

Optimizing the Performance of Multi-threaded Linear Algebra Libraries, a Task Granularity based Approach

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Table of Contents

ACKNOWLEDGMENTS	ii
LIST OF TABLES	v
LIST OF FIGURES	vi
ABSTRACT	viii
CHAPTER	
1. INTRODUCTION	1
1.1. Thesis Statement	1
1.2. Contributions	2
1.3. Document Organization	3
2. BACKGROUND	4
2.1. Asynchronous Many-task Runtime Systems	4
2.2. Blaze	5
2.3. Task Granularity	6
2.4. Analytical modeling of parallel programs	7
2.5. Loop Scheduling	10
3. LITERATURE REVIEW	12
3.1. Literature Review	12
4. METHOD	16
4.1. Parallelization in Blaze	16
4.2. Experiments	17
4.3. Method	24
5. UNDERSTANDING THE EFFECT OF GRAIN SIZE ON CONCURRENCY IN AN ASYNCHRONOUS MANY-TASK RUNTIME SYSTEM	33
5.1. Analytical Modeling	33
6. SETUP	46
7. PROPOSED STUDY	48
7.1. Studying the bathtub model	48
7.2. Generalization for matrix size	48
7.3. Generalization for complex expressions	50
7.4. Generalization for different architectures	50
8. APPENDIX	51
8.1. Bathtub model	51
REFERENCES	56

APPENDIX

List of Tables

4.1.	List of some of the thresholds applied to the operations performed by Blaze, starting from which the operation is executed in parallel	16
4.2.	List of different values used for each variable for running the <i>DMATDMATADD</i> benchmark	20
6.1.	Specifications of the Marvin node from Rostam cluster at CCT.....	47
6.2.	Cache specifications of the Marvin node from Rostam cluster at CCT.....	47
6.3.	Specifications of the libraries used to run our experiments.	47

List of Figures

2.1.	The effect of task size on execution time for Stencil application [1]	7
2.2.	An example of the achievable speedup based on Amdahl's law and USL compared to the ideal linear speedup where $\sigma = 0.04$ and $\kappa = 0.005$	8
4.1.	The results obtained from running <i>DMATDMATADD</i> benchmark through Blazemark for matrix size 690×690 on different number of cores.	20
4.2.	The results obtained from running <i>DMATDMATADD</i> benchmark through Blazemark for matrix of size 690×690 from two different angles	21
4.3.	The results obtained from running <i>DMATDMATADD</i> benchmark through Blazemark for matrix sizes from 200×200 to 1587×1587	22
4.4.	The results obtained from running <i>DMATDMATADD</i> benchmark through Blazemark for matrix size 690×690 with different combinations of block size and chunk size on 4 cores	23
4.5.	The results obtained from running <i>DMATDMATADD</i> benchmark through Blazemark for matrix size 690×690 on 4 cores.	24
4.6.	Throughput vs. grain size graph obtained from running <i>DMATDMATADD</i> benchmark on 4 cores for matrix sizes (a) smaller than 793×793 and (b) larger than 793×793	25
4.7.	The results of fitting the throughput vs grain size data into a 2d polynomial for <i>DMATDMATADD</i> benchmark for matrix size 690×690 with different number of cores on the test data set (a) 1 core, (b) 2 cores, (c) 3 cores, (d) 4 cores, (e) 5 cores, (f) 6 cores, (g) 7 cores, (h) 8 cores.....	26
4.8.	The training and test error for fitting data obtained from the <i>DMATDMATADD</i> benchmark for matrix size 690×690 against different number of cores cores.	27
4.9.	Fitting the parameters of the quadratic function with a 3rd degree polynomial from the <i>DMATDMATADD</i> benchmark for matrix size 690×690 against different number of cores.	28
4.10.	The error in fitting the parameters a , b , and c for matrix size 690×690	28
4.11.	Results of fitting the data from <i>DMATDMATADD</i> benchmark with a polynomial of degree 2 in terms of grain size and of degree 3 in terms of number of cores for matrix size 690×690 for (a) 2 core, (b) 4 cores, (c) 8 cores.	29
4.12.	The training and test error obtained fitting the data to a polynomial of degree 2 in terms of grain size and of degree 3 in terms of number of cores for matrix size 690×690 , for each number of cores. (a) All the data points are include in calculation of error, (b) the leftmost sample was removed from error calculation.	29
4.13.	The range of grain size (shown as the red line) that leads to a performance within 10% of the maximum performance for (a) 2 cores, (b) 4 cores and (b) 8 cores.	30

4.14. The range of grain size within 10% of the maximum performance of the fitted polynomial function for <i>DMATDMATADD</i> benchmark for different number of cores for (a) matrix size 690×690 (b)matrix size 523×523 to 912×912 .	31
4.15. The range of chunk sizes to produce a grain size within 10% of the maximum performance of the fitted quadratic function for <i>DMATDMATADD</i> benchmark for matrix size 690×690 with block size of 4×256 on (a) 2 cores, (b) 4 cores, and (c) 8 cores, and block size of 4×512 on (d) 2 cores, (e) 4 cores, and (f) 8 cores. Silver points denotes the detected range of chunk size, and the red star shows the median point.	32
5.1. The results of running the benchmark in Listing with <i>problem_size</i> = 10000, on different number of cores.	35
5.2. The imbalance ratio calculated for different grain sizes for <i>problem_size</i> = 10000, on 8 cores, where $k = \left\lceil \frac{\text{num_tasks}}{N} \right\rceil$	41
5.3. The imbalance ratio calculated for different grain sizes for <i>problem_size</i> = 10000, on 8 cores, where $k = \left\lceil \frac{\text{num_tasks}}{N} \right\rceil$	42
6.1. An example of the results obtained from running <i>DVECDVECADD</i> benchmark through Blazemark	46
7.1. Throughput vs. grain size graph obtained from running <i>DMATDMATADD</i> benchmark on 4 cores.	49
7.2. The range of grain size within 10% of the maximum performance of the fitted polynomial function for <i>DMATDMATADD</i> benchmark for different number of cores for matrix size 523×523 to 912×912 .	49
8.1. (a)The execution time vs. grain size graph, and (b) execution time vs. number fo tasks graph for <i>DMATDMATADD</i> benchmark for matrix size 690×690 ran on 4 cores.	51
8.2. The prediction of execution time based on grain size using the bathtub model, for (a)4 cores and (b)8 cores for <i>DMATDMATADD</i> benchmark for matrix size 690×690 .	53
8.3. The error in fitting execution time with the bathtub formula for <i>DMATDMATADD</i> benchmark for matrix size 690×690 with different number of cores.	54
8.4. Fitting the three parameters (a) α , (b) t_s , and (c) γ for <i>DMATDMATADD</i> benchmark for matrix size 690×690 .	54
8.5. The error in fitting execution time with the bathtub formula for <i>DMATDMATADD</i> benchmark for matrix size 690×690 with different number of cores.	55

Abstract

Linear algebra libraries play a very important role in HPC applications. In this work we propose a method to tune the performance of a linear algebra library based on a set of compile-time and runtime characteristics including the machine architecture, the expression being evaluated, the number of cores to run the application on, the type of the operation, and also the size of the matrices, to be able to get as close as possible to the highest performance.

There has been an extensive amount of work and study done on how to optimize the compute kernels for matrix-matrix multiplication. In this thesis, we are interested in all types of operations, and our focus is on machine learning applications, where we are potentially dealing with very large matrices and creating temporaries could be very expensive.

For this purpose we decided to use Blaze C++ library, a high performance template-based math library that gives us this option to access the expression tree at compile time, along with HPX, a C++ standard library for concurrency and parallelism, as our runtime system. We propose here that instead of dividing the work equally among the cores and assigning one chunk to each core, we should be able to achieve a higher performance by selecting the right amount of work to be assigned to one core, based on a set of runtime and compile-time parameters.

Studying the significant amount of data we collected with different configurations, we concluded that, grain size, which is the amount of work assigned to each task, is the key factor for the performance when number of cores is fixed. With this assumption, we tried two different approaches to model the relationship between performance and grain size, in order to find a range of grain size that could lead us to the maximum performance.

First model uses a polynomial function to fit the data both for throughput vs. grain size, and throughput vs. number of cores. On the other hand, the second model was developed by studying the behavior of throughput vs. grain size , and integrating it with the an extension of the Universal Scalability Law(USL). This developed model fitted our data reasonably and could be very important in further understanding of the parallelism.

The primary results suggest that, using the data collected for matrix-matrix addition, we are able to improve the performance for matrix-matrix addition, by finding the right range of grain size.

Having the mentioned models, we changed the current implementation of the HPX backend for Blaze by adding two parameters to represent the unit of work and the number of these units included in each task, for fine grained control of the parallelism, which is possible through HPX runtime system. Also, a complexity estimation function has been added to Blaze as an estimate of the number of floating point operations occurring in each unit of work.

Our data was only limited to one specific operation, matrix-matrix addition for two raw-major matrices, one matrix size at a time. In the next step, this problem should be generalized to different matrix sizes, architectures, and arbitrary complex expressions.

Chapter 1. Introduction

The path toward exascale computing, would include more complex machine architectures, deeper memory hierarchies, heterogeneous nodes, complicated networks[2]. Current programming models would not be sufficient. Advanced runtime systems with support for new programming languages and models are needed to manage the huge amount of parallelism that would be available[3]. Asynchronous many-task(AMT) models and their corresponding run-times are the solution to keep application developers safe from the upcoming architectures, by mitigating exascale difficulties to run-time level[4].

On the other hand, the core element of many high performance computing applications is the linear algebra library. The performance of these application heavily relies on the performance of their linear algebra library. BLAS(Basic Linear Algebra Subprograms) are the fundamental routines for basic vector and matrix operations. But in order to tune BLAS for a specific architecture, a lot of effort needs to be made, and requires a deep understanding of memory hierarchy and registers from the programmer[5]. Linear algebra libraries like ATLAS[5], SPIRAL[6] try to use hardware-specific optimizations to improve their performance.

In this work, we are trying to optimize the performance of a linear algebra library based on the application parameters such as matrix size, operation, the expression, data layout, and also the machine architecture. Some of these parameters could be extracted at compile time, while extraction of the others should be postponed to runtime.

1.1. Thesis Statement

The main objective of this thesis is to propose a hybrid runtime and compile-time solution for a linear algebra library to fully take advantage of the available parallelism and resources.

We chose Blaze math library since it is a nice high performance template-based C++ library that allows you to access the expression tree for each assignment at compile time, and we chose HPX as an asynchronous many-task runtime system to manage the parallelism.

HPX makes it possible to create thousands to millions of lightweight user threads, to avoid

expensive context switching. On the other hand, although the overhead of creating one task is negligible, creating millions of tasks when the execution time of the program itself is small, could become significant and cause performance degradation. On the other hand if we create too few tasks it would be very likely for us to not use our resources properly. So in very application in many task systems, it's very important to chose the amount of work assigned to each task, called grain size, properly.

Through analyzing and modeling the relationship between throughput and grain size, we would be able to identify a range of grain size that leads us to maximum performance. Once decided how big one unit of work should be, based on the identified range we would be able to decide on how many units of work should be packed into one task.

On the other hand, there are different models to express the relationship between the throughput and the number of cores. Here, we are interested in developing a model to be as realistic as possible to imitate the behavior of the throughput against both grain size and number of cores. This would help us find how to manage the parallelism in our system to achieve the highest performance possible.

1.2. Contributions

There has been a wide study, mostly by Gunther[7, 8, 9, 10], on different models to represent the relationship between the throughput and the number of cores, for a fixed size problem. Grubel et.al[1] has studied the effect of task granularity on the performance with a fixed number of cores. Our contributions could be summarized into:

- We propose a novel physical model to represent how the execution time is expected to change based on grain size.
- To our knowledge, there has not been a work to create a 3D model of the throughput, grain size, and number of cores.

- We are proposing a method to apply the developed model to a linear algebra library, in a way specific to our application, and the machine architecture.

1.3. Document Organization

In Chapter 2, we will explain briefly the background needed for this thesis, including Blaze and HPX library, the effect of task granularity, and the Universal Scaling Law(USL) method for modeling the throughput based on number of cores. Chapter 3 refers to other works that have been done in our area of our focus. We explain our proposed method to optimize the performance, along with the models we used in Chapter 4. Our work heavily relies on the collected data, the environment we collected the data from and also the library versions are mentioned in Chapter 6. Finally we discuss our concerns and the further steps that needs to be taken in Chapter 7.

Chapter 2. Background

2.1. Asynchronous Many-task Runtime Systems

A parallel programming model includes a programming model, which refers to the mechanism for a program to express the concurrency, and an execution model, indicating how the program creates and controls concurrency[11, 4]. Some of the common existing execution models include fork-join, Communicating Sequential Processes(CSP), event-based models and actor model[4]. Some of the current parallel programming frameworks include, accelerator-based programming models like OpenACC[12], OpenCL[13], and CUDA[14], shared-memory programming models like Intel TBB[15], Cilk[16], OpenMP[17], and distributed-memory programming models like MPI[18], Charm++[19], ParalleX[20], UPC[11].

A runtime system is in charge of creating and managing the concurrency, through implementing parts of an execution model[4]. An asynchronous many-task(AMT) model on the other hand, is a category of programming model and execution model. An AMT programming model breaks the work into small and transferable tasks along with their associated input. In an AMT execution model, tasks are being executed when their inputs are available rather than in a well defined order[4].

Some of well known AMT runtimes include: HPX[3], Charm++[19], Uintah[21], Legion[22].

2.1.1. HPX

HPX[3] is a C++ runtime system for parallel and distributed applications based on ParalleX execution model[20]. HPX provides you with lightweight user-level threads with fast context switching[11]. Whenever one thread is blocked, the scheduler picks a ready thread based on a scheduling policy. This allows you to hide the latency, and avoid starvation while keeping a high utilization of the resources[11].

2.1.1.1. Execution Model

The "SLOW" model identifies four potential sources for performance degradation as: Starvation, Latency, Overheads, and Waiting or contention[20].

Starvation refers to a situation where there is insufficient amount work for the computing resource. this could be due to insufficient total amount of work available, or unbalanced distribution of work among resources[11].

Latency is the time distance, usually measured in processor clock, of accessing remote data or services[23].

Overhead refers to the effort that needs to be taken to manage parallel resources and actions on the critical path[11].

Finally, waiting is the contention of the shared physical and logical resources causing one request to be blocked by another access of the same resource[23]. This could happen due to limited network bandwidth, shared communication channels, memory bank conflicts[11],[23].

2.2. Blaze

Blaze Math Library[24] is a C++ library for linear algebra. Blaze, based upon Expression Templates(ETs)[25], introduces "smart" expression templates(SETs)[24] to optimize the performance for array-based operations. Expression Templates[25] is an abstraction technique that uses overloaded operators in C++ to prevent creation of unnecessary temporaries, while evaluating arithmetic expressions, in order to improve the performance[24]. The ET-based approaches create a parse tree of the expression at compile time and postpone the actual evaluation to when the expression is assigned to a target.

Although being able to achieve promising performances for element-wise operations, these methods are not suitable for high performance computing for the following reasons. Due to their abstraction from both the data type and also the operation itself, they do not allow optimizations specific to the type of the arrays, alongside the operation[24]. As a solu-

tion, Blaze proposes smart ETs with these three main additions: integration with architecture-specific highly optimized compute kernels, creation of intermediate temporaries when needed, and selecting optimal evaluation method automatically for compound expressions[24].

Some of the ET-based linear algebra libraries are: Blitz++[26], Boost uBLAS[27], MTL[28], and Eigen[29]. Among these libraries, Eigen, MTL, alongside Blaze, impose different conceptual changes to ETs in order to make them suitable for HPC. Blaze also makes it possible for programmers to utilize SIMD(Single Instruction Multiple Data) vectorization simply by adding a compile time flag.

