# Adaptive Auto-Tuning of Computations on Heterogeneous Environments

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### **Abstract**

Heterogeneous computing offers increased performance making it popular for mathematically intensive computations. One of the commonly used programming tools for heterogeneous computing is OpenCL, which stands for Open Computing Language. OpenCL has been proposed as an industry standard framework that allows programs to execute on heterogeneous platforms, from multiple processor vendors. OpenCL, like others frameworks similar to it, suffers from poor performance portability. Kernel programs tuned for a particular processing unit need re-tuning on another device to achieve similar performance. Machine learning based auto-tuning provides one possible solution to the performance portability issue. In this paper, we attempt to demonstrate how the Random Forest machine learning model can be applied to the auto-tuning problem. The naive matrix multiplication algorithm is given certain tuning parameters, and the sample data for varying parameters is used to build and train the Random Forest model. The model can then be used to determine suitable tuning parameters for kernel optimization independent of processor model.

**Keywords**: auto-tuning, heterogeneous computing, machine-learning, OpenCL, random forests, SGEMM

### Introduction

For years, central processing units (CPUs) were the only processors capable of performing mathematically intensive computations, but growing demand for faster and cheaper hardware led to the development of accelerated processing units (APUs), and graphics processing units (GPUs). Originally, these APUs and especially GPUs, were designed for a specific purpose and were either incapable or inefficient at performing the numerous functions that CPUs could achieve.

Today, the common heterogeneous system is one with a few CPU cores and thousands of GPU cores that can perform mathematically intensive computations in parallel. While this parallelism is widely advantageous, implementing programs that can take advantage of the entire system remains challenging.

OpenCL was developed to take advantage of the new capabilities of the modern GPU architecture and to attempt to solve the difficulties of CPU-GPU interoperability. While OpenCL is capable of tying programs to heterogeneous systems, it struggles to do so with reliably consistent execution time. OpenCL kernels are programs that can be executed on different devices in a heterogeneous system. These kernels offer portability, but the performance of the kernel can vary dramatically based on the device it is currently executing on.

One common solution to this issue of performance portability is auto tuning. The concept of auto tuning

is to automatically take execution time measurements on various implementations of the kernel and choose the best implementation for that device. Auto tuning can be empirically driven, model driven, or machine learning driven. Empirically driven auto tuning goes through the entire parameter space to find the best implementation, but while it guarantees that the best kernel configuration will be found among the parameter space, it will be a tedious endeavor if the parameter space is large. Model driven auto tuning uses a performance model that is used to find similar kernel candidates to be used. Model driven auto tuning is dependent on the performance model used. If the performance model is inadequate, all possible kernel candidates will also share the inadequacy. Machine learning can be used to generate and tune a performance model. By executing a set of random kernel implementations and calculating their execution times, a prediction can be determined for the best kernel implementation.

The entirety of this paper will demonstrate why auto tuning kernels is necessary and how machine learning could be used in auto tuning to generate performance portable OpenCL kernels without the need to exhaustively search the entire kernel parameter space, or manually without creating a performance model.

### **Related Work**

Other, more successful, research has been done related to heterogeneous computations with matrix multiplication and performance model auto tuning [1]. Additional auto tuning research using Artificial Neural Networks have reported solid outcomes with machine learning applied to auto tuning [2]. The primary differences between what our paper attempts to accomplish and the research of others is the type of kernel used, the kernel parameters used for tuning, and machine learning techniques employed. In other auto tuning research, the most popular machine learning algorithm employed was Artificial Neural Networks. Random Decision Forests is a relatively new machine learning model to be adapted to solve the auto tuning problem.

# **Background**

#### OpenCL

OpenCL is a proposed industry standard framework for heterogeneous computing. It has widely been

implemented into the industry with names like AMD, Apple, Intel, Nvidia, Qualcomm, and over 100 other companies using the framework.

It allows programmers to access the OpenCL compatible devices that include CPUs, GPUs, APUs, and FPGAs. The ability to use the same code across various devices encourages code reuse, but the biggest disadvantage to OpenCL and other heterogeneous APIs is performance variations between devices.

