Towards Transparent and Parsimonious Methods for Automatic Performance Tuning

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The Need for Autotuning

1.1 Introduction

High Performance Computing has been a cornerstone of scientific and industrial progress for at least five decades. By paying the cost of increased complexity, software and hardware engineering advances continue to overcome several challenges on the way of the sustained performance improvements observed during the last fifty years. A consequence of this mounting complexity is that reaching the theoretical peak hardware performance for a given program requires not only expert knowledge of specific hardware architectures, but also mastery of programming models and languages for parallel and distributed computing.

If we state performance optimization problems as *search* or *learning* problems, by converting implementation and configuration choices to *parameters* which might affect performance, we can draw from and adapt proven methods from search, mathematical optimization, and statistical learning. The effectiveness of these adapted methods on performance optimization problems varies greatly, and hinges on practical and mathematical properties of the problem and the corresponding *search space*. The application of such methods to the automation of performance tuning for specific hardware, under a set of *constraints*, is named *autotuning*.

Improving performance also relies on gathering application-specific knowledge, which entails extensive experimental costs since, with the exception of linear algebra routines, theoretical peak performance is not always a reachable comparison baseline. When adapting methods for autotuning we must face challenges emerging from practical properties, such as restricted time and cost budgets, constraints on feasible parameter values, and the need to mix *categorical*, *continuous*, and *discrete* parameters. To achieve useful results we must also choose methods that make hypotheses compatible with problem search spaces, such as the existence of *discoverable*, or at least *exploitable*, relationships between parameters and performance. Choosing an autotuning method requires balancing the exploration of a problem, that is, seeking to discover and explain relationships between parameters and performance, and the exploitation of known or discovered relationships, seeking only to

find the best possible performance.

The contributions of this thesis are strategies to apply to program autotuning the statistical learning methods of *Design of Experiments* [1], or *Experimental Design*, and *Gaussian Process Regression* [2]. This thesis presents background and a high-level view of the theoretical foundations of each method, and detailed discussions of the challenges involved in specializing the general definitions of search heuristics and statistical learning methods to different autotuning problems, as well as what can be *learned* about specific autotuning search spaces, and how that acquired knowledge can be leveraged for further optimization.

This chapter aims to substantiate the claim that autotuning methods have a fundamental role to play on the future of program performance optimization, arguing that the value and the difficulty of the efforts to carefully tune software became more apparent ever since advances in hardware stopped leading to effortless performance improvements, at least from the programmer's perspective. The following sections discuss the historical context for the changes in trends on computer architecture, and characterize the search spaces found when optimizing performance on different domains.

1.2 Historical Trends in Hardware Design

The physical constraints imposed by technological advances on circuit design were evident since the first vacuum tube computers that already spanned entire floors, such as the ENIAC in 1945 [3]. The practical and economical need to fit more computing power into real estate is one force for innovation in hardware design that spans its history, and is echoed in modern supercomputers, such as the *Summit* from *Oak Ridge National Laboratory* [4], which spans an entire room.

Figure 1.1 highlights the unrelenting and so far successful pursuit of smaller transistor fabrication processes, and the resulting capability to fit more computing power on a fixed chip area. This trend was already observed in integrated circuits by Gordon Moore *et al.* in 1965 [5], who also postulated its continuity. The performance improvements produced by the design efforts to make Moore's forecast a self-fulfilling prophecy were boosted until around 2005 by the performance gained from increases in circuit frequency.

Robert Dennard *et al.* remarked in 1974 [8] that smaller transistors, in part because they generate shorter circuit delays, decrease the energy required to power a circuit and enable an increase in operation frequency without breaking power usage constraints. This scaling effect, named *Dennard's scaling*, is hindered primarily by leakage current, caused by quantum tunneling effects in small transistors. Figure 1.1 shows a marked stagnation on frequency increase after around 2005, as transistors crossed the 10²nm fabrication process. It was expected that leakage due to tunneling would limit frequency scaling strongly, even before the transistor fabrication process reached 10nm [9].

