

Data Distribution Equivalence for Data Intensive Interconnection Networks

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Abstract of the Dissertation

Data Distribution Equivalence for Data Intensive Interconnection Networks

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This thesis considers two problems. One problem is finding closed-form solutions for equivalence computation of divisible workload origins from a single source in mesh and toroidal networks and the other problem is scheduling divisible workloads from multiple sources in mesh and toroidal networks of processors.

We propose a flow matrix closed-form equation to present the equivalence, which allows a characterization of the nature of minimal time solution and a simple method to determine when and how much load to distribute for processors. Also, we propose the use of a reduced Manhattan distance Voronoi diagram algorithm (RMDVDA) to minimize the overall processing time of these workloads by taking advantage of the processor equivalence technique. Three case studies with 10 sources of workloads are presented to illustrate the general approach for multiple sources of workloads. In the first phase, a Manhattan distance Voronoi diagram is used to obtain a network cluster division. In the second phase, we propose an efficient algorithm to obtain near-optimal load distribution among processors represented by equivalent processors. The algorithm minimizes the number of processors utilized. Experimental evaluation through simulations demonstrates that a task can be finished in the same suboptimal time and yet save over 30% of processor resources.

*Key Words : Divisible Load Theory, Voronoi Diagram, Multi-source, Network on Chip
(NOC) Interconnection Network, Mesh, Data Intensive Load*

To my Parents and my loved ones

Table of Contents

List of Figures	xii
List of Tables	xiii
Acknowledgements	xiv
1 Introduction	1
1.1 Divisible Load Theory	1
1.2 Processor Equivalence	2
1.3 Voronoi Diagram, Constrained Voronoi Diagram, Centroidal Voronoi Diagram	2
1.4 Networks on Chip	3
1.4.1 Topologies Used In This Thesis	3
1.4.2 Routing (Related Work)	4
1.4.3 Flow Control (Related Work)	4
1.5 Scientific Computation (Related Work)	5
1.6 Related Work on Single-source Assignment	6
1.7 Related Work on Multi-source Assignment	6
1.8 This Thesis	8
2 Problem Description	9
2.1 Problem Background	9
2.2 Definitions and Assumption	11
2.2.1 Notations	12

3 Mesh Network Virtual Cut-through Switching	14
3.1 Virtual Cut-through Switching Scenario	14
3.1.1 Data Injection on The Corner Processor	14
3.1.2 Data Injection On The Boundary Processor	32
3.1.3 Data Injection On The Inner Grid Processor	36
3.1.4 Sensitivity Analysis Virtual Cut-through Switching Scenario	42
3.1.5 Multi-source Uniform Data Fraction	48
3.1.6 Multi-source Non-uniform Data Injection	70
4 Mesh Network Modified Store and Forward Switching	74
4.1 Modified Store and Forward Switching Scenario	74
4.1.1 Data Injection on The Corner Processor	75
4.1.2 Data Injection on The Boundary Processor	87
4.1.3 Data Injection on The Inner Grid Processor	91
4.1.4 Sensitivity Analysis Modified Store and Forward Scenario	93
4.1.5 Multi-source Uniform Data Fraction	99
4.1.6 Multi-source Non-uniform Data Injection	109
5 Toroidal Rectangle Network	110
5.1 Toroidal Rectangle Network Notions	110
5.2 Virtual Cut-through Switching Scenario	112
5.2.1 Data Injection On The Grid Processor	112
5.2.2 Sensitivity Analysis of Toroidal Rectangle Network	115
5.2.3 Multi-source Uniform Data Fraction	118
5.2.4 Multi-source Non-uniform Data Injection	134
5.3 Modified Store and Forward Switching Scenario	134
5.3.1 Data Injection On The Grid Processor	134
5.3.2 Sensitivity Analysis of Toroidal Rectangle Network	136
5.3.3 Multi-source Uniform Data Injection	138
5.3.4 Multi-source Non-uniform Data Injection	138
5.4 Comparison Result	138
5.4.1 Comparison Result Between Mesh Network and Toroidal With Same Number of Processors	138
5.4.2 Comparison Result With Corner Processor and Inner Grid Processor .	139

5.4.3	Comparison Result Between Virtual Cut-through Switching Scenario and Modified Store and Forward Scenario	140
6	Hypercube Network and General Network	146
6.1	Hypercube Network Virtual Cut-through Switching Scenario	146
6.1.1	Data Injection On The Grid Processor	147
6.2	General Case Virtual Cut-through Switching Scenario	149
6.3	Hypercube Network Modified Store and Forward Switching Scenario	150
6.4	General Case Store and Forward Switching Scenario	151
7	Conclusion and Future Work	152
7.1	Conclusion	152
7.2	Future Work	153
	Bibliography	154

List of Figures

2.1	A m^*n mesh network($m = 5, n = 5$)	10
2.2	A toroidal rectangle network with grid unit cores	10
3.1	The 2^*2 mesh network and the root processor is P_0	15
3.2	The timing diagram for 2^*2 mesh network and the root processor is P_0	16
3.3	2^*2 mesh network. $\alpha_0, \alpha_1, \alpha_2, \alpha_3$ value curve	18
3.4	The 2^*3 mesh network and the data injection happens on corner processor P_0 .	19
3.5	The timing diagram for a 2^*3 mesh network and the data injection happens on processor P_0	20
3.6	2^*3 mesh network. $\alpha_0, \alpha_1, \alpha_3, \alpha_5$ value curve	22
3.7	2^*n ($n = 10$) mesh network and the workload happens on P_0	23
3.8	The timing diagram for 2^*10 mesh network and the data injection happens on P_0	25
3.9	3^*8 mesh network. The data injection position is P_0	29
3.10	The 3^*3 mesh network and the root processor is P_0	32
3.11	The timing diagram for 3^*3 mesh network and the data injection occurs on P_0 .	33
3.12	The data fraction simulation result of 3^*3 mesh network and the data injection happens on the boundary P_0	36
3.13	3^*3 mesh network. The data injection position is inner grid point P_0	37
3.14	The timing diagram for 3^*3 mesh network and the data injection is inner grid P_0 .	38
3.15	3^*3 mesh network. The data injection position is inner grid point P_0	40
3.16	Sensitivity analysis result of 2^*n mesh network (where $n = 1 \cdots 10$)	42
3.17	Speedup curves of 2^*n mesh network (where $n = 1 \cdots 10$)	43
3.18	Sensitivity analysis result of 3^*n mesh network and the injection position on boundary processor P_2 (where $n = 1 \cdots 8$)	44
3.19	Speedup curve of 3^*n mesh network (where $n = 1 \cdots 8$)	45

3.20	Sensitivity analysis result of $5 * n$ mesh network and the injection position on inner processor P_{12} (where $n = 1 \dots 5$)	46
3.21	Speedup curves of $5 * n$ mesh network Fig. 2.1 (where $n = 1 \dots 5$)	47
3.22	Example 1: Data injection consists of a connected subgraph of G	48
3.23	Example 2: Data injection consists of a connected subgraph of G	49
3.24	Connected subgraph Fig. 3.22 speedup vs σ	52
3.25	Connected subgraph Fig. 3.22's processor data fraction vs σ	53
3.26	Connected subgraph Fig. 3.23 speedup vs σ	54
3.27	Connected subgraph Fig. 3.23's processor data fraction vs σ	55
3.28	Reduced Voronoi diagram algorithm average saved processors' percentage . . .	61
3.29	10 reduced Voronoi cells	65
3.30	Reduced constrained Voronoi diagram for connected subgraphs and individual data injections on mesh network	67
3.31	Speedup curves of connected subgraphs and individual data injection on mesh network	68
3.32	Maximal speedup for $6 * 6$ mesh network and 4 data injections	69
3.33	Maximal speedup for $16 * 16$ mesh network and 1 data injections	69
4.1	The timing diagram for $2 * 2$ mesh network store and forward scenario	75
4.2	Data fraction curves of mesh network $2 * 2$ vs σ	77
4.3	The timing diagram for store and forward switching $2 * 3$ mesh network.	79
4.4	Data fraction curves of mesh network $2 * 3$ vs σ	81
4.5	The timing diagram for $3 * 3$ boundary data injection on P_0	87
4.6	The fraction curve for $3 * 3$ boundary data injection on P_0	89
4.7	The timing diagram for $3 * 3$ inner grid injection P_0	92
4.8	Sensitivity analysis result of $2 * n$ mesh network result ($n = 1 \dots 10$)	93
4.9	Speedup curve of $2 * n$ mesh network result ($n = 1 \dots 10$)	94
4.10	Sensitivity analysis result of $3 * n$ mesh network result ($n = 1 \dots 8$)	95
4.11	Speedup curve of $3 * n$ mesh network result ($n = 1 \dots 8$)	96
4.12	Sensitivity analysis result of data injection position on inner grid processor of $5 * 5$ mesh network	97
4.13	Speedup curve of $5 * n$ mesh network result ($n = 1 \dots 5$)	98
4.14	Connected subgraph Fig. 3.22 speedup vs σ	99
4.15	Connected subgraph Fig. 3.22's data injection processor workload fraction vs σ	100
4.16	Connected subgraph Fig. 3.23 speedup vs σ	101

4.17	Connected subgraph Fig. 3.23's data injection processor workload fraction vs σ	102
4.18	10 Manhattan distance Voronoi Cells	103
4.19	10 Manhattan distance Voronoi cells modified store and forward switching speedup curves	104
4.20	10 reduced Manhattan distance modified store and forward switching Voronoi cells	105
4.21	10 reduced Voronoi cells' store and forward switching speedup curves	106
4.22	10 reduced Manhattan distance modified store and forward switching Voronoi cells	107
4.23	10 reduced Voronoi cells' store and forward switching speedup curves	108
5.1	The rectangular toroidal network. Figure citing from [1]	110
5.2	The rectangular toroidal network. Figure citing from [1]	111
5.3	The $m \times n$ toroidal rectangle network and the data injection is $P_{4,2}$	112
5.4	Data fraction curve of toroidal rectangle network Fig. 5.3	114
5.5	Sensitivity analysis of $5 \times n$ torus network (where $n = 1 \dots 5$)	116
5.6	Data fraction curves of a $5 \times n$ torus network (where $n = 1 \dots 5$) vs σ	117
5.7	Toroidal mesh network subgraph data injection	119
5.8	Data fraction curves of connected subgraph Fig. 5.7	120
5.9	Toroidal mesh network subgraph data injection	121
5.10	Data fraction on different type processors for Fig. 5.9	122
5.11	Copy the original domain to 8 copies	123
5.12	How to calculate torus Voronoi Diagram	123
5.13	Initial Manhattan distance Voronoi Diagram	124
5.14	Torus Manhattan distance Voronoi diagram	125
5.15	Voronoi diagram casting to the torus	126
5.16	Toroidal Voronoi cells' speedup curves	127
5.17	Torus reduced Voronoi diagram	128
5.18	Torus reduced Voronoi diagram casting to a toroidal rectangle network	129
5.19	Torus reduced Voronoi diagram speedup curves	130
5.20	Torus reduced Voronoi diagram average saved processors	131
5.21	The data fraction curve of 6×6 torus network Fig. 5.3	135
5.22	Sensitivity analysis result of $5 \times n$ torus network ($n = 1 \dots 5$)	137
5.23	Speedup difference between inner grid injection and corner injection in $5 \times n$ network ($n = 1 \dots 5$)	139

5.24	Comparing result between virtual cut-through switching and store and forward switching in $2 * 2$ mesh network	141
5.25	Comparing result between virtual cut-through switching and store and forward switching in $2 * 3$ mesh network	142
5.26	Comparing result between virtual cut-through switching scenario and store and forward switching scenario. The data injection is boundary processor of a $3 * 3$ mesh network.	143
5.27	The comparing result between virtual cut-through switching processor with without virtual cut-through switching processor in $3*3$ mesh network injection on inner grid processor	144
5.28	Speedup difference between modified store and forward switching and virtual cut-through switching in $5 * n$ mesh network ($n = 1 \cdots 5$)	145
6.1	Hypercube in 0, 1, 2, 3 dimension. [2]	146
6.2	A hypercube network	147

List of Tables

3.1	The 10 Manhattan distance Voronoi Cells and speedup curves	57
3.2	The 10 Reduced Voronoi Cells and speedup curves	60
3.3	The 10 Accelerated Reduced Voronoi Cells and Speedup Curves	64
3.4	The 10 Constrained Voronoi Cells and Speedup Curves	73
5.1	D_i vs Number	113
5.2	D_i vs Number of processor on each D_i	115
5.3	Constrained Voronoi Diagram and RCVDA	132
5.4	Speedup curves Vs RCVDA speedup curves	133
5.5	D_i vs Number of processor on D_i	136

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Chapter 1

Introduction

A number of technological areas related to this thesis are discussed in this chapter. This includes modeling/evaluation techniques such as divisible load theory and Voronoi diagram. It includes concepts such as topology, routing, flow control and processor equivalence. Finally, application areas such as networks on chips and scientific computing are discussed.

1.1 Divisible Load Theory

This thesis makes heavy use of Divisible Load Theory concepts for modeling and solution.

During the past decades, divisible load scheduling and distribution techniques have been widely studied forming the Divisible Load Theory (DLT). The load is defined to be “divisible” for computation and communication if it can be partitioned arbitrarily among processors. Divisible Load Theory [3] [4] [5] is characterized by the fine granularity and large size of loads. There are also no precedence relations among the data elements. Such a load may be arbitrarily partitioned and distributed among processors and links in a system.

The divisible load model is a good approximation of tasks that require a large number of identical, low-granularity computations, thus has been proposed for a wide range of scientific and engineering data processing, such as image processing, matrix multiplication, fast-Fourier-transformation (FFT), video encoding/decoding, stereo matching, etc [6] [7] [8] [9] [10]. Various constraints on different interconnection topology [11] [12] [13] scheduling problems have been studied by the researchers.

1.2 Processor Equivalence

The concept of equivalent networks [14] was presented for complex networks such as multilevel tree networks.

The concept of processor equivalence was first proposed in [15]. A linear daisy chain of processors where processor load is divisible and shared among the processors is examined. It is shown that two or more processors can be collapsed into a single equivalent processor. This equivalence allows a characterization of the nature of the minimal time solution, a simple method to determine when to distribute load for linear daisy chain networks of processors without front end communication sub-processors and closed form expressions for the equivalent processing speed of infinitely large daisy chains of processors.

- By solving a series of layer and layer meshes and torus networks, we find the problem formulation and solution for single-source assignment for meshes and torus networks of arbitrary sites. In addition, the distribution and routing policy used in this thesis is represented in many other variations are can be considered.
- We propose a flow matrix closed-form equation to present the equivalence for single source assignment, which allows a characterization of the nature of minimal time solution and a simple method to determine when and how much load to distribute for processors.

This [15] [16] inspires us to adopt this model to tackle multi-source workload scheduling problems

1.3 Voronoi Diagram, Constrained Voronoi Diagram, Centroidal Voronoi Diagram

Voronoi diagrams are used in this thesis for multi-sources assignment.

Voronoi diagrams are induced by a set of points (called sites) : Subdivision of the plane where the faces correspond to the regions where one site is closest. Voronoi diagram is comprehensively utilized in network optimization application [17] [18] [19].

A constrained Voronoi diagram [20] [21] is that given a set of n vertices in the plane together with a set of noncrossing, straight-line edges, the constrained Delaunay triangulation (CDT) is the triangulation of the vertices with the following properties:

- (1) the prespecified edges are included in the triangulation
- (2) it is as close as possible to the Delaunay triangulation.

Applications also include motion planning in the presence of polygonal obstacles and constrained Euclidean minimum spanning trees, spanning trees subject to the restriction that some edges are prespecified.

A centroidal Voronoi tessellation [22] is a Voronoi tessellation whose generating points are the centroids (centers of mass) of the corresponding Voronoi regions. Some applications for example image compression, quadrature, finite difference methods, distribution of resources, cellular biology, statistics, and the territorial behavior of animals are investigated in Professor Qiang's paper [23] [22]. We discuss methods for computing these tessellations, provide some analyses concerning both the tessellations and the methods for their determination, and, finally, present the results of some numerical experiments. It can be viewed as an optimal partition corresponding to an optimal distribution of generators. A number of algorithms can be used to generate centroidal Voronoi tessellations, including Lloyd's algorithm [24] for K-means clustering.

1.4 Networks on Chip

This work has implications for network on chip design.

As the number of cores and Intellectual property (IP) increase on a single chip, conventional bus technology can no longer satisfy nowadays requirements of throughput and latency. In today's and future's electronic technology, Networks on Chip (NOC) have become important and even essential [25]. This paradigm has emerged as a communication backbone to enable a high degree of integration in multi-core System-on-chip (SOC) [26] [27] [28].

The distance between nodes, referred to as the hop count, is measured as the number of channels and nodes a message must traverse on average to reach its destination.

1.4.1 Topologies Used In This Thesis

The most basic structured interconnection network for on-chip networking: the mesh and toroidal network will be considered in detail in our discussion.

Mesh Network

Meshes are commonly used in network on chips.

A mesh is basically a chessboard (Manhattan streets) patterns of interconnection. Intellectual property (IP) are connected to their north, south, east and west neighbors through a network interface (NI). A network interface at the boundary of each core is used to package any data generated by the core. The topology is just like a chessboard, the lines are channels, and crossover points represent routers and processing elements. The most significant proposition of mesh topology is its modularization and total symmetry which reduce the hardware design complexity. However, congestion is easily formed in the center of mesh network. Simulation shows that total bandwidth is proportional to the square root of the number of processors, thus, the network has low average bandwidth and high latency when core number is large.

Torus

In short, torus is a 2D-mesh with successive boundary condition, thus, each router is connected to 5 ports. For example, if we describe a torus network with a 5-by-5 metrics, router $(0, 0)$ is connected to $(4, 0)$, $(0, 4)$, $(0, 1)$ and $(1, 0)$. According to the torus network, the congestion is less likely to happen in torus than mesh since torus has higher channel density.

1.4.2 Routing (Related Work)

Once a topology has been chosen, there can be many possible paths (sequences of nodes and channel) that a message could take through the network to reach its destination. A basic routing method is dimension-order routing [29], routing first in the x -dimension and then in the y -dimension. However dimension-order routing can produce significant load imbalance for some traffic patterns. We also have oblivious algorithms which include deterministic as a subset, and adaptive algorithms [30], adapt to the state of the network, using this state information in making routing decisions.

1.4.3 Flow Control (Related Work)

Flow control dictates which messages get access to particular network resources over time. The simplest flow-control mechanisms are buffer-less and, rather than temporarily

storing-blocked packets, they either drop or misroute these packets.

The next step up in complexity and efficiency is circuit switching, where only packet headers are buffered. There are two main switching strategies.

- One is circuit switching
 - In circuit switching, the header of a packet traverses the network ahead of any packet payload, reserving the appropriate resources along the path.
- Another is packet switching, containing store and forward switching, virtual cut-through switching and wormhole. Further, switching on a packet basis can either be done using *store and forward* [31] or *virtual cut-through* [32] switching approaches.
 - By breaking large, variable-length packets into smaller, fixed-sized flits, the amount of storage needed at any particular node can be greatly reduced.
 - Allocating resources in units of flits also facilitates creating multiple virtual channels per physical channel in the network, which can alleviate blocking and increase throughput.

These techniques confront with four problems, for example faults in network, congestion, deadlock [33] and livelock [34] issues.

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1.5 Scientific Computation (Related Work)

Besides networks on chips, another application area for this thesis is high performance large scale scientific computing.

Scientific computation applications include computational grids, a network of large number of loads and load sources with large number of transmission, processing and storage resources. Computational grids aim at exploiting synergies that result from cooperation by sharing and aggregating distributed computational capabilities and delivering them as service.

Another example of grid problems is the case of high energy and nuclear physics experiments. Here large amounts of data originate from distributed researchers who must work closely together. The analysis of data in this type of experiments requires an increased computational power and network based computing platforms such as Globus [35] and Condor

[36]. In such recently emerging platforms, scheduling and performance evaluation analysis for efficient use of distributed resources are important but challenging tasks. Effective scheduling of jobs in such a system is complicated by a number of factors including data locality, network topology and latency.

1.6 Related Work on Single-source Assignment

The mathematical theory of divisible loads is uniquely suited to serve as a basis for analytically modeling and solving grid scheduling problems in its ability to capture both computing and communication in a single model. The original work of Cheng and Robertazzi [37] [38] was proposed in 1988, after that there has been an increasing amount of study in Divisible Load Theory. In a single root processor assumption, the efficient load distribution strategy and protocol are developed by the majority of these studies.

The optimal solution is obtained by forcing the processors over a network to all stop processing simultaneously. Otherwise, the efficiency of the whole network system could be improved by transferring load if some processors were idle while other are still busy [39]. The various network topologies include linear daisy chains, tree and bus networks using a set of recursive equations which were presented in [40] [41] [42], respectively. Different load distribution policies for hypercubes [43] and mesh networks [44] are further studied.

Work has also been considered scheduling policy with multi-installment [45], multi-round algorithms [46], independent task scheduling [47], fixed communication charges [48], detailed parameterization and solution reporting time optimization [49] and combinatorial optimization [50]. Recently, though Divisible Load Theory (DLT) is fundamentally a deterministic theory, a study has been done to show some equivalence to Markov chain models [13].

1.7 Related Work on Multi-source Assignment

Most of the earlier studies in the literature so far have assumed that the processing load originates from a single root processor. In a practical scenario, this need not be always true. The problem of minimizing the processing time of extensive processing loads originated from various sources presents a great challenge that, if successfully met, could foster a range of new creative applications.

Inspired by this challenge, we sought to apply Divisible Load Theory to the problem of grid computing involving multiple sources connected to multiple sinks. So far research in this area includes [51] where tasks arrive according to a basic stochastic process to multiple nodes and [52] presents a first step technique for scheduling divisible loads from multiple sources to multiple sinks, with and without buffer capacity constraints. This paper, unlike the previous research papers, presents closed form solutions using Divisible Load Theory to mesh networks, toroidal or hypercube networks with multi-load originating (root) processors.

There is a limited amount of literature on divisible load modeling with multiple sources. A 2002 paper on multi-source load distribution combining Markovian queuing theory and divisible load scheduling theory is by Ko and Robertazzi [51]. In 2003 Wong, Yu, Veeravalli and Robertazzi [52] examined two source grid scheduling with memory capacity constraints. Marchal, Yang, Casanova and Robert [10] in 2005 studied the use of linear programming to maximize throughput for large grids with multiple loads/sources. In 2005, Lammie and Robertazzi [53] presented a numerical solution for a linear daisy chain network with load originating at both ends of the chain. Finally, Yu and Robertazzi examined mathematical programming solutions and flow structure in multi-source problems in 2006 [54].

Jia [55] propose a general method to address the multi-source scheduling problem. Yet, the algorithm has three issues that need more attention. First, his algorithm need to discard the part of previous computation result and do the global re-distribution. Then, his method doesn't give a quantitative model to represent the data fraction on each processor. Third, the algorithm doesn't propose a general equivalence computation power, which is very useful for user to estimate the makespan and help me to allocate the data source workload. In this paper, we utilize the Voronoi diagram to partition the NOCs cores.

Data flow analysis and optimization is considered for homogeneous rectangular mesh networks. Firstly, closed-form solutions are found for the equivalence computation [15] of divisible workload originating from a single data injection point in a homogeneous mesh network. We propose a flow matrix closed-form equation to present the equivalence, which allows a characterization of the nature of minimal time solution and a simple method to determine when and how much load to distribute to processors. We also propose a rigorous mathematical proof about the flow matrix optimal solution existence and that the solution is unique.

Secondly, load originating at multiple data injection points is considered. A reduced Manhattan distance Voronoi diagram algorithm (RMDVDA) is used to minimize the overall processing time of these workloads by taking advantage of a processor equivalence technique.

In the first phase, a Voronoi Manhattan distance diagram is used to obtain a network cluster division. In the second phase, an efficient algorithm to obtain near-optimal load distribution among processors represented by equivalent processors is proposed. The algorithm minimizes the number of processors utilized. Experimental evaluation through simulations demonstrates that a load can be processed in about the same suboptimal time and yet save about 35% - 40% of processor resources. Further, the lower bound of intuitive and heuristic algorithms are also investigated.

This work has important consequences for networks on chips where the demonstrated ability to reduce the number of processors needed for a computational problem has significant consequences for chip area and chip scalability.

1.8 This Thesis

Four chapters are organized as follows :

- A problem introduction, notation and definitions are shown in Chapter II.
- Chapter III considers mesh networks. We solve the processor equivalence problem with virtual cut-through scenario and modified store and forward distribution model. Sensitivity analysis is another topic. In considering the multi-source workload assignment problem, we propose a reduced Manhattan distance Voronoi diagram algorithm. We propose two objective functions. First, we finish the load in the same makespan, yet utilizing less processors. Another one is changing the cluster partitioning to increasing the bottleneck's capacity to increase the whole cluster's equivalence computation.
- Chapter IV investigates the toroidal network situation. The virtual cut-through switching and modified store and forward switching assumption are discussed. Sensitivity analysis is examined and the multi-source assignment in torus network environment is also discussed. Two objective function, finishing the load with less sources and increasing the total equivalence computation, of torus network are also implemented in this chapter.
- Chapter V mentioned some general case for example, hypercube network and general network. We give two equivalence computation flow matrix in virtual cut-through switching and store and forward switching.
- Chapter VI discuss the conclusion and future work.

