees. After generation 19 the best tree in every generation has a fitness of -0.0462157 (4 hits) (tree returns 0.0769947 regardless of test case, cf. Figure 4). In 90% of this run, the whole population has identical fitness (y=0). (Mean number of poor fitness children is 0.1315.) Mean prog size (linear scale, black) and best fitness (log, blue).

Figure 17: Fitness convergence in last Sextic polynomial run with population of 48 trees. (Run 295 in Figure 2.) In this run 67% of generations all 48 trees have identical fitness. The last improvement is found in generation 752 071 (5759 generation before the end of the run) in a tree of 13 196 331 nodes. It takes over the whole population in 4 generations. However in 21% of the remaining generations it does not totally dominate (mean number trees with lower fitness 0.394 per generation). Mean prog size (linear scale, black) and best fitness (log, blue).
5 Is there a Limit to Evolution?

In the Sextic Polynomial experiments with larger populations there is no hint of either evolution of fitness or bloat totally stopping. In the smaller populations, it is both possible to run evolution for longer and to allow trees to bloat to be even larger. Four of the eleven extended runs reached a million generations but in the remaining seven, bloat ran into memory limits and halted the run. Only in one run did we see anti-bloat, in which the population converged in a few generation on a small high fitness tree which crossover was able to replicate across a million generations. Interestingly two other runs found similar solution but after thousands of generations crossover found bloated version of them.

In the binary 6-Mux Boolean problem [Langdon (2017)] there are only 65 different fitness values. Therefore the number of fitness improvements is very limited. An end to bloat was found. By which we mean it was possible for trees to grow so large that crossover was unable to disrupt the important part of their calculation next to the root node and many generations were evolved where everyone had identical fitness. This lead to random selection and random fluctuations in tree size. (I.e. enormous trees but without a tendency for progressive end less growth.)

This did not happen here. Even in some of the smallest Sextic polynomials runs, we are still seeing innovation in the second half of the run, with tiny fitness improvements being created by crossover between enormous parents. Also we are still slightly short of total fitness convergence. Even with populations containing Sextic polynomial trees of hundreds of millions of nodes, crossover can still be disruptive and frequently even tiny populations can contain a tree of lower fitness. This is sufficient to provide some pressure (over thousands of generations) for tree size to increase on average.

Can bloat continue forever? It is still difficult to be definitive in our answer. We have seen cases where it does not and of course there are plenty of engineering techniques to prevent bloat. But we see other cases where crossover over thousands of generations can create an innovative child which allows bloat into a converged population of small trees. Perhaps more interestingly, we see crossover finding fitness improvement in bloated trees after many thousand of generations.

It is still an open question in continuous domains if, given sufficient memory and computational resources, bloat will always stifle innovation so completely that crossover will always only reproduce children of exactly the same fitness for long enough that the lack of selection pressure [Langdon and Poli (1997)] will in turn stifle bloat.

6 Conclusions

The availability of mult-core SIMD capable hardware has allowed us to push GP performance on single computers with floating point problems to that previously only approached with sub-machine code GP operating in discreet domains [Poli and Langdon (1999); Poli and Page (2000)]. This in turn has allowed GP runs far longer than anything previously attempted whilst evolving far bigger programs.

As expected, without size or depth limits or biases crossover with brutal selection pressure tends to evolve very large non-parsimonious programs. Known in the GP community as bloat (page 2 [Koza 1992, page 617]). After a few initial generations, GP tree bloat typically follows a sub-quadratic power law Langdon (2000a). But eventually effective selection pressure (Nordin, 1997, sec. 14.2), (Banzhaf et al., 1998, page 187), Stephens and Waelbroeck (1999), Langdon and Poli (2002) within highly evolved populations falls, leading to bloat at a reduced rate. However we only see the chaotic lack of bloat found in long running Boolean problems [Langdon (2017)] in a few unsuccessful runs with tiny populations (red plots in Figure 2 page 3). Nevertheless in all cases bloated binary trees evolve to be randomly shaped and lie close to Flajolet’s square root limit (Section 2.2).

Evolving binary Sextic polynomial trees for up to a million generations, during which some programs grow to four hundred million nodes, suggests even a simple floating point benchmark allows long term fitness improvement over thousands of generations.
Acknowledgements

This work was inspired by conversations at Dagstuhl Seminar 18052 on Genetic Improvement of Software.

The new parallel GPQuick code is available via http://www.cs.ucl.ac.uk/staff/W.Langdon/ftp/gp-code/GPavx.tar.gz

References

Altenberg, L. (1994). The evolution of evolvability in genetic programming. In Kinnear, Jr., K. E., editor, Advances in Genetic Programming, chapter 3, pages 47–74. MIT Press.

Angeline, P. J. (1994). Genetic programming and emergent intelligence. In Kinnear, Jr., K. E., editor, Advances in Genetic Programming, chapter 4, pages 75–98. MIT Press.

Banzhaf, W., Nordin, P., Keller, R. E., and Francone, F. D. (1998). Genetic Programming – An Introduction; On the Automatic Evolution of Computer Programs and its Applications. Morgan Kaufmann, San Francisco, CA, USA.