2.3. Task Granularity

Defining the grain size as the amount of work assigned to one HPX thread, Grubel[1] studies the effect of grain size on the execution time for a fixed number of cores. The results show that, for small grain sizes the overhead of creating the tasks, and for large grain sizes the starvation, is the dominant factor affecting the execution time[1]. When grain size is small, to perform same amount of work, higher number of tasks is created, and there is an overhead associated with creation of each task. Although this overhead is very small (order of microseconds), when the amount of work performed by each thread is also small, this overhead becomes significant. As the grain size increases, these overheads are amortized by the time it takes to execute the task.

On the other hand, when grain size is increases, the number of tasks being created decreases, up until a point where the number of tasks being created is smaller than the number of cores. At this point another factor would interfere with the performance, which is referred to as starvation. Starvation happens where a large amount of work is assigned to some of the cores while the other cores are idling. At this point we are not using our resources efficiently.

While overheads of creating tasks degrades the performance for small grain sizes and starvation causes the execution time to increase for large grain sizes, there is a region in between where changing the grain size does not affect the performance.

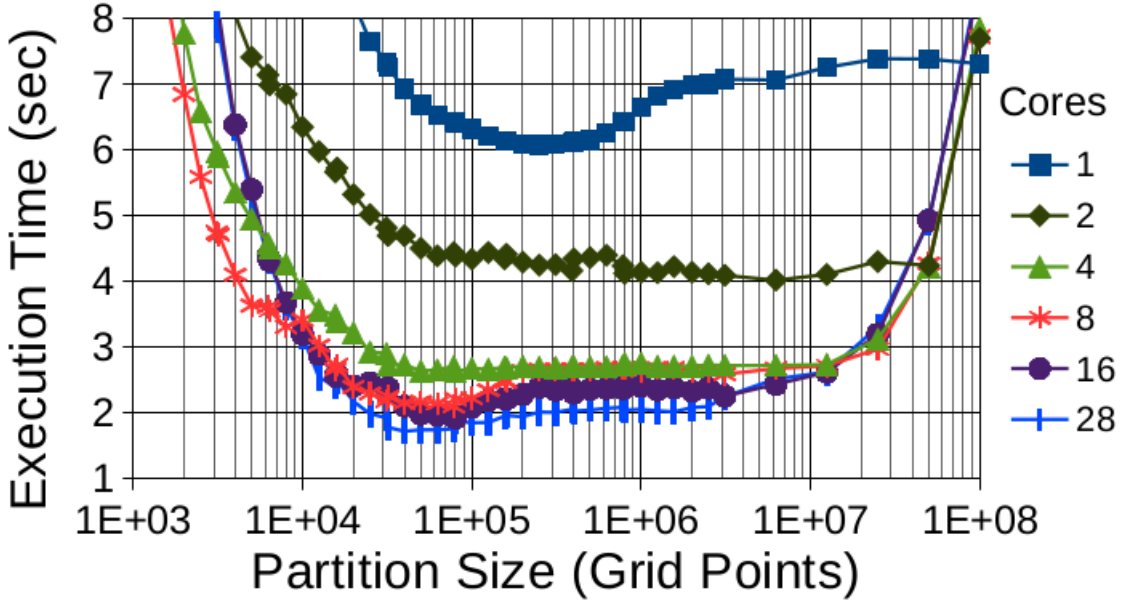


Figure 2.1. The effect of task size on execution time for Stencil application [1]

2.4. Analytical modeling of parallel programs

The execution time of a parallel program is highly dependent of the parallel algorithm to be evaluated, alongside the architecture it is implemented on[30].

2.4.1. Universal Scalability Law

Amdahl's law[31], states that the amount of achievable speed up by adding more processors when running a parallel application, is restricted by the amount of code that could actually be parallelized. Equation 2.1, shows the relationship between speedup and number of processors, where σ is the serial fraction of the execution time, based on Amdahl's law[9].

$$S(p) = \frac{p}{1 + \sigma(p - 1)} \quad (2.1)$$

On the other hand, Gunther[9] extends Amdahl's law by incorporating the effect of three factors, namely concurrency, contention, and coherency, as shown in Equation 2.2.

$$S(p) = \frac{p}{1 + \sigma(p - 1) + \kappa p(p - 1)} \quad (2.2)$$

Concurrency(p) represents the linear speedup that could have been achieved if no interaction existed among the processors, contention(σ) represents the serialization effect of shared writable data, and finally coherency or data consistency(κ) represents the effort that needs to be made for keeping shared writable data consistent[9].

Figure 2.2 shows an example of the ideal linear speedup we expect to see when increasing the number of the processors, against the actual achievable speedup based on Amdahl's law and USL.

Equation 2.3 generalizes Equation 2.2 to represent the throughput by adding another parameter(γ) to represent the serial throughput.

$$X(p) = \frac{\gamma p}{1 + \sigma(p - 1) + \kappa p(p - 1)} \quad (2.3)$$

Universal scalability law also suggests that for some values of σ and κ there could be a certain number of processors that yield to maximum performance[9]. Increasing the number of processor beyond that point would only cause performance degradation.

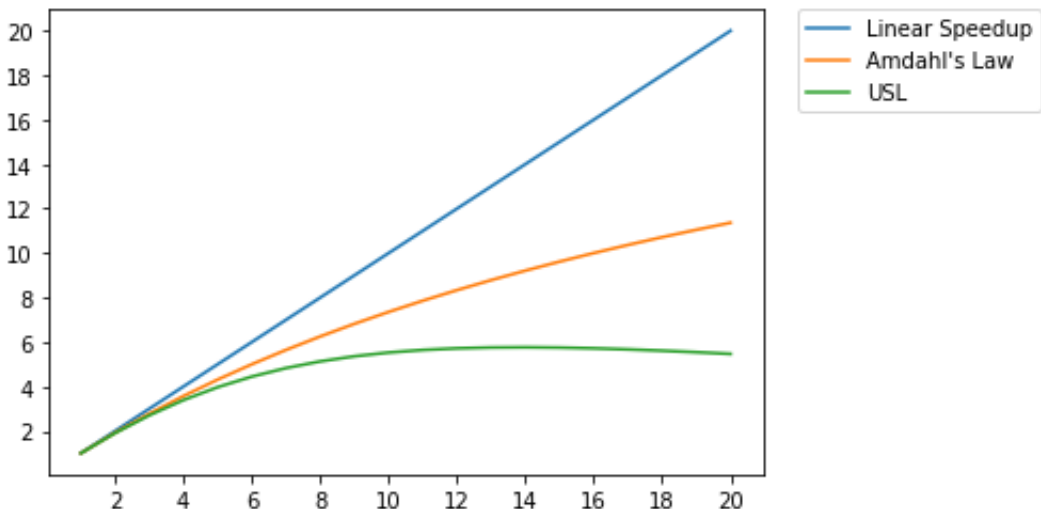


Figure 2.2. An example of the achievable speedup based on Amdahl's law and USL compared to the ideal linear speedup where $\sigma = 0.04$ and $\kappa = 0.005$.

2.4.2. Other Models

There are a few other models that have also been suggested to simulate the scalability. Geometric model is a one-parameter model, in which speedup has the following relationship with the number of processors:

$$S(p) = \frac{1 - \phi^p}{1 - \phi} \quad (2.4)$$

The parameter ϕ , where $0 < \phi \leq 1$, is called the MP factor, and it represents the remaining section of the processor capacity after deducting overheads. The geometric model is non-physical for large number of processors, due to inconsistency with Coxian queuing model[8].

Quadratic model[7], with overhead parameter γ , where $0 \leq \gamma < 1$, is represented in Equation quad.

$$S(p) = p - \gamma p(p - 1) \quad (2.5)$$

The quadratic model has a critical point at:

$$p^* = \lfloor \frac{1 + \gamma}{2\gamma} \rfloor \quad (2.6)$$

The problem with this model is that its not physical. This model represents an inverted parabola that will intersect the x axis at two points, implying that there will be a certain number of processors starting from which the speedup would be negative.

Exponential model, is also a single parameter model α where $0 < \alpha \leq 1$. This parameter is a combination of coherency and contention.

$$S(p) = p(1 - \alpha)^{(p-1)} \quad (2.7)$$

This model also has a critical point, but this point is very sensitive to α . Although this model works very well for small number of processors, it imposes a severe capacity degradation for large number of processors.

2.5. Loop Scheduling

Loop scheduling refers to different ways iterations could be assigned to the processors and the order of their execution. The main reason for performance degradation in loop scheduling is load imbalance, which refers to situations where different amount of work is assigned to different processors[32].

The simplest loop scheduling method is static scheduling, in which, the iterations are divided evenly among all the processors statically, either as a consecutive block -also called cyclic- or in a round-robin manner[33]. Since all the assignments happen at compile time or before execution of the application, this method imposes no runtime scheduling overhead. Several factors including interprocessor communication, cache misses, and page faults can lead to different execution times for different iterations, leading to load imbalance among the processors[34].

In the meanwhile, dynamic scheduling methods postpone the assignment to runtime, which tends to improve load balancing, at the cost of higher scheduling overhead. Some of dynamic scheduling methods include: Pure Self-scheduling, Chunk Self-scheduling, Guided Self-scheduling[35], Factoring[36] and Trapazoid Self-scheduling[37],[33]. We briefly go over some of these loop scheduling techniques here.

In Pure Self-scheduling every time a processor becomes idle, it fetches one loop iteration. This approach, while achieving a high load balance, imposes a considerable amount of scheduling overhead when we are dealing with a fine-grain workload, and a large number of iterations. Also frequent access to shared variables like loop index could lead to memory contention[33].

In order to decrease the high scheduling overhead of Pure Self-scheduling methods, Chunk Self-scheduling method assigns a certain number of iterations(called chunk size) to each idle processor. This method trades lower scheduling overhead with higher load imbalance. Selection of the chunk size plays a very important role in the performance, as so a large chunk size increases the scheduling overhead decreases and causes load imbalance, while a small chunk

size increases memory contention and scheduling overhead[33].

As an adaptive loop scheduling technique, Guided Self-scheduling[35] divides the remaining number of iterations at each request evenly among the processors, and assigns it to the processor that made the request, while updating the number of remaining iterations. This causes larger number of iterations to be assigned to the processors at the beginning of the loop execution, which results in lower scheduling overhead. The number of iterations assigned to each processor decreases as it approaches to the end of the execution, generating tasks containing only one or two iterations, causing an increase in the scheduling overhead. In order to tackle this issue, a minimum number of chunks could be set to avoid creation of very small chunks[38].

Very similar to Guided Self-scheduling, Factoring[36] also decreases the chunk size as the loop execution proceeds, with this difference that it does it in batches of equal sized chunks. If the first iterations of the loop are more time consuming than the rest of the iterations, Factoring performs better than Guided scheduling[39].

Along with the mentioned loop scheduling approaches, two other scheduling techniques can be utilized for load balancing. Work stealing[40] lets the processors to steal work from other processors queue, resulting in a more balanced load distribution. In work sharing on the other hand, each time a processor creates new threads, the scheduler would try to migrate some of them to other processors for a more balanced load distribution[40].

But each of these methods work well for specific problem. We are looking for a general solution which can automatically decide on the chunk size parameter to achieve the best performance.

Chapter 3. Literature Review

3.1. Literature Review

Loop scheduling techniques has been extensively studied by different researchers. In [41] the authors propose a hybrid static/dynamic method for loop scheduling that improves the performance of dense matrix factorization, compared to both fully static and fully dynamic scheduling. The authors of [41], divide the dependency graph into two subgraphs, one of which is scheduled dynamically and the other one is scheduled statically. The tasks on the critical path are scheduled statically and each thread is forced to prioritize the static tasks[41]. They were able to improve data locality and scheduling overhead, while creating a more balanced workload.

The previous work on predicting the performance of a parallel application mainly focuses on three major types of models: analytical, trace-based, and empirical models[42].

The analytical models[43],[44],[45], while providing an arithmetic formula to represent the execution time of an application, require a deep understanding of the application, to apply platform-specific optimizations, and can not be generalized to different domains and architectures[46],[47],[48]. Traced-based models, on the other hand, use the traces collected through instrumentation, to predict the performance. These models, opposed to analytical models, do not rely on an expert's knowledge of the application, but while adding some overhead to the runtime, these models require a large storage space to save the traces, and are hard to interpret[47]. In empirical modeling, the results obtained from running an application with a set of parameters on a specific set of machines to build a model for unknown set of application and system parameters[42]. This type of modeling includes machine learning based approaches.

In [49], the authors use neural networks to predict the performance focusing on SMG2000 application, a parallel multigrid solver for linear systems[50], on two different platforms. Defining application parameters N_x , N_y , N_z , representing the working set size per processor, and P_x , P_y , P_z , describing the three-dimension processor topology, as the features, [49] uses a

fully connected neural network to learn the model. Since they use absolute mean square error as the loss function, they use stratification to replicate samples with lower values by a factor which is proportional to their target value. They also apply bagging technique to decrease the variance in the model. As they increase the size of the training set to 5K points, they reach an error rate of 4.9%.

As a trace-based model, [47] analyzes the abstract syntax tree of the code and collects data through inserting special code for instrumentation when encounters 4 different situations, namely, assignments, branches, loops, and MPI communications. The authors then use 5 different machine learning methods including random forests, support vector machine, and ridge regression to build a prediction model from the collected data. Through applying two filtration processes, they were able to decrease the amount of overhead introduced along with the storage space requirement. Their results were inclined towards random forest, mainly because of the lower impact of categorical features on it, which is helpful in general cases where we do not have any knowledge about the type of features[47].

In [42] the authors investigate a set of machine learning techniques, including deep neural networks, support vector machine, decision tree, random forest, and k-nearest neighbor to predict the execution time of 4 different applications. Each of these applications require a certain set of features as input, for example, for the miniMD application in molecular dynamics, the number of processes and the number of atoms were considered as the input features, while for miniAMR, an application for studying adaptive mesh refinement, number of processes and also block sizes in x , y , and z direction, where used as the input features. While achieving promising results especially for deep neural networks, bagging, and boosting methods, [42] suggest utilizing transfer learning through deep neural networks to predict performance on other platforms.

Although concentrating on GPUs, [51] proposes a lightweight machine learning based performance model to choose the number of threads to use for parallelization for a specific data size and operation. With the final goal of improving the training time in a neural network,

[51] selects 4 performance features collected by hardware counters namely, number of CPU cycles, number of cache misses, cache accesses for the last cache level, and number of level 1 cache hits. Then they take two different approaches to build their model. In the first one they try 10 different regression models including random forest, and in the second one they use hill climbing algorithm to choose the number of threads. In addition to hardware independent, and not requiring the training process, hill climbing algorithm achieves a much higher accuracy compared to the best performing regression model.

In this paper, we suggest using machine learning to directly predict the optimal chunk size to achieve the best performance instead of predicting the execution time or the optimal number of cores to run the application on. For this purpose, we have offered a set of general features that are not specific to an application and could easily be extracted at compile time or at run time. Once the data has been collected and our model has been created, the prediction results could be easily applied to a new application with a negligible overhead.

As another field to use machine learning, [52] collects seven runtime events and uses machine learning not to predict the performance, but to schedule the tasks. These events include, task creation, suspension, execution, completion, implicit/explicit barrier, parallel region, and finally loop/master/single region runtime events, collected through the OMPT using ORA API. Experimenting with four different machine learning techniques, including support vector machine, random forest, neural networks, and naive bayes, they would select one specific task pool configuration out of the three pre-defined options as the final classification result. Testing this framework on a real life molecular dynamics application, they observed an up to 31% improvement in performance.