Chapter 2

Problem Description

2.1 Problem Background

In large-scale data intensive problems with geographically distributed or compact resources, load is generated from multiple sources [56] for a class of problems. It is assumed that the problem representation can be arbitrarily divided amongst the processors. Thus the problem representation is said to be “divisible”. The processing of massive amounts of data on distributed and parallel networks is becoming more and more common. The problem of minimizing the processing time of extensive loads originating from a multiplicity of sources and being processed on a multiplicity of nodes presents a challenge.

In this chapter, the closed-form processor equivalence [15] [16] problem in the grid networks of mesh network and toroidal rectangle network is discussed. Also, the multi-source workload assignment is also taken into account.

In this thesis, we investigate two problems. One is the processor equivalence problem for single sources. The other one is scheduling divisible workloads from multiple sources in mesh networks Fig. 2.1, toroidal rectangle networks Fig. 2.2.

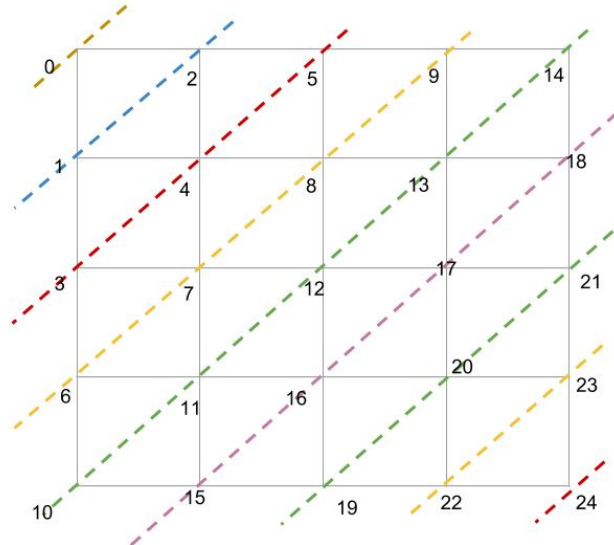


Figure 2.1: A $m \times n$ mesh network ($m = 5$, $n = 5$)

Torus Network

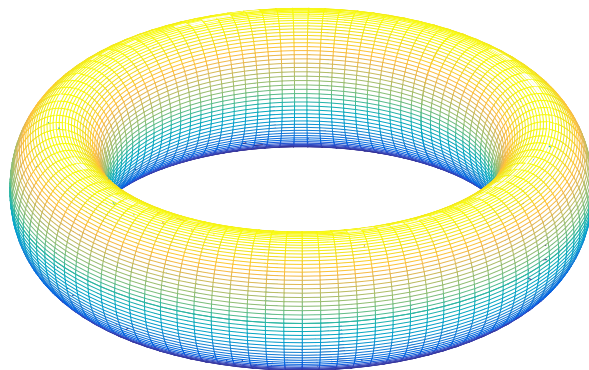


Figure 2.2: A toroidal rectangle network with grid unit cores

2.2 Definitions and Assumption

Definition 2.2.1. *Equivalence Computation*

Equivalence computation is a technique, which consists of combining a cluster of processors as one whole equivalent processor to process a unit 1 workload.

The following assumptions are used throughout the paper:

- Virtual cut-through [32] switching and store and forward switching is used to transmit the assigned workload between processors.
 - Under virtual cut-through switching, a node can relay the beginning bit of a message (packet) before the entire message is received.
 - Under store and forward switching, a message must be completely received by a node before it can be relayed to the next node along their transmission path.
- For simplicity, return communication is not considered.
- The communication delays are taken into consideration.
- The time costs of computation and communication are assumed to be linear function of the data size.
- The network environment is homogeneous, that is, all the processors have the same computation capacity. The link speeds between any two unit cores are identical.
- The number of outgoing ports in each processor is limited. In NOC (network on chip), the port number is fixed 4 or 5.
- The general graph's grid node's in-degree and out-degree is 5, .

The optimization objective functions are as follows :

- Single Path Communication : data transfer between two nodes follows a single path
- Homogeneous : a homogeneous (all link and processors speed are identical) is assumed.
- Equivalence computation : the problem's objective function is how to partition and schedule the workloads amongst the processors to obtain the minimum makespan (finish time).

- Multi-source assignment : how all processors can finish processing a unit 1 workload at the same time utilizing fewer processors.

To achieve the minimum time solution is obtained by forcing the processors over a network to stop processing simultaneously. Intuitively, this is because the solution could be improved by transfer load from some busy processors to idle ones [3] [57].

2.2.1 Notations

The following notations and definitions are utilized:

- P_i : The i th processor. $0 \leq i \leq m * n - 1$.
- L_i : The i th work load. $1 \leq i \leq k$.
- D_i : The minimum number of hops from the processor P_i to the data load injection site L .
- α_0 : The load fraction assigned to the root processor.
- α_i : The load fraction assigned to the i th processor.
- $\hat{\alpha}_i$: The load fraction assigned to each professor on the i th layer $i \in 0 \dots (k - 1)$.
- ω_i : The inverse computing speed on the i th processor.
- ω_{eq} : The inverse computing speed on an equivalent node collapsed from a cluster of processors.
- r : The rank of flow matrix.
- z_i : The inverse link speed on the i th link.
- T_{cp} : Computing intensity constant. The entire load is processed in time $\omega_i T_{cp}$ seconds on the i th processor.
- Sp_i : The speedup of i th Voronoi cell $i \in 0 \dots (k - 1)$.
- Sp_b : The speedup of the bottleneck cell.
- T_{cm} : Communication intensity constant. The entire load is transmitted in time $z_i T_{cm}$ seconds over the i th link.
- \hat{T}_f : The finish time of the whole processor network. Here \hat{T}_f is equal to $\omega_{eq} T_{cp}$.

- T_b : The finish time for the bottleneck cell.
- T_f : The finish time for the entire divisible load solved on the root processor. Here T_f is equal to $1 \times \omega_0 T_{cp}$, that is $\omega_0 T_{cp}$.
- α_{j,l_j} : The data fraction of each processor on the l_j layer of j th Voronoi cell.
- $T_{f,i}$: The finish time for the i th processor, $i \in 0 \cdots (m * n - 1)$.
- S_j : The processor's number of j th cell.
- $\sigma = \frac{zT_{cm}}{\omega T_{cp}}$: The ratio between the communication speed to the computation speed, $0 < \sigma < 1$ [3] [58].
- $\hat{T}_{j,f}$: The finish time of j th cell.
- In multi-source situation, $\sum_{i=1}^k L_i = 1$
- $\sum_{i=0}^{m*n-1} \alpha_i = 1$
- $Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$

Chapter 3

Mesh Network Virtual Cut-through Switching

3.1 Virtual Cut-through Switching Scenario

In the virtual cut-through environment, a node can begin relaying the first part of a message (packet) along a transmission path as soon as it starts to arrive at the node , that is, it doesn't have to wait to receive the entire message before it can begin forwarding the message.

First we consider about the $2 * 2$ mesh network, which can be generalized to a $2 * n$ mesh network. After, we analyze a more general case $m * n$ mesh network and obtain a general closed-form matrix presentation. Finally, we give a key methodology to address this type of question. In addition, different single data injection position, such as the corner, boundary and inner grid are also discussed.

3.1.1 Data Injection on The Corner Processor

2*2 Mesh Network

The load L is assigned on the corner processor P_0 Fig. 3.1. The whole task is tackled by four processors P_0, P_1, P_2, P_3 together.

The processor P_0, P_1 and P_2 start to process its respective fraction at the same time. This includes P_1 and P_2 as they are relayed load in virtual cut-through mode at $t = 0$. Because we assume a homogeneous network (in processing speed and communication speed), $\alpha_1 = \alpha_2$ and P_1 and P_2 stop processing at the same time. The processor P_3 starts to work when the α_1 and α_2 complete transmission. That is, the link $0 - 1$ and $0 - 2$ are occupied

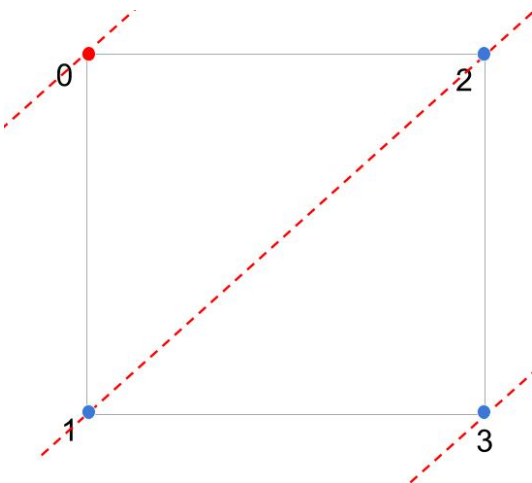


Figure 3.1: The 2×2 mesh network and the root processor is P_0

transmitting load to processor 1 and 2, respectively and only transmission to 3 when that is finished.

According to the divisible load theory [4], we obtain the timing diagram Fig. 3.2.

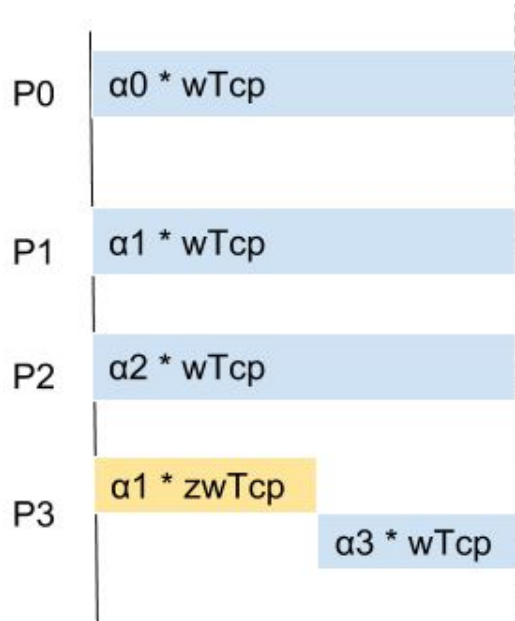


Figure 3.2: The timing diagram for 2×2 mesh network and the root processor is P_0

Here in the Gantt-like timing diagram communication appears above each axis and computations appears below the each axis. Let's assume that all processors stop computing at the same time in order to minimize the makespan [57].

Based on the timing diagram, we obtain a group of linear equations to find the fraction workload assigned to each processor α_i :

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ \alpha_0 + \alpha_1 + \alpha_2 + \alpha_3 = 1 \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1, \alpha_2, \alpha_3 < 1 \end{array} \right. \quad \begin{array}{l} (3.1) \\ (3.2) \\ (3.3) \\ (3.4) \\ (3.5) \\ (3.6) \\ (3.7) \\ (3.8) \\ (3.9) \end{array}$$

The group of equations are represented by the matrix form:

$$\begin{bmatrix} 1 & 2 & 1 \\ 1 & -1 & 0 \\ 0 & \sigma - 1 & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix} \quad (3.10)$$

The matrix is represented as $A \times \alpha = b$. A is named as the ***flow matrix***. Here because of symmetry $\alpha_1 = \alpha_2$, so α_2 is not listed in the matrix equations.

Finally, the explicit solution is:

$$\sigma = \frac{z T_{cm}}{\omega T_{cp}} \quad (3.11)$$

$$\alpha_0 = \frac{1}{4 - \sigma} \quad (3.12)$$

$$\alpha_1 = \frac{1}{4 - \sigma} \quad (3.13)$$

$$\alpha_3 = \frac{1 - \sigma}{4 - \sigma} \quad (3.14)$$

The simulation result is illustrated:

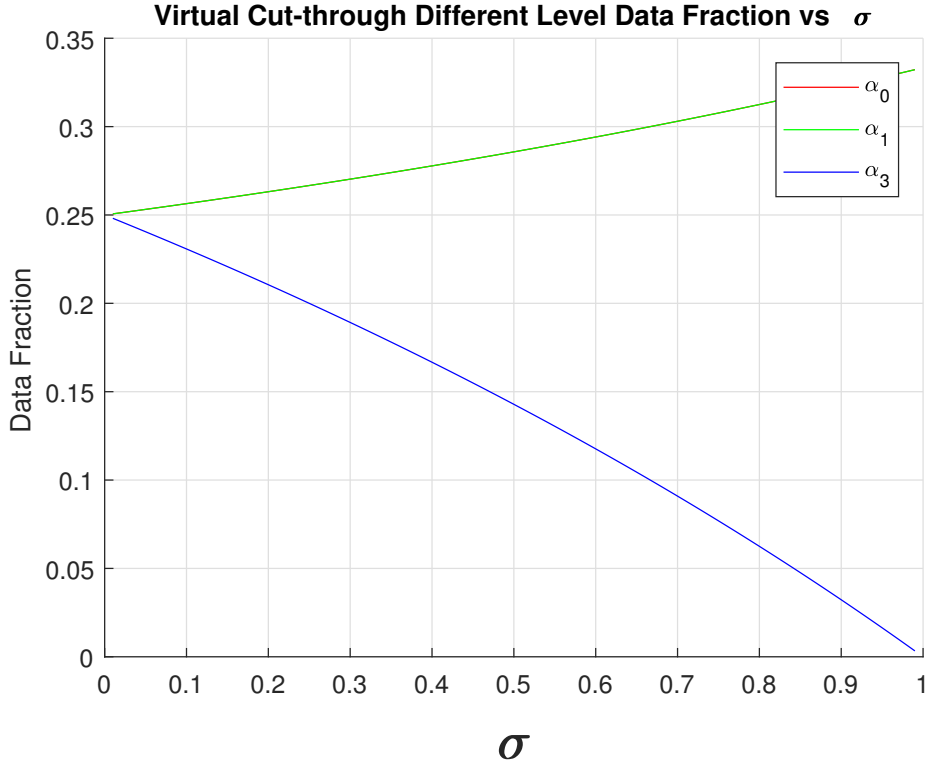


Figure 3.3: 2*2 mesh network. α_0 , α_1 , α_2 , α_3 value curve

In Fig. 3.3, the three processors P_0 , P_1 , P_2 three processors have the same data fraction workload, so the curve of α_0 and α_1 coincide.

The figure says that as σ grows, the value α_3 drops. In other words, as the communication capacity decreases, there is less data workload assigned to P_3 . Further, it means it will be economical to keep the load local on P_0 P_1 P_2 and not distribute it, to other processors.

The equivalence inverse speed of a single processor is w_{eq} , that can replace the original network as

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = 4 - \sigma$$

2*3 Mesh Network

In Fig. 3.4 mesh network, L originates on processor P_0 . There are 6 processors in this network.

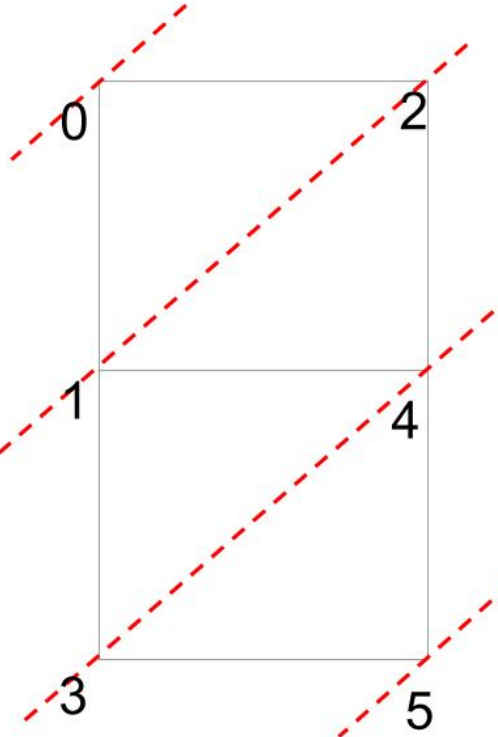


Figure 3.4: The 2*3 mesh network and the data injection happens on corner processor P_0

Here P_0 , P_1 and P_2 start processing at the same time. Processor P_3 and P_4 start to work when they receive the data from processor P_1 , P_2 , after the $0 - 1$ and $0 - 2$ link become available to transfer load to P_3 and P_4 .

The last processor P_5 doesn't start to execute until the work load fraction α_0 , α_1 , α_2 , α_3 , α_4 are transmitted completed. According to the divisible load theory [4], we obtain the timing diagram Fig. 3.5.

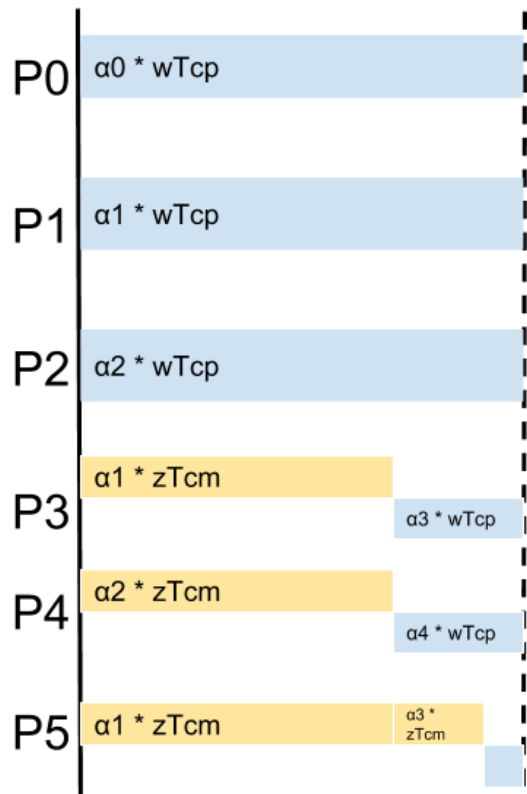


Figure 3.5: The timing diagram for a 2×3 mesh network and the data injection happens on processor P_0

The timing diagram equations as follows:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_3) z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ \alpha_0 + \alpha_1 + \alpha_2 + \alpha_3 + \alpha_4 + \alpha_5 = 1 \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1, \alpha_2, \alpha_3, \alpha_4, \alpha_5 < 1 \end{array} \right. \quad \begin{array}{l} (3.15) \\ (3.16) \\ (3.17) \\ (3.18) \\ (3.19) \\ (3.20) \\ (3.21) \\ (3.22) \\ (3.23) \\ (3.24) \\ (3.25) \end{array}$$

The flow matrix closed-form formula is:

$$\begin{bmatrix} 1 & 2 & 2 & 1 \\ 1 & -1 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_5 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (3.26)$$

The explicit solution is:

$$\sigma = \frac{z T_{cm}}{\omega T_{cp}} \quad (3.27)$$

$$\alpha_0 = \frac{1}{\sigma^2 - 4 \times \sigma + 6} \quad (3.28)$$

$$\alpha_1 = \frac{1}{\sigma^2 - 4 \times \sigma + 6} \quad (3.29)$$

$$\alpha_3 = \frac{1 - \sigma}{\sigma^2 - 4 \times \sigma + 6} \quad (3.30)$$

$$\alpha_5 = \frac{\sigma^2 - 2 \times \sigma + 1}{\sigma^2 - 4 \times \sigma + 6} \quad (3.31)$$

The α calculation result are shown in Fig. 3.6. P_0 , P_1 and P_2 have the same fraction so

the curve of α_0 and α_1 coincide. Because of symmetry, $\alpha_1 = \alpha_2$ and $\alpha_3 = \alpha_4$. If the communication time is zero, each of the six processors receive $\frac{1}{6}$ of the load which is in the figure. If the communication delay equals computation time $\frac{1}{3}$ of the load goes to each of P_0 , P_1 and P_2 . It is uneconomical to distribute load to P_3 , P_4 and P_5 .

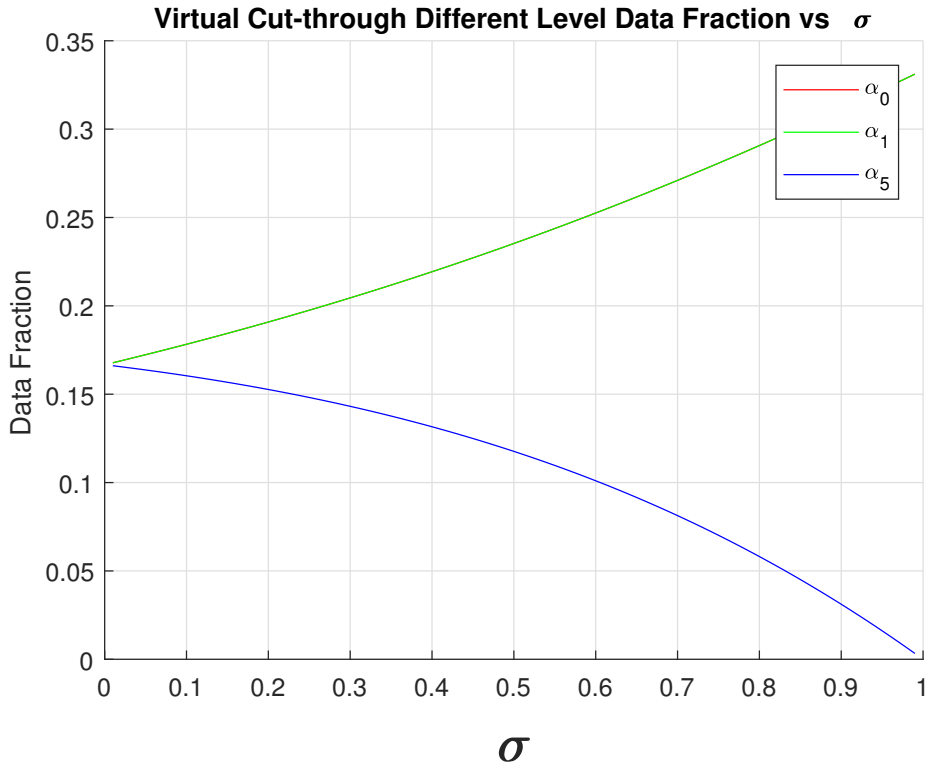


Figure 3.6: 2×3 mesh network. α_0 , α_1 , α_3 , α_5 value curve.

One also has for the equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

For speedup:

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

2*N Mesh Network

The $2 * n$ Fig. 3.7 homogeneous mesh network processes load L and L originates P_0 .

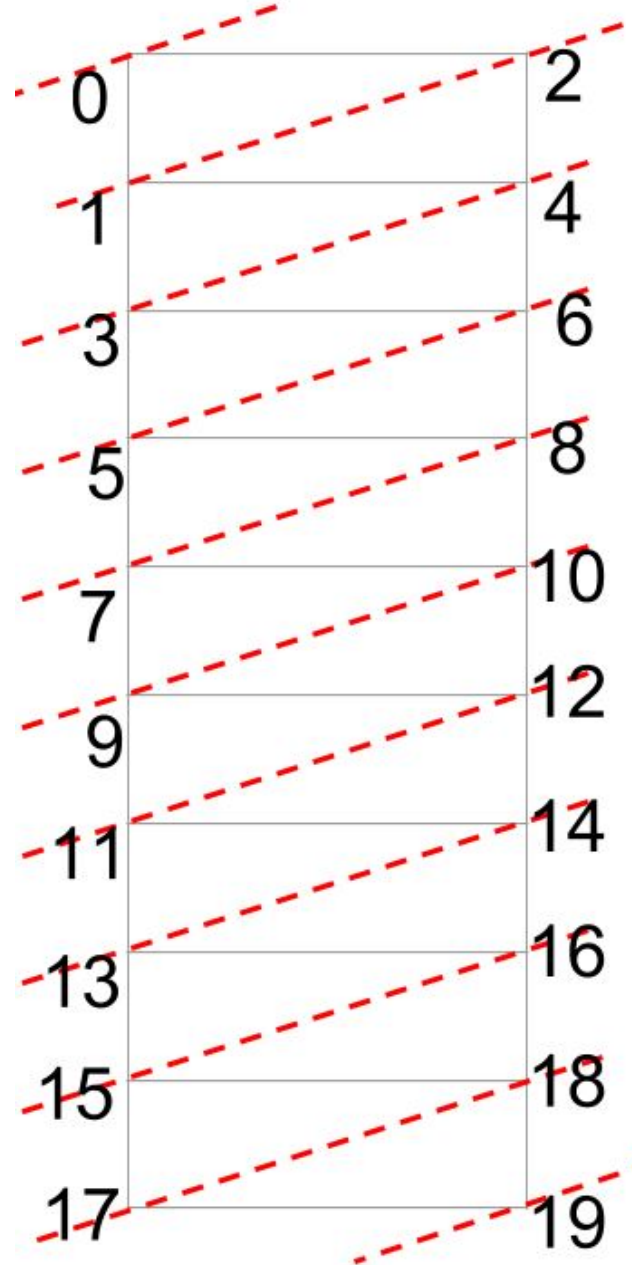


Figure 3.7: $2 * n$ ($n = 10$) mesh network and the workload happens on P_0

Load a distribution from P_0 to P_1 and P_2 via virtual cut-through. After P_1 and P_2 finish receiving load from link $0 - 1$ and $0 - 2$, they will be used to forward load to P_3 and P_4 and so on.