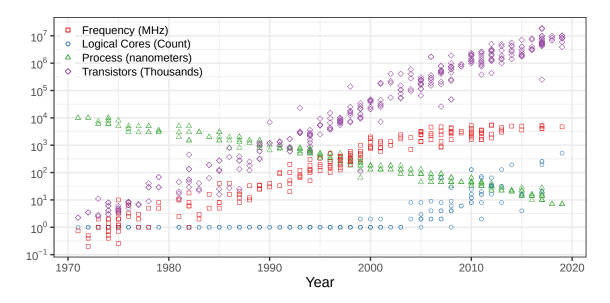


Figure 1.1: 49 years of microprocessor data, highlighting the sustained exponential increases and reductions on transistor counts and fabrication processes, the stagnation of frequency scaling around 2005, and one solution found for it, the simultaneous exponential increase on logical core count. Data from Wikipedia [6, 7]

Current hardware is now past the effects of Dennard's scaling. The increase in logical cores around 2015 can be interpreted as preparation for and mitigation of the end of frequency scaling, and ushered in an age of multicore scaling. Still, in order to meet power consumption constraints, up to half of a multicore processor could have to be powered down, at all times. This phenomenon is named *Dark Silicon* [10], and presents significant challenges to current hardware designers and programmers [11, 12, 13].

The Top500 [14] list gathers information about commercially available supercomputers, and ranks them by performance on the LINPACK benchmark [15]. Figure 1.2 shows the peak theoretical performance RPeak, and the maximum performance achieved on the LINPACK benchmark RMax, in Tflops/s, for the top-ranked supercomputers on TOP500. Despite the smaller performance gains from hardware design that are to be expected for post-Dennard's scaling processors, the increase in computer performance has sustained an exponential climb, sustained mostly by software improvements.

Although *hardware accelerators* such as GPUs and FPGAs, have also helped to support exponential performance increases, their use is not an escape from the fundamental scaling constraints imposed by current semiconductor design. Figure 1.3 shows the increase in processor and accelerator core count on the top-ranked supercomputers on Top500. Half of the top-ranked supercomputers in the last decade had accelerator cores and, of those, all had around ten times more accelerator than processor cores. The apparent stagnation of core count in top-ranked supercomputers, even considering accelerators, highlights the crucial impact software optimization has on performance.

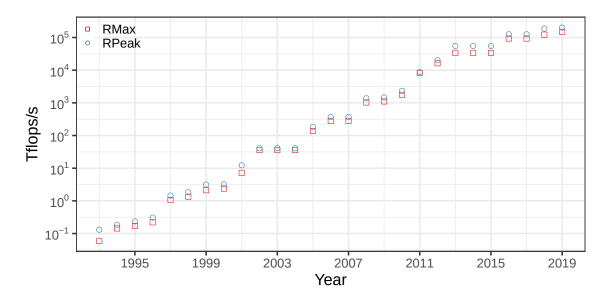


Figure 1.2: Sustained exponential increase of theoretical RPeak and achieved RMax performance for the supercomputer ranked 1^{st} on TOP500 [14]

Advances in hardware design are currently not capable of providing performance improvements via frequency scaling without dissipating more power than the processor was designed to support, which violates power constraints and risks damaging the circuit. From the programmer's perspective, effortless performance improvements from hardware have not been expected for quite some time, and the key to sustaining historical trends in performance scaling has lied in accelerators, parallel and distributed programming libraries, and fine tuning of several stages of the software stack, from instruction selection to the layout of neural networks.

The problem of optimizing software for performance presents its own challenges. The search spaces that emerge from autotuning problems grow quickly to a size for which it would take a prohibitive amount of time to determine the best configuration by exhaustively evaluating all possibilities. Although this means we must seek to decrease the amount of possibilities, by restricting allowed parameter values, or dropping parameters completely, it is often unclear how to decide which parameters should be restricted or dropped. The next section introduces a simple autotuning problem, presents an overview of the magnitude of the dimension of autotuning search spaces, and briefly introduces the methods commonly used to explore search spaces, some of which are discussed in detail in Chapter 3.