The authors of [53] propose using machine learning to predict the optimal number of threads, and also the optimal scheduling policy for running an OpenMP application. Through that, they were able to develop an automatic compiler-based method to map a parallel application to a multicore processor. They collect three type of features namely, code, data, and runtime features. Code features are extracted from the code directly, and they include cycles

per instruction, number of branches, load and store instructions, and computations per instruction. While the code features could be collected statically at compile time, the data and run-time features are collected through low-cost profiling runs. This group of features include loop iteration count, branch miss rate, and $L1$ data cache miss rate. The authors of [53] then use an artificial neural network to predict the speedup achieved for a program with certain number of threads, and at the same time they use a support vector machine model to predict the best scheduling policy, out of block, cyclic, dynamic, and guided scheduling policies, for an unseen program.

Chapter 4. Method

4.1. Parallelization in Blaze

Depending on the operation and the size of operands, this assignment could be parallelized through four different backends, namely, HPX, OpenMP[17], C++ threads, and Boost[54]. Table 4.1 shows the default value for some of the threshold for parallelization applied to operations performed in Blaze. It should be noted that these thresholds should be tuned based on the parallelization backend and also the system architecture.

Benchmark	Array size
<i>DVECDVCEADD, DVECDVCECMULT</i>	38000
<i>DMATDMATADD</i>	36100 elements equivalent to a 175×175 matrix
<i>DMATDMATMULT</i>	3025 elements equivalent to a 55×55 matrix

Table 4.1. List of some of the thresholds applied to the operations performed by Blaze, starting from which the operation is executed in parallel

4.1.1. Implementation of HPX Backend

As stated earlier, as an ET-based library, blaze performs the calculations when an expression is assigned to a target, which is implemented through the *blaze::Assign* function.

The four mentioned backends, parallelize this assignment process through a parallel for-loop, in which at each iteration a specific section of each of the vectors or matrices(called a block) is selected and assigned to a core. Each core then performs the operation on the block they have been assigned to.

Each backend uses their own method for parallelizing this for loop. For HPX backend, current implementation uses a HPX *parallel::for_loop* with static chunking policy and chunk size of 1. This way, knowing the number of cores to run the application on, we can divide the

original matrix equally among the cores, while the order of assignment of blocks to the cores is known at compile time. Listings4.1 shows the current implementation of the HPX backend in Blaze.

What we suggest here is that, some prior knowledge for example, architecture of the system we are running the application on, the expression that has to be executed, number of cores of the system, size and type of the arrays we are dealing with, and etc. should be able to help us to achieve a higher performance. For this purpose we introduced two parameters `block_size` and `chunk_size`.

4.2. Experiments

In order to capture the relationship between number of cores, `chunk_size`, `block_size`, and the performance, we ran a series of experiments with different of these parameters and measured the number of floating point operations per second performed.

For these experiments ,at the first step we selected the *DMatDMatADD* benchmark which was implemented in Blazemark. *DMatDMatADD* benchmark is a level 3 BLAS function to perform matrix-matrix addition in the form of $A = B + C$, where A , B , C are square matrices of the same size. For simplification we are only studying raw-major matrices at this point. Our final goal is to extend the work to cover arbitrary data layouts for arrays.

To avoid adding the scheduling overhead for small matrix sizes, Blaze uses a threshold to start parallelization, which is specific to the type of operation. For matrix-matrix addition, if the number of elements in the matrix is greater than 36100 elements(which is equivalent to a square matrix of size 190×190) Blaze uses the configured backend to parallelize the assignment operation. For this reason, we start our experiments with matrix size of 200x200 and gradually increase the size to 1587×1587 . Table 4.2 show the matrix sizes and the number of cores chosen for our experiments with *DMATDMATADD* benchmark.

Listing 4.1: Previous implementation of Assign function for HPX backend in Blaze.

```

1 template< typename MT1 // Type of the left-hand side dense matrix
2 , bool SO1 // Storage order of the left-hand side dense matrix
3 , typename MT2 // Type of the right-hand side dense matrix
4 , bool SO2 // Storage order of the right-hand side dense matrix
5 , typename OP > // Type of the assignment operation
6 void hpxAssign( DenseMatrix<MT1,SO1>& lhs , const DenseMatrix<MT2,SO2>& rhs , OP op )
7 {
8     using hpx::parallel::for_loop;
9     using hpx::parallel::execution::par;
10
11    BLAZE_FUNCTION_TRACE;
12
13    using ET1 = ElementType_t<MT1>;
14    using ET2 = ElementType_t<MT2>;
15
16    constexpr bool simdEnabled( MT1::simdEnabled && MT2::simdEnabled && IsSIMDCombinable_v<ET1,ET2> );
17    constexpr size_t SIMDSize( SIMDTrait< ElementType_t<MT1> >::size );
18
19    const bool lhsAligned( (~lhs).isAligned() );
20    const bool rhsAligned( (~rhs).isAligned() );
21
22    const size_t threads ( getNumThreads() );
23    ThreadMapping threadmap( createThreadMapping( threads , ~rhs ) );
24
25    const size_t addon1 ( ( ( (~rhs).rows() % threadmap.first ) != 0UL )? 1UL : 0UL );
26    const size_t equalShare1( (~rhs).rows() / threadmap.first + addon1 );
27    const size_t rest1 ( equalShare1 & ( SIMDSize - 1UL ) );
28    const size_t rowsPerThread( ( simdEnabled && rest1 )?( equalShare1 - rest1 + SIMDSize ):( equalShare1 ) );
29
30    const size_t addon2 ( ( ( (~rhs).columns() % threadmap.second ) != 0UL )? 1UL : 0UL );
31    const size_t equalShare2( (~rhs).columns() / threadmap.second + addon2 );
32    const size_t rest2 ( equalShare2 & ( SIMDSize - 1UL ) );
33    const size_t colsPerThread( ( simdEnabled && rest2 )?( equalShare2 - rest2 + SIMDSize ):( equalShare2 ) );
34
35    for_loop( par , size_t(0) , threads , [&](int i)
36    {
37        const size_t row ( ( i / threadmap.second ) * rowsPerThread );
38        const size_t column( ( i % threadmap.second ) * colsPerThread );
39
40        if( row >= (~rhs).rows() || column >= (~rhs).columns() )
41            return;
42
43        const size_t m( min( rowsPerThread , (~rhs).rows() - row ) );
44        const size_t n( min( colsPerThread , (~rhs).columns() - column ) );
45
46        if( simdEnabled && lhsAligned && rhsAligned ) {
47            auto target( submatrix<aligned>( ~lhs , row , column , m , n ) );
48            const auto source( submatrix<aligned>( ~rhs , row , column , m , n ) );
49            op( target , source );
50        }
51        else if( simdEnabled && lhsAligned ) {
52            auto target( submatrix<aligned>( ~lhs , row , column , m , n ) );
53            const auto source( submatrix<unaligned>( ~rhs , row , column , m , n ) );
54            op( target , source );
55        }
56        else if( simdEnabled && rhsAligned ) {
57            auto target( submatrix<unaligned>( ~lhs , row , column , m , n ) );
58            const auto source( submatrix<aligned>( ~rhs , row , column , m , n ) );
59            op( target , source );
60        }
61        else {
62            auto target( submatrix<unaligned>( ~lhs , row , column , m , n ) );
63            const auto source( submatrix<unaligned>( ~rhs , row , column , m , n ) );
64            op( target , source );
65        }
66    } );
67 }
```

Listing 4.2: New implementation of Assign function for HPX backend in Blaze.

```

1 template< typename MT1    // Type of the left-hand side dense matrix
2 , bool SO1      // Storage order of the left-hand side dense matrix
3 , typename MT2    // Type of the right-hand side dense matrix
4 , bool SO2      // Storage order of the right-hand side dense matrix
5 , typename OP > // Type of the assignment operation
6 void hpxAssign( DenseMatrix<MT1,SO1>& lhs , const DenseMatrix<MT2,SO2>& rhs , OP op )
7 {
8     using hpx::parallel::for_loop;
9     using hpx::parallel::execution::par;
10
11    BLAZE_FUNCTION_TRACE;
12
13    using ET1 = ElementType_t<MT1>;
14    using ET2 = ElementType_t<MT2>;
15
16    constexpr bool simdEnabled( MT1::simdEnabled && MT2::simdEnabled && IsSIMDCombinable_v<ET1,ET2> );
17    constexpr size_t SIMDSize( SIMDTrait< ElementType_t<MT1> >::size );
18
19    const bool lhsAligned( (~lhs).isAligned() );
20    const bool rhsAligned( (~rhs).isAligned() );
21
22    const size_t threads   ( getNumThreads() );
23    const size_t numRows ( min( static_cast<std::size_t>( BLAZE_HPX_MATRIX_BLOCK_SIZE_ROW ), (~rhs).rows() ) );
24    const size_t numCols ( min( static_cast<std::size_t>( BLAZE_HPX_MATRIX_BLOCK_SIZE_COLUMN ), (~rhs).columns()
25        ) );
26
27    const size_t rest1      ( numRows & ( SIMDSize - 1UL ) );
28    const size_t rowsPerIter( ( simdEnabled && rest1 )?( numRows - rest1 + SIMDSize ):( numRows ) );
29    const size_t addon1     ( ( ( (~rhs).rows() % rowsPerIter ) != 0UL )? 1UL : 0UL );
30    const size_t equalShare1( (~rhs).rows() / rowsPerIter + addon1 );
31
32    const size_t rest2      ( numCols & ( SIMDSize - 1UL ) );
33    const size_t colsPerIter( ( simdEnabled && rest2 )?( numCols - rest2 + SIMDSize ):( numCols ) );
34    const size_t addon2     ( ( ( (~rhs).columns() % colsPerIter ) != 0UL )? 1UL : 0UL );
35    const size_t equalShare2( (~rhs).columns() / colsPerIter + addon2 );
36
37    hpx::parallel::execution::dynamic_chunk_size chunkSize ( BLAZE_HPX_MATRIX_CHUNK_SIZE );
38
39    for_loop( par.with( chunkSize ), size_t(0), equalShare1 * equalShare2, [&](int i)
40    {
41        const size_t row   ( ( i / equalShare2 ) * rowsPerIter );
42        const size_t column( ( i % equalShare2 ) * colsPerIter );
43
44        if( row >= (~rhs).rows() || column >= (~rhs).columns() )
45            return;
46
47        const size_t m( min( rowsPerIter, (~rhs).rows() - row ) );
48        const size_t n( min( colsPerIter, (~rhs).columns() - column ) );
49
50        if( simdEnabled && lhsAligned && rhsAligned ) {
51            auto target( submatrix<aligned>( ~lhs, row, column, m, n ) );
52            const auto source( submatrix<aligned>( ~rhs, row, column, m, n ) );
53            op( target, source );
54        }
55        else if( simdEnabled && lhsAligned ) {
56            auto target( submatrix<aligned>( ~lhs, row, column, m, n ) );
57            const auto source( submatrix<unaligned>( ~rhs, row, column, m, n ) );
58            op( target, source );
59        }
60        else if( simdEnabled && rhsAligned ) {
61            auto target( submatrix<unaligned>( ~lhs, row, column, m, n ) );
62            const auto source( submatrix<aligned>( ~rhs, row, column, m, n ) );
63            op( target, source );
64        }
65        else {
66            auto target( submatrix<unaligned>( ~lhs, row, column, m, n ) );
67            const auto source( submatrix<unaligned>( ~rhs, row, column, m, n ) );
68            op( target, source );
69        }
70    });

```

Category	Configuration
Matrix sizes	200, 230, 264, 300, 396, 455, 523, 600, 690, 793, 912, 1048, 1200, 1380, 1587
Number of cores	1, 2, 3, 4, 5, 6, 7, 8
Number of rows in the block	4, 8, 12, 16, 20, 32
Number of columns in the block	64, 128, 256, 512, 1024
Chunk size	Between 1 and total number of blocks (logarithmic increase)

Table 4.2. List of different values used for each variable for running the *DMATDMATADD* benchmark

Figure 4.2 shows the results of running *DMatDMatADD* benchmark for matrix sizes and number of cores listed in Table 4.2 based on grain size.

On the other hand, Figure 4.3 integrates the results obtained from running the same benchmark with different matrix sizes. Each color in this graph represents a specific matrix size.

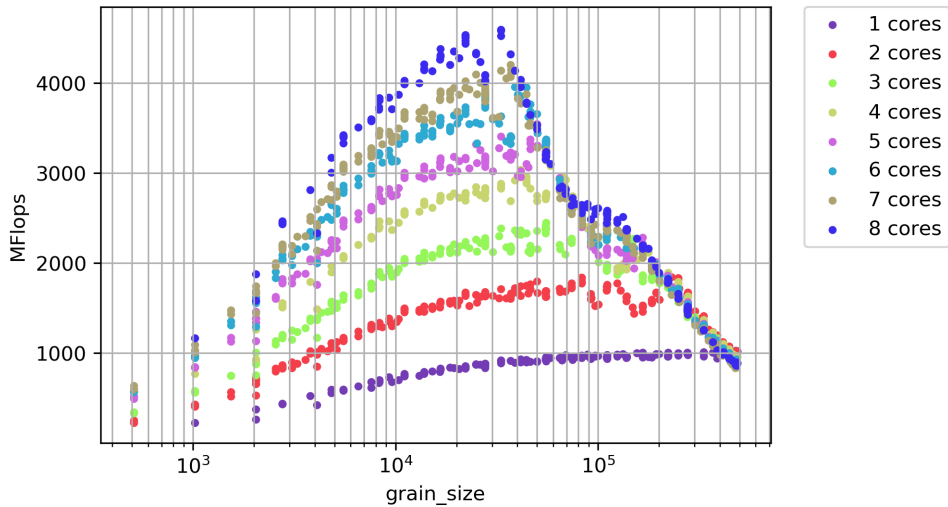


Figure 4.1. The results obtained from running *DMATDMATADD* benchmark through Blaze-mark for matrix size 690×690 on different number of cores.

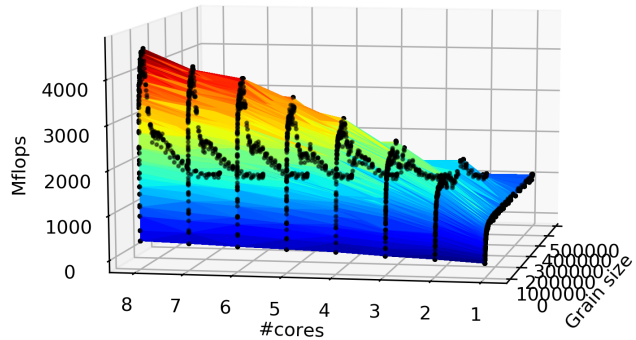
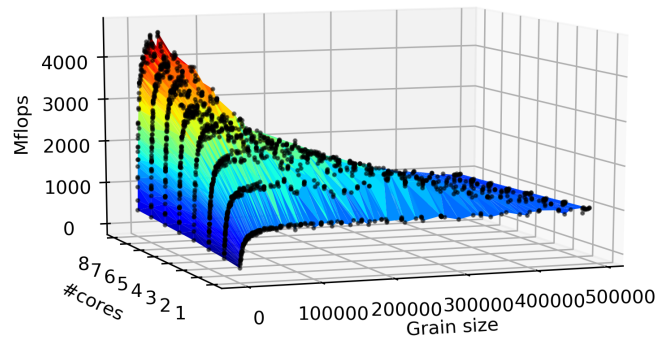


Figure 4.2. The results obtained from running *DMATDMATADD* benchmark through Blaze-mark for matrix of size 690×690 from two different angles

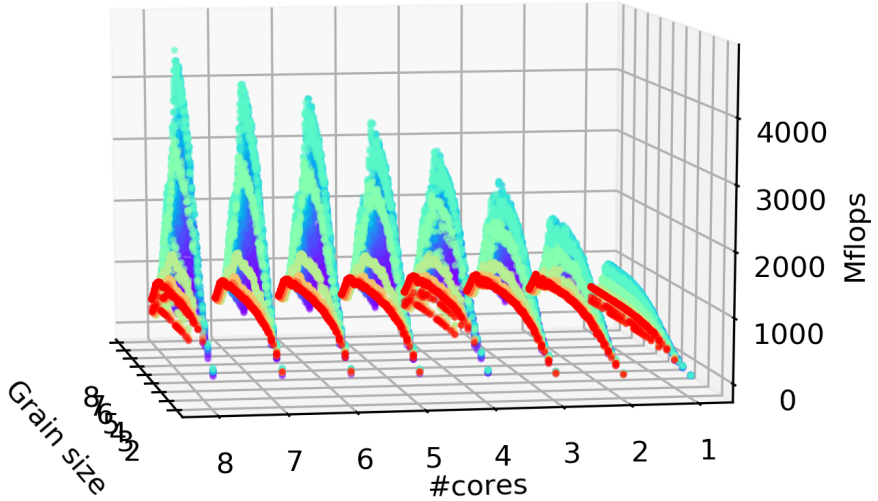


Figure 4.3. The results obtained from running *DMATDMATADD* benchmark through Blaze-mark for matrix sizes from 200×200 to 1587×1587

4.2.1. Observation

The final purpose of our experiments is to find a chunk size that gives us the best performance for a given matrix size on a given machine. This chunk size should also be tailored to the expression being executed, and this all is based on assuming that we have already fixed the block size. So the first step appeared to be selecting the block size. For this purpose, we ran the experiments with a selection of block sizes as shown in Table 4.2.