Similarly to the analysis of Fig. 3.2 and Fig. 3.5, the timing diagram for Fig. 3.7 is shown in Fig. 3.8

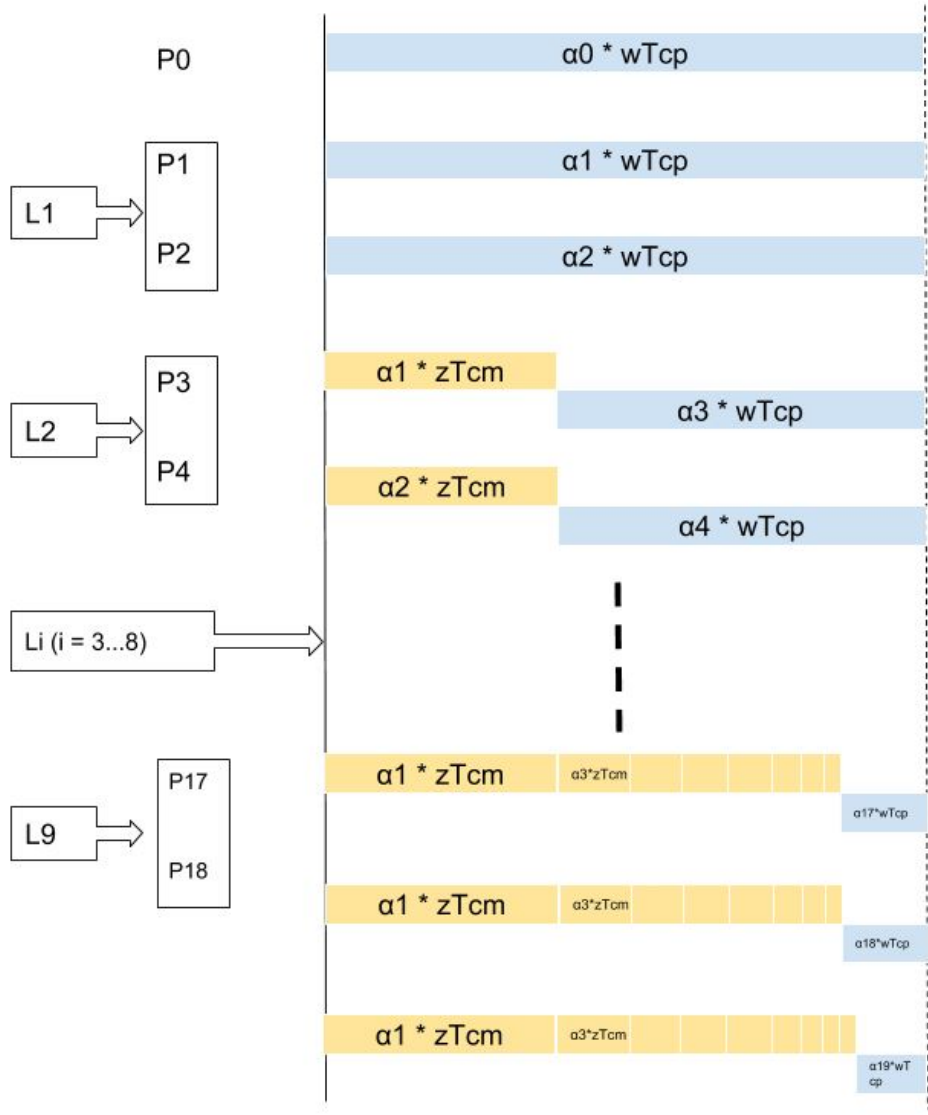


Figure 3.8: The timing diagram for 2*10 mesh network and the data injection happens on P_0

The equations are presented as:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_3) z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ \vdots \\ (\alpha_1 + \alpha_3 + \cdots + \alpha_{2 \times n-1}) z T_{cm} + \alpha_{2 \times n-1} \omega T_{cp} = T_{f,m} \\ \alpha_0 + \cdots + \alpha_{2 \times n-1} = 1 \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1 \alpha_2 \cdots \alpha_{2 \times n-1} < 1 \end{array} \right. \quad \begin{array}{l} (3.32) \\ (3.33) \\ (3.34) \\ (3.35) \\ (3.36) \\ (3.37) \\ (3.38) \\ (3.39) \\ (3.40) \\ (3.41) \\ (3.42) \\ (3.43) \\ (3.44) \end{array}$$

The flow matrix closed-form is shown:

$$\begin{bmatrix} 1 & 2 & 2 & \cdots & 2 & 2 & 1 \\ 1 & -1 & 0 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_5 \\ \vdots \\ \alpha_{2 \times n-3} \\ \alpha_{2 \times n-1} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ \vdots \\ 0 \\ 0 \end{bmatrix} \quad (3.45)$$

According to the **Cramer's rule**, the explicit solution for the group of equations is:

$$\left\{ \alpha_i = \left| \frac{\det A_i^*}{\det A} \right| \right. \quad (3.46)$$

where A_i^* is the matrix formed by replacing the i -th column of A by the column vector b .

Specifically,

$$A_0^* = \begin{bmatrix} 1 & 2 & 2 & \cdots & 2 & 2 & 1 \\ 0 & -1 & 0 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \ddots \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{bmatrix} \quad (3.47)$$

$$\alpha_0 = \left| \frac{\det A_0^*}{\det A} \right|$$

$$\det A_0^* = -1$$

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

Finally, the speedup is:

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = |-\det A|$$

Further, we prove the matrix $\det A \neq 0$.

$$C = \begin{bmatrix} -1 & 0 & \cdots & 0 & 0 & 0 \\ \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots \\ \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{bmatrix} \quad (3.48)$$

C is a lower triangular matrix and the diagonal elements are not 0. So C is non-degenerate, that is, the matrix is column linear independence.

After a series of column reduction and row reduction actions, we get

$$A = \left[\begin{array}{ccccccc} 1 & 2 & 2 & \cdots & 2 & 2 & 1 \\ 1 & -1 & 0 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{array} \right] \xrightarrow{\substack{\text{Column} \\ \text{Reduction}}} \left[\begin{array}{ccccccc} 1 & 0 & 0 & \cdots & 0 & 0 & 0 \\ 1 & -3 & -2 & \cdots & -2 & -2 & -1 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{array} \right]$$

$$\xrightarrow{\substack{\text{Row} \\ \text{Reduction}}} \left[\begin{array}{ccccccc} 1 & 0 & 0 & \cdots & 0 & 0 & 0 \\ 0 & -3 & -2 & \cdots & -2 & -2 & -1 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{array} \right]$$

Considering the matrix \hat{C}

$$\hat{C} = \left[\begin{array}{ccccccc} -3 & -2 & \cdots & -2 & -2 & -1 \\ \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots \\ \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{array} \right] \quad (3.49)$$

, which is still column linear independence. Considering $0 < \sigma < 1$, the flow matrix is full rank. So $\det A \neq 0$.

After three user cases' investigation, we find a crucial methodology:

$$\forall D_i = D_j, \quad \text{then} \quad \alpha_i = \alpha_j, \quad 0 \leq i, j \leq m * n - 1$$

$m*n$ Mesh Network

Considering a general $m * n$ mesh network, such as Fig. 3.9 Fig. 2.1.

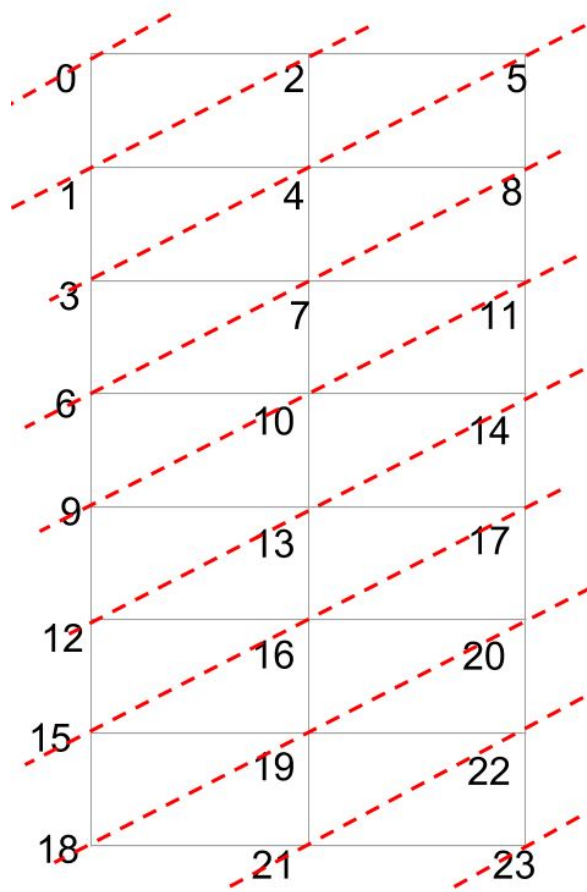


Figure 3.9: $3*8$ mesh network. The data injection position is P_0

Utilizing the previous methodology, we obtain the closed-form flow matrix equations

for Fig. 3.9:

$$\begin{bmatrix} 1 & 2 & 3 & 3 & 3 & 3 & 3 & 3 & 2 & 1 \\ 1 & -1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & 1 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & 1 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & \sigma & 1 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & \sigma & \sigma & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_6 \\ \alpha_9 \\ \alpha_{12} \\ \alpha_{15} \\ \alpha_{18} \\ \alpha_{21} \\ \alpha_{23} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ \vdots \\ 0 \end{bmatrix} \quad (3.50)$$

Also, the flow matrix equations for Fig. 2.1:

$$\begin{bmatrix} 1 & 2 & 3 & 4 & 5 & 4 & 3 & 2 & 1 \\ 1 & -1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & 1 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & 1 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & \sigma & 1 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & \sigma & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_6 \\ \alpha_{10} \\ \alpha_{15} \\ \alpha_{19} \\ \alpha_{22} \\ \alpha_{24} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ \vdots \\ 0 \end{bmatrix} \quad (3.51)$$

We use the similar method to prove $\det A \neq 0$. The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

so the speedup is:

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = |-\det A|$$

The first row in flow matrix describe the number of cores on each D_i . For example,

there is 1 core with 0 hop distance (D_0) with load site L . There are 2 cores with 1 hop distance (D_1) with load site L . There are 3 cores with 2 hops distance (D_2) with load site L , and so on.

The number of rows means the number of different type processor data fraction.

3.1.2 Data Injection On The Boundary Processor

After the corner scenario, we extend the methodology to boundary processor load origination.

A single data injection root-node on the boundary processor, as shown in Fig. 3.10.

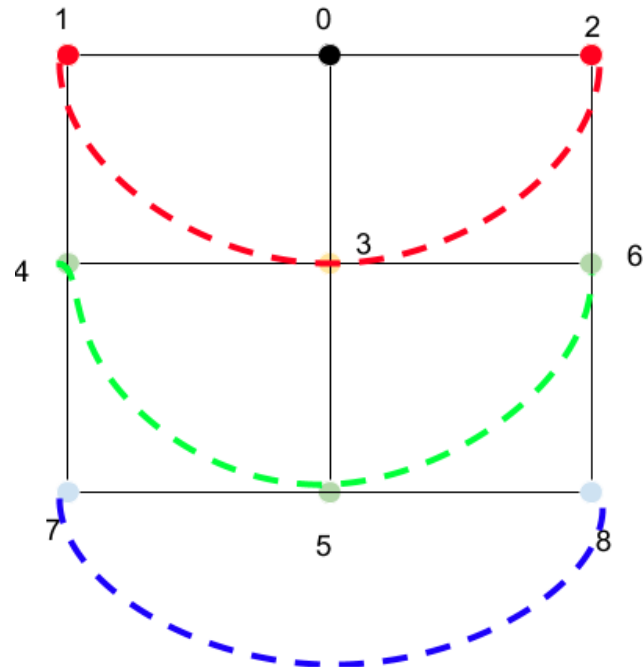


Figure 3.10: The 3×3 mesh network and the root processor is P_0

The timing diagram is Fig. 3.11:

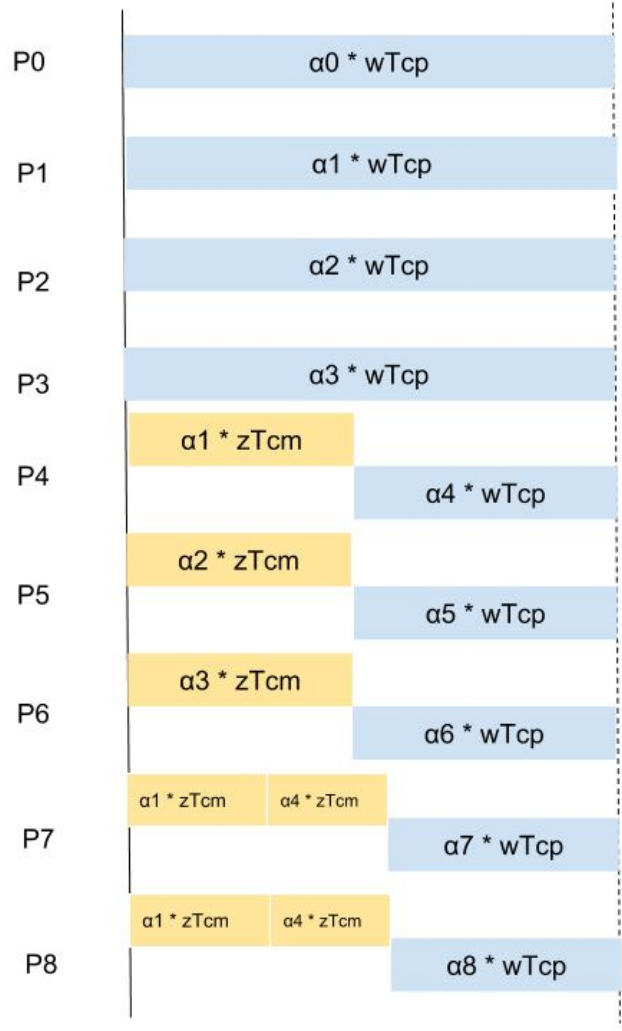


Figure 3.11: The timing diagram for 3×3 mesh network and the data injection occurs on P_0

P_0 , P_1 , P_2 and P_3 commence processing at $t = 0$ axis timing diagram. P_1 , P_2 and P_3 receive load in virtual cut-through mode from P_0 . Once that is finished, link $0 - 1$, $0 - 2$ and $0 - 3$ is used to forward load to nodes P_4 , P_5 and P_6 in virtual cut-through mode. Once that is finished, the link that have been so far are used to forward load to P_7 and P_8 in virtual cut-through mode.

The equations are:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_3 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ \alpha_3 z T_{cm} + \alpha_6 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_4) z T_{cm} + \alpha_7 \omega T_{cp} = T_{f,m} \\ (\alpha_2 + \alpha_5) z T_{cm} + \alpha_8 \omega T_{cp} = T_{f,m} \\ \alpha_0 + \cdots + \alpha_8 = 1 \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1 \alpha_2 \cdots \alpha_8 < 1 \end{array} \right. \begin{array}{l} (3.52) \\ (3.53) \\ (3.54) \\ (3.55) \\ (3.56) \\ (3.57) \\ (3.58) \\ (3.59) \\ (3.60) \\ (3.61) \\ (3.62) \\ (3.63) \\ (3.64) \\ (3.65) \\ (3.66) \end{array}$$

And the flow matrix form is :

$$\begin{bmatrix} 1 & 3 & 3 & 2 \\ 1 & -1 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_4 \\ \alpha_7 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (3.67)$$

The explicit solution is:

$$\left\{ \begin{array}{l} \alpha_0 = \frac{1}{9 - 7 \times \sigma + 2 \times \sigma^2} \\ \alpha_1 = \frac{1}{9 - 7 \times \sigma + 2 \times \sigma^2} \end{array} \right. \quad (3.68)$$

$$\left\{ \begin{array}{l} \alpha_4 = \frac{1 - \sigma}{9 - 7 \times \sigma + 2 \times \sigma^2} \\ \alpha_7 = \frac{(1 - \sigma)^2}{9 - 7 \times \sigma + 2 \times \sigma^2} \end{array} \right. \quad (3.70)$$

$$\left\{ \begin{array}{l} \alpha_0 = \frac{1}{9 - 7 \times \sigma + 2 \times \sigma^2} \\ \alpha_1 = \frac{1}{9 - 7 \times \sigma + 2 \times \sigma^2} \end{array} \right. \quad (3.71)$$

$$\left\{ \begin{array}{l} \alpha_4 = \frac{1 - \sigma}{9 - 7 \times \sigma + 2 \times \sigma^2} \\ \alpha_7 = \frac{(1 - \sigma)^2}{9 - 7 \times \sigma + 2 \times \sigma^2} \end{array} \right. \quad (3.72)$$

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

The simulation result is shown: The curve α_0 is identified to the curve for α_1 , α_2 and

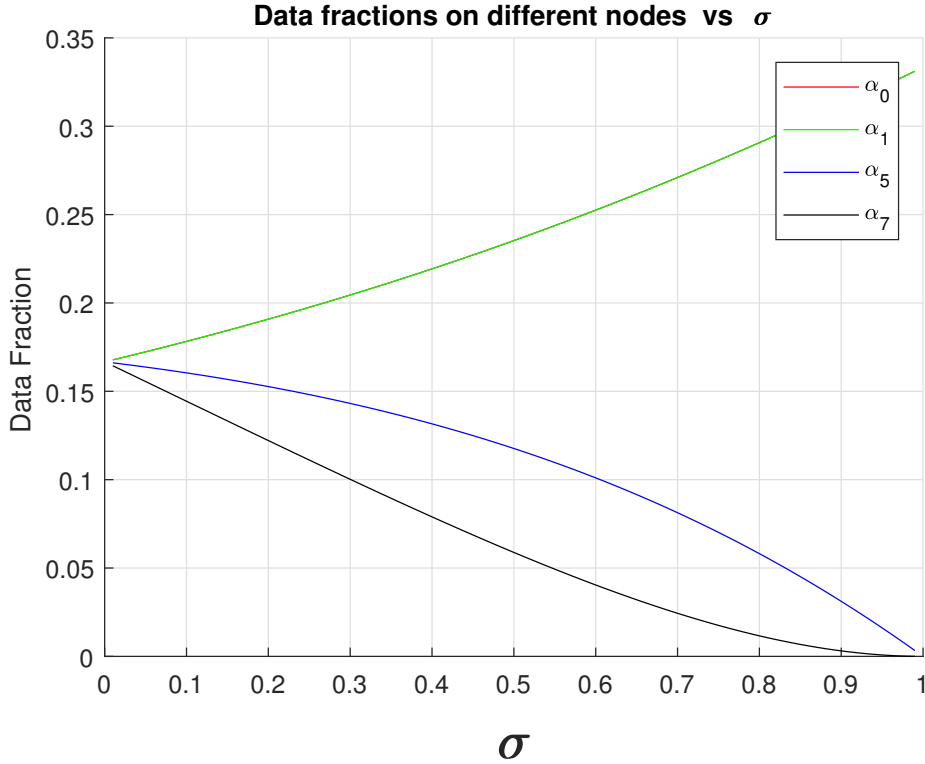


Figure 3.12: The data fraction simulation result of 3×3 mesh network and the data injection happens on the boundary P_0

α_3 . The curve α_4 is identified to the curves of α_5 and α_6 . The curve α_7 is identified to the curve for α_8 . If communication time is zero (extremely fast communication), each processor get $\frac{1}{9}$ of the load. If communication time equals computation time, each of the four start stage nodes (P_0, P_1, P_2, P_3) get $\frac{1}{4}$ of the load fraction.

3.1.3 Data Injection On The Inner Grid Processor

Fig. 3.13 shows that L load on the inner grid processor P_0 , P_0 , P_1 , P_2 , P_3 and P_4 commence processing at time = 0 through virtual cut-through switching. Once this is finished links $0 - 1$, $0 - 2$, $0 - 3$ and $0 - 4$ are released to forward load to P_5 , P_6 , P_7 and P_8 in virtual cut-through mode.

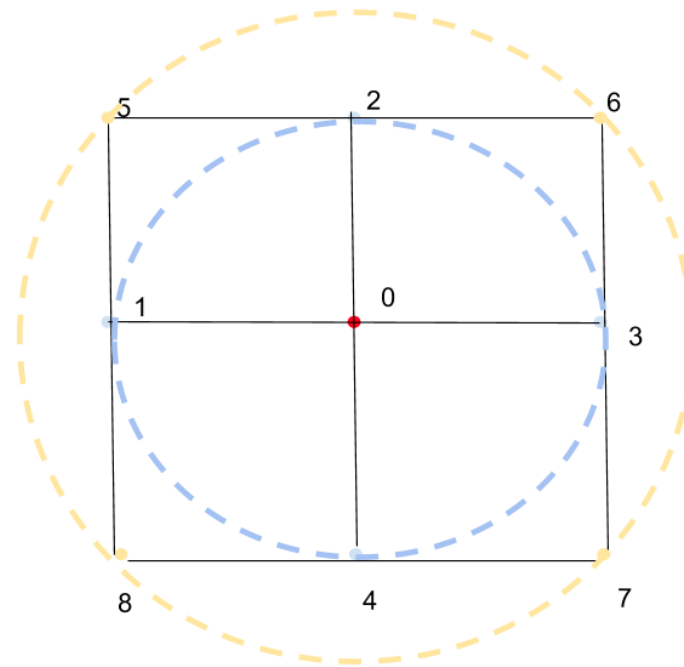


Figure 3.13: 3×3 mesh network. The data injection position is inner grid point P_0

The timing diagram for this user case is illustrated as Fig. 3.14:

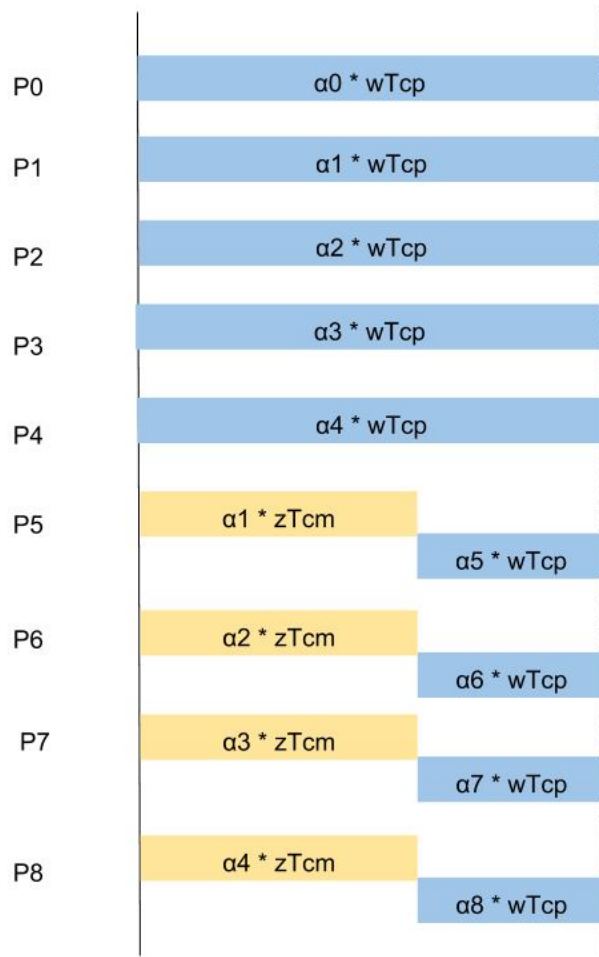


Figure 3.14: The timing diagram for 3*3 mesh network and the data injection is inner grid P_0

The group of equations are:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_3 \omega T_{cp} = T_{f,m} \\ \alpha_4 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_6 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_7 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_8 \omega T_{cp} = T_{f,m} \\ \sigma = \frac{z T_{cm}}{w T_{cp}} \end{array} \right. \quad \begin{array}{l} (3.73) \\ (3.74) \\ (3.75) \\ (3.76) \\ (3.77) \\ (3.78) \\ (3.79) \\ (3.80) \\ (3.81) \\ (3.82) \end{array}$$

The flow matrix form is :

$$\begin{bmatrix} 1 & 4 & 4 \\ 1 & -1 & 0 \\ 0 & \sigma - 1 & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_5 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix} \quad (3.86)$$

The simulation result is Fig. 3.15: P_0 and P_1 have the same α value, so the curve of α_0 and α_1 coincide.