Keith, M. J. and Martin, M. C. (1994). Genetic programming in C++: Implementation issues. In Kinnear, Jr., K. E., editor, Advances in Genetic Programming, chapter 13, pages 285–310. MIT Press.

Koza, J. R. (1992). Genetic Programming: On the Programming of Computers by Means of Natural Selection. MIT Press, Cambridge, MA, USA.

Koza, J. R. (1994). Genetic Programming II: Automatic Discovery of Reusable Programs. MIT Press, Cambridge Massachusetts.

Langdon, W. B. (1998). Genetic Programming and Data Structures: Genetic Programming + Data Structures = Automatic Programming!, volume 1 of Genetic Programming. Kluwer, Boston.

Langdon, W. B. (1999a). Linear increase in tree height leads to sub-quadratic bloat. In Haynes, T. et al., editors, Foundations of Genetic Programming, pages 55–56, Orlando, Florida, USA.

Langdon, W. B. (1999b). Scaling of program tree fitness spaces. Evolutionary Computation, 7(4):399–428.

Langdon, W. B. (1999c). Size fair and homologous tree genetic programming crossovers. In Banzhaf, W. et al., editors, Proceedings of the Genetic and Evolutionary Computation Conference, volume 2, pages 1092–1097, Orlando, Florida, USA. Morgan Kaufmann.

Langdon, W. B. (2000a). Quadratic bloat in genetic programming. In Whitley, D. et al., editors, Proceedings of the Genetic and Evolutionary Computation Conference (GECCO-2000), pages 451–458, Las Vegas, Nevada, USA. Morgan Kaufmann.

Langdon, W. B. (2000b). Size fair and homologous tree genetic programming crossovers. Genetic Programming and Evolvable Machines, 1(1/2):95–119.

Langdon, W. B. (2013). Large scale bioinformatics data mining with parallel genetic programming on graphics processing units. In Tsutsui, S. and Collet, P., editors, Massively Parallel Evolutionary Computation on GPGPUs, Natural Computing Series, chapter 15, pages 311–347. Springer.

Langdon, W. B. (2017). Long-term evolution of genetic programming populations. In GECCO 2017: The Genetic and Evolutionary Computation Conference, pages 235–236, Berlin. ACM.

Langdon, W. B. and Poli, R. (1997). Fitness causes bloat. In Chawdhry, P. K. et al., editors, Soft Computing in Engineering Design and Manufacturing, pages 13–22. Springer-Verlag London.

Langdon, W. B. and Poli, R. (2002). Foundations of Genetic Programming. Springer-Verlag.

Langdon, W. B., Soule, T., Poli, R., and Foster, J. A. (1999). The evolution of size and shape. In Spector, L. et al., editors, Advances in Genetic Programming 3, chapter 8, pages 163–190. MIT Press, Cambridge, MA, USA.

Lenski, R. E. et al. (2015). Sustained fitness gains and variability in fitness trajectories in the long-term evolution experiment with Escherichia coli. Proceedings of the Royal Society B, 282(1821).

McPhee, N. F. and Poli, R. (2001). A schema theory analysis of the evolution of size in genetic programming with linear representations. In Miller, J. F. et al., editors, Genetic Programming, Proceedings of EuroGP’2001, volume 2038 of LNCS, pages 108–125, Lake Como, Italy. Springer-Verlag.

Nordin, P. (1997). Evolutionary Program Induction of Binary Machine Code and its Applications. PhD thesis, der Universitat Dortmund am Fachbereich Informatik.
Poli, R. and Langdon, W. B. (1999). Sub-machine-code genetic programming. In Spector, L. et al., editors, Advances in Genetic Programming 3, chapter 13, pages 301–323. MIT Press, Cambridge, MA, USA.

Poli, R., Langdon, W. B., and McPhee, N. F. (2008). A field guide to genetic programming. Published via http://lulu.com and freely available at http://www.gp-field-guide.org.uk. (With contributions by J. R. Koza).

Poli, R. and Page, J. (2000). Solving high-order Boolean parity problems with smooth uniform crossover, sub-machine code GP and demes. Genetic Programming and Evolvable Machines, 1(1/2):37–56.

Sedgewick, R. and Flajolet, P. (1996). An Introduction to the Analysis of Algorithms. Addison-Wesley.

Singleton, A. (1994). Genetic programming with C++. BYTE, pages 171–176.

Stephens, C. and Waelbroeck, H. (1999). Schemata evolution and building blocks. Evolutionary Computation, 7(2):109–124.

Syswerda, G. (1990). A study of reproduction in generational and steady state genetic algorithms. In Rawlings, G. J. E., editor, Foundations of genetic algorithms, pages 94–101. Morgan Kaufmann, Indiana University. Published 1991.

Tackett, W. A. (1994). Recombination, Selection, and the Genetic Construction of Computer Programs. PhD thesis, University of Southern California, Department of Electrical Engineering Systems, USA.