### Random Decision Forests

The algorithm was first implemented by Tin Kam Ho of AT&T Bell Labs [3]. Random decision forests are collections of random decision trees. A single decision tree can provide insight for machine learning, but the tree is prone to overfitting. Random decision forests can reduce overfitting with multiple decision trees that are smaller and contain randomized subsets of the dataset features. Random decision forests are nonlinear and work well with features that have varying ranges from one another. Later implementations of random decision forests were developed such as Leo Breiman's trademarked Random Forest that uses bagging techniques alongside Ho's original algorithm [4].

# Single Precision Floating Point General Matrix Multiplication

Single Precision Floating Point General Matrix Multiplication or SGEMM is any matrix multiplication algorithm that processes single precision floating point elements. Matrix multiplication is a commonly used mathematical operation that has applications across all science and engineering disciplines. Matrix multiplication in its naive form is calculated as:

$$c_{ij} = a_{i1}b_{1j} + ... + a_{im}b_{mj} = \sum_{k=1}^{m} a_{ik}b_{kj}$$
  
where  $i = 1 ... n$  and  $j = 1 ... p$ 

The computational complexity is  $O(n^3)$  for the naive matrix multiplication algorithm. Divide and conquer variations of the algorithm that take advantage of parallel processing and/or heterogeneous computing has allowed for faster execute times.

# **Implementation**

Our benchmark test uses the classic single precision general purpose matrix multiplication (SGEMM). We placed a constraint for the dimensions of the matrices to be square, 2^N, and within the range up to 512x512. Input matrices A and B, as well as the output matrix C, have the same row and column dimensions.

Benchmark Kernel	Description
Single Precision Floating Point General Matrix Multiplication (SGEMM)	Matrix Multiplication of several sizes of square input matrices. 1x1, 2x2, 4x4, 8x8, 16x16, 32x32, 64x64, 128x128, 256x256, 512x512

Table 1: Benchmark Kernel

Our parameter space consists of the following values:

Parameters	Test Values
MATRIX DIMENSION (GLOBAL WORK SIZE)	1, 2, 4, 8, 16, 32, 64, 128, 256, 512
USE_LOCAL_MEMORY	0, 1
BLOCK SIZE (LOCAL WORK SIZE)	1, 2, 4, 8, 16

Table 2: Parameters

Originally the parameter space was designed to include global work group size x, global work group size y, local work group size x, and local work group size y. We had to limit the parameter space down to BLOCK SIZE (local work size) and MATRIX DIMENSION (global work size) because our matrices were specified as square matrices of  $2^N$  x  $2^N$  in the range where  $2^N = \{1, 2, 4, 8, 16, 32, 64, 128, 256, 512\}$ . The parameter space was also limited up to 512 due to the OpenCL constraints of our test hardware. OpenCL global work size must the size of the job size, in our case the size of the matrix.

According to the OpenCL specification [5] the number of work groups can be computed as:

$$(W_x, W_y) = (G_x / S_x, G_y / S_y)$$

where  $W_x$  is the number of work groups in the x-dimension,  $W_y$  is the number of work groups in the y-dimension,  $G_x$  is the index space of the work items in the x-dimension,  $G_y$  is the index space of the work items in the y-dimension,  $S_x$  is the size of the work

group in the x-dimension,  $S_y$  is the size of the work group in the y-dimension. The OpenCL specification uses this equation as part of its execution model. This creates a limitation to what combination of kernel parameters can properly execute without fault.

We build our kernel using the OpenCL API for the C language. The C program is the OpenCL host code that sets up the execution environment for the kernel on a particular device. The host code contains functions to establish an OpenCL context, OpenCL command queue, OpenCL program, and OpenCL kernel. The kernel will operate on a work environment that is defined by clEnqueueNDRangeKernel. This function takes the global and local work sizes as parameters, and these parameters depend on the size of the matrix, as well as the maximum allowed work size for the device.

The host function contains an OpenCL function for event profiling that determines the kernel execution start and finish times. The difference between the two determines the execution time of the kernel in nanoseconds. For convenience, we shifted the results from nanoseconds to milliseconds. The kernel parameters and their corresponding execution times are then emitted to a CSV (comma separated values) file that will be parsed by a Python3 script. This Python script contains the Random Forest algorithm, which was taken from a popular module called Scikit Learn.