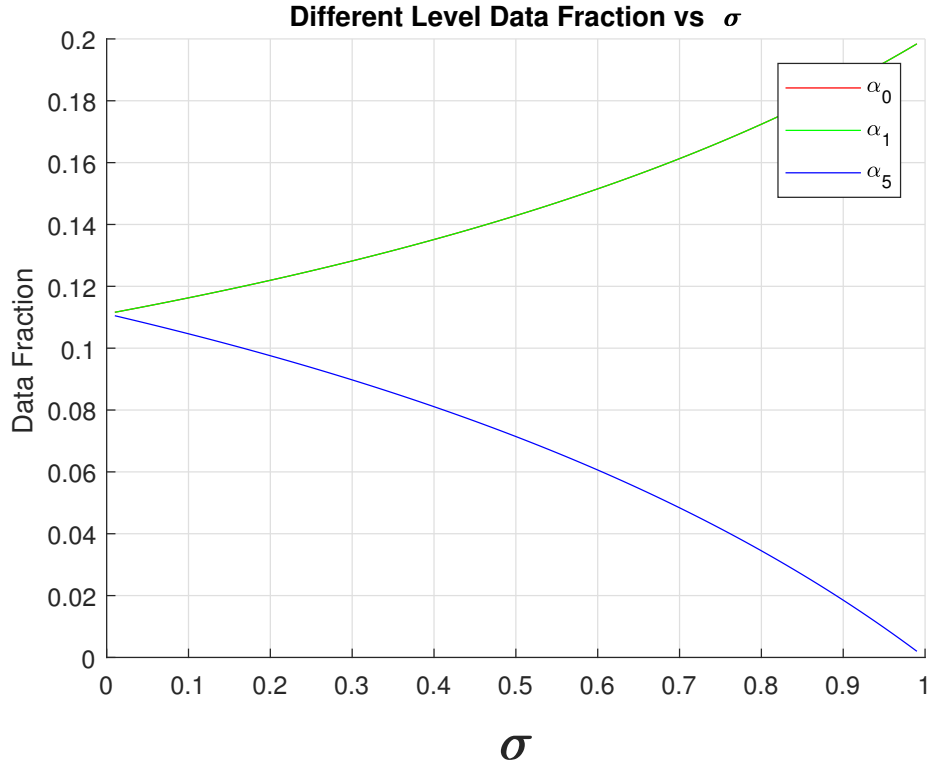


Figure 3.15: 3*3 mesh network. The data injection position is inner grid point P_0

The explicit solution is:

$$\left\{ \begin{array}{l} \alpha_0 = \frac{1}{9 - 4 \times \sigma} \\ \alpha_1 = \frac{1}{9 - 4 \times \sigma} \end{array} \right. \quad (3.87)$$

$$\left\{ \begin{array}{l} \alpha_0 = \frac{1}{9 - 4 \times \sigma} \\ \alpha_1 = \frac{1}{9 - 4 \times \sigma} \end{array} \right. \quad (3.88)$$

$$\left\{ \begin{array}{l} \alpha_0 = \frac{1 - \sigma}{9 - 4 \times \sigma} \\ \alpha_1 = \frac{1 - \sigma}{9 - 4 \times \sigma} \end{array} \right. \quad (3.89)$$

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

If the communication time is zero (very fast communication), each of nine processors get $\frac{1}{9}$ of the load. If communication time equals computation time, each of processors P_0 , P_1 , P_2 , P_3 and P_4 get $\frac{1}{5}$ of the load.

3.1.4 Sensitivity Analysis Virtual Cut-through Switching Scenario

From Chapter 2, we know the speedup is :

$$\text{Speedup} = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = |\det A|$$

Data Injection On The Corner Processor

The simulation result of sensitivity analysis of $2 * n$ mesh network Fig. 3.7 is as follows:

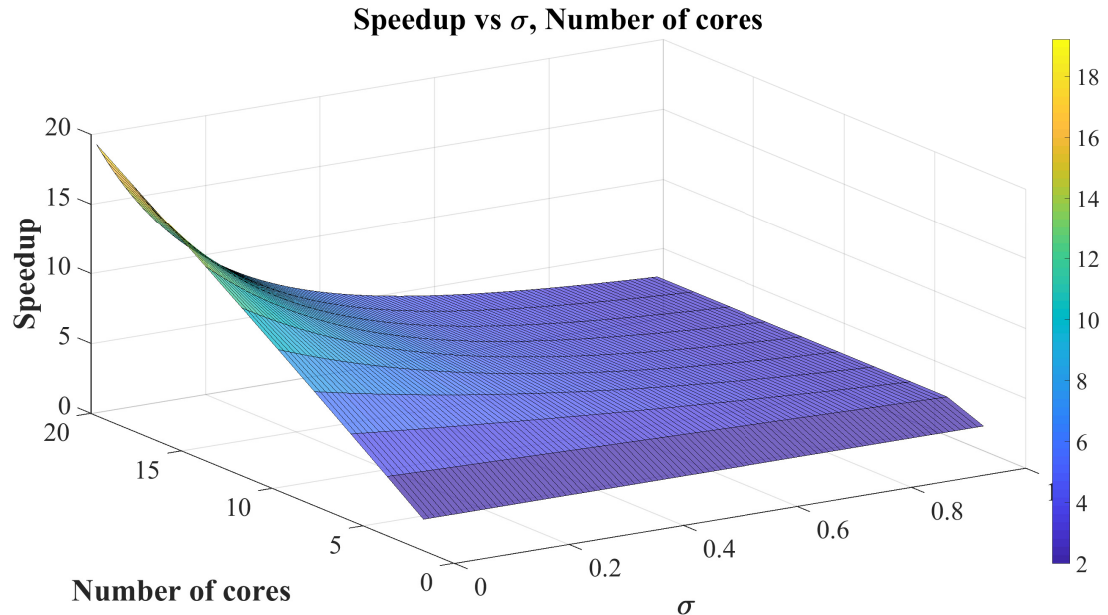


Figure 3.16: Sensitivity analysis result of $2 * n$ mesh network (where $n = 1 \dots 10$)

One can see speedup increases with an increasing number of core (i.e. processor) and also increases with decreasing σ (that is increasing communication speed relative to computation speed).

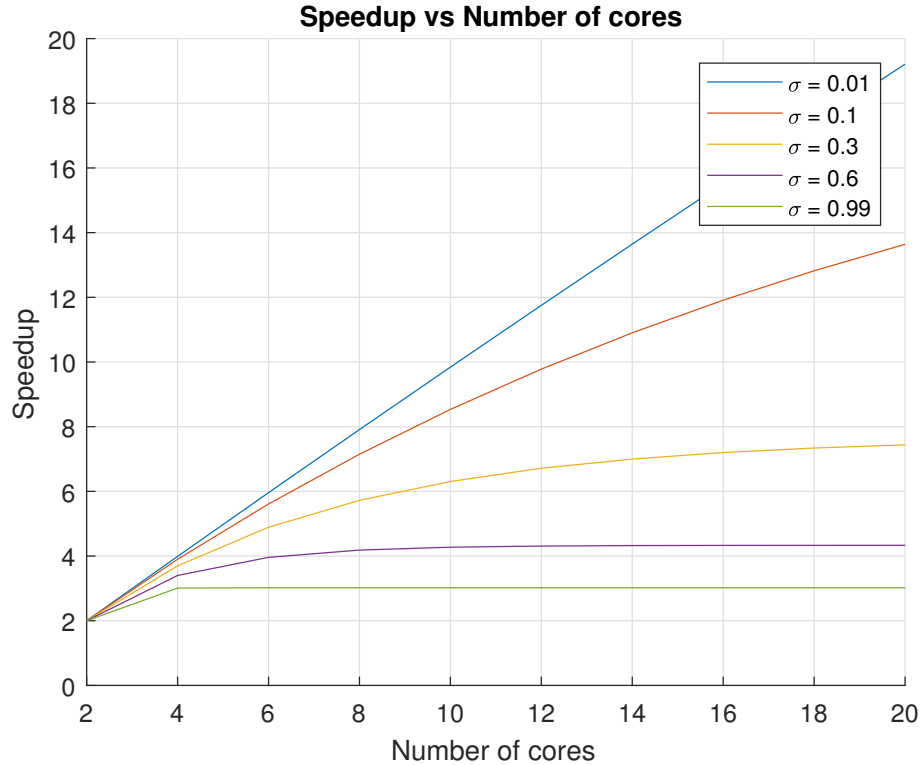


Figure 3.17: Speedup curves of 2^*n mesh network (where $n = 1 \cdots 10$)

For a large number of cores and σ close to one, speedup is about three. P_0 and two adjacent neighbor processor do most of the processing.

Data Injection On The Boundary Processor

For a $3 * n$ mesh network Fig. 3.9, the data injection happens on boundary processor P_2 .

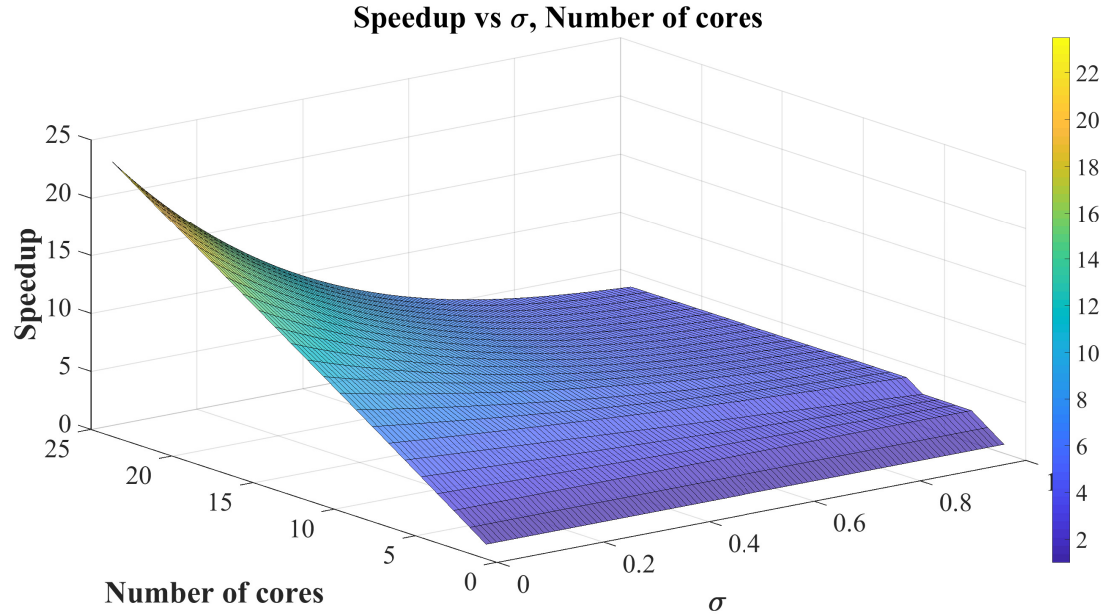


Figure 3.18: Sensitivity analysis result of $3*n$ mesh network and the injection position on boundary processor P_2 (where $n = 1 \dots 8$)

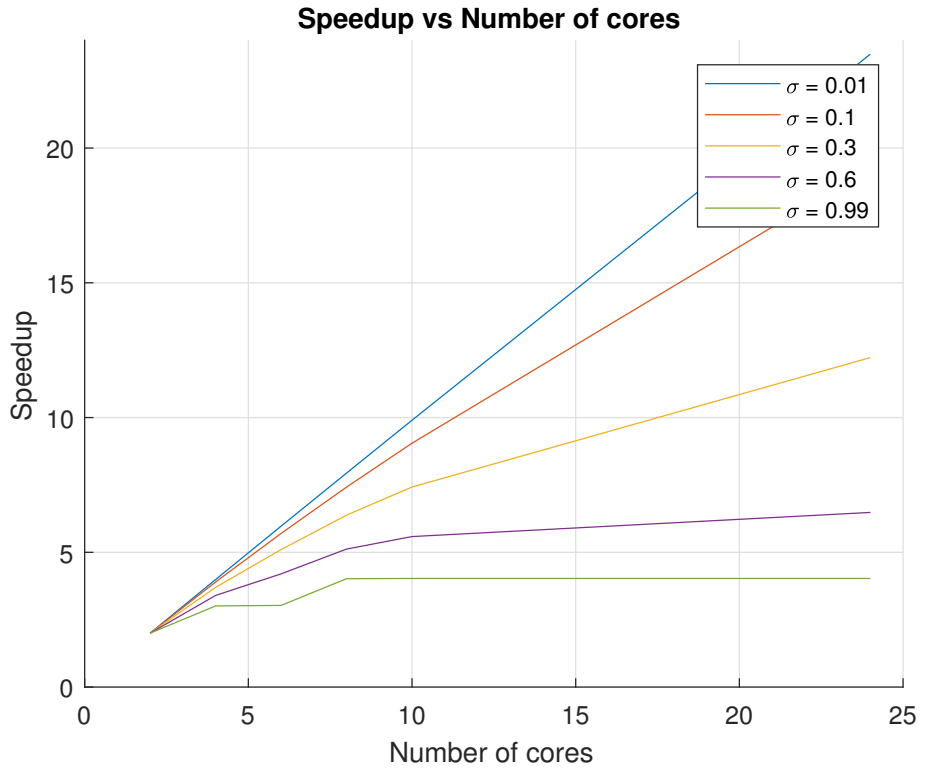


Figure 3.19: Speedup curve of 3^*n mesh network (where $n = 1 \cdots 8$)

Fig. 3.18 and Fig. 3.19 show that if the value $\sigma > 0.2$, the speedup increases rapidly as a function of the number of cores. If the value $\sigma < 0.1$, the number of cores has linear impact on the speedup performance. If the number of cores is for σ close to one, the speedup is about four since only the three processors close to P_0 and P_0 do almost all the processing.

Data Injection On The Inner Grid Processor

For a $5 * n$ mesh network is in Fig. 2.1, L originate on the inner grid P_{12} and the simulation result says:

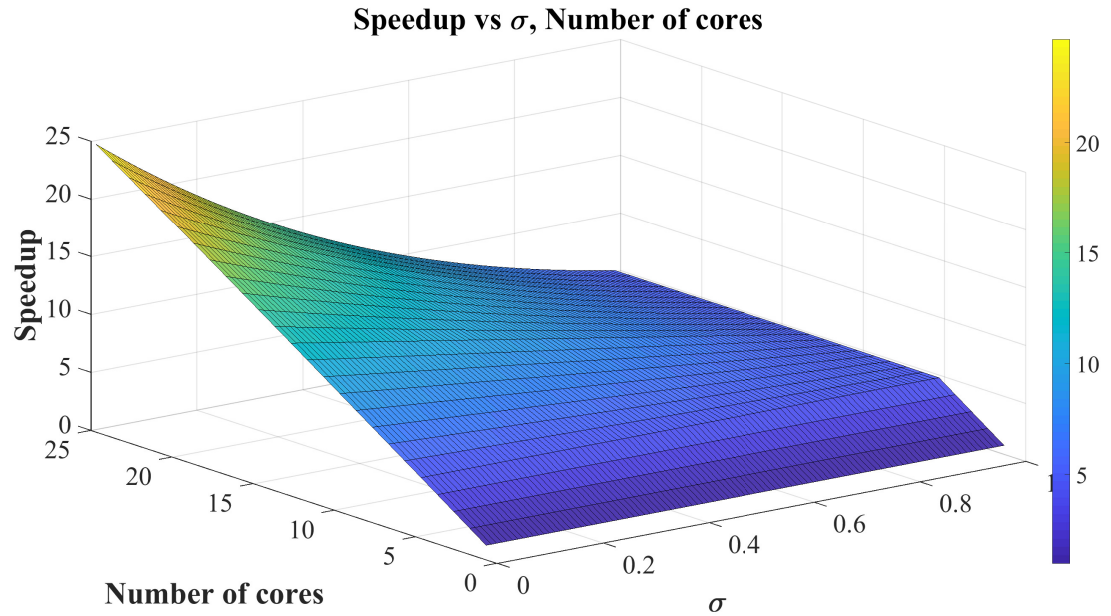


Figure 3.20: Sensitivity analysis result of $5 * n$ mesh network and the injection position on inner processor P_{12} (where $n = 1 \dots 5$)

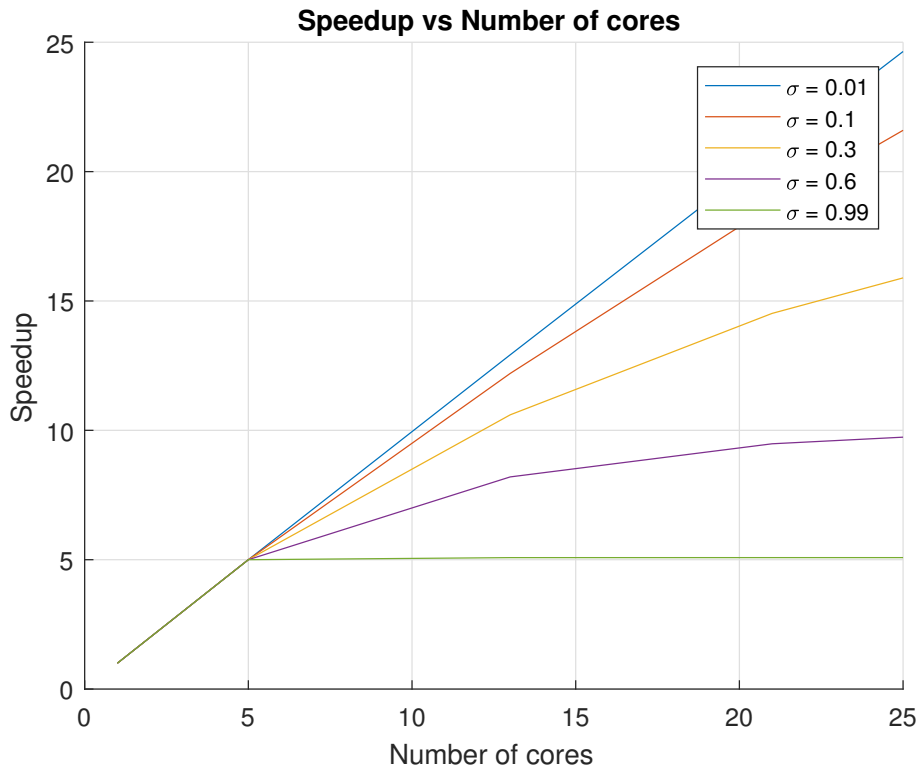


Figure 3.21: Speedup curves of $5 \times n$ mesh network Fig. 2.1 (where $n = 1 \dots 5$)

If the number of processor > 5 , the cluster equivalence computation ability is at least 5 time speedup. There are 5 processors in the first stage. For small σ speedup growth is linear. For large σ and a large number of cores, speedup is about five as only P_0 and four adjacent processors do almost all the processing.

3.1.5 Multi-source Uniform Data Fraction

We extend the single source assignment to the multi-source assignment problem [55] [16]. For each processor, we concentrate on the processors' geographical location P_i , and assigned data fraction α_i .

Assume the data fractions are distributed uniformly. For example, the workload is unit 1 and there are k different data injection options. So each data injection is assigned $\frac{1}{k}$ workload.

From the data injection position relationship we consider three different situations :

- Data injection positions consist of a connected subgraph G_L of G .
- Data injection processors don't connect with each other.
- Some data injection positions consist of some connected subgraphs and some are individual injection points.

Model I

If the data injection positions consist of a connected subgraph of G , we use G_L to present it.

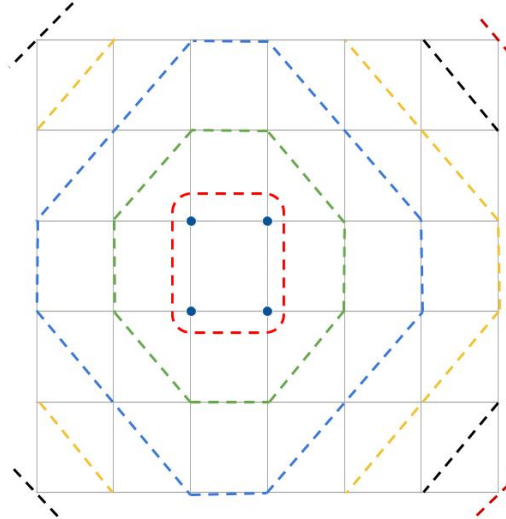


Figure 3.22: Example 1: Data injection consists of a connected subgraph of G

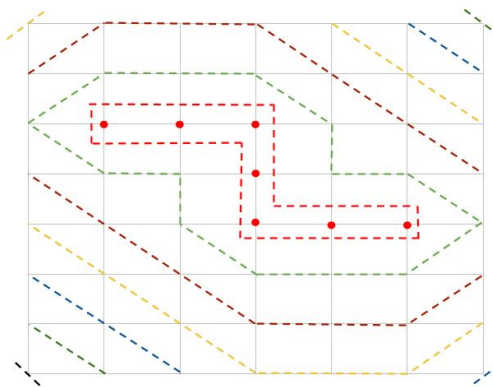


Figure 3.23: Example 2: Data injection consists of a connected subgraph of G

Connected subgraph Fig. 3.22 and connected subgraph Fig. 3.23 illustrate two examples that the data injections consist of connected subgraphs of G .

Our objective is to propose a general algorithm framework to minimize the makespan and give a quantitative model analysis utilizing the flow matrix.

$$\min \quad \hat{T}_f \quad (3.90)$$

$$\text{s.t.} \quad \sum_{j=0}^{k-1} \sum_{i=0}^{S_j-1} \alpha_{j,i}^* = 1 \quad (3.91)$$

$$1 \geq \alpha_{j,i}^* \geq 0, \quad (3.92)$$

$$T_{f,i} = T_{f,0}, \quad i \in 1 \cdots (m * n - 1) \quad (3.93)$$

$$\alpha_{j,0}^* w T_{cp} = \hat{T}_{j,f}, \quad j \in 0 \cdots (k - 1) \quad (3.94)$$

$$\vdots, \quad (3.95)$$

$$(\alpha_{j,1}^* + \cdots + \alpha_{j,\hat{l}_j-2}) z T_{cm} + \alpha_{j,\hat{l}_j-1}^* w T_{cp} = \hat{T}_{j,f}, \quad (3.96)$$

- Equation 3.90 is the objective function, minimizing the makespan. \hat{T}_f is the finish time of the whole processor network. Here \hat{T}_f is equal to $w_{eq} T_{cp}$.
- Equation 3.91 means the total fraction of each processor is unit 1. $\alpha_{j,i}^*$ is the load fraction assigned to i th processor of j th layer.
- Equation 3.92 means each processor's data fraction is nonnegative.
- Equation 3.94 means each processor has the same finish time.
- Equation 3.96 means the j th layer's finish time is consisted of previous $(j - 1)$ layers' transmitting time and j th computation time.

This algorithm is named as ***Equivalence Processor Scheduling Algorithm (EPSA)***.

Algorithm 1 Equivalence Processor Scheduling Algorithm (EPSA)

Input: k data injection positions

Output: $m * n$ processor data fractions α_i

Collapse the data injection processors into one “big” equivalent processor [15].

Calculate $m * n$ processor’s D_i .

Obtain the flow matrix A .

Calculate the determinant of flow matrix.

Calculate $m * n$ processors’ data fraction α_i .

In term of the time complexity :

- r is the rank of flow matrix.
- The time complexity of calculating the determinant is $O(r^3)$ with Gaussian elimination or LU decomposition.
- The time complexity of calculating the flow matrix A_i is $O(k * m * n)$. k is the number of data injection location.
- The total time complexity is $O(r^3 + k * m * n)$.

For example, connected subgraph Fig. 3.22’s ***flow matrix*** is :

$$\begin{bmatrix} 4 & 8 & 12 & 10 & 6 & 2 \\ 1 & -1 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_2 \\ \alpha_3 \\ \alpha_4 \\ \alpha_5 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (3.97)$$

The simulation result illustrates as follows:

This simulation says the best performance happens on value $\sigma \leq 0.05$, which hits about 42 times speedup as all 42 processors in the network are engaged in processing networks. There are 12 processors at the start stage. So if $\sigma \approx 1$, that is, communication time equals computation time, the network achieves about 12 times speedup performance, only those 12 processors do processing.

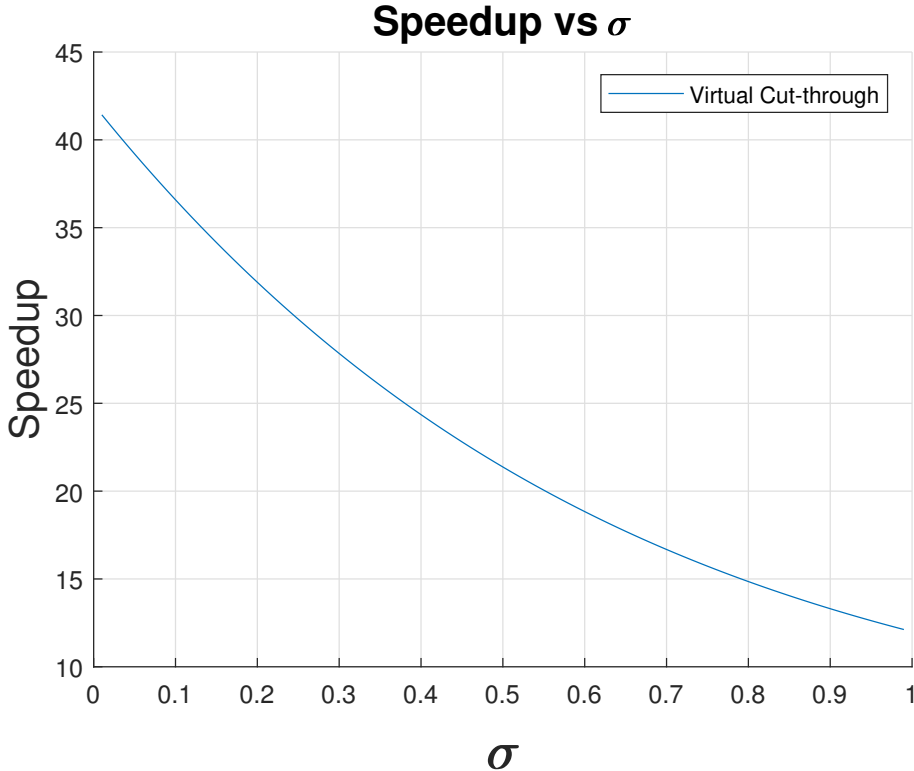


Figure 3.24: Connected subgraph Fig. 3.22 speedup vs σ

For example, connected subgraph Fig. 3.23's **flow matrix** is :

$$\begin{bmatrix} 7 & 14 & 15 & 10 & 6 & 3 & 1 \\ 1 & -1 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & 1 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_2 \\ \alpha_3 \\ \alpha_4 \\ \alpha_5 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (3.98)$$

The simulation result illustrates as follows:

This simulation says the best performance happens on value $\sigma \leq 0.05$, which hits about 56 times speedup as there are 56 processors engaged over the entire network. There are 21 processors in the start stage. So if $\sigma \approx 1$, the network achieves 21 times speedup performance as only those 21 processors do processing.