1.3 Characterizing Search Spaces

Algorithms for linear algebra problems are fundamental to scientific computing and statistics. Therefore, decreasing the execution time of algorithms such as general matrix multiplication (GEMM) [16], and others from the original BLAS [17], is an interesting and well

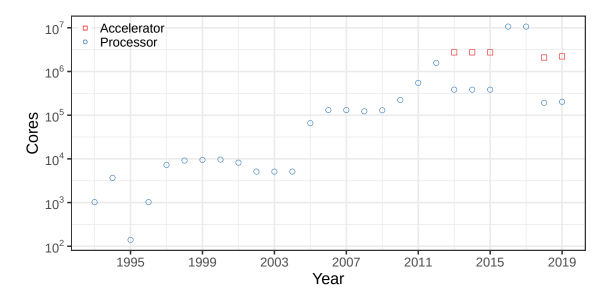


Figure 1.3: Processor and accelerator core count in supercomputers ranked 1st on TOP500 [14]. Core count trends for supercomputers are not necessarily bound to processor trends observed on Figure 1.1.

motivated example to introduce autotuning.

One way to improve the performance of such linear algebra programs is to exploit cache locality by reordering and organizing loop iterations, using source code transformation methods such as loop *tiling*, or *blocking*, and *unrolling*. We will now briefly describe loop tiling and unrolling for a simple linear algebra problem. After, we will discuss an autotuning search space for blocking and unrolling applied to GEMM, and how these transformations generate a relatively large and complex search space, which we can explore with autotuning methods.

For three square matrices A, B, and C, we wish to compute $C = C + A + B^{T}$.

Figure 1.6 shows three GEMM implementations in C. Figure 1.6a shows a naive implementation, with three nested loops spanning the size of the square matrices, accessing elements of the *destination* matrix *C* in the order shown on the leftmost panel of Figure 1.5. If loop indices are split into blocks, as shown in Figure 1.6b, memory accesses will be performed in *tiles*, shown on the center panel of Figure 1.5, which can improve performance by using only elements loaded into the cache. Loop unrolling, shown conceptually in Figure 1.6c, is an additional source transformation that can be performed independently, or in addition to tiling, and that can improve performance by grouping iterations of a loop, as represented on the rightmost panel of Figure 1.5, ensuring that processor registers are used.

The two *parameters* involved in this example are the *block size*, which controls the stride, and the *unrolling factor*, which controls the number of unrolled iterations. Larger block sizes are desirable, because we want to avoid extra comparisons, but blocks should be

```
int N = 256;
int B size = 4;
int A[N][N], B[N][N], C[N][N];
int i, j, x, y;
// Initialize A, B, C
for(i = 0; i < N; i++){
    // Load line i of A to fast memory
    for(j = 0; j < N; j++){
        // Load c(i)[j] to fast memory
        // (Load column j of B to fast memory
        // write C[i][j] to main memory
    }
}

(a) Regular implementation

(b) Blocked, or tiled

int N = 256;
int B size = 4;
int A[N][N], B[N][N], C[N][N];
int i, j, x,
// Initialize A, B, C
for(i = 0; i < N; i++){
        // Load block (i, j) of C to fast memory
        // Load block (i, j) of C to fast memory
        // Load cill[j] to fast memory
        // Write C[i][j] to main memory
    }
}

(a) Regular implementation

(b) Blocked, or tiled

int N = 256;
int B size = 4;
int A[N][N], B[N][N], C[N][N];
int i, j, x,
// Initialize A, B, C
for(i = 0; i < N; i+= B size, N); x++){
    for(y = 0; j < N; i+= B size, N); x++){
        for(y = j; y < min(j + B size, N); y++){
            // Load block (i, j) of C to main memory
            // Write block (i, j) of C to main memory
            // Load block (j, j) of C to fast memory
            // Load block (j, j) of A to fast memory
            // Load block (j, j) of A to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Load block (j, j) of B to fast memory
            // Write block (i, j) of C to fast memory
            // B size - 1][j] * B size - 2];
            // B size - 1][j] * B size - 1][j] * B size - 1][j] * B size - 1][j] *
```

Figure 1.4: Loop nest optimizations for $C = C + A + B^{T}$, in C

(c) Tiled and unrolled

small enough to ensure access to as few as possible out-of-cache elements. Likewise, the unrolling factor should be large, to leverage available registers, but not so large that it forces memory to the stack.