We choose Random Forest as our machine learning algorithm because the nature of Random Forest fits the problem well. Our parameter space consists of features that have varying ranges from 1 to 512, 0 to 1, and 1 to 16. Random Forest is a nonlinear machine learning algorithm and responds well with features that have variable ranges. The Random Forest algorithm also doesn't need to be normalized between 0 and 1. In fact, normalization would smooth out the nonlinearity of the model. If normalization were to occur with our parameters, it would be difficult to denormalize back to the original values. Normalization would make sense if all the parameters operated within the same range of values.

Once the set of kernel implementations is fed into the random forest regressor, a prediction model is generated and undergoes training to obtain accurate predictions for kernel parameters that have not been provided in the original dataset.

The Python script then generates predictions from the kernel parameters of the test set and with the actual execution times. The error is generated between the predicted and actual. Then, the best actual kernel implementation is found. Once we collect information about the best kernel implementations found in each matrix dimension, another Python script would take all the information about these kernels and plot a performance curve.

Testing was performed on the following devices:

Test Hardware / OS	OpenCL Device Specs
Intel i5-3317U / macOS High Sierra 10.13.4	Max Compute Units: 4 Max Work Group Size: 1024 Max Work Item Dimension 3 Max Work Item Sizes: 1024 / 1 / 1 Max Clock Frequency: 1700 MHz
Intel i7-4578U / macOS High Sierra 10.13.4	Max Compute Units: 4 Max Work Group Size: 1024 Max Work Item Dimension: 3 Max Work Item Sizes: 1024 / 1 / 1 Max Clock Frequency: 3000 MHz
Intel HD Graphics 4000 / macOS High Sierra 10.13.4	Max Compute Units: 16 Max Work Group Size: 512 Max Work Item Dimension: 3 Max Work Item Sizes: 512 / 512 / 512 Max Clock Frequency: 1050 MHz
Intel Iris 1536 MB / macOS High Sierra 10.13.4	Max Compute Units: 40 Max Work Group Size: 512 Max Work Item Dimension: 3 Max Work Item Sizes: 512 / 512 / 512 Max Clock Frequency: 1200 MHz

Table 3: Test Hardware

Due to resource constraints, the amount of available hardware to test and compare kernel performance was limited to the devices shown in Table 3.

### **Results**

We performed various tests on our method by using the matrix multiplication kernel for varying dimensions for the input matrices. See Table 1 for the list of different input dimensions.

The OpenCL kernel was sampled for the different matrix multiplication dimensions on the Intel Iris GPU, and the Intel HD Graphics 4000. The Intel i5-3317U CPU was running the naive matrix multiplication with no OpenCL to compare the

performance of the kernels to the naive implementation. The results below illustrate that, for matrix dimension 512x512, the worst kernel parameters can execute at least 2x slower than a naive matrix multiplication on a CPU.

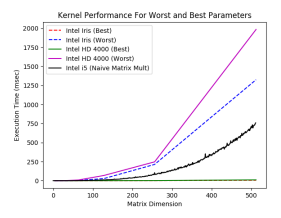


Figure 1: Kernel Performance

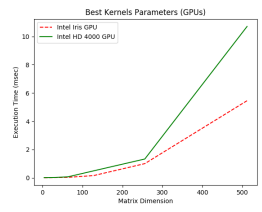


Figure 2: Best Kernel Parameters (GPUs)

The best kernel parameters on the GPUs show that for matrices of 512 x 512, the execution times are 10 milliseconds or less. In figure 1, the best kernel execution times appear linear, relative to the worst execution times. Figure 2 shows that they are not linear, but rather they execute at least 75x faster than the naive matrix multiplication on CPU.

Looking solely at the best kernel implementation in Figure 2, the Intel Iris GPU exhibits faster, about 2x faster, execution times than the Intel HD Graphics 4000. This is most likely due to the fact that the Intel Iris has 2.5x more compute units than the Intel HD Graphics 4000, and the Iris also has a max clock frequency that is 14.28% faster. This combination of more compute units and high clock frequency means

that Iris would theoretically have greater parallelism and execute more instructions faster than the HD Graphics 4000 counterpart.

Unfortunately, our dataset was extremely small due to the fact that only 3 kernel parameters were used with values of limited range. Going through every possible iteration of the parameter space results in a dataset of 66 possible samples, and some of these kernel parameters might not execute because local work size cannot be larger than the global work size. Machine learning algorithms are built with large datasets in mind, on the order of millions of samples. This means that it is extremely difficult to generate accurate models for small datasets. The alternative method to find the best kernels for small datasets was to perform an empirical auto tune. Through an exhaustive search the results were placed into a separate Python script to illustrate the kernel comparisons.