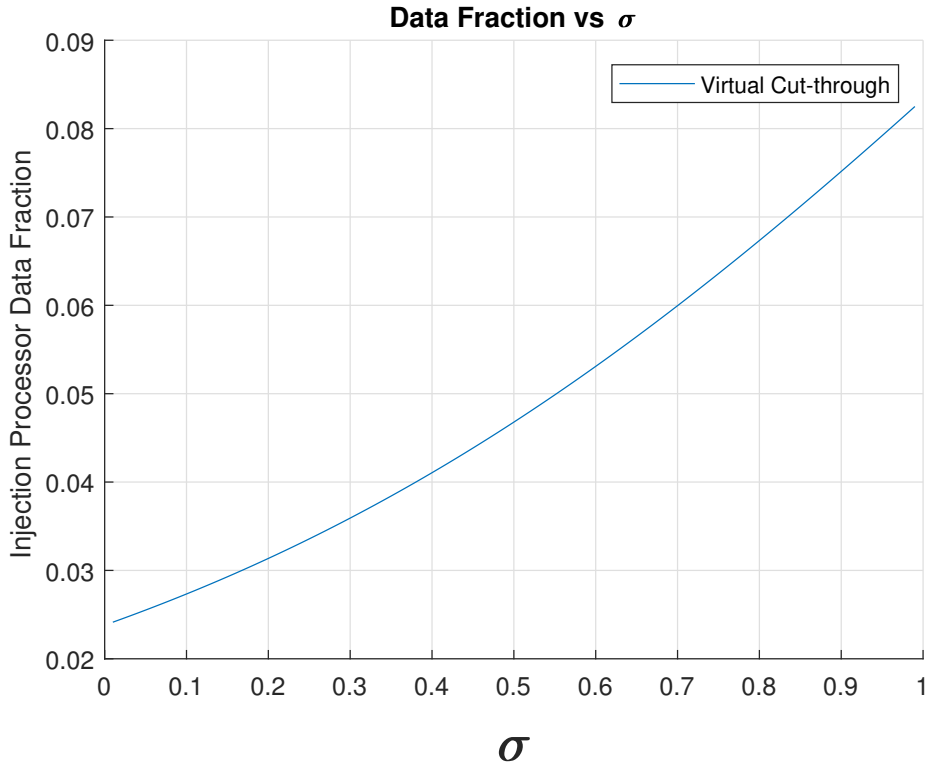


Figure 3.25: Connected subgraph Fig. 3.22's processor data fraction vs σ

Model II

Now the data injections don't consist of a connected subgraph of G . Jia [55] proposes a genetic algorithm, which utilize a novel Graph Partitioning (GP) scheme to partition the network such that each source in the network gains a portion of network resources and then these sources cooperate to process their loads.

Relevant divisible load applications include, for example, big file transmission, HPC scientific computation, like BlueGene [59] or Hadoop jobs.

Our objective is to propose a general algorithm framework to utilize less processors and achieve the same finish time. This is a critical issue for networks on chip (NOC) where the number of active processor is closely related to the chip power computation chip. Power computation is a limiting factor to scalability in such NOCs.

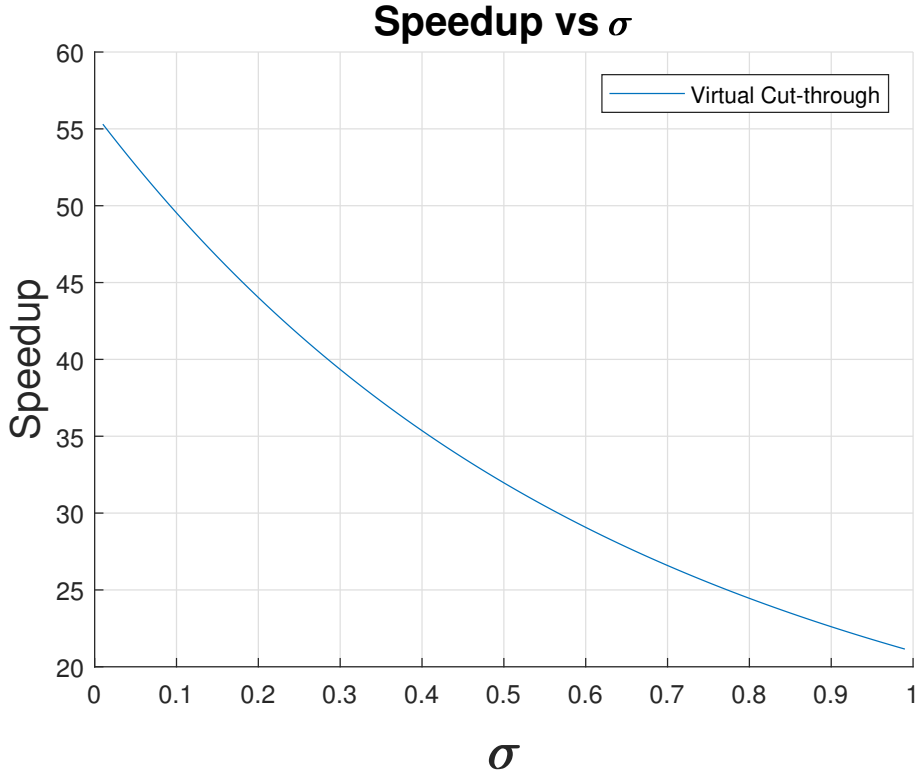


Figure 3.26: Connected subgraph Fig. 3.23 speedup vs σ

Definition 3.1.1. Voronoi diagram [60]

“Let S denote a set of n points (called sites) in the plane. For two distinct sites $p, q \in S$, the dominance of p over q is defined as the subset of the plane being at least as close to p as to q . Formally,

$$dom(p, q) = \{x \in R^2 \mid \delta(x, p) \leq \delta(x, q)\}$$

for δ denoting the euclidean distance function. Clearly, $dom(p, q)$ is a closed half plane bounded by the perpendicular bisector of p and q .

This bisector separates all points of the plane closer to p from those closer to q and will be termed the separator of p and q . The region of a site $p \in S$ is the portion of the plane lying in all of the dominances of p over the remaining sites in S . Formally,

$$reg(p) = \bigcap_{q \in S - p} dom(p, q)$$

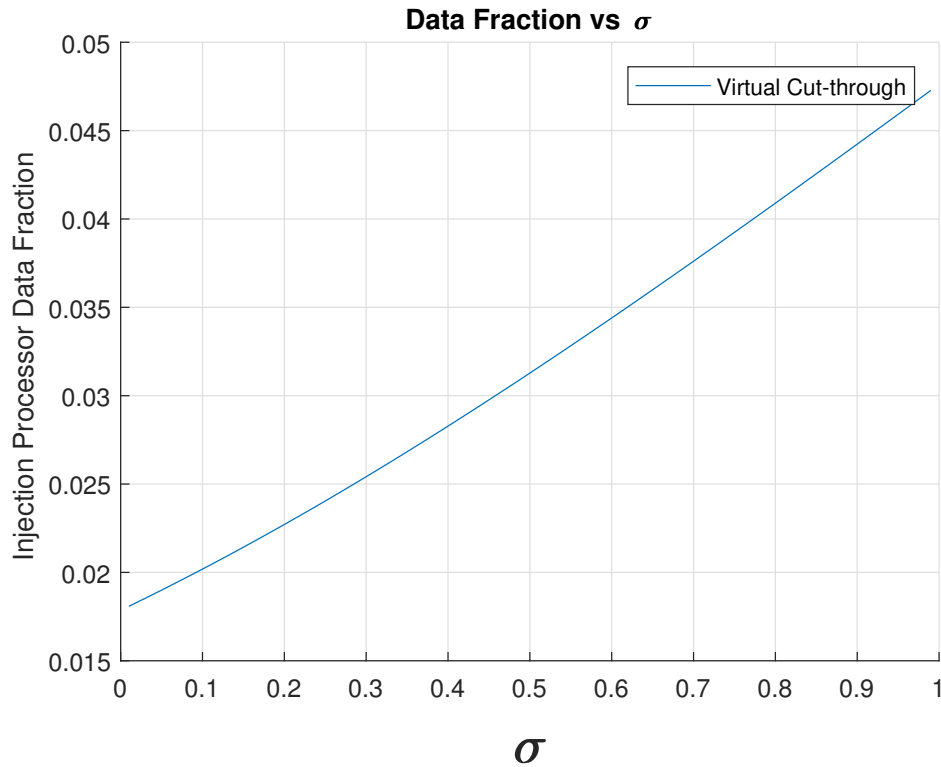


Figure 3.27: Connected subgraph Fig. 3.23's processor data fraction vs σ

"

For each seed there is a corresponding region consisting of all points closer to that seed than to any other.

Definition 3.1.2. *Manhattan distance Voronoi diagram* Unlike most Voronoi diagrams, which use the L2 (Euclidean distance) metric, **Manhattan distance Voronoi diagrams** use the L1 metric.

In this thesis, the intuitive algorithm is to implement the graph partitioning by the Manhattan distance Voronoi diagram.

Manhattan Distance Voronoi Diagram Algorithm

Our objective is to propose an intuitive algorithm to minimize the makespan and give quantitative model analysis utilizing the flow matrix.

The objective functions and constraints are the same with Equivalence Processor Scheduling Algorithm (EPSA), equations 3.90, 3.91, 3.92, 3.94, 3.96.

The intuitive algorithm is named as ***Manhattan Distance Voronoi Diagram Algorithm***:

Algorithm 2 Manhattan Distance Voronoi Diagram Algorithm (RMDVDA)

Input: k data injection positions

Output: $m * n$ processor data fractions α_j .

Calculate k Voronoi cells with Manhattan distance.

Calculate k flow matrix A_i .

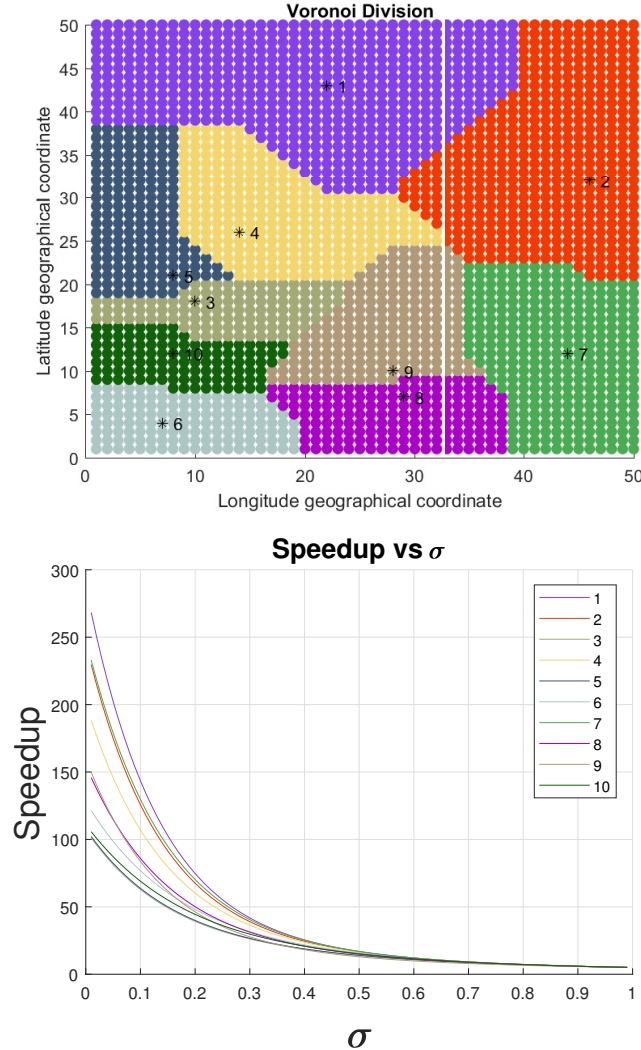
Calculate k the determinant of flow matrix A_i .

Calculate $m * n$ processors' data fraction α_j .

The time complexity is the same with EPSA.

Manhattan distance Voronoi diagram division shows 10 Voronoi cells division.

Table 3.1: The 10 Manhattan distance Voronoi Cells and speedup curves



After investigation, we find the makespan depends on the bottleneck makespan. In other words, if other cells consist of more processors than the bottleneck cell, it does not help to minimize the makespan.

Reduced Manhattan Distance Voronoi Diagram Algorithm

Our objective is to propose a heuristic algorithm to minimize the makespan and give quantitative model analysis utilizing the flow matrix. The merits of new algorithm is finishing the task within the same makespan as MDVDA, yet utilizing less processor resources.

$$\min \hat{T}_f, \quad (3.99)$$

$$\text{s.t. } \sum_{j=0}^{k-1} \sum_{i=0}^{S_j-1} \alpha_{j,i}^* = 1 \quad (3.100)$$

$$1 \geq \alpha_{j,i}^* \geq 0, \quad (3.101)$$

$$T_{f,i} = \hat{T}_{j,f}, \quad i \in 0 \cdots (S_j - 1) \quad (3.102)$$

$$\alpha_{j,0}^* w T_{cp} = \hat{T}_{j,f}, \quad (3.103)$$

$$\vdots, \quad (3.104)$$

$$(\alpha_{j,1}^* + \cdots + \alpha_{j,\hat{l}_j-2}^*) z T_{cm} + \alpha_{j,\hat{l}_j-1}^* w T_{cp} = \hat{T}_{j,f}, \quad j \in 0 \cdots (k-1) \quad (3.105)$$

$$Sp_i = Sp_b + \epsilon, \quad i \in 0 \cdots (k-1) \quad (3.106)$$

$$\epsilon \geq 0, \quad (3.107)$$

- $\hat{T}_f = \max T_{f,i}, i \in 0 \cdots (k-1)$. Equation 3.99 means minimizing the makespan of the whole mesh network.
- Equation 3.106 means each Voronoi equivalence speedup is approximate same with the bottleneck cell's equivalence speedup. $\max T_{f,i}$ is the speedup of bottleneck cell's finish time.
- ϵ is a small enough nonnegative real number.

The heuristic algorithm is named as *Reduced Manhattan Distance Voronoi Diagram Algorithm*:

Algorithm 3 Reduced Manhattan Distance Voronoi Diagram Algorithm (RMDVDA)

Input: k data injection positions

Output: $m * n$ processor data fractions α_i .

Calculate k Voronoi cells with Manhattan distance.

Calculate k Voronoi cells' radius R_i .

Calculate k flow matrix A_i .

$depth_{min} = \min(Sp_i)$'s R_i .

Calculate the reduced Voronoi cells by setting the $depth_i = depth_{min}$ in each Voronoi cell.

Calculate reduced Voronoi cell's flow matrix \hat{A}_i .

Calculate $m * n$ processors' data fraction α_i .

“Reduced” means the Voronoi cell shrinks from the outside layer to interior direction layer by layer, which keeps the equivalence computation faster than the bottleneck’s computation. The analysis of the time complexity is the same as EPSA.

Table 3.2: The 10 Reduced Voronoi Cells and speedup curves

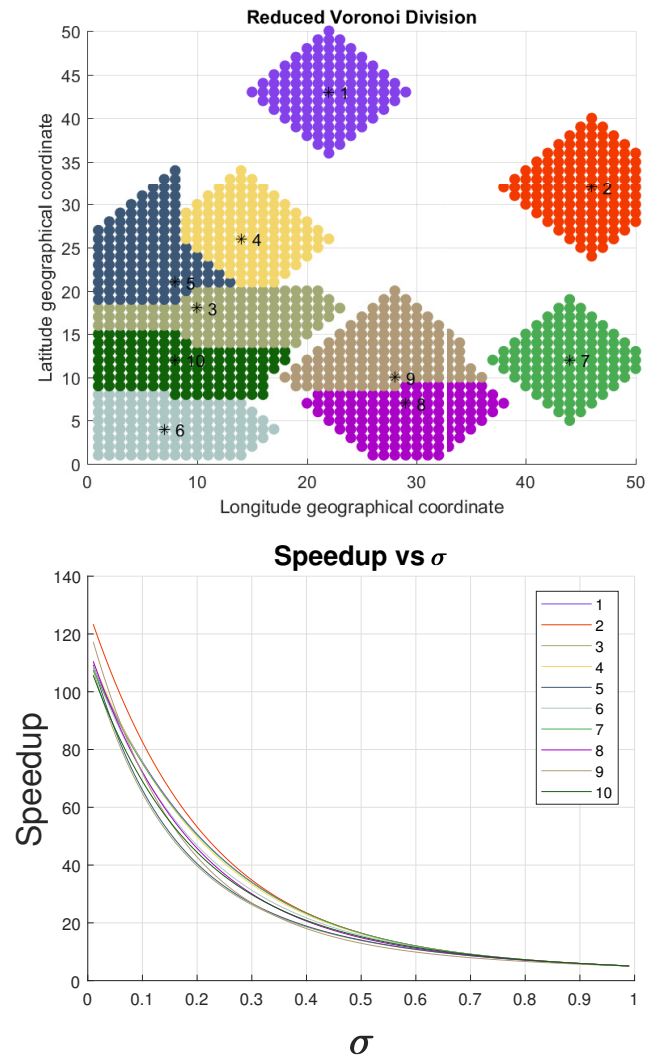


Table 3.2 shows reduced Voronoi cells and reduced Voronoi speedup curves, which saves about 30% processors within the same makespan.

- Voronoi speedup curves show that $\sigma < 0.2$, the ratio $\max(\frac{Sp_{max}}{Sp_{min}}) = \frac{271}{107} \approx 2.533$.
- Reduced Voronoi speedup curves show that $\sigma < 0.2$, the ratio is $\max(\frac{Sp_{max}}{Sp_{min}}) = \frac{122}{107} = 1.14$.

It displays that 10 cells' equivalence computation is more balanced than the initial setting, and the whole cluster finishes processing load within the same time by less processors. After 1000 round random sampling experiments, we obtain the average saved processors ratio in Fig. 3.28.

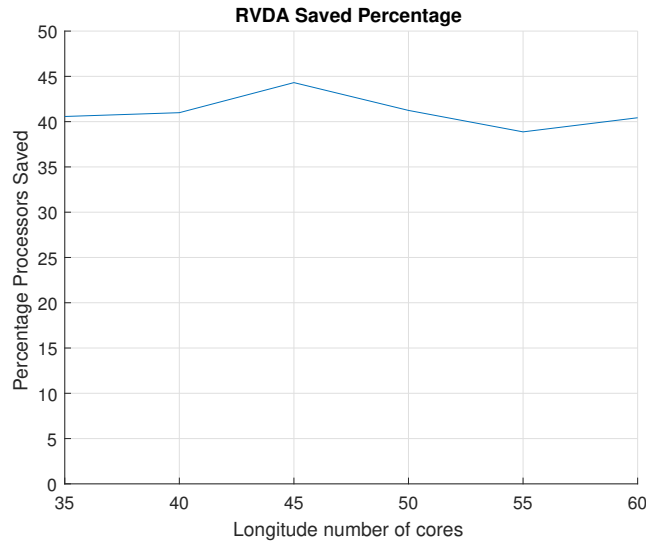


Figure 3.28: Reduced Voronoi diagram algorithm average saved processors' percentage

From the average saved processors ratio Fig. 3.28, it shows the average percentage of saved processor is about 40%.

Accelerated Reduced Manhattan Distance Voronoi Diagram Algorithm

In order to accelerate the bottleneck division computation, we propose a heuristic algorithm to minimize the makespan and give quantitative model analysis utilizing the flow matrix.

The merits of new algorithm is finishing the task within less makespan as MDVDA and also utilizing less processor resources.

$$\max \quad Sp_b, \quad (3.108)$$

$$\text{s.t.} \quad \sum_{j=0}^{k-1} \sum_{i=0}^{S_j-1} \hat{\alpha}_{j,i} = 1 \quad (3.109)$$

$$1 \geq \hat{\alpha}_{j,i} \geq 0, \quad (3.110)$$

$$T_{f,i} = \hat{T}_{j,f}, \quad i \in 0 \cdots (S_j - 1) \quad (3.111)$$

$$\hat{\alpha}_{j,0} w T_{cp} = \hat{T}_{j,f}, \quad j \in 0 \cdots (k - 1) \quad (3.112)$$

$$\vdots, \quad (3.113)$$

$$(\hat{\alpha}_{j,1} + \cdots + \hat{\alpha}_{j,l_j-2}) z T_{cm} + \hat{\alpha}_{j,l_j-1} w T_{cp} = \hat{T}_{j,f}, \quad j \in 0 \cdots (k - 1) \quad (3.114)$$

$$Sp_i = Sp_b + \epsilon, \quad i \in 0 \cdots (k - 1) \quad (3.115)$$

$$\epsilon \geq 0, \quad (3.116)$$

- Equation 3.108 is the objective function. Through maximizing the bottleneck cell's speedup, we minimize the makespan.

Algorithm 4 Accelerated Reduced Manhattan Distance Voronoi Diagram Algorithm (AR-MDVDA)

Input:

- k data injection positions
- $\sigma = \frac{zT_{cm}}{wT_{cp}}$
- processors and links' parameters
- f_T : the requirement of finish time
- r_{save} : the requirement ratio of saved processors
- tmp_{save} : the ratio of saved processors
- Sp_i : the speedup of i Voronoi cell.
- $Sp_{min} : \min(Sp_i) \quad i \in 1 \cdots k$
- $index_{min}$: the index of Sp_{min}

Output:

- $m * n$ processor data fractions α_i .

Calculate k Voronoi cells with Manhattan distance and k Voronoi cells' radius R_i .

Calculate k flow matrix A_i and set $depth_{min} = \min(Sp_i)$'s R_i .

Calculate the reduced Voronoi cells by setting the $depth_i = depth_{min}$ in each Voronoi cell.

Calculate the determinant of flow matrix.

Calculate reduced Voronoi cell's flow matrix \hat{A}_i and Sp_i of each cell.

while $\frac{1}{Sp_{min}} > f_T$ and $tmp_{save} > r_{save}$ **do**

 merge processors to $index_{min}$ Voronoi cell one by one

 by increasing order of D_i .

end while

Calculate $m * n$ processors' data fraction α_i .

Comparing with RMDVDA, ARMDVDA accelerates the speedup about 20%.

Table 3.3: The 10 Accelerated Reduced Voronoi Cells and Speedup Curves

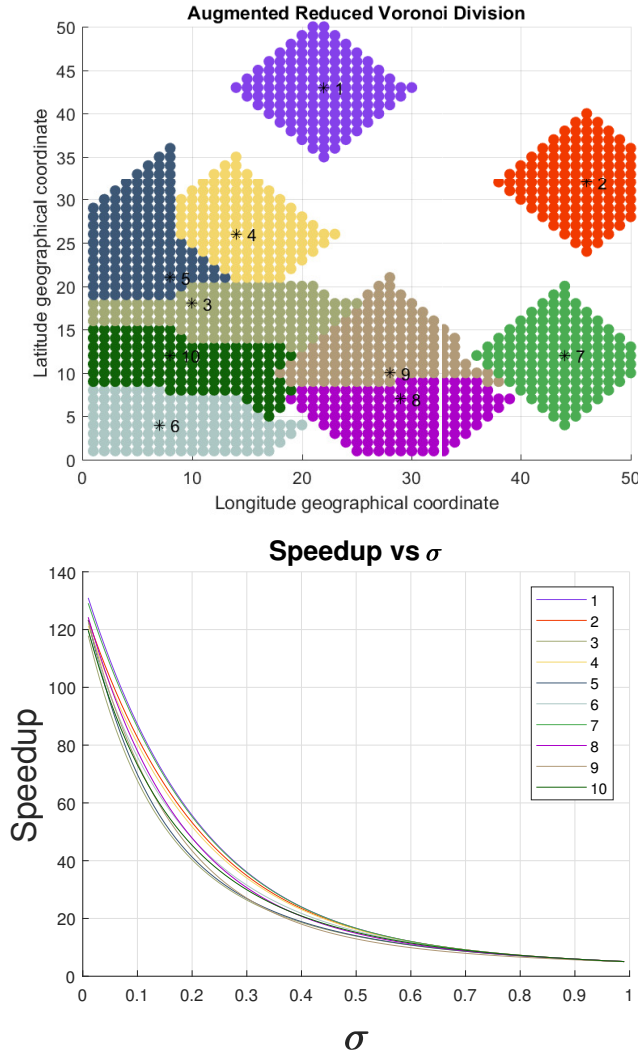


Fig. 3.29 shows the Voronoi cell 3 is the bottleneck division and there is no other economical way to enlarge this constraint cell.