It should be mentioned that there were three constraints on selecting the block sizes. First, Blaze forces the number of columns in a raw-major matrix to be divisible to SIMD register size in order to be able to take advantage of vectorization. Second, we have selected the number of columns in our blocks to be either divisible by cache line or to contain all the columns of the matrix.

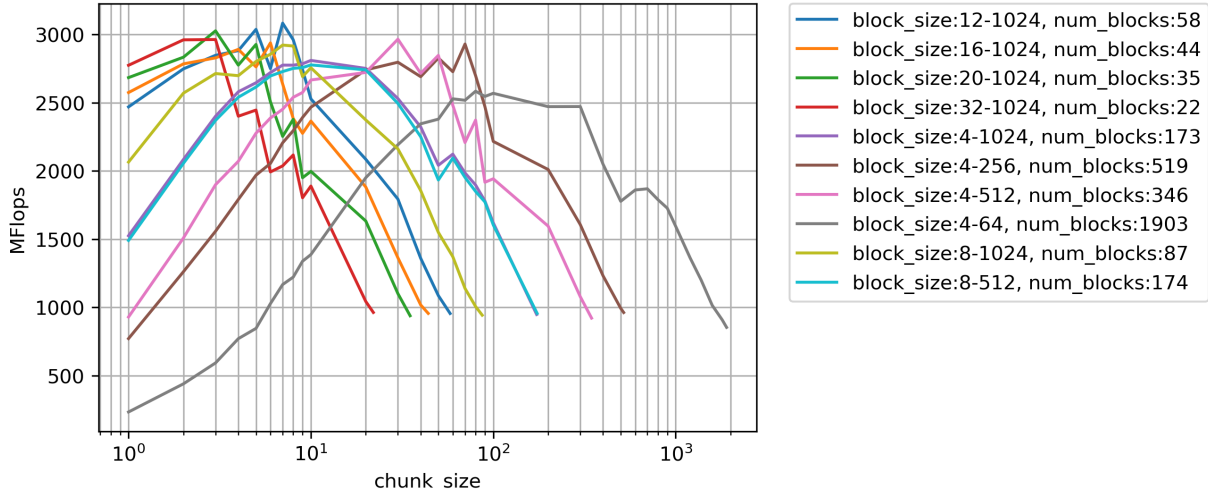


Figure 4.4. The results obtained from running *DMATDMATADD* benchmark through Blaze-mark for matrix size 690×690 with different combinations of block size and chunk size on 4 cores

The collected data, as seen in Figure 4.4, suggests two main points:

- For each selected block size, there is a range of chunk sizes that gives us the best performance.
- Except for some uncommon cases, no matter which block size we choose, we are able to achieve the maximum performance if we select the right chunk size.

This motivated us to move our search parameter from chunk size to grain size. As stated earlier, grain size is the amount of work assigned to one HPX thread. Here we represent grain size by number of floating point operations performed by a HPX thread. For example, performing addition among two matrices, if we choose the block size as 4×64 and chunk size as 3, the grain size would be $3 \times 4 \times 64 = 768$. Note that in our experiments whenever the number of columns of the original matrix is not divisible to the selected number of columns for block size, there would be a set of blocks with less number of elements than the selected block size, this has been considered when calculating the grain size.

By changing our focus to the grain size instead of the block size and the chunk size, Figure 4.5 shows how the throughput changes with regards to the grain size for the *DMATDMATADD*

benchmark, for each specific block size. Each combination of block size and chunk size generates a point in the graph. On the other hand, Figure 4.1 looks at these graphs from another aspect, keeping the problem size constant but changing the number of the cores to run the benchmark on, instead.

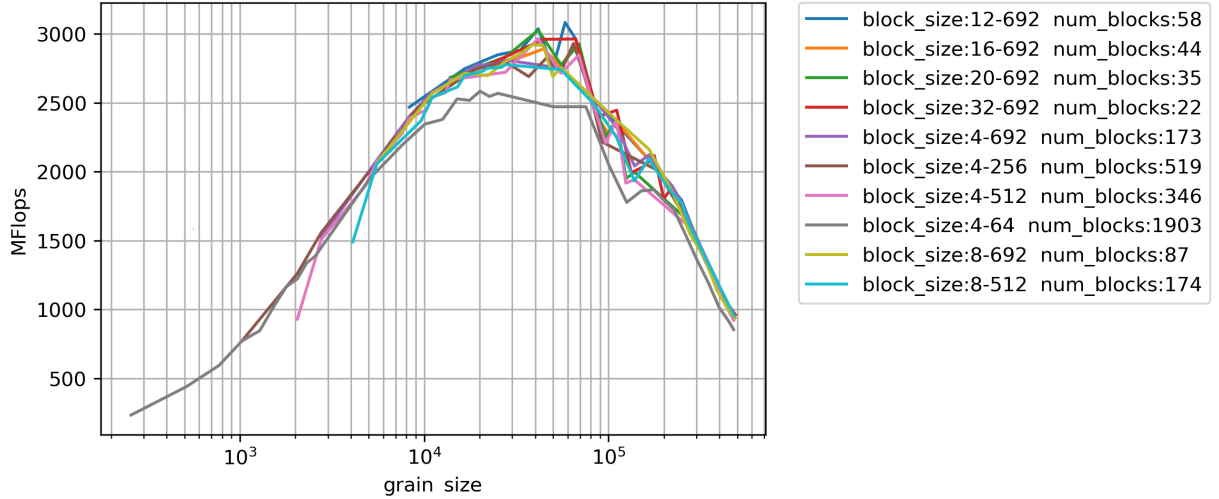


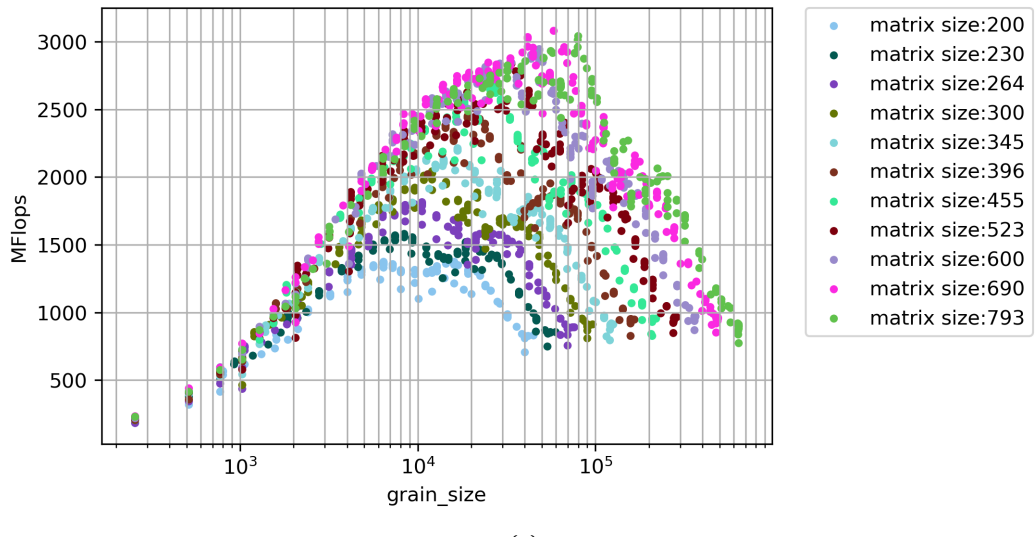
Figure 4.5. The results obtained from running *DMATDMATADD* benchmark through Blaze-mark for matrix size 690×690 on 4 cores.

4.3. Method

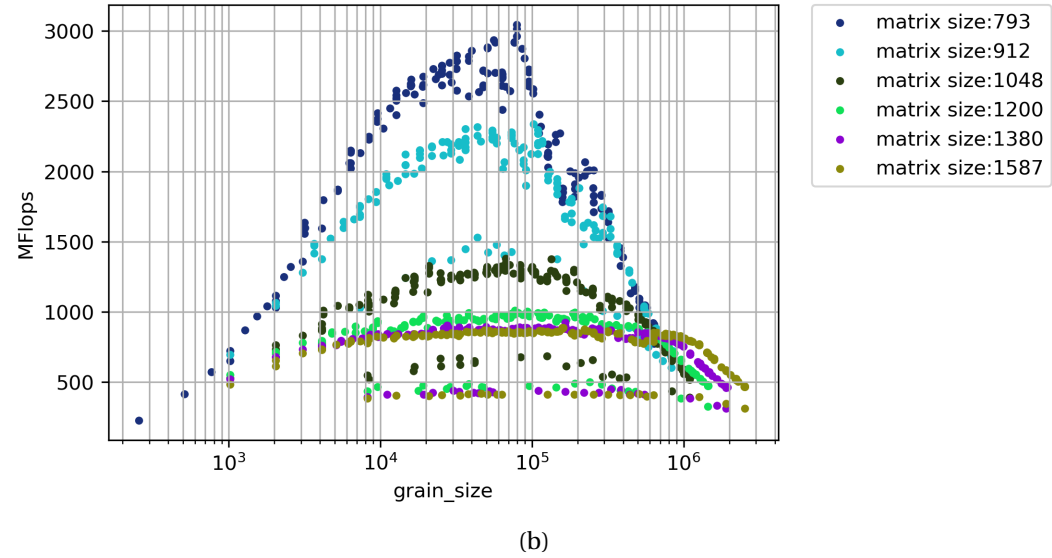
Looking at the throughput vs. grain size graphs and the consistent pattern observable motivated us to try to model the relationship between throughput and grain size. In order to simplify the process and eliminate the effect of different possible factors, we started with limiting the problem to a fixed matrix size.

4.3.1. Polynomial Fit

In our first attempt we used a 2nd degree polynomial to model throughput against grain size. For each matrix size, we fitted the corresponding graphs shown in Figure 4.6 to a second degree polynomial.



(a)



(b)

Figure 4.6. Throughput vs. grain size graph obtained from running *DMATDMATADD* benchmark on 4 cores for matrix sizes (a) smaller than 793×793 and (b) larger than 793×793 .

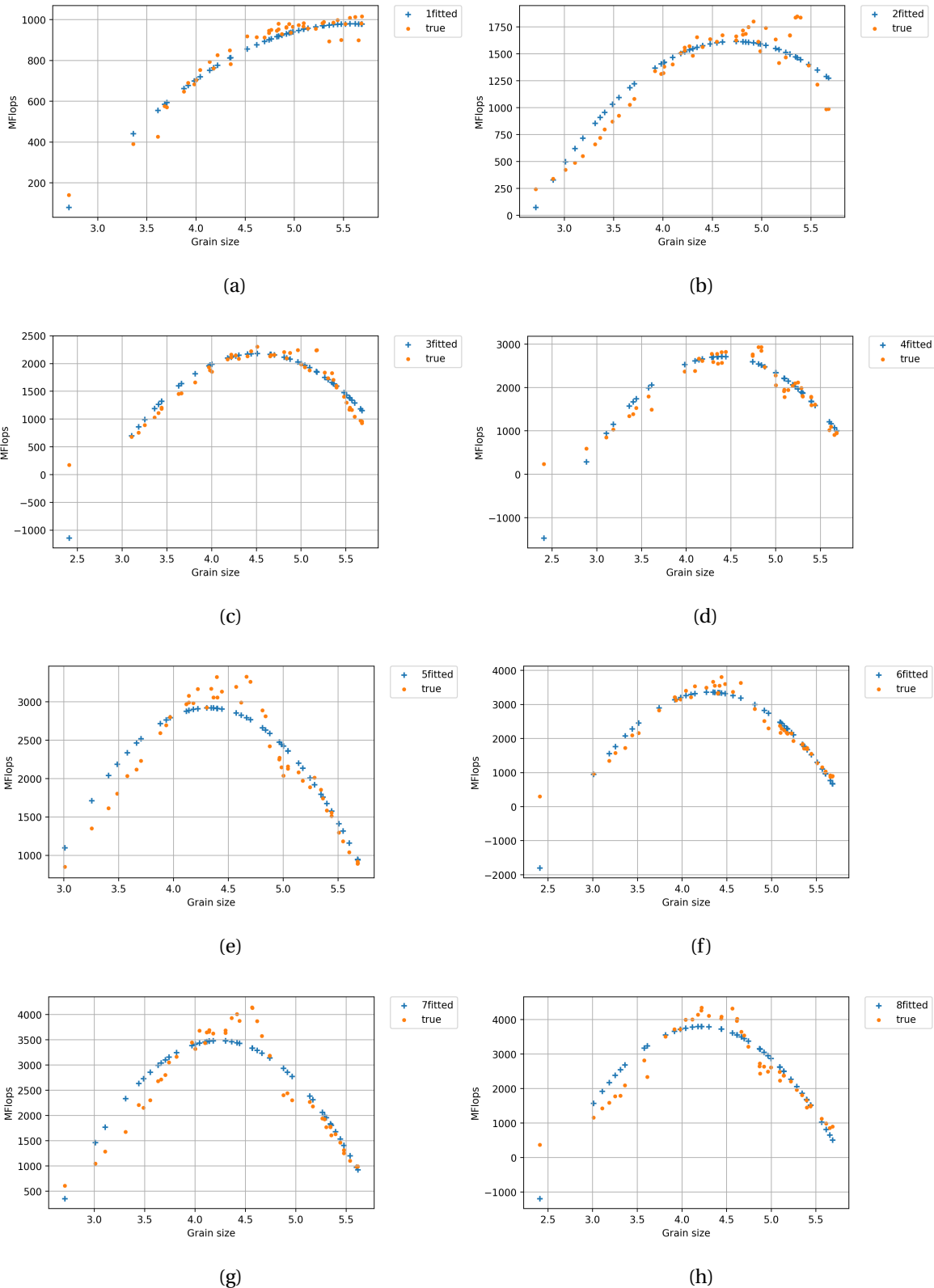


Figure 4.7. The results of fitting the throughput vs grain size data into a 2d polynomial for *DMATDMATADD* benchmark for matrix size 690×690 with different number of cores on the test data set (a) 1 core, (b) 2 cores, (c) 3 cores, (d) 4 cores, (e) 5 cores, (f) 6 cores, (g) 7 cores, (h) 8 cores.