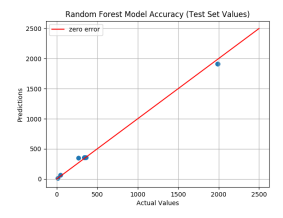


Figure 3:Random Forest Accuracy

The Random Forest machine learning prediction model was executed 100 times to obtain an average prediction accuracy of 95%, but because the dataset was limited, we believe these values are due our test cases being closely related to our training cases. For each matrix dimension we ran 50 samples for the kernel parameters. These combinations are likely to repeat and appear in both the training and testing cases. At this time, we cannot guarantee that the model prediction accuracy is truly 95% due to the highly likelihood of training and testing similarities.

# **Conclusions and Future Work**

This paper set out to prove that adaptive auto tuning is necessary in heterogeneous computing and that

Random Forest machine learning could be a useful tool for auto tuning. Our results demonstrate that the way the OpenCL host code sets up the kernel for execution can dramatically affect the performance of the kernel. If the kernel parameters are set incorrectly, the execution time can perform 2x worst than the naive matrix multiplication running on a CPU. On the positive side, if you were to tune the kernels to quality parameters, the kernels obtain an estimated 75x speedup factor over the same naive matrix multiplication.

When it comes to the machine learning aspect of this project, it is hard to say if Random Forest machine learning can effectively be used in auto tuning. The only way to determine effectiveness of Random Forest is to generate larger datasets for the test cases, so that we can evaluate our model on more unseen samples. Future work could be done to increase the amount of samples in the dataset. The best way to do this would be to increase the number of features of the Random Forest. Besides global work size, local work size, local memory, and global memory, additional kernel parameters could be added, like OpenCL texture memory usage, and loop unroll factors for matrix multiplication. Also, opening the matrix dimensions to different sizes besides 2<sup>N</sup> could open more options for the global work size and local work size. We also need to keep in mind that some additional kernel parameters, such as loop unrolling, are features unique to the matrix multiplication kernel. Developing other kernels besides matrix multiplication would open up new features unique to that problem.

Another possible issue that could arise in the random forest auto tuning model is the presence of local maxima and minima. In each case instance of the machine learning, the model could be finding kernel implementations that are not the global execution time minimum, but actually a local minima for those samples applied to the current instance of the model. In order to solve this issue, we would likely need two types of machine learning models in our program. The first machine learning instance would be the random forest model that would evaluate the local minima at each matrix dimension for several executions of the model. The second instance would be an artificial neural network, or other type of machine learning algorithm, that would take the interpolation of all the random forests executions in order to find the global kernel execution time minimum. This could be a possible goal for any future work on this project.

The execution time of the program could be dramatically improved in two areas. First, the

comparison check to see if the kernel returns the correct result could be improved. The check is needed to ensure that kernels not only run fast but return the right result. There is likely a more efficient method to perform kernel error checking. Second, the Random Forest prediction generation could also be improved. For our purposes, Python and the SciKit Learn module was sufficient enough to generate a prediction model, but if this were to be used in real time applications, the program would be extremely slow. Possibly the best solution to improve the prediction generation would be to implement the Random Forest algorithm using C/C++ instead of Python.

Finally, the OpenCL specification is being restructured into a new API called Vulkan Compute. Vulkan combines the heterogeneous capabilities of OpenCL with the graphics capabilities of OpenGL, and OpenCL is being moved into the Vulkan Compute section of Vulkan. This decision to restructure OpenCL to Vulkan Compute means that our program has a high risk of containing deprecated functions with one to two years from the date of this paper. Any future work directly related to this project will have to refactor the code to meet the new Vulkan specification.

# **Acknowledgments**

The authors would like to thank Carlos Reyes, the technical mentor and sponsor for this project. He provided a wonderful and challenging project to address. This was a project that none of us had prior knowledge about, and we like to thank Carlos for the new knowledge we gained by tackling this problem. The authors would also extend thanks to the wonderful employees at Stellar Science for allowing project discussions to take place in their office.

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