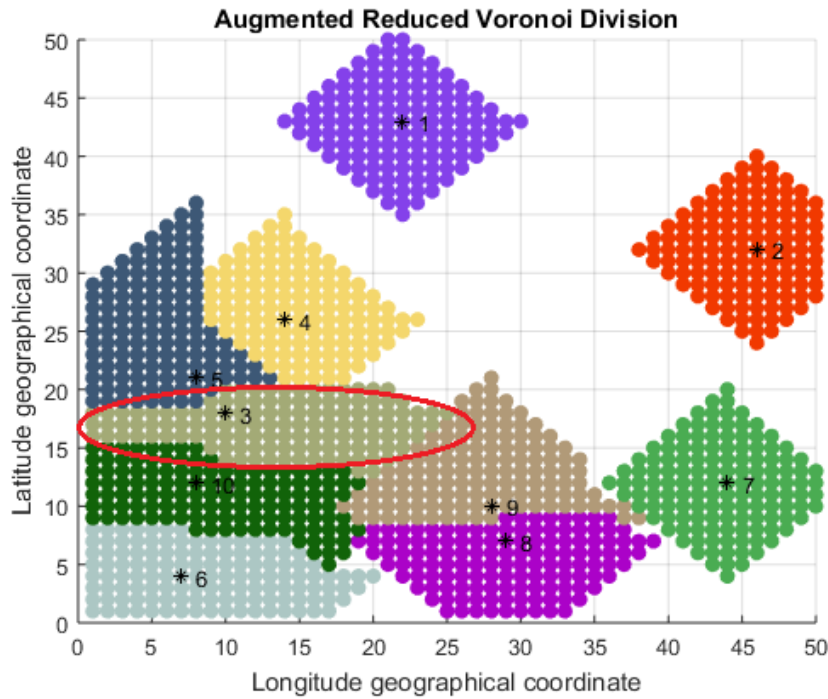


Figure 3.29: 10 reduced Voronoi cells

Model III

If there are some nodes consisting some connected subgraphs and there are some nodes individual processors, our objective to finish the load in the same time and save processors. We process an Reduced Constrained Manhattan Distance Voronoi Diagram Algorithm (RCMDVDA) to tackle this situation.

In term of the time complexity, it's the same with EPSA.

Algorithm 5 Reduced Constrained Manhattan Distance Voronoi Diagram Algorithm (RCMDVDA)

Input: \hat{k} data injection positions

Output: $m * n$ processor data fractions

Collapse the connected data injection processors into some “big” equivalent processors, individually.

Calculate k constrained Manhattan distance Voronoi cells [21].

Calculate k constrained Voronoi cells’ radius R_i .

Calculate k constrained Voronoi cells’ flow matrix A_i .

Calculate k constrained Voronoi cells’ Sp_i .

Set the $depth_{min} = \min(Sp_i)$ ’s radius.

Calculate k reduced constrained Voronoi cells by setting the $depth_i = depth_{min}$ in each Voronoi cell.

Calculate Voronoi cell’s flow matrix \hat{A}_i .

Calculate the determinant of flow matrix.

Calculate $m * n$ processors’ data fraction α_i .

- Voronoi speedup curves show that $\sigma < 0.2$, the ratio $\max(\frac{Sp_{max}}{Sp_{min}}) = \frac{360}{98} \approx 3.67$
- Reduced Voronoi speedup curves show that $\sigma < 0.2$, the ratio is $\max(\frac{Sp_{max}}{Sp_{min}}) = \frac{125}{98} = 1.276$.

It displays that 10 cells’ equivalence computation is more balanced than the initial setting, and the whole cluster finishes processing load within the same time by less processors.

In this user case, we save about $294/2500 \approx 11.7\%$ processors.

Lower Bound of Intuitive and Heuristic Algorithms

- Sp : Speedup of an equivalence computation cell.
- $optimal_p$: practical optimal speedup of a load data injection, which is NP hard [16].
- $optimal_i$: optimal speedup of a load data injection in ideal situation
- β : $\frac{Sp}{optimal_i}$ is the ratio between the speedup of an algorithm (i.e. EPSA or RMDVDA) over the practical ideal situation speedup.
- lower bound : β is the value of lower bound.

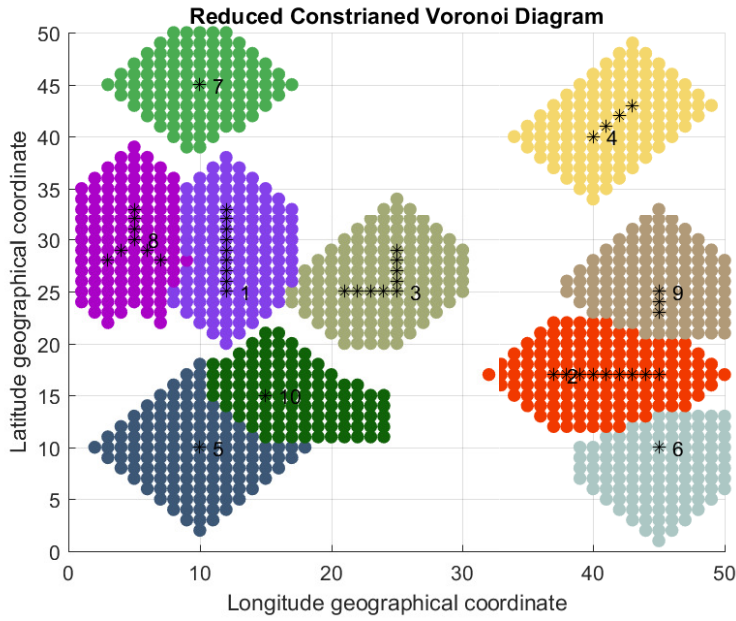


Figure 3.30: Reduced constrained Voronoi diagram for connected subgraphs and individual data injections on mesh network

Considering a $n * n$ mesh network, there are l data load injections and the makespan is constrained by the makespan of bottleneck. So the best solution is that each injection deploys data to its community and the size of community is $\frac{n*n}{l}$.

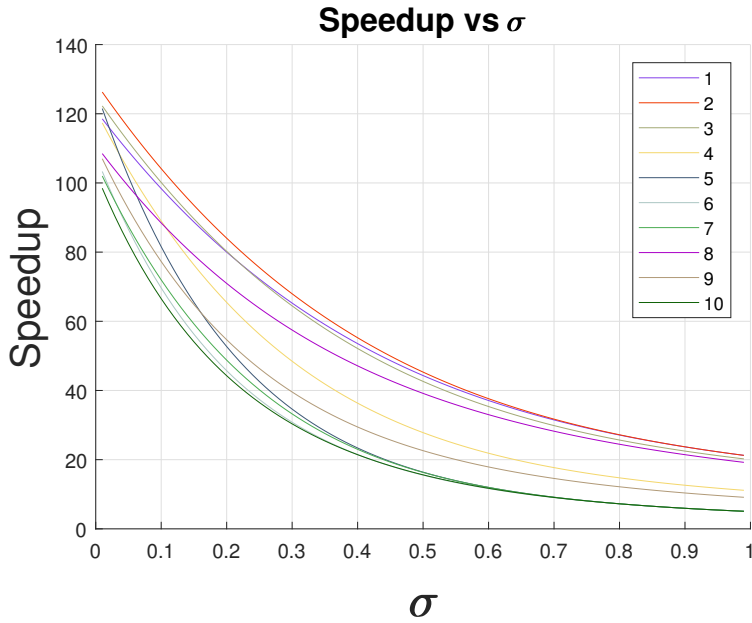


Figure 3.31: Speedup curves of connected subgraphs and individual data injection on mesh network

According to the intuitive and heuristic algorithms, we obtain a lower band of these algorithms.

$$\beta = \frac{Sp}{optimal_i} \leq \frac{Sp}{optimal_p}$$

$$Sp \geq \beta * optimal_p$$

For example, we calculate a $50 * 50$ mesh network and there are $l = 10$ load data injections. Each node transmits data to $\frac{50*50}{10} = 250$ cores.

In ideal assumption, the first row of flow matrix is $row_i = [1\ 4\ 8\ 12\ 16\ 20\ 24\ 28\ 32\ 36\ 40\ 29]$;

We calculate the speedup of a community and the speedup of ideal situation. In other words, we guarantee obtain a $\beta * optimal_i$ times approximation speedup of our algorithms.

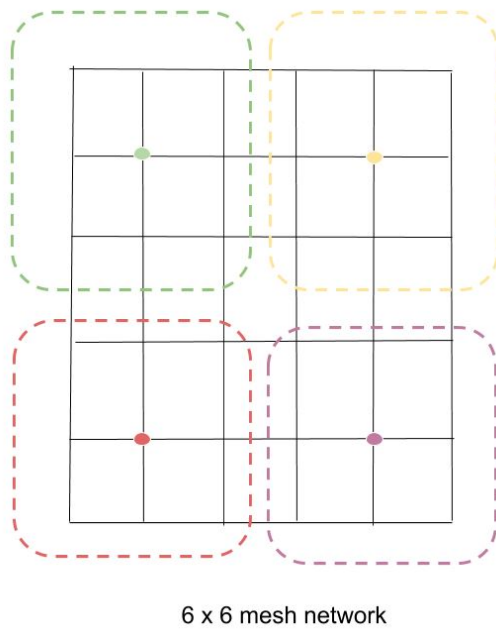


Figure 3.32: Maximal speedup for $6 * 6$ mesh network and 4 data injections

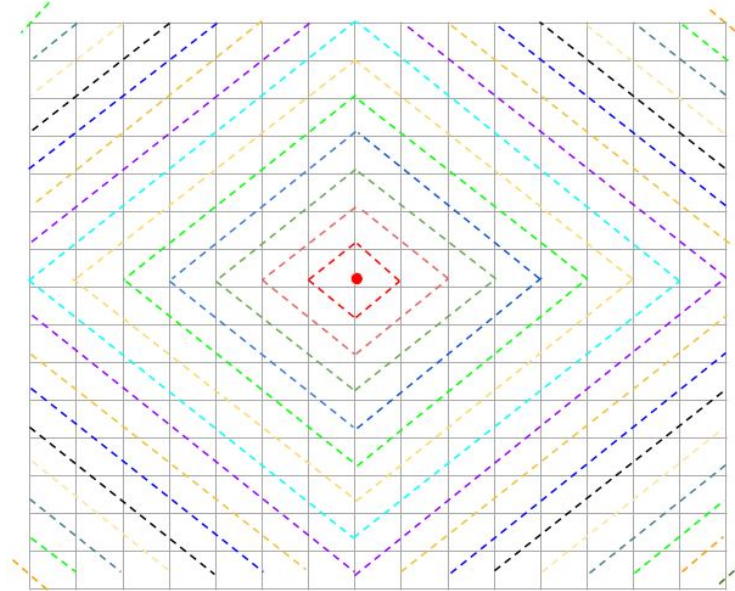


Figure 3.33: Maximal speedup for $16 * 16$ mesh network and 1 data injections

3.1.6 Multi-source Non-uniform Data Injection

From the data injection position relationship, we consider three different situations :

- Data injection positions consist of a subgraph G_L of G .
- Data injection processors don't connect with each other.
- Some data injection positions consist of some connected subgraphs and some are individual injection.

Model I

We propose a suboptimal algorithm framework to minimize the makespan and give quantitative model analysis utilizing the flow matrix. The constraint comes from the Divisible Load Theory linear equations. The objective function and the constraints are the same with EPSA.

The main idea is divided into two stages.

- Re-balance the data fraction between the nodes in connected subgraph first.
- Run EPSA.

Algorithm 6 Equivalence Processor Scheduling Extended Algorithm (EPSEA)

Input: k data injection positions

Output: $m * n$ processor data fractions

$local_s$:

Re-balance the connected subgraph G_L node's fraction distribution to be uniform.

$global_s$:

Collapse the data injection processors into one “big” equivalent processor [15].

Calculate $m * n$ processor's D_i

Obtain the flow matrix A_i .

Calculate $m * n$ processors data fraction α_i .

Calculate the determinant of flow matrix.

Calculate $m * n$ processors' data fraction α_i .

In $local_s$ stage, our target is from a data fraction distribution to a uniform distribution and the minimum cost is called Earth Mover Distance [61].

The time complexity is the same with EPSA.

Model II

We utilize the Reduced Manhattan Distance Voronoi Diagram Algorithm (RMDVDA) and Accelerated Reduced Manhattan Distance Voronoi Diagram Algorithm (ARMDVDA) to address this problem.

Model III

We utilize the same idea of Situation I.

The constraint comes from the Divisible Load Theory linear equations.

$$\min \hat{T}_f \quad (3.117)$$

$$\text{s.t. } \sum_{j=0}^{k-1} \sum_{i=0}^{S_j-1} \hat{\alpha}_{j,i} = 1 \quad (3.118)$$

$$1 \geq \hat{\alpha}_{j,i} \geq 0, \quad (3.119)$$

$$T_{f,i} = \hat{T}_{j,f}, \quad i \in 0 \cdots (S_j - 1) \quad (3.120)$$

$$\hat{\alpha}_{j,0} w T_{cp} = \hat{T}_{j,f}, \quad j \in 0 \cdots (k - 1) \quad (3.121)$$

$$\vdots, \quad (3.122)$$

$$(\hat{\alpha}_{j,1} + \cdots + \hat{\alpha}_{j,\hat{l}_j-2}) z T_{cm} + \hat{\alpha}_{j,\hat{l}_j-1} w T_{cp} = \hat{T}_{j,f}, \quad j \in 0 \cdots (k - 1) \quad (3.123)$$

$$Sp_i = Sp_b + \epsilon, \quad i \in 0 \cdots (k - 1) \quad (3.124)$$

$$\epsilon \geq 0, \quad (3.125)$$

Algorithm 7 Reduced Constrained Manhattan Distance Voronoi Diagram Extended Algorithm (RCMDVDEA)

Input: \hat{k} data injection positions

Output: $m * n$ processor data fractions

local_s :

Collapse the connected data injection processors into some “big” equivalent processors, individually.

Redistribute the data fraction to uniform fraction within each connected subgraph processors.

global_s :

Calculate k constrained Manhattan distance Voronoi cells [21].

Calculate k constrained Voronoi cells’ radius R_i .

Calculate k constrained Voronoi cells’ flow matrix A_i .

Calculate k constrained Voronoi cells’ Sp_i .

Set the $depth_{min} = \min(Sp_i)$ ’s radius.

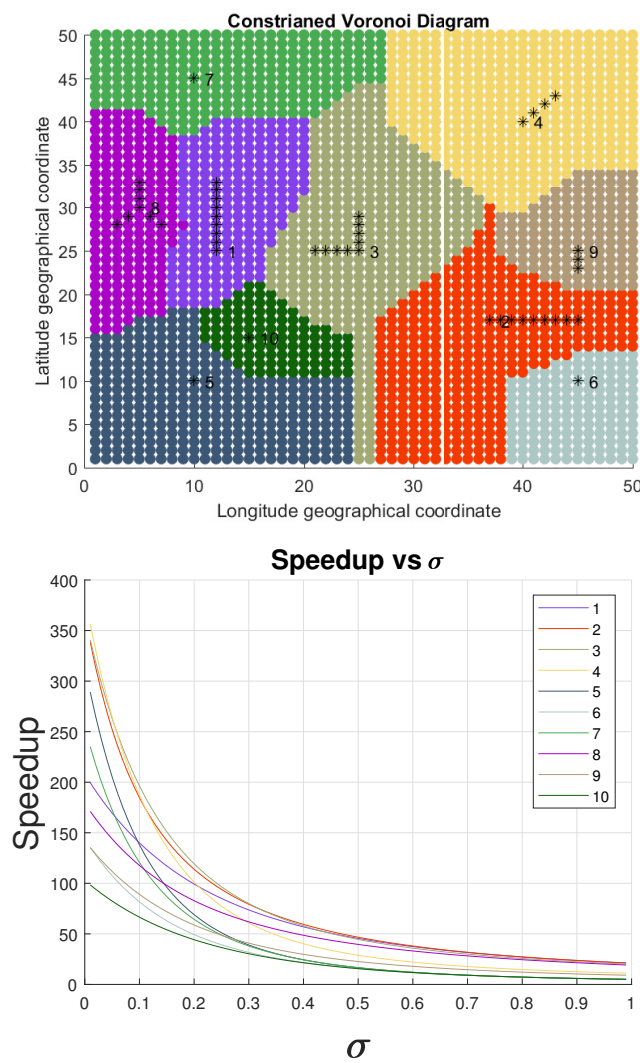
Calculate the reduced constrained Voronoi cells by setting the $depth_i = depth_{min}$ in each Voronoi cell.

Calculate Voronoi cell’s flow matrix \hat{A}_i .

Calculate the determinant of flow matrix.

Calculate $m * n$ processors’ data fraction α_i .

Table 3.4: The 10 Constrained Voronoi Cells and Speedup Curves



Chapter 4

Mesh Network Modified Store and Forward Switching

4.1 Modified Store and Forward Switching Scenario

In the modified store and forward scenario, the processors start to process as soon as each processor receives its entire load assignment [62]. However it is assumed that once a load assignment is received by a node, it can serve as zero delay (virtual cut-through) relay for load to successive processors. Pure store and forward could also be modeled, though it is not discussed in this thesis.

4.1.1 Data Injection on The Corner Processor

2*2 Mesh Network

The timing diagram of Fig. 4.1 is shown:

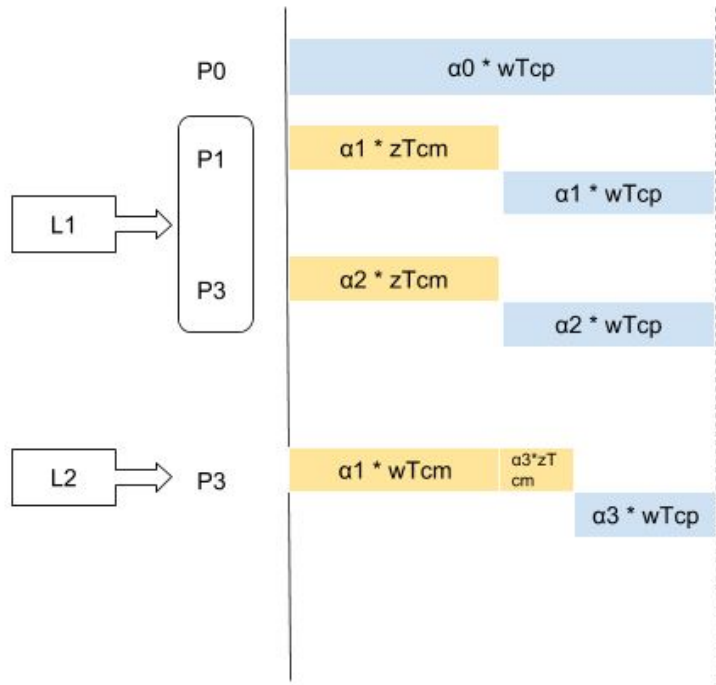


Figure 4.1: The timing diagram for 2*2 mesh network store and forward scenario

The corresponding group of equations are as follows:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_2 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_3) z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ \alpha_0 + \alpha_1 + \alpha_2 + \alpha_3 = 1 \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1, \alpha_2, \alpha_3 < 1 \end{array} \right. \quad \begin{array}{l} (4.1) \\ (4.2) \\ (4.3) \\ (4.4) \\ (4.5) \\ (4.6) \\ (4.7) \\ (4.8) \\ (4.9) \end{array}$$

The matrix closed-form is presented as:

$$\begin{bmatrix} 1 & 2 & 1 \\ 1 & -(\sigma+1) & 0 \\ 1 & -\sigma & -(\sigma+1) \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix} \quad (4.10)$$

The explicit solution is:

$$\alpha_0 = \left(\frac{\sigma+1}{\sigma+2} \right)^2 \quad (4.11)$$

$$\alpha_1 = \frac{\sigma+1}{(\sigma+2)^2} \quad (4.12)$$

$$\alpha_3 = \frac{1}{(\sigma+2)^2} \quad (4.13)$$

The simulation result for Fig. 3.1 is provided in Fig. 4.2:

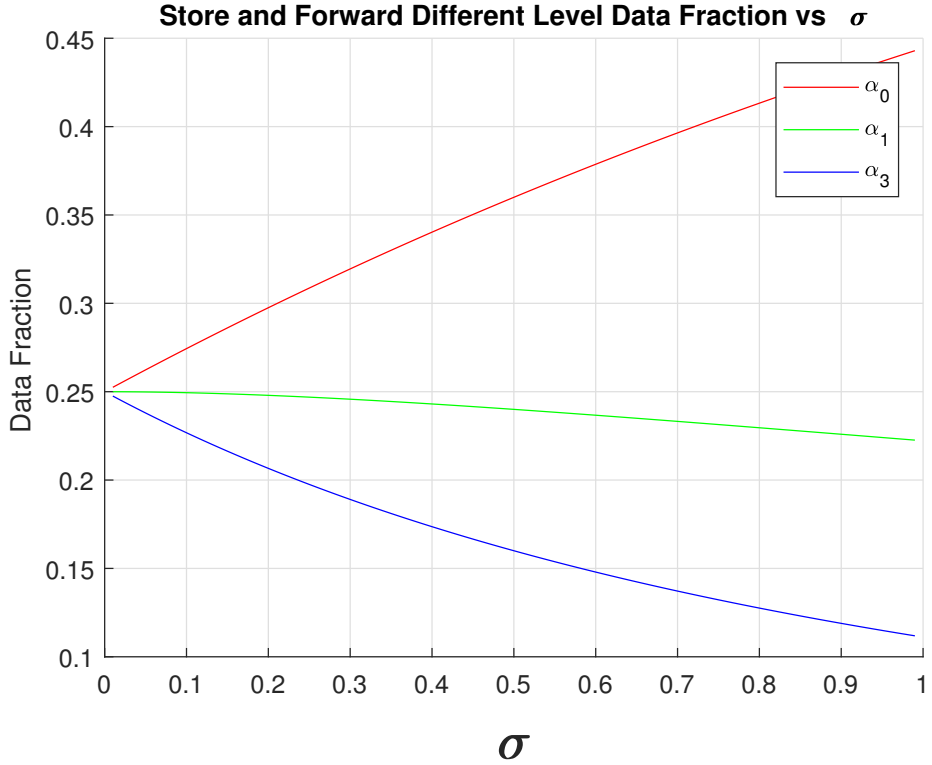


Figure 4.2: Data fraction curves of mesh network $2 * 2$ vs σ

A $2 * 2$ mesh network Fig. 3.1's data fraction curves Fig. 4.2 shows that as the value σ increases, the fraction assigned to P_0 increases, the fractions distributed to $level_1$ and $level_2$ reduces. In other words, if the communication capability decreases, there are more data processed locally, which is reasonable. If the ability of the link decreases, asymptotically equaling to the processor computation capacity, there is only 11% data is deployed to the $level_2$. In addition, if the $\sigma > 1$, it means that the transmitting time is greater than the processor's processing time. In this scenario, keeping the data locally is more economical than transmitting it. If σ is small (fast communication), $\alpha_0 = \alpha_1 = \alpha_2 = \alpha_3$, that is, all four processors are engaged in processing equally.

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

2*3 Mesh Network

P_0 starts to process the assigned workload and it starts to transfer the data. P_1 and P_2 are idle until they get their data fraction from P_0 through the link 0 – 1 and 0 – 2.

According to the $level_2$, the similar situation happens to P_3 and P_4 . They both start to transmit to α_5 after they receive those workload. In other words, P_5 has to wait until the previous two layers processors obtain their own data.

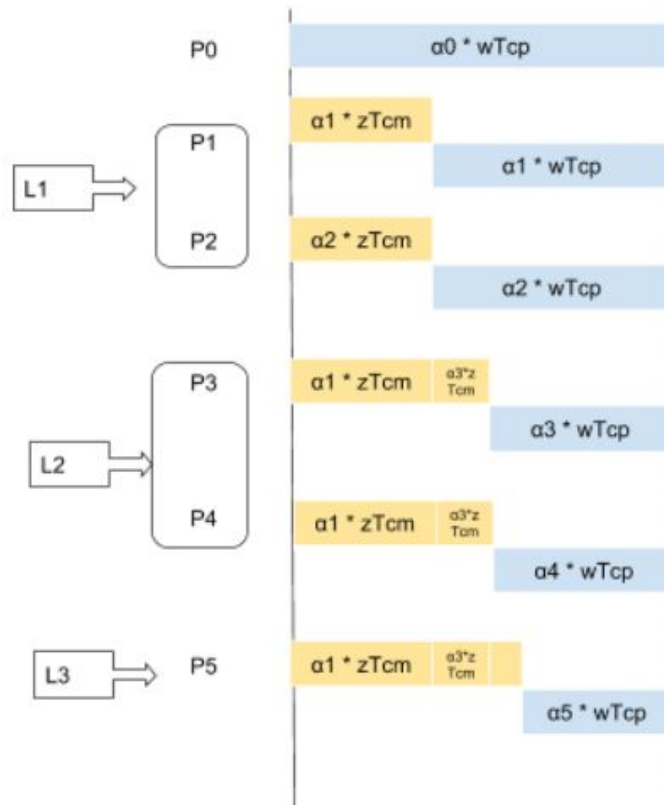


Figure 4.3: The timing diagram for store and forward switching 2*3 mesh network.

In addition, the group of equations are as follows:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_2 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_3) z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_4) z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_3 + \alpha_5) z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ \alpha_0 + \alpha_1 + \alpha_2 + \alpha_3 + \alpha_4 + \alpha_5 = 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1 \quad \alpha_2 \quad \alpha_3 \quad \alpha_4 \quad \alpha_5 < 1 \end{array} \right. \begin{array}{l} (4.14) \\ (4.15) \\ (4.16) \\ (4.17) \\ (4.18) \\ (4.19) \\ (4.20) \\ (4.21) \\ (4.22) \\ (4.23) \end{array}$$

The flow matrix is :

$$\begin{bmatrix} 1 & 2 & 2 & 1 \\ 1 & -(\sigma + 1) & 0 & 0 \\ 1 & -\sigma & -(\sigma + 1) & 0 \\ 1 & -\sigma & -\sigma & -(\sigma + 1) \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_5 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (4.24)$$

The explicit solution is:

$$\alpha_0 = \left(\frac{\sigma + 1}{\sigma + 2} \right)^3 \quad (4.25)$$

$$\alpha_1 = \frac{(\sigma + 1)^2}{(\sigma + 2)^3} \quad (4.26)$$

$$\alpha_3 = \frac{\sigma + 1}{(\sigma + 2)^3} \quad (4.27)$$

$$\alpha_5 = \frac{1}{(\sigma + 2)^3} \quad (4.28)$$

The simulation result is:

If σ is small (fast communication), each of six processor gets $\frac{1}{6}$ of the load.