Figure 4.7 shows the results of using a quadratic function to fit the data for one matrix size with different number of threads. We used the *polyfit* package from *numpy* library in *python* for this purpose, which tries to minimize the least-square error over all the samples.

For our experiment, we divided the data into two sections, training and test. 60% of the data was randomly chosen for the training part and the rest was considered as the test set. The training set was used to find the best 2nd degree polynomial for the data, and once the parameters were identified, the generated 2nd degree polynomial was applied to the test set to measure how good our fit was performing.

For the matrix size 690×690 our dataset contained 117 data points, 72 of which were randomly selected to build the model. The mean relative error for each number of cores, calculated using Equation 4.1, is represented in Figure 4.8 for training and test set. In this equation, t_i and p_i denote the true value and the predicted value of the i th sample respectively, where n is the number of samples with the particular number of cores.

$$\text{Relative Error} = \frac{1}{n} \sum_{i=1}^n 1 - p_i / t_i \quad (4.1)$$

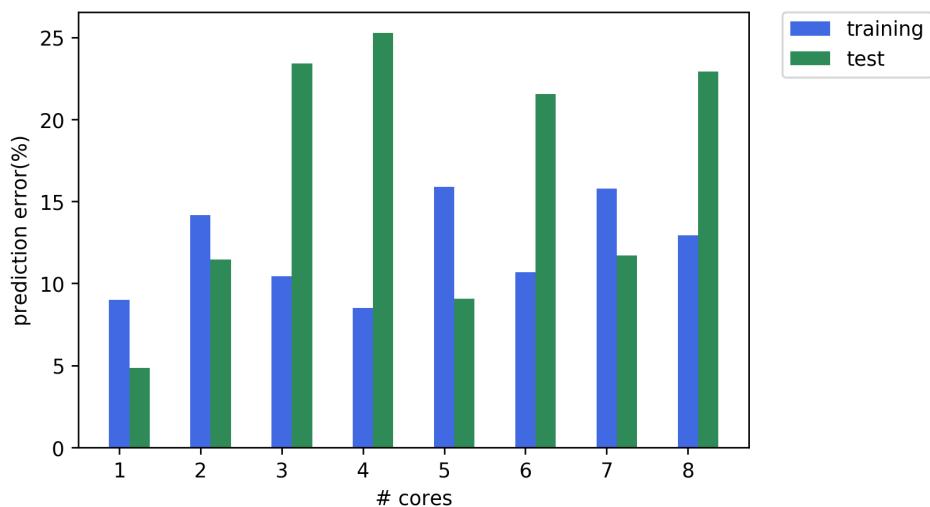


Figure 4.8. The training and test error for fitting data obtained from the *DMATDMATADD* benchmark for matrix size 690×690 against different number of cores cores.

4.3.1.1. Generalizing the fitted function to include number of cores

In this step, we try to generalize the fitted 2nd degree polynomial obtained from the previous step, represented by $P = ag^2 + bg + c$, where P is the throughput and g is the grain size, by looking at how the three parameters a , b , and c change when number of cores changes. A 3rd degree polynomial seems to a reasonable fit for each of these parameters, in regards to number of cores. In order to avoid overfitting, we excluded two of the data points(2 and 5) from the data points used for fitting the polynomial and tested the fitted function on those two points to see how well the function is working on unseen data points.

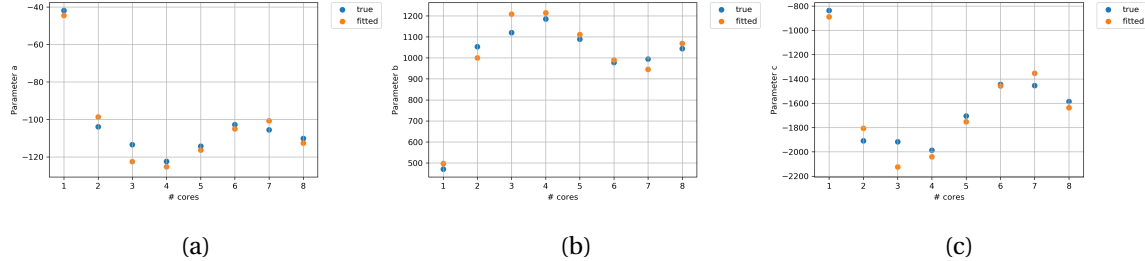


Figure 4.9. Fitting the parameters of the quadratic function with a 3rd degree polynomial from the *DMATDMATADD* benchmark for matrix size 690×690 against different number of cores.

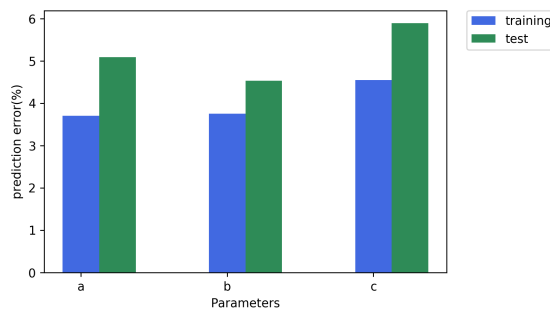


Figure 4.10. The error in fitting the parameters a , b , and c for matrix size 690×690 .

Using this 3rd degree polynomial to fit the parameters, we can generalize the relationship between throughput and grain size in the following equation:

$$P = a_{11}g^3 + a_{10}g^2N^2 + \dots + a_1N + a_0 \quad (4.2)$$

where P is the throughput, g is the grain size, and N is the number of cores and coefficients a_{11}, \dots, a_0 are the real values.

Knowing that a polynomial of degree 2 in terms of grain size and of degree 3 in terms of number of cores, we can try to fit our original data directly to the above mentioned formula (Equation 4.2). The results of the original data obtained from running *DMATDMATADD* benchmark, the fitted polynomial based on Equation 4.2, is represented in Figure 4.11, for 2,4, and 8 cores for a matrix of size 690×690 .

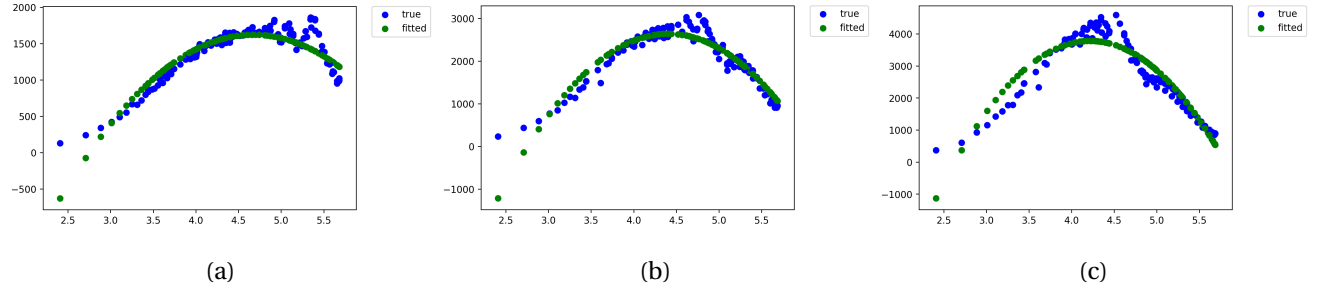


Figure 4.11. Results of fitting the data from *DMATDMATADD* benchmark with a polynomial of degree 2 in terms of grain size and of degree 3 in terms of number of cores for matrix size 690×690 for (a) 2 core, (b) 4 cores, (c) 8 cores.

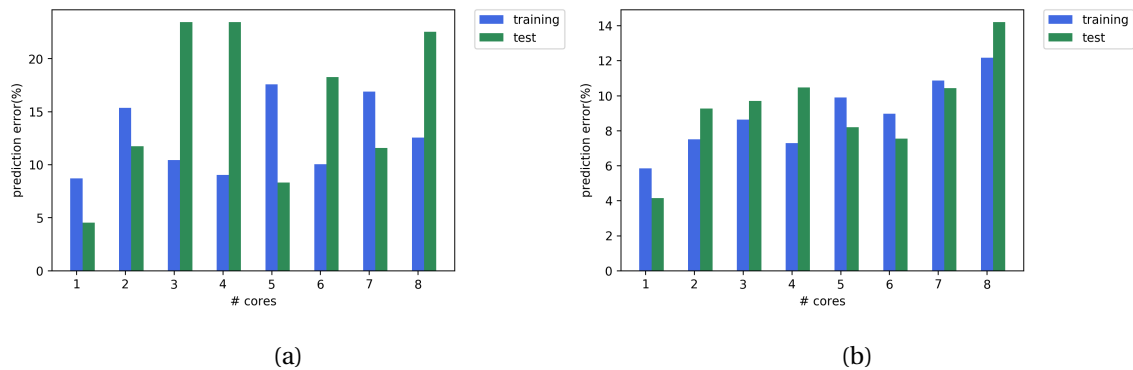


Figure 4.12. The training and test error obtained fitting the data to a polynomial of degree 2 in terms of grain size and of degree 3 in terms of number of cores for matrix size 690×690 , for each number of cores. (a) All the data points are include in calculation of error, (b) the leftmost sample was removed from error calculation.

Figure 4.12a shows the obtained relative error on both training and test sets. The graph

suggests a higher test error compared to the training error, mostly caused by the left hand side of the graph. The effect of removing the leftmost sample from error calculations is depicted in Figure 4.12b.

Although we are interested in finding a model that results in a low training and test error, our purpose is mainly finding the region that generates the highest performance. So, even though our model might not match the original data in all data points, due to having a different nature than a quadratic function, our focus would be on how this fit can help us to find which range of grain sizes, or how big the task sizes should be, to achieve the highest performance.

4.3.1.2. Finding the range of grain size to achieve the highest performance

The major advantage of using a quadratic function to fit the data in terms of grain size, when number of cores is fixed, is the simplicity of the formula, which makes it possible for us to find the peak of the graph very easily. In order to add some uncertainty to our prediction, instead of finding the maximum of the quadratic function, we identified the range of grain size that results in a performance within 10% of the maximum performance. For a second degree polynomial in terms of g , $P = ag^2 + bg + c$, the minimum or maximum of the polynomial is located at $p^* = \frac{-b}{2a}$, and a, b, c are 3rd degree polynomials of number of cores.

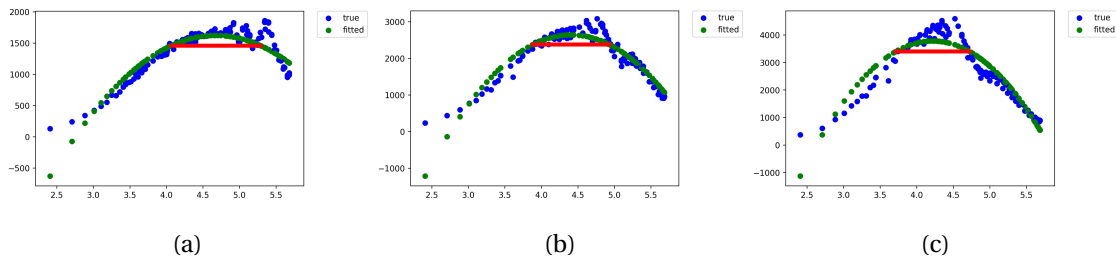


Figure 4.13. The range of grain size (shown as the red line) that leads to a performance within 10% of the maximum performance for (a) 2 cores, (b) 4 cores and (b) 8 cores.

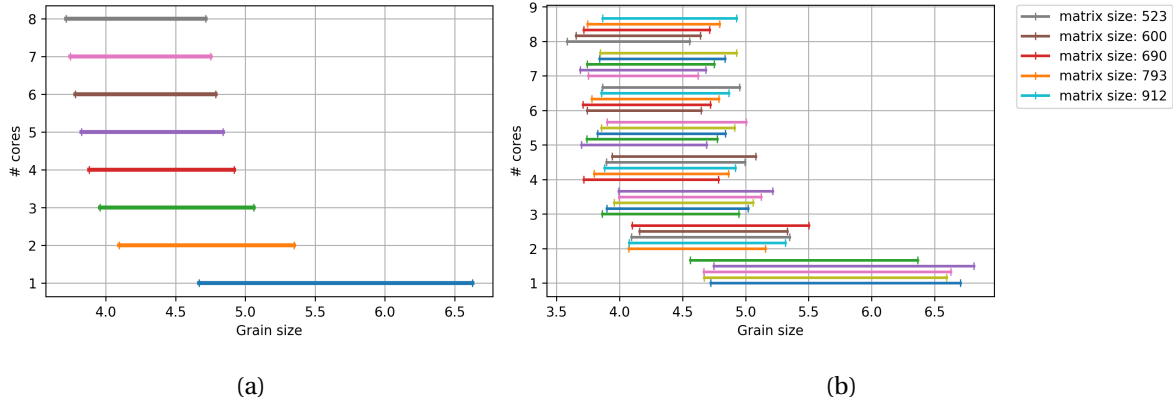


Figure 4.14. The range of grain size within 10% of the maximum performance of the fitted polynomial function for *DMATDMATADD* benchmark for different number of cores for (a) matrix size 690×690 (b)matrix size 523×523 to 912×912 .

Figure 4.14a shows the calculated range for matrix size 690×690 for each specific number of threads, while Figure 4.14b compares the range for different matrix sizes.

4.3.1.3. Estimating the chunk size

Once we identified a range of grain sizes that is expected to lead us to highest achievable performances for a specific matrix size and a specific number of cores, the next step is finding the possible combinations of block size and chunk size to achieve that range of grain sizes. As stated earlier in this chapter, results obtained from Figure 4.5 suggests that with a fixed grain size, our choice of block size does not affect the performance directly, as long as there exist a chunk size that when combined by the block size could result in the specified grain size.

In our experiment, we selected our block size to be 4×256 . With this assumption, in order for the grain size to be within the specified range for each matrix size, chunk size has to be within a specific range size too.

For example, for a 690×690 matrix we calculated the range of maximum performance for 4 cores to be $[3.88, 4.92]$ in logarithmic scale which is equivalent to $[7586, 83176]$. Setting the block size to 4×256 , this range forces the chunk size to be within the range $[9, 90]$. The range of chunk sizes to match the range of grain sizes identified, and their corresponding throughput

is shown in Figure 4.15, for matrix size 690×690 and block size 4×256 . The green line is the throughput achieved by the current implementation of HPX backend. Since the graph from the original data is skewed to right, we selected the point after the median of all the chunk sizes in the range, as our candidate chunk size for this specific configuration.