The values of block size and unrolling factor that optimize performance depend on the cache hierarchy and register layout of the target processor, and on the memory access pattern of the target algorithm. An autotuner should also aim to account for the *interaction* between parameters, that is, for the fact that the best value for each parameter also depends on the value chosen for the other.

It is straightforward to change the block size of the implementations from Figure 1.6, but the unrolling factor is not exposed as a parameter. To test different unrolling values we need to generate new versions of the source code with different numbers of unrolled iterations. We can do that with code generators or with *source-to-source transformation* tools [19, 20, 21]. It is often necessary to modify the program we wish to optimize in order to provide a configuration interface and expose its implicit parameters.

Once we are able to control the block size and the loop unrolling factor, we determine the target search space by choosing the values to be explored. The loop nest optimization example in this section comes from Seymour *et al.* [18], and considers 128 integer values in

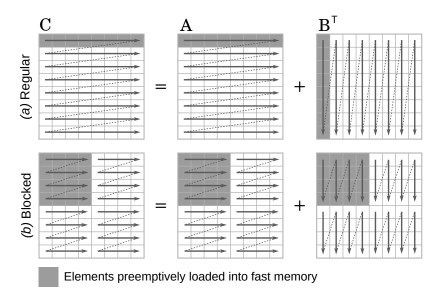


Figure 1.5: Access patterns for matrices in $C = A + B^{T}$, with loop nest optimizations. Panel (a) shows the access order of a regular implementation, and panel (b) shows the effect of loop *tiling*, or *blocking*

the interval [0, 127]. A block size of zero results in the implementation from Figure 1.6a, and an unrolling factor of zero performs a single iteration per condition check.

The performance of each of the $128^2 = 16384$ combinations of blocking and unrolling factor, shown in Mflops/s in Figure 1.7, was measured for a sequential matrix multiplication algorithm, using square matrices of size 400 [18]. We can represent this autotuning search space as a 3D landscape, since we have two configurable parameters and a single target performance metric. An autotuner's objective is to find the *highest* point of this landscape, since the objective is to *maximize* Mflops/s, although it is more usual to transform the performance metric so that the objective is to *minimize* it.

On a first look, there seems to be no apparent global search space structure in the landscape on Figure 1.7, but local features jump to the eyes, such as the "valley" across all block sizes for low unrolling factors, the "ramp" across all unrolling factors for low block sizes, and the series of jagged "plateaus" across the middle regions, with ridges for identical or divisible block sizes and unrolling factors. A careful look reveals a curvature along the unrolling factor axis. The best performance on this program was achieved with a block size of 80 and an unrolling factor of 2.

In this conceptual example, all $\approx 1.64 \times 10^4$ configurations were exhaustively evaluated, but in most settings where autotuning methods are useful it is impossible to do so.

(c) Tiled and unrolled

Figure 1.6: Loop nest optimizations for GEMM, in C

| System | Domain | Method | Year |
|-----------------------------|---------------------------|------------------------|------|
| PhiPAC [42] | Linear Algebra | SH (Exhaustive) | 1997 |
| ATLAS [43] | Linear Algebra | SH (Exhaustive) | 1998 |
| FFTW [44] | Digital Signal Processing | SH (Exhaustive) | 1998 |
| Active Harmony [45] | Domain-Specific Language | SH | 2002 |
| OSKI [46] | Linear Algebra | SH | 2005 |
| Seymour, K. et al. [18] | Linear Algebra | SH | 2008 |
| PRO [33] | Linear Algebra | SH | 2009 |
| ParamILS [34] | Combinatorial Auctions | SH | 2009 |
| PetaBricks [21] | Domain-Specific Language | SH (Genetic Algorithm) | 2009 |
| MILEPOST GCC [47] | Compiler Parameters | ML | 2011 |
| Orio [22] | Linear Algebra | ML (Decision Trees) | 2012 |
| pOSKI [24] | Linear Algebra | SH | 2012 |
| INSIEME [48] | Compiler Parameters | SH (Genetic Algorithm) | 2012 |
| OpenTuner [23] | Compiler Parameters | SH | 2014 |
| Lgen [49] | Linear Algebra | SH | 2014 |
| OPAL [50] | Parallel Computing | SH | 2014 |
| Mametjanov, A. et al. [30] | High-Level Synthesis | ML (Decision Trees) | 2015 |
| CLTune [51] | Parallel Computing | SH | 2015 |
| Guerreirro, J. et al. [52] | Parallel Computing | SH | 2015 |
| Collective Mind [53] | Compiler Parameters | ML | 2015 |
| Abdelfattah, A. et al. [31] | Linear Algebra | SH (Exhaustive) | 2016 |