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

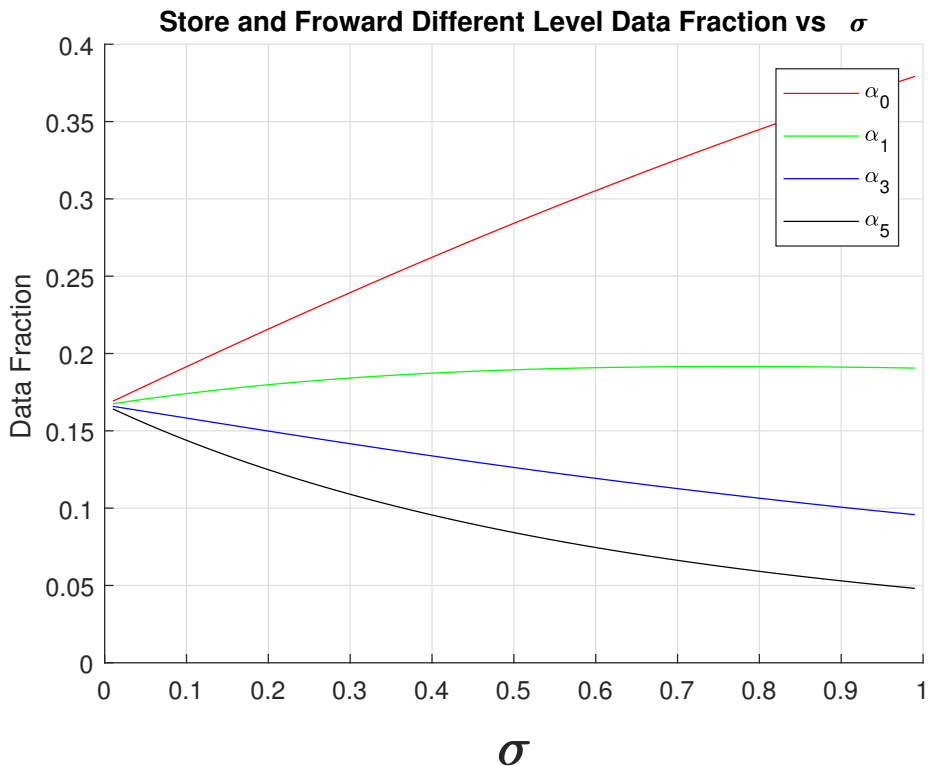


Figure 4.4: Data fraction curves of mesh network $2 * 3$ vs σ

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$\text{Speedup} = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

2*n Mesh Network

Considering a $2 * 10$ mesh network Fig. 3.7, the equations are demonstrated as follows:

$$\left\{
\begin{array}{ll}
\alpha_0 \omega T_{cp} = T_{f,m} & (4.29) \\
\alpha_1 z T_{cm} + \alpha_1 \omega T_{cp} = T_{f,m} & (4.30) \\
\alpha_2 z T_{cm} + \alpha_2 \omega T_{cp} = T_{f,m} & (4.31) \\
(\alpha_1 + \alpha_3) z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} & (4.32) \\
(\alpha_1 + \alpha_4) z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} & (4.33) \\
(\alpha_1 + \alpha_3 + \alpha_5) z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} & (4.34) \\
\vdots & (4.35) \\
(\alpha_1 + \alpha_3 + \cdots + \alpha_{2 \times n - 1}) z T_{cm} + \alpha_{2 \times n - 1} \omega T_{cp} = T_{f,m} & (4.36) \\
\sigma = \frac{z T_{cm}}{\omega T_{cp}} & (4.37) \\
0 < \sigma < 1 & (4.38) \\
0 < \alpha_0 \leq 1 & (4.39) \\
0 \leq \alpha_1 \alpha_3 \cdots \alpha_{2 \times n - 1} < 1 & (4.40)
\end{array}
\right.$$

Utilize σ^* to present $-(\sigma + 1)$ and then the flow matrix form for the group of equations is :

$$\begin{bmatrix} 1 & 2 & 2 & \cdots & 2 & 2 & 1 \\ 1 & \sigma^* & 0 & \cdots & 0 & 0 & 0 \\ 1 & -\sigma & \sigma^* & \cdots & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & \sigma^* & 0 & \cdots & 0 \\ 1 & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \\ 1 & -\sigma & -\sigma & \cdots & -\sigma & -\sigma & \sigma^* \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_5 \\ \vdots \\ \alpha_{2 \times n-3} \\ \alpha_{2 \times n-1} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ \vdots \\ 0 \end{bmatrix} \quad (4.41)$$

According to the **Cramer's rule**, the explicit solution for the group of equations is:

$$\left\{ \alpha_i = \left| \frac{\det A_i^*}{\det A} \right| \right. \quad (4.42)$$

where A_i^* is the matrix formed by replacing the i -th column of A by the column vector b.

We use $-\sigma - 2 = \epsilon$ and $\sigma^* - 2 = \beta$.

After a series of column reduction and row reduction, the flow matrix changes as follows

:

$$A = \left[\begin{array}{ccccccc} 1 & 2 & 2 & \cdots & 2 & 2 & 1 \\ 1 & \sigma^* & 0 & 0 & 0 & 0 & 0 \\ 1 & -\sigma & \sigma^* & 0 & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & \sigma^* & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* \end{array} \right] \xrightarrow{\text{Column Reduction}} \left[\begin{array}{ccccccc} 1 & 0 & 0 & \cdots & 0 & 0 & 0 \\ 1 & \beta & -2 & \cdots & -2 & -2 & -1 \\ 1 & \epsilon & \beta & \cdots & 0 & 0 & 0 \\ 1 & \epsilon & \epsilon & \beta & 0 & \cdots & 0 \\ 1 & \epsilon & \epsilon & \epsilon & \beta & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \ddots \\ 1 & \epsilon & \epsilon & \cdots & \epsilon & \epsilon & \beta \end{array} \right]$$

$$\xrightarrow{\text{Row Reduction}} \left[\begin{array}{ccccccc} 1 & 0 & 0 & \cdots & 0 & 0 & 0 \\ 0 & \beta & -2 & \cdots & -2 & -2 & -1 \\ 0 & \epsilon & \beta & \cdots & 0 & 0 & 0 \\ 0 & \epsilon & \epsilon & \beta & 0 & \cdots & 0 \\ 0 & \epsilon & \epsilon & \epsilon & \beta & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \ddots \\ 0 & \epsilon & \epsilon & \cdots & \epsilon & \epsilon & \beta \end{array} \right]$$

We define

$$C = \left[\begin{array}{cccccc} \sigma^* & 0 & 0 & 0 & 0 & 0 \\ -\sigma & \sigma^* & 0 & 0 & 0 & 0 \\ -\sigma & -\sigma & \sigma^* & 0 & 0 & 0 \\ -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 \\ -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* & 0 \\ -\sigma & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* \end{array} \right]$$

$0 < \sigma < 1$, then $-2 < \sigma^* < -1$, which means C is column linear independent, after column and row reduction.

Further, we define

$$\hat{C} = \begin{bmatrix} \beta & -2 & \cdots & -2 & -2 & -1 \\ \epsilon & \beta & \cdots & 0 & 0 & 0 \\ \epsilon & \epsilon & \beta & 0 & \cdots & 0 \\ \epsilon & \epsilon & \epsilon & \beta & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots \\ \epsilon & \epsilon & \cdots & \epsilon & \epsilon & \beta \end{bmatrix}$$

\hat{C} is also full rank. So the flow matrix A is full rank, that is, $\det A \neq 0$ and $\det A^* \neq 0$.

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = \frac{\det A}{\det A^*} = \left| \frac{\det A}{(\sigma^*)^{n-1}} \right|$$

m*n Mesh Network

Referring to a $5 * 5$ mesh network Fig. 2.1, we utilize σ^* to represent the $-(\sigma + 1)$.

The flow matrix closed-form is:

$$\begin{bmatrix} 1 & 2 & 3 & 4 & 5 & 4 & 3 & 2 & 1 \\ 1 & \sigma^* & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 1 & -\sigma & \sigma^* & 0 & 0 & 0 & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & \sigma^* & 0 & 0 & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* & 0 \\ 1 & -\sigma & \sigma^* \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_3 \\ \alpha_6 \\ \alpha_{10} \\ \alpha_{15} \\ \alpha_{19} \\ \alpha_{22} \\ \alpha_{24} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (4.43)$$

The first row in flow matrix describe the number of cores on each D_i . For example, there is 1 core with 0 hop distance (D_0) with load site L . There are 2 cores with 1 hop distance (D_1) with load site L . There are 3 cores with 2 hops distance (D_2) with load site L , and so on.

The number of rows means the number of different type processor data fraction.

4.1.2 Data Injection on The Boundary Processor

A 3×3 mesh network Fig. 3.10 shows an example of boundary processor P_0 receiving L .

The timing diagram for Fig. 3.10 is Fig. 4.5.

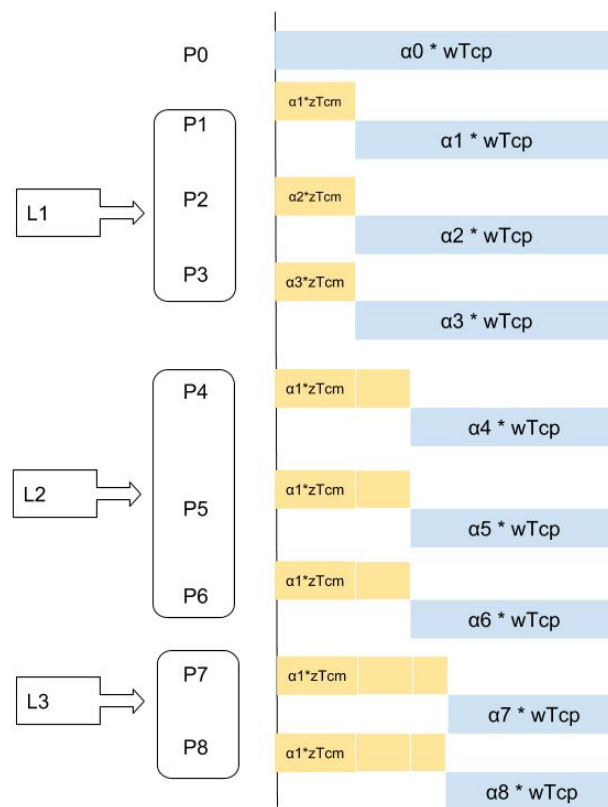


Figure 4.5: The timing diagram for 3×3 boundary data injection on P_0

The equations are:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_3 z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_4) z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} \\ (\alpha_2 + \alpha_5) z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ (\alpha_3 + \alpha_6) z T_{cm} + \alpha_6 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_4 + \alpha_7) z T_{cm} + \alpha_7 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_4 + \alpha_8) z T_{cm} + \alpha_8 \omega T_{cp} = T_{f,m} \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1 \quad \alpha_3 \quad \alpha_4 \quad \alpha_5 \quad \alpha_6 \quad \alpha_7 \quad \alpha_8 < 1 \end{array} \right. \begin{array}{l} (4.44) \\ (4.45) \\ (4.46) \\ (4.47) \\ (4.48) \\ (4.49) \\ (4.50) \\ (4.51) \\ (4.52) \\ (4.53) \\ (4.54) \\ (4.55) \\ (4.56) \end{array}$$

The flow matrix is :

$$\begin{bmatrix} 1 & 3 & 3 & 2 \\ 1 & -(\sigma+1) & 0 & 0 \\ 1 & -\sigma & -(\sigma+1) & 0 \\ 1 & -\sigma & -\sigma & -(\sigma+1) \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_4 \\ \alpha_7 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (4.57)$$

The simulation result is shown in :

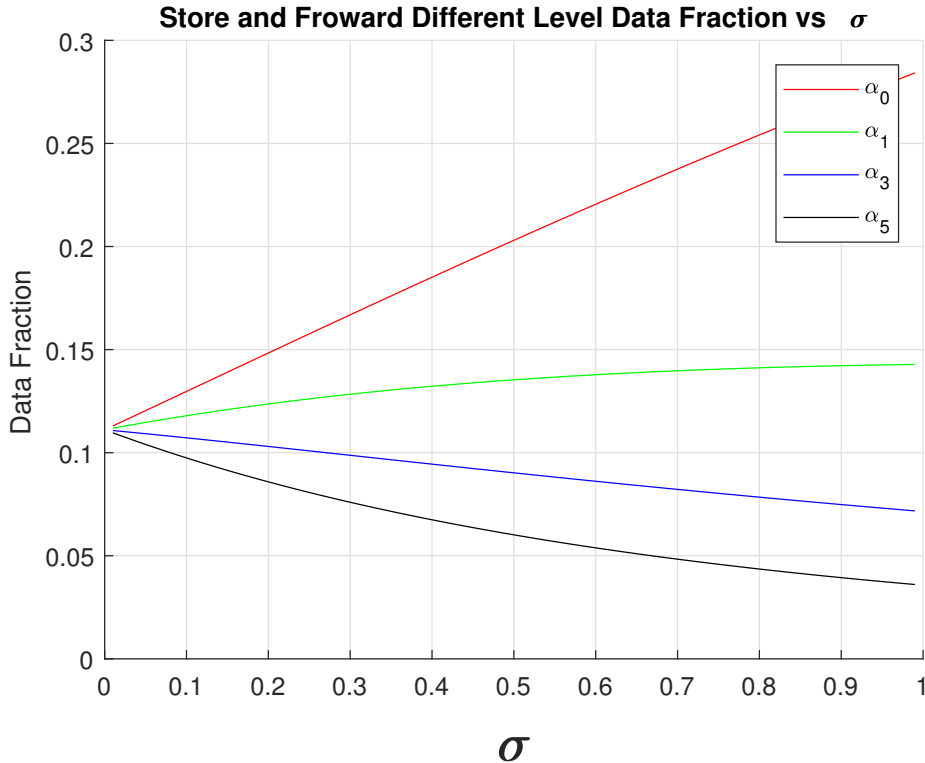


Figure 4.6: The fraction curve for 3*3 boundary data injection on P_0

A 3*3 mesh network appears in Fig. 3.10 and its data fraction curves Fig. 4.6 shows that as the value σ increases, the fraction assigned to P_0 increases and the fractions distributed to $level_2$ and $level_3$ decreases. In other words, if the communication capability decreases, there are more data processed locally, which is reasonable.

If the ability of the link degrades, asymptotically equaling to the processor computation capacity, there is solely 4% data is deployed to the $level_3$. In addition, if the $\sigma > 1$, it means that the transmitting time is greater than the processor's processing time. In this scenario, keeping the data locally is more economical than transmitting it.

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

4.1.3 Data Injection on The Inner Grid Processor

The mesh network considered is $3 * 3$.

The equations are:

$$\left\{ \begin{array}{l} \alpha_0 \omega T_{cp} = T_{f,m} \\ \alpha_1 z T_{cm} + \alpha_1 \omega T_{cp} = T_{f,m} \\ \alpha_2 z T_{cm} + \alpha_2 \omega T_{cp} = T_{f,m} \\ \alpha_3 z T_{cm} + \alpha_3 \omega T_{cp} = T_{f,m} \\ \alpha_4 z T_{cm} + \alpha_4 \omega T_{cp} = T_{f,m} \\ (\alpha_1 + \alpha_5) z T_{cm} + \alpha_5 \omega T_{cp} = T_{f,m} \\ (\alpha_2 + \alpha_6) z T_{cm} + \alpha_6 \omega T_{cp} = T_{f,m} \\ (\alpha_3 + \alpha_7) z T_{cm} + \alpha_7 \omega T_{cp} = T_{f,m} \\ (\alpha_4 + \alpha_8) z T_{cm} + \alpha_8 \omega T_{cp} = T_{f,m} \\ \sigma = \frac{z T_{cm}}{\omega T_{cp}} \\ 0 < \sigma < 1 \\ 0 < \alpha_0 \leq 1 \\ 0 \leq \alpha_1 \quad \alpha_3 \quad \alpha_4 \quad \alpha_5 \quad \alpha_6 \quad \alpha_7 \quad \alpha_8 < 1 \end{array} \right. \begin{array}{l} (4.58) \\ (4.59) \\ (4.60) \\ (4.61) \\ (4.62) \\ (4.63) \\ (4.64) \\ (4.65) \\ (4.66) \\ (4.67) \\ (4.68) \\ (4.69) \\ (4.70) \end{array}$$

The flow matrix closed-form is:

$$\begin{bmatrix} 1 & 4 & 4 \\ 1 & -(\sigma + 1) & 0 \\ 1 & -\sigma & -(\sigma + 1) \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_5 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix} \quad (4.71)$$

The simulation result shows:

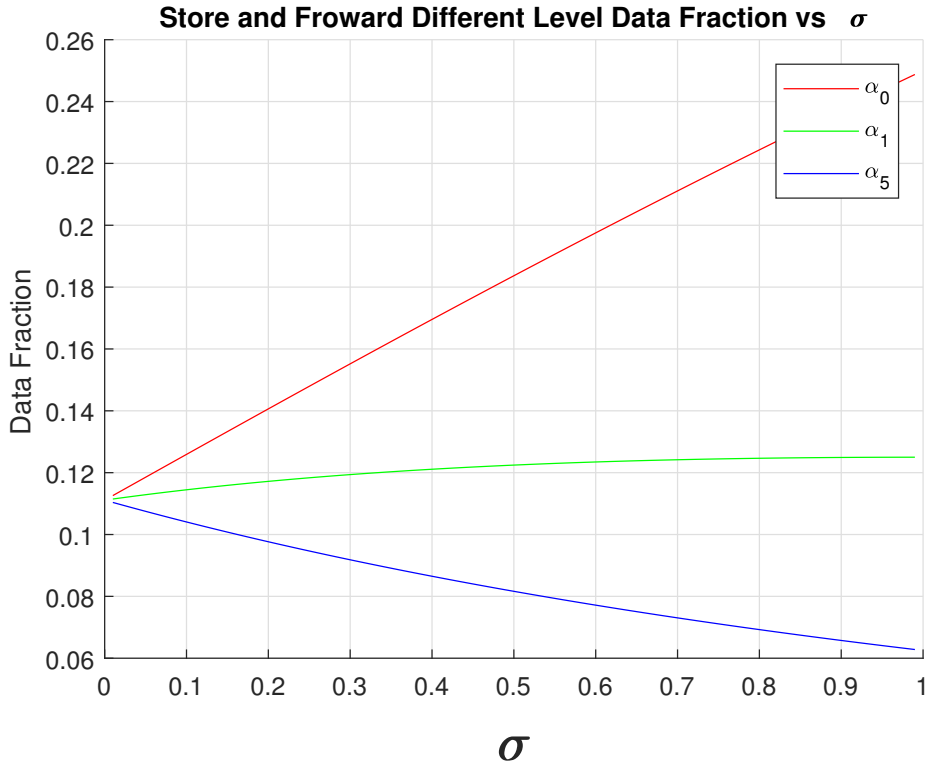


Figure 4.7: The timing diagram for 3×3 inner grid injection P_0

A 3×3 mesh network appears in Fig. 3.13 and its data fraction curves Fig. 4.7 illustrates that as the value σ increases, the fraction assigned to P_0 increases and the fractions distributed to $level_2$ reduces. In other words, if the communication capability decreases, there are more data processed locally, which is reasonable.

If the ability of the link degrades, asymptotically equaling to the processor computation capacity, there is only 6% data is deployed to the $level_2$.

If σ is very small (fast communication), each of nine processors process $\frac{1}{9}$ of the load

4.1.4 Sensitivity Analysis Modified Store and Forward Scenario

Data Injection on The Corner Processor

The simulation result of sensitivity analysis of $2 * n$ mesh network Fig. 3.7 is as follows:

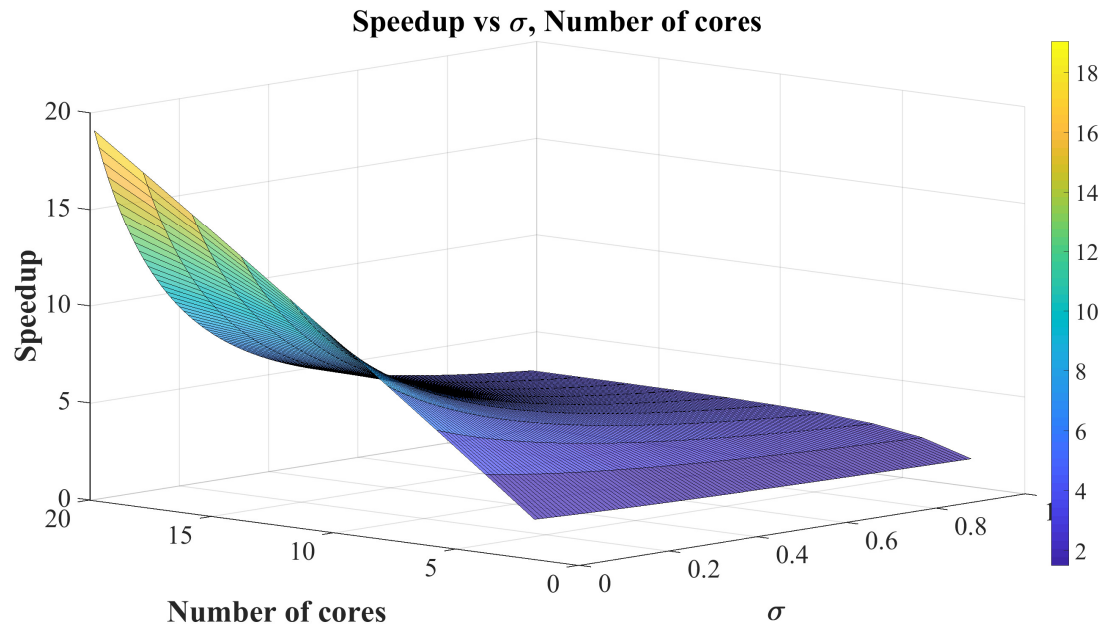


Figure 4.8: Sensitivity analysis result of $2 * n$ mesh network result ($n = 1 \dots 10$)

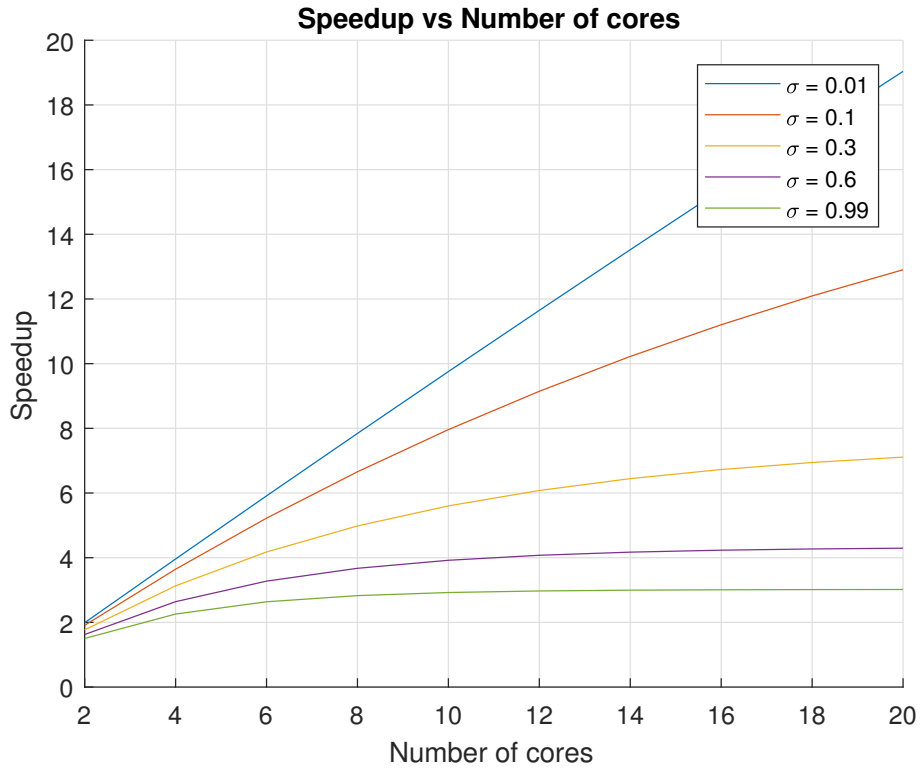


Figure 4.9: Speedup curve of 2^*n mesh network result ($n = 1 \dots 10$)

The figure illustrates that if $\sigma < 0.1$, the number of cores increases, the speedup efficiency is linearly increasing. Alternatively speaking, if $\sigma < 0.1$, the core's number is the most important factor for efficiency. If the $\sigma > 0.3$, the efficiency drops dramatically. That is, the σ value plays more critical role in the speedup simulation.

If σ is close to one (slow communication) and the number of cores is large, then speedup is three times since P_0 and P_0 two neighbors do almost all the processing.