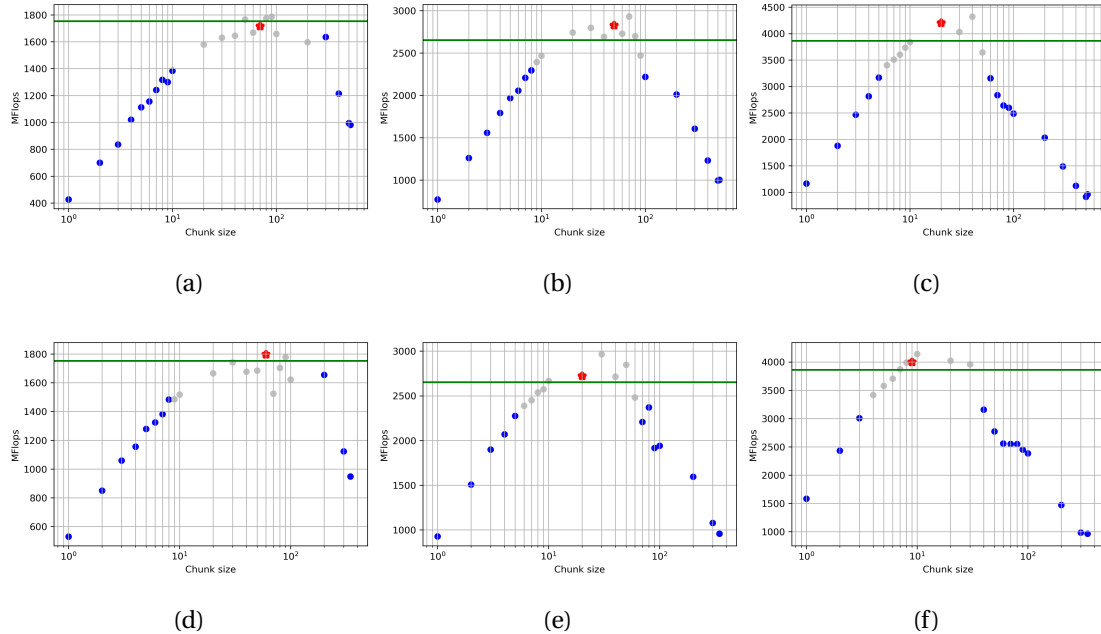


Figure 4.15. The range of chunk sizes to produce a grain size within 10% of the maximum performance of the fitted quadratic function for *DMATDMATADD* benchmark for matrix size 690×690 with block size of 4×256 on (a) 2 cores, (b) 4 cores, and (c) 8 cores, and block size of 4×512 on (d) 2 cores, (e) 4 cores, and (f) 8 cores. Silver points denotes the detected range of chunk size, and the red star shows the median point.

Chapter 5. Understanding the effect of grain size on concurrency in an asynchronous many-task runtime system

5.1. Analytical Modeling

In the previous section we studied the possibility of using a polynomial to capture the relationship between grain size, number of cores, and throughput for a fixed matrix size, with the purpose of finding a range of grain size that leads us to maximum performance. Although the polynomial function was helpful in directing us toward our objective, it does not have a physical implication.

This motivated us to change our view, and instead of looking just at the data and trying to find a function to fit the data, study the behavior of the data, and then find a function that would be likely to fit the data. That function would be a good fit mostly because that's how we expect the throughput to change with grain size, and not just how the data looks like.

In this chapter we attempt to understand the effect of grain size on the achievable speedup in an asynchronous many-task runtime system. As discussed in chapter 2, our knowledge here comes from Amdahl's law and Universal Scalability Law, which is an extension to the Amdahl's law. What these models suggest is that as we increase the number of cores in a multicore system, we do not observe a linear speedup, and they hold two major factors accountable for that, latency.

Here we are interested in developing an analytical model for predicting the execution time in an asynchronous many task runtime system. Even if we were able to determine all the factors affecting the execution time, it is still very hard to find an analytical model describing the relationship between these factors and the execution time.

Definitions: We represent the overhead of creating one task with α , the number of tasks created with num_tasks , the maximum amount of work assigned to one core as w_c , the number of cores that are actually doing the work as M , the total amount of work available as $problem_size$, and the sequential execution time as t_{seq} .

In an attempt to find this analytical model, we started with looking into two major factors,

the overhead of creating tasks, and the maximum amount of work assigned to one core. In order to understand how these factors contribute to the execution time, we created a benchmark based on a simple *for_loop* with different number of iterations, iteration lengths, and chunk sizes, as shown in Listing 5.1. Each iteration consists of a while loop that makes sure the iteration lasts a certain amount of time. This way, knowing how long it would take to execute one iteration(denoted as *iter_length*), how many of iterations are executed by one HPX thread(denoted as *chunk_size*), and finally how many iterations there are(denoted as *num_iterations*), we can see how the execution time changes when the problem is executed on different number of cores. Here, we define *problem_size* as the time it takes to execute all the iterations, which is:

$$problem_size = iter_length \times num_iterations \quad (5.1)$$

The total execution time could be assumed to be the maximum amount of time it takes for one of the cores to finish its work. With this assumption, the execution time for this simple problem, could be estimated as summation of maximum overhead of creating tasks on one core and the time it takes to run *w_c* amount of work on *N* cores, shown in Formula 5.2.

$$\begin{aligned} execution_time &= \alpha \times \left\lceil \frac{num_tasks}{M} \right\rceil + (1 + \gamma \times (M - 1)) \times w_c \\ &= \alpha \times \left\lceil \frac{num_tasks}{M} \right\rceil + w_c + \gamma \times w_c \times (M - 1) \end{aligned} \quad (5.2)$$

Where,

$$w_c = \begin{cases} problem_size & \text{if } N = 1 \\ problem_size - g \times (N - 1) \times (\left\lceil \frac{num_tasks}{N} \right\rceil - 1) & \text{or} \\ g \times (\left\lceil \frac{num_tasks}{M} \right\rceil - 1) + (num_iterations \% chunk_size) \times (iter_length) & (5.3) \\ \text{if } num_tasks \% N = 1 \text{ and } num_iterations \% chunk_size \neq 0 \\ g \times \left\lceil \frac{num_tasks}{N} \right\rceil & \text{otherwise} \end{cases}$$

The first part of Formula 5.2 is associated with the overhead of creating num_tasks tasks on N cores, and the second part is deducted from the Universal Scalability Law[9], that suggests an overhead associated with the number of cores should be added to the expected execution time. Here, we adjusted the total number of cores (N) to the number of cores that are actually executing the work (M).

Assuming we are running our application on N cores, with a grain size equal to g , num_tasks tasks are being created, and M cores are actually doing the work. If $num_tasks < N$, M would be equal to num_tasks , otherwise $M = N$.

$$M = \begin{cases} num_tasks & \text{if } num_tasks < N \\ N & \text{otherwise} \end{cases} \quad (5.4)$$

Figure shows an example of the results obtained from running the benchmark, for $problem_size = 10000$, on different number of cores.

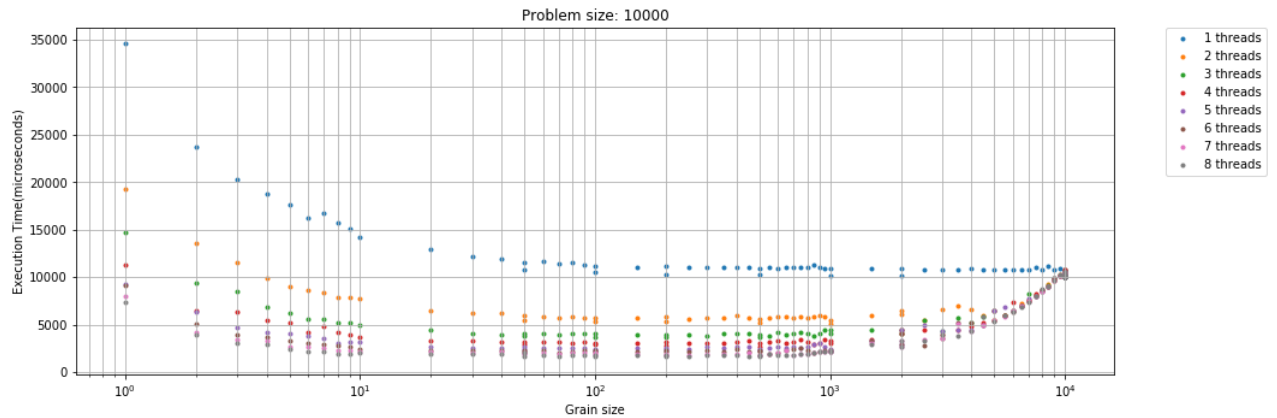


Figure 5.1. The results of running the benchmark in Listing with $problem_size = 10000$, on different number of cores.

As stated in Chapter 2, at the right hand side of the graph in Figure 5.1, the number of tasks created is smaller than the number of cores which results in making at least one of the cores idle, while the other cores are assigned a rather big chunk of work. The performance degradation we observe in that points is associated with starvation, meaning that we are not

utilizing our computation resources to the full extent. In these points, the number of cores actually doing the work is equal to the number of the tasks, since each core gets to execute at most one task. At this region of the graph, the maximum assigned work to a core (w_c) is the dominant factor.

On the other hand, on the left hand side of the graph, since the grain size is very small we end up with creating a large number of tasks. Since there is an overhead associated with each created task, we observe a performance degradation in that region. As the grain size increases, the number of created tasks, and the overhead associated to that decreases consequently. At this region of the graph, the overhead of creating the tasks($\alpha \times num_tasks$) is the dominant factor.

From overhead point of view though, if we represent the overhead of creating one task on a particular machine as α , the overhead of creating num_tasks tasks would be $\alpha \times num_tasks$, but this overhead is divided between the M cores actually doing the work.

To summarize, for this simple experiment, where we do not have to deal with the cash effects, we believe the important factors are: number of HPX threads being created, number of cores the program is ran on, the maximum amount of work one core has to perform. The maximum number of tasks assigned to one core, and the number of cores that are actually performing the work, are two other important factors that can be deducted from the aforementioned factors.

5.1.1. Analyzing the Data

Assuming the proposed formula is a good fit for this problem, we can estimate the range of grain sizes for which we achieve the lowest execution time, for a specific problem size, run on specific number of cores.

5.1.1.1. Left side of the graph

As stated earlier, in Formula 5.2, for small grain sizes the first term is the dominant factor while the second term roughly stays constant. Same way, for large grain sizes the second factor

is the dominant factor.

In order to find the lower-bound of the range for which the execution time stays constant, we assume the second factor is constant in that region. Taking the derivative of the function based on the grain size then leads to:

$$\begin{aligned}
 \frac{\partial \text{execution_time}}{\partial g} &= \frac{\alpha}{M} \times \frac{\partial \text{num_tasks}}{\partial g} \\
 &= \frac{\alpha}{M} \times \frac{\partial \text{problem_size}}{\partial g} \\
 &= \frac{\alpha \times \text{problem_size}}{M} \times \frac{-1}{g^2} \\
 &= \frac{\alpha \times \text{problem_size}}{N} \times \frac{-1}{g^2}
 \end{aligned} \tag{5.5}$$

From Formula 5.5, it can be observed that for the left hand side of the graph the rate of changes is negative and decreases as the grain size increases. Here we are looking for the value of the grain size for which the rate of change becomes very small (we introduce a threshold λ_b for this purpose).

$$\begin{aligned}
 \frac{\alpha \times \text{problem_size}}{N} \times \frac{1}{g^2} &\leq \lambda_b \\
 g^2 &\geq \frac{\alpha \times \text{problem_size}}{N \times \lambda_b} \\
 g &\geq \sqrt{\frac{\alpha \times \text{problem_size}}{N \times \lambda_b}}
 \end{aligned} \tag{5.6}$$

Formula 5.6 can also be represented as shown in Formula 5.7. This representation shows that when the ratio of the time it takes to execute one task to the total overhead of creating num_tasks tasks on N core, is greater than a threshold, we will end up in the flat region of the graph, close to the left hand side.

$$\begin{aligned}
\frac{\alpha}{N} \times \frac{\text{problem_size}}{g} \times \frac{1}{g} &\leq \lambda_b \\
\frac{\alpha}{N} \times \text{num_tasks} \times \frac{1}{g} &\leq \lambda_b \\
\frac{\alpha \times \text{num_tasks}}{N} &\leq g \times \lambda_b \\
\frac{g}{\frac{\alpha \times \text{num_tasks}}{N}} &\geq \frac{1}{\lambda_b}
\end{aligned} \tag{5.7}$$

5.1.1.2. Right side of the graph

Now looking at the right hand side of the graph, the overhead of creating the tasks becomes negligible on that side, since only few tasks are being created and the overhead of creating these many tasks is not significant compared to the execution time. On this side, the maximum amount of work executed by one core (w_c) is the dominant factor.

Formula 5.3 shows how w_c is calculated for different cases, but in general we can estimate w_c with $g \times \left\lceil \frac{\text{num_tasks}}{N} \right\rceil$.

$$\begin{aligned}
w_c &\approx g \times \left\lceil \frac{\text{num_tasks}}{N} \right\rceil \\
&\approx g \times \left\lceil \frac{\frac{\text{problem_size}}{g}}{N} \right\rceil \\
&\approx \frac{\text{problem_size}}{N}
\end{aligned} \tag{5.8}$$

What happens here is that as the grain size changes, there are points for which $\left\lceil \frac{\text{num_tasks}}{N} \right\rceil$ is the same but since the grain size is different, a different w_c would be resulted. For all the values of g that create the same $\left\lceil \frac{\text{num_tasks}}{N} \right\rceil$, as g increases the difference between w_c and $\frac{\text{problem_size}}{N}$ increases.

For example, considering a case where $\text{problem_size} = 100,000$, and $N = 8$, for the grain sizes in range of $[4167, 6249]$ would result in creating between $\left\lceil \frac{100,000}{6,249} \right\rceil = 17$ and $\left\lceil \frac{100,000}{4,167} \right\rceil = 24$ tasks. This amount of tasks created itself would result in $\lceil \frac{17}{8} \rceil = 3$ and $\lceil \frac{24}{8} \rceil = 3$ tasks. On the other hand, $w_c = g \times \left\lceil \frac{\text{num_tasks}}{N} \right\rceil = 3 \times g$, would have a value in range of $[3 \times 4167, 3 \times 6249] =$

[12501, 18747], where the average amount of work per core is $\frac{\text{problem_size}}{M} = 12500$. This means that for grain sizes closer to the end of the range, we are observing that a much bigger amount of work is assigned to the core with maximum amount of work, which would result in a higher execution time.

In the general case, if we denote $\left\lceil \frac{\text{num_tasks}}{N} \right\rceil$ as k , then:

$$\begin{aligned} k - 1 &< \frac{\text{num_tasks}}{N} \leq k \\ (k - 1) \times N &< \text{num_tasks} \leq k \times N \\ (k - 1) \times N &< \left\lceil \frac{\text{problem_size}}{g} \right\rceil \leq k \times N \\ (k - 1) \times N &< \frac{\text{problem_size}}{g} \leq k \times N \end{aligned} \tag{5.9}$$

If $k = 1$, then,

$$\begin{aligned} 0 &< \frac{\text{problem_size}}{g} \leq N \\ \frac{\text{problem_size}}{N} &\leq g \leq \text{problem_size}. \end{aligned} \tag{5.10}$$

Otherwise, when $k > 1$,

$$\frac{\text{problem_size}}{k \times N} \leq g < \frac{\text{problem_size}}{(k - 1) \times N}. \tag{5.11}$$

Since $\left\lceil \frac{\text{num_tasks}}{N} \right\rceil = k$, and $w_c = g \times \left\lceil \frac{\text{num_tasks}}{N} \right\rceil = k \times g$ if $\text{num_tasks \% } N \neq 1$, we can

conclude for $k > 1$:

$$\begin{aligned} k \times \frac{\text{problem_size}}{k \times N} &\leq w_c < k \times \frac{\text{problem_size}}{(k - 1) \times N} \\ \frac{\text{problem_size}}{N} &\leq w_c < \frac{k}{k - 1} \times \frac{\text{problem_size}}{N} \\ 0 \leq w_c - \frac{\text{problem_size}}{N} &< \frac{1}{k - 1} \times \frac{\text{problem_size}}{N} \end{aligned} \tag{5.12}$$

For the cases where $k > 1$ and $\text{num_tasks \% } N = 1$, there could be a change in w_c if $\text{problem_size \% } g \neq 0$. For these cases:

$$\begin{aligned}
\left\lceil \frac{\text{num_tasks}}{N} \right\rceil = k \quad & \& \quad \text{num_tasks} \% N = 1 \Rightarrow \\
\text{num_tasks} = (k-1) \times N + 1 \Rightarrow \\
(k-1) \times N < \frac{\text{problem_size}}{g} \leq (k-1) \times N + 1 \Rightarrow \\
\frac{\text{problem_size}}{(k-1) \times N + 1} \leq g < \frac{\text{problem_size}}{(k-1) \times N}
\end{aligned} \tag{5.13}$$

From Formula 5.3 we know,

$$w_c = \text{problem_size} - (k-1) \times (N-1) \times g. \tag{5.14}$$

Therefore,

$$\begin{aligned}
(k-1)(N-1) \frac{\text{problem_size}}{(k-1)N+1} \leq (k-1)(N-1)g < (k-1)(N-1) \frac{\text{problem_size}}{(k-1)N} \Rightarrow \\
\frac{\text{problem_size}}{N} < \text{problem_size} - (k-1)(N-1)g \leq k \frac{\text{problem_size}}{(k-1)N+1} \\
\frac{\text{problem_size}}{N} < w_c \leq k \times \frac{\text{problem_size}}{(k-1)N+1}
\end{aligned} \tag{5.15}$$

And for $k = 1$, where $\text{num_tasks} \leq N$,

$$\begin{aligned}
w_c &= g \\
\frac{\text{problem_size}}{N} &\leq g \leq \text{problem_size} \Rightarrow
\end{aligned}$$

$$0 \leq w_c - \frac{\text{problem_size}}{N} = g - \frac{\text{problem_size}}{N} \leq (N-1) \times \frac{\text{problem_size}}{N} \tag{5.16}$$

Defining $imbalance_ratio = \frac{w_c - \frac{\text{problem_size}}{N}}{\frac{\text{problem_size}}{N}}$, then,

$$\begin{aligned}
0 \leq imbalance_ratio &\leq N-1 && \text{for } k=1 \\
0 \leq imbalance_ratio &< \frac{1}{k-1} && \text{for } k>1 \text{ and } num_tasks \% N \neq 1 \\
0 \leq imbalance_ratio &< \frac{N-1}{N(k-1)+1} = \frac{1}{k-1 + \frac{k}{N-1}} && \text{otherwise}
\end{aligned} \tag{5.17}$$

Formula 5.17 shows that as number of created tasks increases, as long as number of tasks per core is the same, the imbalance factor decreases.