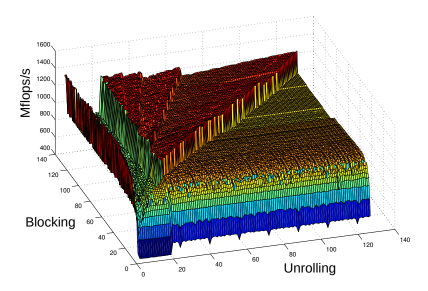


Figure 1.7: A search space defined for loop blocking and unrolling parameters, in a matrix multiplication kernel [18]

1.4 Thesis Contributions

1.5 Text Structure

Chapter 3 presents background for the application of methods derived from search heuristics, mathematical optimization, and statistical learning to autotuning.

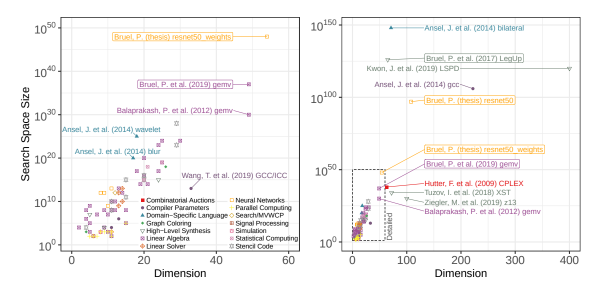


Figure 1.8: Dimension and search space size for autotuning problems from different domains [22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 18] The left panel shows a zoomed view of the right panel

Efforts for Reproducible Science

- 2.1 Introduction
- 2.2 Computational Documents with Emacs
- 2.3 Versioning for Code, Text, and Data

Search Heuristics and Statistical Learning for Autotuning

3.1 Introduction

• Methods Tree

3.2 Search Heuristics

3.2.1 Introduction

- Implicit hypotheses
- No explicit surrogate models

3.2.2 Software for Autotuning with Heuristics

- 3.2.3 Results with GPU Compiler Parameters
- 3.2.4 Results with High-Level Synthesis for FPGAs

3.3 Statistical Learning

3.3.1 Introduction

- Explicit surrogate models
- The Bias-Variance Trade-off
- Inference and prediction
- Supervised and unsupervised learning
- Parametric and Nonparametric Methods

3.3.2 Supervised Methods for Regression and Classification

- KNN, Linear Regression, Logistic Regression, Neural Networks
- Briefly mention others

3.3.3 Model Assessment

- Train and Test Sets
- Train and Test Mean Squared Error
- Cross Validation, Bootstrapping
- The question of choosing experiments *X* is never posed

3.3.4 Linear Regression

- Ordinary Least Squares: Gauss-Markov (BLUE)
- Ridge, Lasso
- Transformations of *X*

3.3.5 Software for Autotuning with Statistical Learning

3.3.6 Results

• Briefly mention results, link to following chapters

A Design of Experiments Methodology for Autotuning

4.1 Introduction

- Poses the question of choosing the experiments *X*
- Mixing categorical and numerical (continuous and discrete) factors

4.2 Inference with Analysis of Variance

- 4.3 Autotuning under a Tight Budget
- 4.3.1 Screening for Main Effects
- 4.3.2 Fractional Factorial Designs
- 4.3.3 Optimal Designs
- 4.4 Optimal Design
- 4.4.1 The KL Exchange Algorithm
- 4.5 Results
- 4.5.1 Results in a GPU Laplacian Kernel
- 4.5.2 Results with SPAPT
 - 1. CCGRID Paper

2. Later developments

Gaussian Process Regression

5.1 Introduction

- Sampling functions
- Covariance Kernels: modeling uncertainty regions
- Expressing structure?

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