Figure 5.2 shows the imbalance ratio calculated for different grain sizes for $problem_size = 10000$, on 8 cores. Each of the regions between two dashed green lines correspond to a specific value for $k = \left\lceil \frac{num_tasks}{N} \right\rceil$. At each of the regions with $k > 1$, $\left\lceil \frac{num_tasks}{N} \right\rceil = k$, $imbalance_ratio$ starts from 0 and approaches $\frac{1}{k-1}$ ($\frac{1}{k-1 + \frac{k}{N-1}}$ for regions where $num_tasks \% N \neq 1$) at the end of the region. When $k = 1$, $imbalance_ratio$ increases linearly starting from 0 and reaching the maximum of $N-1$ when $g = problem_size$. As we move to larger grain sizes, $\left\lceil \frac{num_tasks}{N} \right\rceil$ decreases, therefore the upper-bound for $imbalance_ratio$ increases.

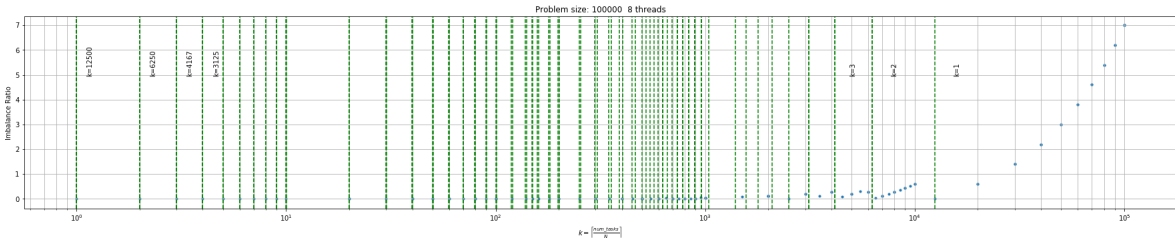


Figure 5.2. The imbalance ratio calculated for different grain sizes for $problem_size = 10000$, on 8 cores, where $k = \left\lceil \frac{num_tasks}{N} \right\rceil$.

Figure 5.1 represents the imbalance ratio, along with the ratio of the sequential execution time over execution time(speed-up) against grain size for $problem_size = 10000$, ran on 8 cores. As it can be observed, as the $imbalance_ratio$ increases, the speed-up decreases.

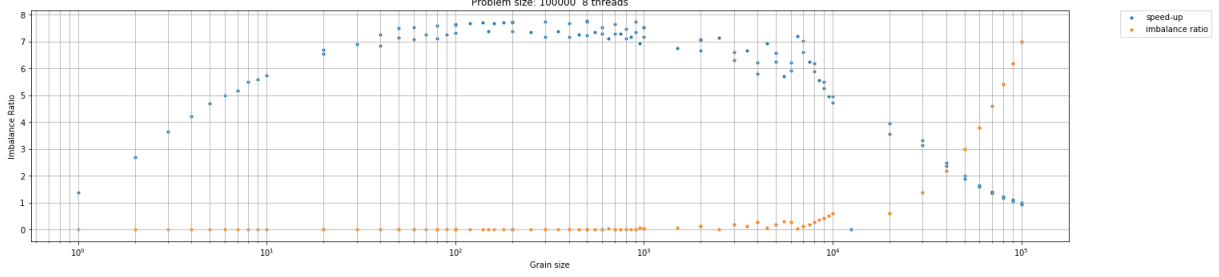


Figure 5.3. The imbalance ratio calculated for different grain sizes for $problem_size = 10000$, on 8 cores, where $k = \left\lceil \frac{num_tasks}{N} \right\rceil$.

To summarize, as the grain size increases the maximum imbalance in the loads assigned to the cores also increases, and some point on, this imbalance has a significant affect in the execution time. We define a threshold, λ_s ($0 < \lambda_s < 1$), where for $imbalance_ratios$ smaller than this threshold the imbalance effect is not significant. As we get close to this threshold, we are likely to reach the right hand side of the flat region of the bathtub curve of the execution time against grain size.

We are interested in finding the maximum grain size that would generate a reasonable imbalance ($imbalance_ratio \leq \lambda_s$), to make sure we should stay in the flat region of the bathtub curve of execution time against grain size, from load imbalance point of view.

Formula 5.16 states that for grain sizes greater than $problem_size$, $imbalance_ratio$ increases linearly with grain size from 0 to $N-1$. While for grain sizes smaller than $problem_size$, the maximum $imbalance_ratio$ depends on $k = \left\lceil \frac{num_tasks}{N} \right\rceil$. So, in order to assure $imbalance_ratio$ is smaller than or equal to a threshold (λ_s), first we search the grain sizes smaller than $\frac{problem_size}{N}$.

Since $0 < \lambda_s < 1$, and $k \geq 2$ in this region, there exists a k such that $\frac{1}{k-1} \leq \lambda_s$. If there exists a k_{min} (creating an imbalance ratio between 0 and $\frac{1}{k_{min}-1}$), where $\frac{1}{k_{min}-1} \leq \lambda_s$, $\forall k < k_{min}$ maximum value of $imbalance_ratio$ would be greater than λ_s . So in order to find the grain size that would create maximum $imbalance_ratio$ of λ_s :

$$\begin{aligned}
imbalance_ratio &\leq \lambda_s \Rightarrow \\
\frac{1}{k-1} &\leq \lambda_s \\
k &\geq 1 + \frac{1}{\lambda_s} \\
k_{min} &= \left\lceil 1 + \frac{1}{\lambda_s} \right\rceil + 1 \\
g &< \frac{problem_size}{(k_{min}-1) \times N} \\
g_{max} &= \frac{problem_size}{(k_{min}-1) \times N} - 1 = \frac{problem_size}{(1 + \left\lceil \frac{1}{\lambda_s} \right\rceil) \times N}
\end{aligned} \tag{5.18}$$

If $g < g_{max}$, we can assure that $imbalance_ratio$ never exceeds λ_s . Since we already found a match at grain sizes smaller than $\frac{problem_size}{N}$, checking the rest of grain sizes would not be necessary.

5.1.2. Identifying the range of grain size for maximum speedup

In the previous section, we proposed a method to identify the lower-bound and the upper-bound of the grain sizes for which we observe the minimum execution time. Integrating Formula 5.6 and Formula 5.18 suggests the following range for maximum speedup:

$$\sqrt{\frac{\alpha \times problem_size}{N \times \lambda_b}} \leq g \leq \frac{problem_size}{(1 + \left\lceil \frac{1}{\lambda_s} \right\rceil) \times N} \tag{5.19}$$

In this section we represent the identified range for a number of problem sizes, with different number of cores.

5.1.3. Step 1: Single-task, single-core runs

At the first step, we look at the cases where only one task has been created, the program is run on only one core. This problem could be assumed to be equivalent to running the same amount of work sequentially with an additional cost of creating just one task.

Listing 5.1: A simple hpx for_loop used to study the effect of grain size on the achieved parallelism.

```

1 //////////////////////////////////////////////////////////////////
2 void measure_function_futures_for_loop(std::uint64_t count, bool csv, std::uint64_t chunk_size, std::uint64_t
3     iter_length)
4 {
5 // start the clock
6 high_resolution_timer waltime;
7 hpx::parallel::for_loop(hpx::parallel::execution::par.with(
8     hpx::parallel::execution::dynamic_chunk_size(chunk_size)),
9     0, count, [&](std::uint64_t) { worker_timed(iter_length*1000); });
10 // stop the clock
11 const double duration = waltime.elapsed();
12 print_stats("for_loop", "par", "parallel_executor", count, duration, csv);
13 }
14
15 //////////////////////////////////////////////////////////////////
16 int hpx_main(variables_map& vm)
17 {
18 {
19     const int repetitions = vm["repetitions"].as<int>();
20     num_threads = hpx::get_num_worker_threads();
21     const std::uint64_t chunk_size = vm["chunk_size"].as<std::uint64_t>();
22     const std::uint64_t iter_length = vm["iter_length"].as<std::uint64_t>();
23     const std::uint64_t count = vm["num_iterations"].as<std::uint64_t>();
24     bool csv = vm.count("csv") != 0;
25     if (HPX_UNLIKELY(0 == count))
26         throw std::logic_error("error: count of 0 futures specified\n");
27     for (int i = 0; i < repetitions; i++)
28     {
29         measure_function_futures_for_loop(count, csv, chunk_size, iter_length);
30     }
31 }
32 return hpx::finalize();
33 }
34 //////////////////////////////////////////////////////////////////
35 inline void worker_timed(std::uint64_t delay_ns)
36 {
37     if (delay_ns == 0)
38         return;
39     std::uint64_t start = hpx::util::high_resolution_clock::now();
40     while (true)
41     {
42 // Check if we've reached the specified delay.
43     if ((hpx::util::high_resolution_clock::now() - start) >= delay_ns)
44         break;chunk
45     }
46 }
47 //////////////////////////////////////////////////////////////////
48 int main(int argc, char* argv[])
49 {
50 // Configure application-specific options.
51     options_description cmdline("usage: " HPX_APPLICATION_STRING " [options]");
52     cmdline.add_options()("num_iterations",
53     value<std::uint64_t>()>default_value(500000),
54     "number of iterations to invoke")
55     ("repetitions", value<int>()>default_value(1),
56     "number of repetitions of the full benchmark")
57     ("iter_length", value<std::uint64_t>()>default_value(1), "length of each iteration")
58     ("chunk_size", value<std::uint64_t>()>default_value(1), "chunk size");
59 // Initialize and run HPX.
60     return init(cmdline, argc, argv);
61 }
```

5.1.3.1. Expected Model

With this assumption we expect the execution time to be summation of the time it takes to perform the total amount of work(*problem_size*) and the overhead of creating one HPX task(α). Formula 5.20 shows the expected formula.

$$\text{execution time} = \alpha + \text{problem size} \quad (5.20)$$

5.1.3.2. Original Data

In order to check our proposed model for this simplified problem, we collected data from running the program, setting the *chunk_size* to 1, *num_iterations* to 1, and changing the *iter_length* from 1 to 10,000,000.

Chapter 6. Setup

6.0.1. Blazemark

Blazemark is a benchmark suite provided by Blaze to compare the performance of Blaze with other linear algebra libraries including Blitz++[26], Boost uBLAS[54], GMM++[55], Armadillo[56], MTL4[28], and Eigen3[57], alongside plain BLAS libraries like Atlas[58], Goto[59], and Intel MKL.[60]

```
Dense Vector/Dense Vector Addition:  
C-like implementation [MFlop/s]:  
    100      1115.44  
    10000000  206.317  
Classic operator overloading [MFlop/s]:  
    100      415.703  
    10000000  112.557  
Blaze [MFlop/s]:  
    100      2602.56  
    10000000  292.569  
Boost uBLAS [MFlop/s]:  
    100      1056.75  
    10000000  208.639  
Blitz++ [MFlop/s]:  
    100      1011.1  
    10000000  207.855  
GMM++ [MFlop/s]:  
    100      1115.42  
    10000000  207.699  
Armadillo [MFlop/s]:  
    100      1095.86  
    10000000  208.658  
MTL [MFlop/s]:  
    100      1018.47  
    10000000  209.065  
Eigen [MFlop/s]:  
    100      2173.48  
    10000000  209.899  
  
N=100, steps=55116257  
C-like      = 2.33322 (4.94123)  
Classic     = 6.26062 (13.2586)  
Blaze       = 1 (2.11777)  
Boost uBLAS = 2.4628 (5.21565)  
Blitz++     = 2.57398 (5.4511)  
GMM++      = 2.33325 (4.94129)  
Armadillo   = 2.3749 (5.0295)  
MTL         = 2.55537 (5.41168)  
Eigen        = 1.19742 (2.53585)  
N=10000000, steps=8  
C-like      = 1.41805 (0.387753)  
Classic     = 2.5993 (0.710753)  
Blaze       = 1 (0.27344)  
Boost uBLAS = 1.40227 (0.383437)  
Blitz++     = 1.40756 (0.384884)  
GMM++      = 1.40862 (0.385172)  
Armadillo   = 1.40215 (0.383403)  
MTL         = 1.39941 (0.382656)  
Eigen        = 1.39386 (0.381136)
```

Figure 6.1. An example of the results obtained from running *DVECDVECADD* benchmark through Blazemark

6.0.2. Configurations

Our experiments were run on Marvin nodes of Rostam cluster at Center for Computation and Technology(CCT) at Louisiana State University. Table 6.1 and Table 6.2 show some of the specifications of this node.

Category	Specification
CPU	2 x Intel(R) Xeon(R) CPU E5-2450 0 @ 2.10GHz
RAM	48 GB
Number of Cores	16
Hyperthreading	Off

Table 6.1. Specifications of the Marvin node from Rostam cluster at CCT.

Cache Level	Coherency Line Size	Number of Sets	Ways of Associativity	Size
1	64	512	8	32KB
2	64	512	8	256KB
3	64	512	20	20480KB

Table 6.2. Cache specifications of the Marvin node from Rostam cluster at CCT.

Library	Version
HPX	1.3.0
Blaze	3.5

Table 6.3. Specifications of the libraries used to run our experiments.

Chapter 7. Proposed Study

In Chapter 4 two different functions were proposed to model the relationship between grain size, number of cores, and performance (execution time or throughput). Using these function, for a specific matrix size, we were able to identify a reasonable value for grain size in order to be able to amortize the overhead of task creation with execution time on one hand, and utilize our resources properly on another hand. This value(range of values) helped us to select a reasonable value(range of values) for chunk size, for a fixed block size. In this section we will state the next steps that needs to be taken.

7.1. Studying the bathtub model

The bathtub model needs to be studied more to check for the missing factor. We suspect this factor to be relevant to load imbalance, where some cores end up to be idle while the others are still doing the work.

Also, we need to study the formula in terms of throughput instead of execution time, since small changes in execution time could make a large difference in throughput, so it would be easier to spot it.

7.2. Generalization for matrix size

As stated earlier, the suggested functions were fitted for each matrix size individually for simplification, but this assumption is not practical, since even though the characteristics of the function is the same for all matrix sizes, the fitted parameters vary from one matrix size to another.

The matrix size should somehow be integrated into the model itself. For this purpose, first we need to understand how changing the matrix size affects the relationship between grain size and performance. One immediate effect of increasing the matrix size is an increase in maximum possible grain size (right hand side of Figure 7.1), while the minimum possible

grain size is the same.

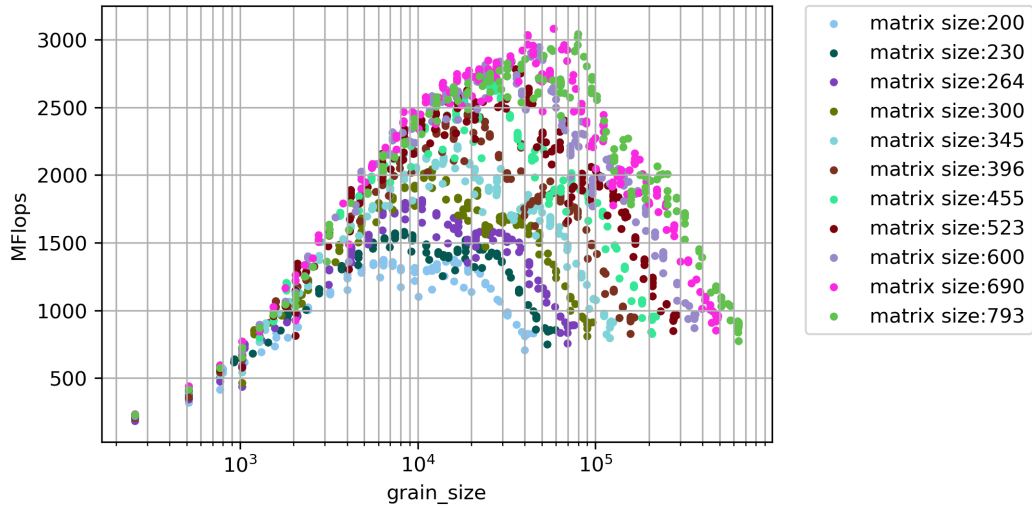


Figure 7.1. Throughput vs. grain size graph obtained from running *DMATDMATADD* benchmark on 4 cores.

Moreover, Figure 7.2 shows how the predicted grain size range changes for different matrix sizes for each number of cores.

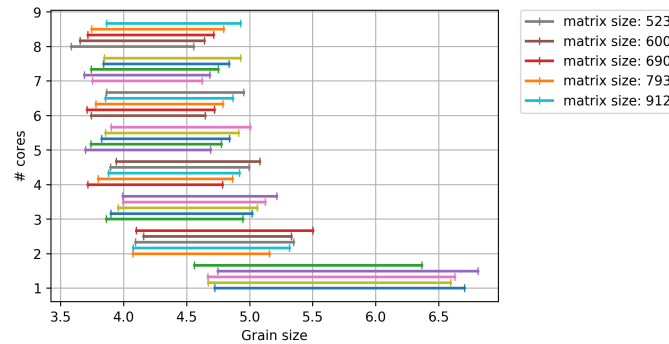


Figure 7.2. The range of grain size within 10% of the maximum performance of the fitted polynomial function for *DMATDMATADD* benchmark for different number of cores for matrix size 523×523 to 912×912 .

Also, larger matrix sizes should be added to the experiments to validate the current results.

7.3. Generalization for complex expressions

The whole study here was based on a simple matrix-matrix addition, and was validated on the same operation. We have been looking at the grain size as the key factor here, while the complexity of the operation has been included in the grain size. In the next step we need to add more benchmarks and study if the behavior is consistent for the other benchmarks. In addition to the operation, the number of matrices involved in the operation is also important, since it affects the threshold from which the matrices would not fit into the cache, which itself results in a performance degradation.

7.4. Generalization for different architectures

We are proposing here that once we have a generalized model, the parameters of the model are the factors that would change from one architecture to another. This, along with the cache specifications of the machines, should direct us toward fine tuning the performance for each architecture. One option could be running a set of benchmarks at build time to find these parameters for each machine. Once these parameters are found, we would be able to use our models to find the best grain size for the specific expression we are evaluating, considering the matrix size and the number of cores.

Chapter 8. Appendix

8.1. Bathtub model

In the previous section we studied the possibility of using a polynomial to capture the relationship between grain size, number of cores, and throughput for a fixed matrix size, with the purpose of finding a range of grain size that leads us to maximum performance. Although the polynomial function was helpful in directing us toward our objective, it does not have a physical implication.

This motivated us to change our view, and instead of looking just at the data and trying to find a function to fit the data, study the behavior of the data, and then find a function that would be likely to fit the data. That function would be a good fit mostly because that's how we expect the throughput to change with grain size, and not just how the data looks like.

As stated in Section 2.3, Grubel et.al.[1] has studied the task granularity for a specific problem(1D stencil). Looking at the graph representing the execution time based on the grain size, which resembles a bathtub, we are interested in formulating this graph based on our understanding of the effect of task granularity. Figure 8.1 shows the execution time in terms of grain size for *DMATDMATADD* benchmark for matrix size 690×690 on 4 cores.

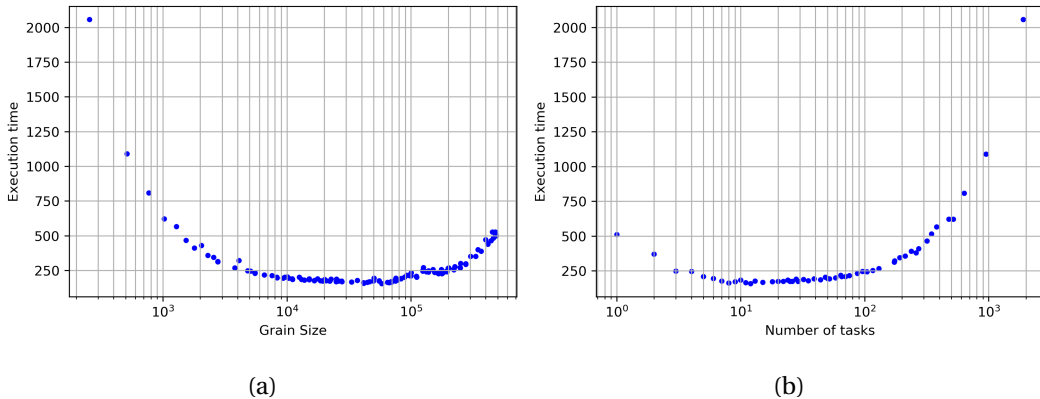


Figure 8.1. (a)The execution time vs. grain size graph, and (b) execution time vs. number fo tasks graph for *DMATDMATADD* benchmark for matrix size 690×690 ran on 4 cores.

For the sake of simplicity, we change the x axis from grain size to number of tasks. Each specific grain size would create a specific number of tasks(HPX threads), since the parameters we are interested in are directly associated with the number of tasks, we represent execution time based on number of tasks, as shown in Figure 8.1b.

Looking at the left hand side of the graph in Figure 8.1b, we can observe that for the first three points, the number of tasks created is smaller than the number of cores(which is 4 in this example). This means that in any of these cases there is at least one idle core, while the other cores are assigned a rather big chunk of work. The performance degradation we observe in that points is associated with starvation, meaning that we are not utilizing our computation resources to the full extent. In these three points, the number of cores actually doing the work is equal to the number of the tasks, since each core gets to execute at most one task.

To generalize the problem, assuming we are running our application on N cores, with a grain size equal to g , n_t tasks are being created, and M cores are actually doing the work. If $n_t < N$, M would be equal to n_t , otherwise $M = N$.

From overhead point of view though, if we represent the overhead of creating one task on a particular machine as α , the overhead of creating d tasks would be $n_t \alpha$, but this overhead is divided between the M cores actually doing the work.

To summarize, knowing the grain size, we are expecting the execution time in a many-task runtime system to be mainly affected by these factors, the overhead of creating one task(α), the number of cores that are actually doing the work(M), the sequential execution time(t_s), and finally the portion of the program that could actually be parallelized(γ).

If we try to integrate these information into a formula, we would expect the relation between execution time(t) and number of tasks(n_t) as follows:

$$t = \frac{\alpha n_t + t_s}{M} + \gamma \quad (8.1)$$

which could be decomposed into these two equations:

$$t = \begin{cases} \alpha + \frac{t_s}{n_t} + \gamma & \text{if } n_t < N \\ \frac{\alpha n_t + t_s}{N} + \gamma & \text{otherwise} \end{cases} \quad (8.2)$$

Now we use this function to find the best three parameters α , t_s , and γ so that the collected data would fit this model. For this purpose we used the *curve_fit* package from *SciPy* library in *python*.

In order to make Equation 8.1 differentiable, we used the softplus function(Equation 8.3) to represent M based on n_t .

$$f(x) = \ln(1 + e^x) \quad (8.3)$$

Which results in Equation 8.4:

$$t = \frac{\alpha n_t + t_s}{(N-1) - \ln(1 + (e^{N-1} - 1)e^{-n_t})} + \gamma \quad (8.4)$$

Here again we limited our problem to one specific matrix size at a time, and divided the whole data for each matrix size and number of cores into two parts, 60% for training and 40% for testing.

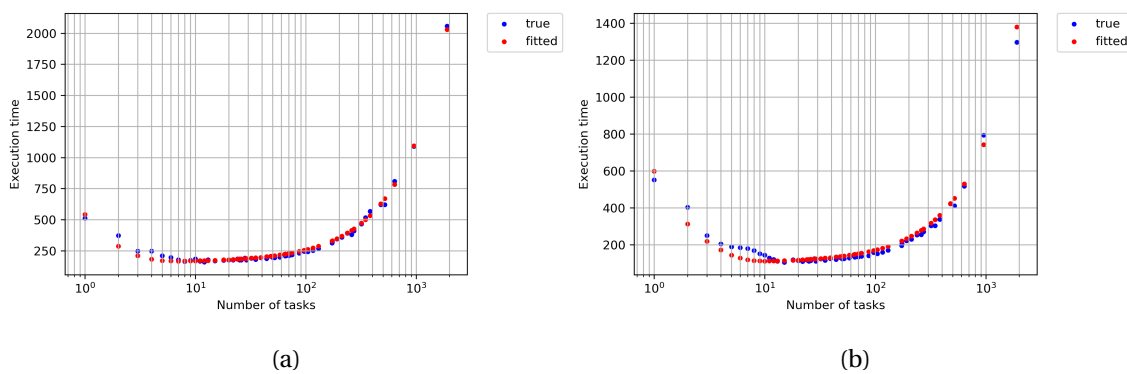


Figure 8.2. The prediction of execution time based on grain size using the bathtub model, for (a)4 cores and (b)8 cores for *DMATDMATADD* benchmark for matrix size 690×690 .

Figure 8.3 represents the relative error calculated for both training and test set, which is less than 5%.

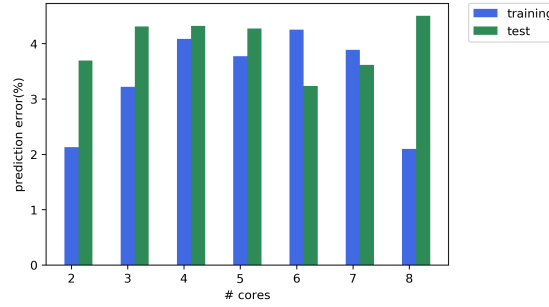


Figure 8.3. The error in fitting execution time with the bathtub formula for *DMATDMATADD* benchmark for matrix size 690×690 with different number of cores.

Assuming that this function fits our data in acceptable manner, next step would be to check how these three parameters change with the number of cores.

$$f(x) = \frac{m_0}{x} + \frac{m_1(x-1)}{x} + m_2(x-1) + m_3(x)(x-1) \quad (8.5)$$

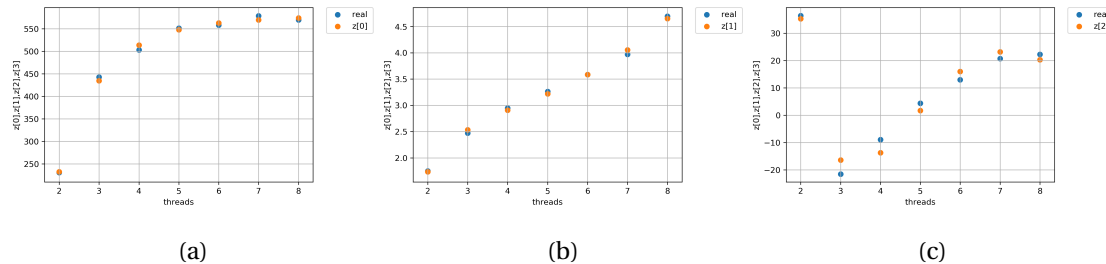


Figure 8.4. Fitting the three parameters (a) α , (b) t_s , and (c) γ for *DMATDMATADD* benchmark for matrix size 690×690 .

We can integrate Equation 8.1 and Equation 8.5 to predict the execution time for a given matrix size and number of cores. For each matrix size having found the parameters m_0 to m_3 , we can find α , t_s , and γ for the particular number of cores requested through Equation 8.5.

Then we can plug in the calculated values for α , t_s , and γ into Equation 8.1 to predict the execution time.

Figure 8.5 shows the prediction error on the test set for *DMATDMATADD* benchmark for matrix size 690×690 . The axis shows the different samples in the test set, and the label of each point represents the number of tasks created for that particular data point. As it could be seen, certain number of tasks result in higher prediction error, which needs to be studied.

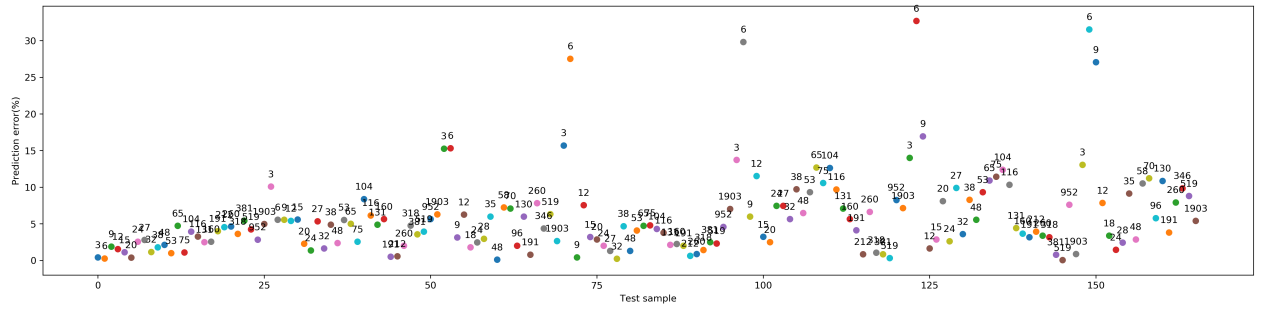


Figure 8.5. The error in fitting execution time with the bathtub formula for *DMATDMATADD* benchmark for matrix size 690×690 with different number of cores.

The problem with the current model is that with this formula we know that the minimum occurs at $n_t = N$, but that's not usually the case. This inspires us to check for a missing factor. This model still needs to be studied. The estimate that we have is for execution time, which is in our experiments very small. Changing the graphs from execution time to throughput can help us find the location of the maximum throughput easier compared to minimum of the execution time.

8.1.1. Studying the effect of matrix size

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