Data Injection on The Boundary Processor

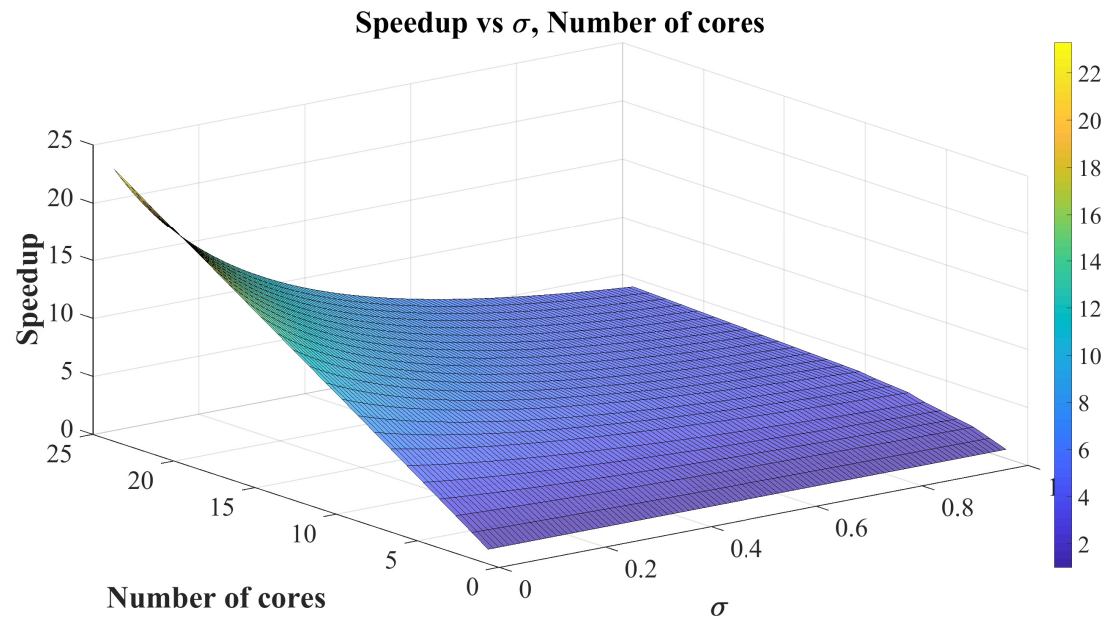


Figure 4.10: Sensitivity analysis result of $3*n$ mesh network result ($n = 1 \cdots 8$)

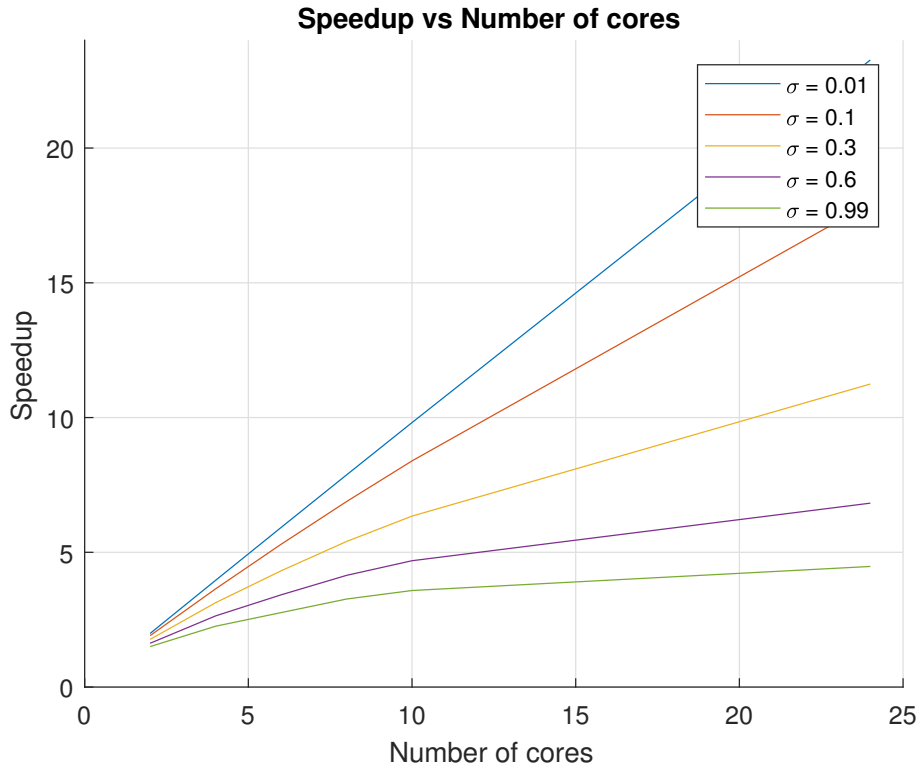


Figure 4.11: Speedup curve of $3*n$ mesh network result ($n = 1 \cdots 8$)

Sensitivity analysis result Fig. 4.10 and speedup curve Fig. 4.11 show the speedup efficiency for the data injection on the boundary scenario.

The figure illustrates that if $\sigma < 0.1$ and the number of cores increases, the speedup efficiency is linearly increasing. The maximum speedup is 24, the number of the cores in a $3 * 8$ mesh network. Alternatively speaking, if $\sigma < 0.1$, the number of cores is the most important factor for improving speedup. If the $\sigma > 0.3$, the speedup drops dramatically. That is, the σ value plays more critical role in the speedup simulation.

Data Injection on The Inner Grid Processor

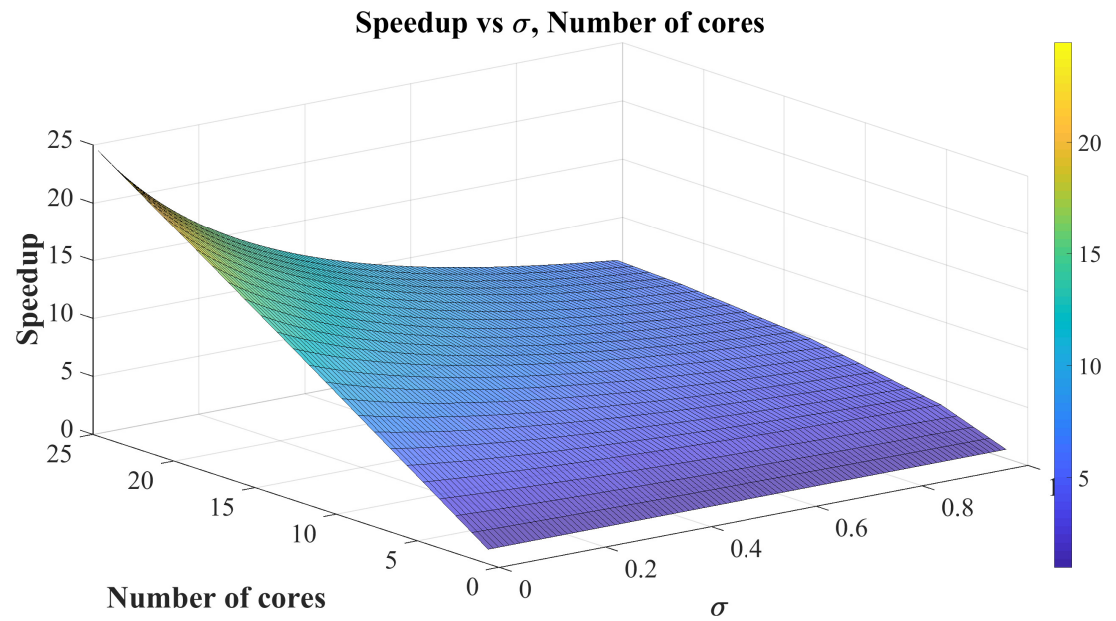


Figure 4.12: Sensitivity analysis result of data injection position on inner grid processor of $5 * 5$ mesh network

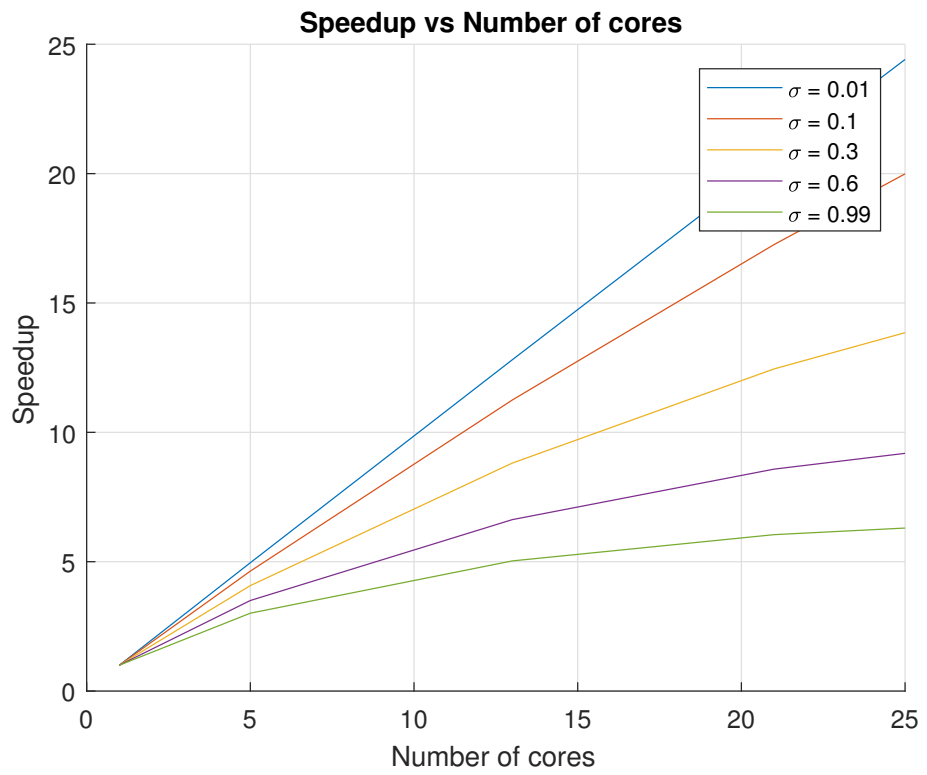


Figure 4.13: Speedup curve of $5 * n$ mesh network result ($n = 1 \dots 5$)

Sensitivity analysis Fig. 4.12 and speedup curves Fig. 4.13 display the simulation result for the data injection inner position P_{12} . If $\sigma < 0.1$, the speedup linear increases and the maximum speedup is 24.

4.1.5 Multi-source Uniform Data Fraction

We utilize the similar methodology and algorithms and in virtual cut-through situation to deal with these problems. The critical point is that the flow matrix formula is different, which is replaced by the modified store and forward flow matrix formula.

Model I

We utilize the Equivalence Processor Scheduling Algorithm to address this problem.

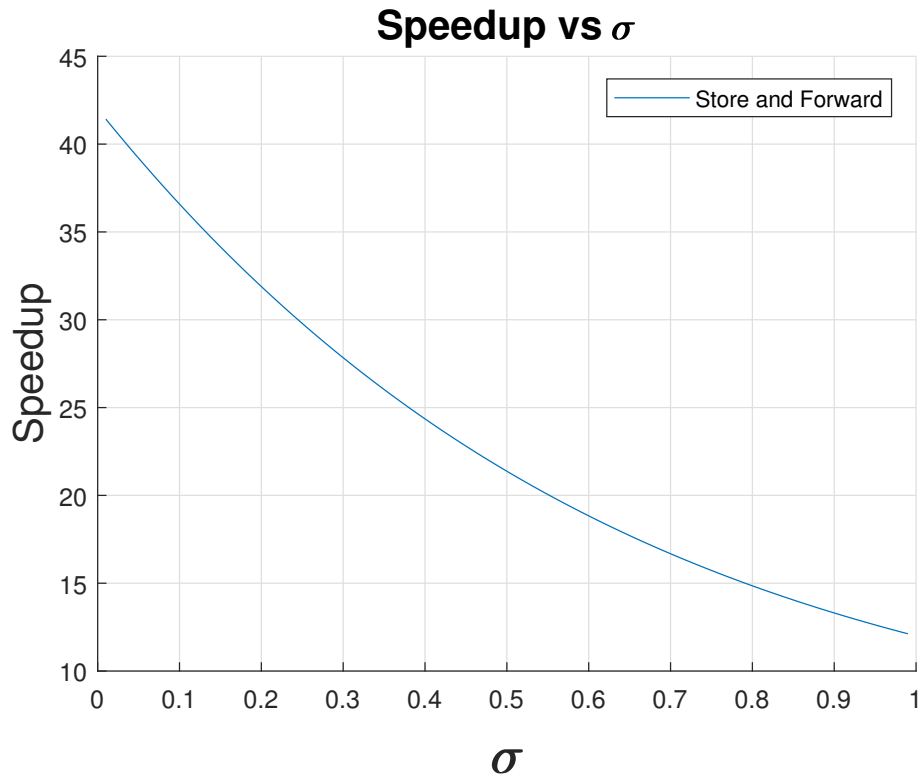


Figure 4.14: Connected subgraph Fig. 3.22 speedup vs σ

For fast communication ($\sigma \approx 0$), the maximum speedup is 42.

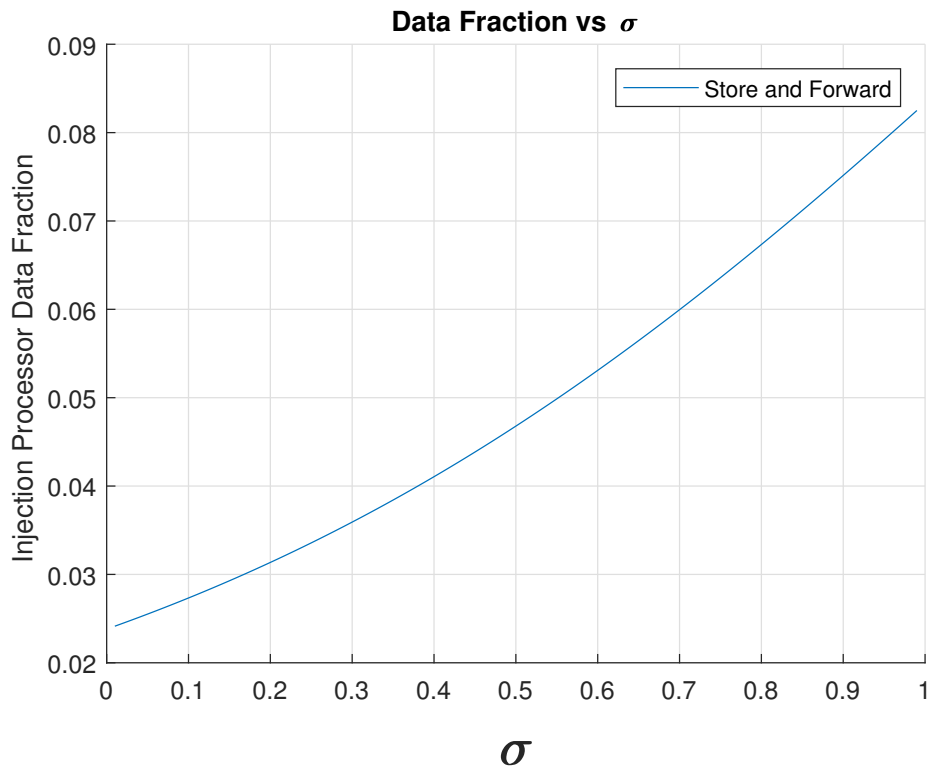


Figure 4.15: Connected subgraph Fig. 3.22's data injection processor workload fraction vs σ

For fast communication ($\sigma \approx 0$), each of 42 processors in Fig. 3.22 gets $\frac{1}{42}$ of the load since they are all equals engaged in processing.

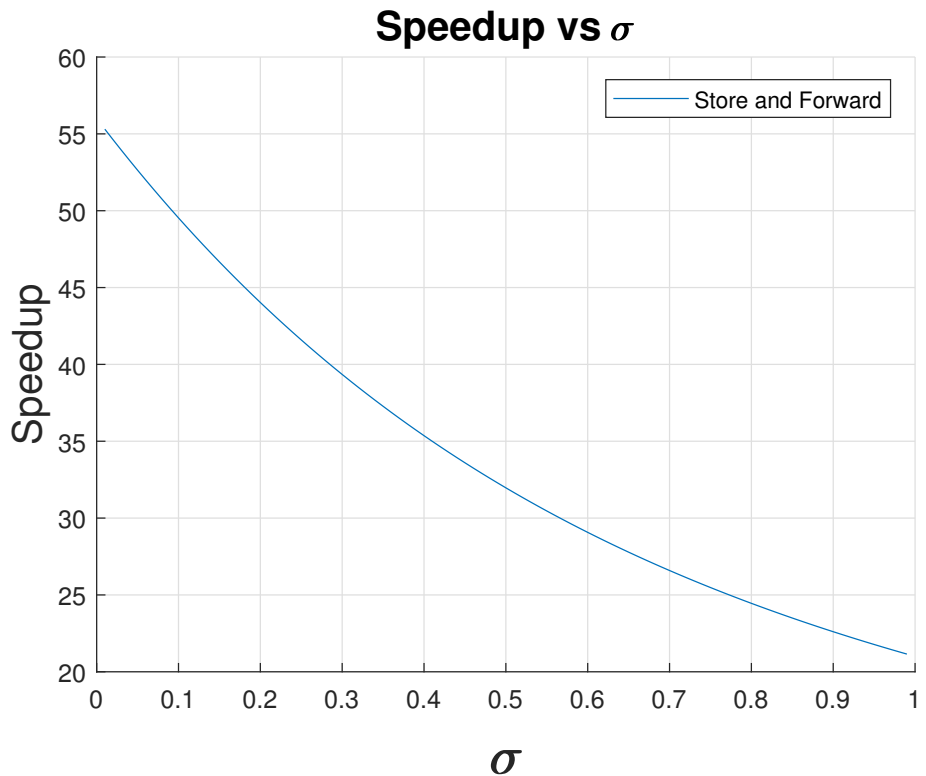


Figure 4.16: Connected subgraph Fig. 3.23 speedup vs σ

For fast communication $\sigma \approx 0$, the maximum speedup is 56 (there are $7 * 8 = 56$ processors in Fig. 3.23).

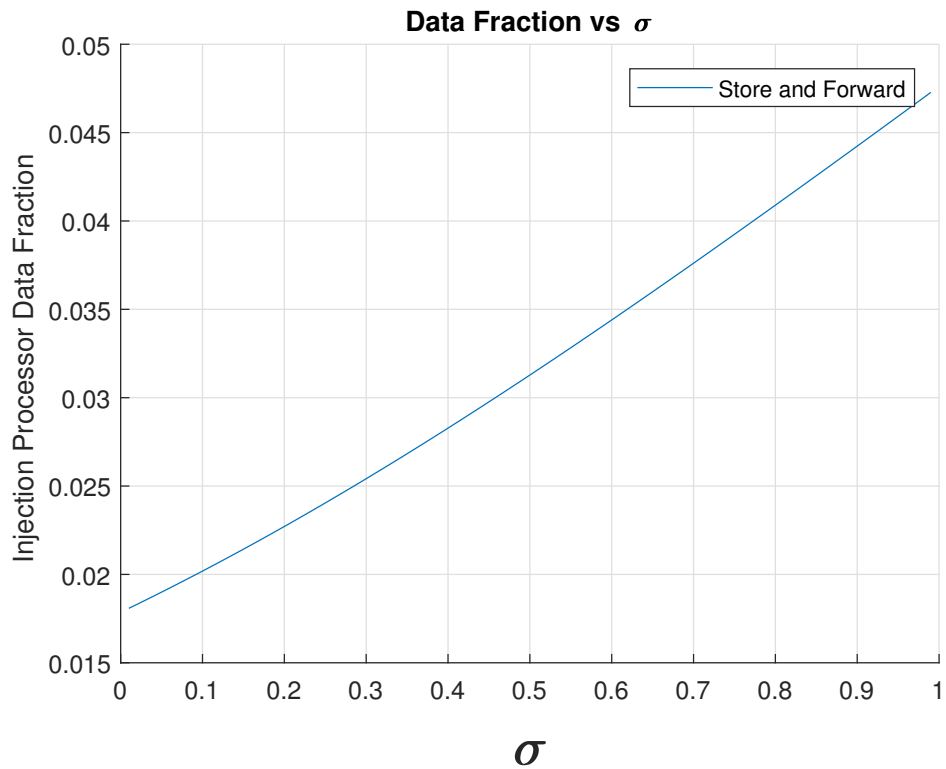


Figure 4.17: Connected subgraph Fig. 3.23's data injection processor workload fraction vs σ

For fast communication ($\sigma \approx 0$), each of 56 processors in Fig. 3.23 gets $\frac{1}{56}$ of the load.

Model II

We utilize the Reduced Voronoi Diagram Algorithm to address this problem. The simulation result is as follows :

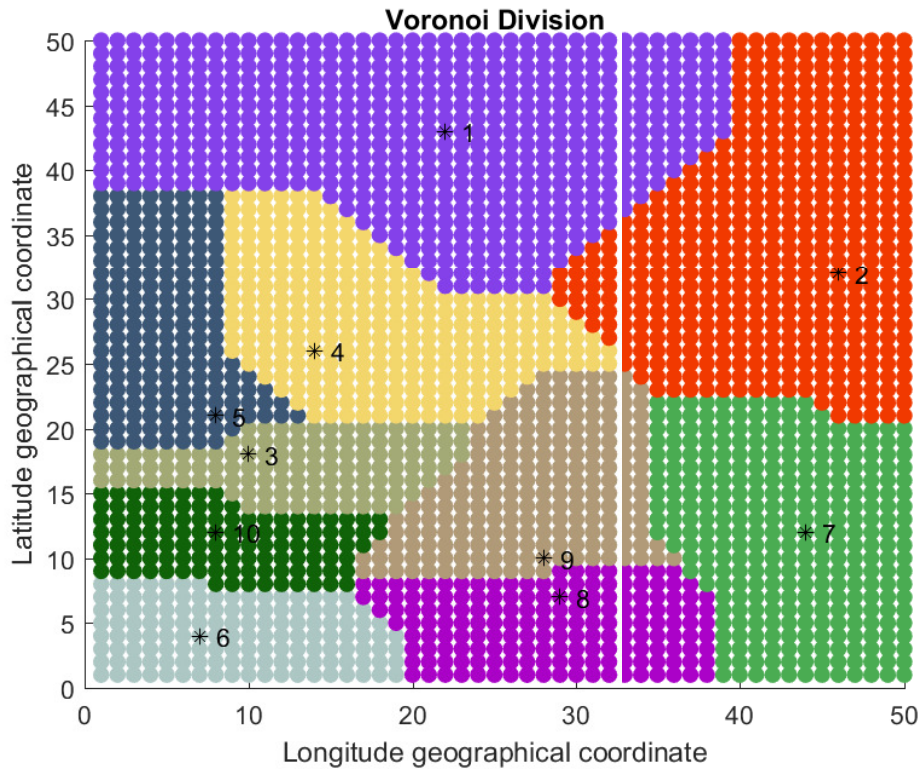


Figure 4.18: 10 Manhattan distance Voronoi Cells

Fig. 4.18 shows an intuitive algorithm to minimizing the makespan. There are 10 bases and the largest cell contains about 495 cores. There are about 107 cores in the smallest cell. So the ratio is $\frac{495}{107} \approx 4.62$.

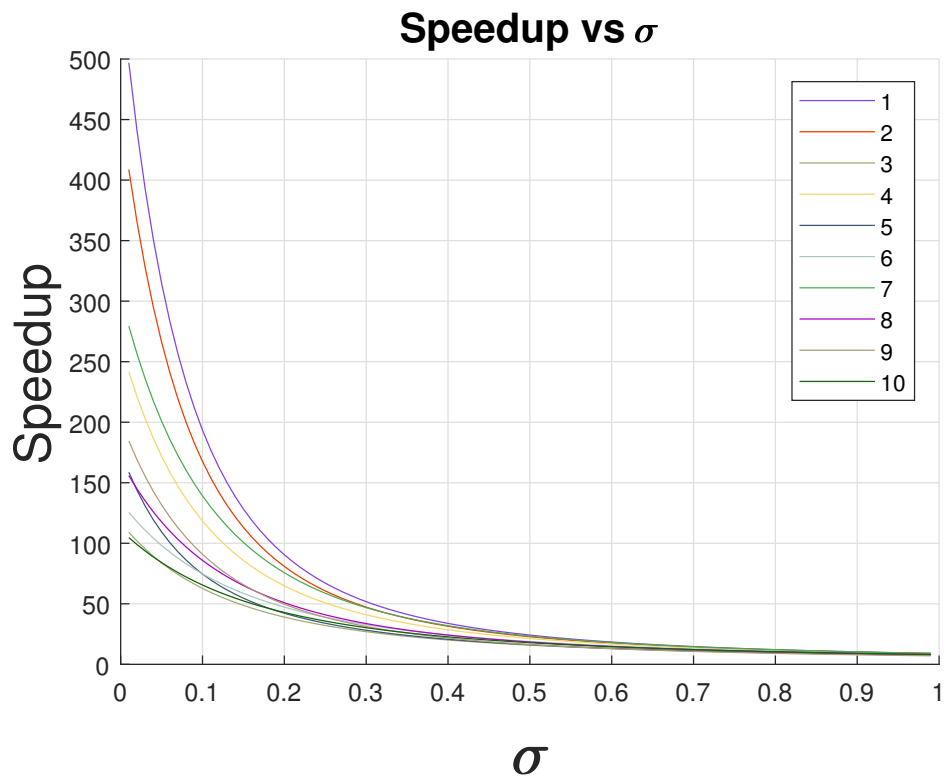


Figure 4.19: 10 Manhattan distance Voronoi cells modified store and forward switching speedup curves

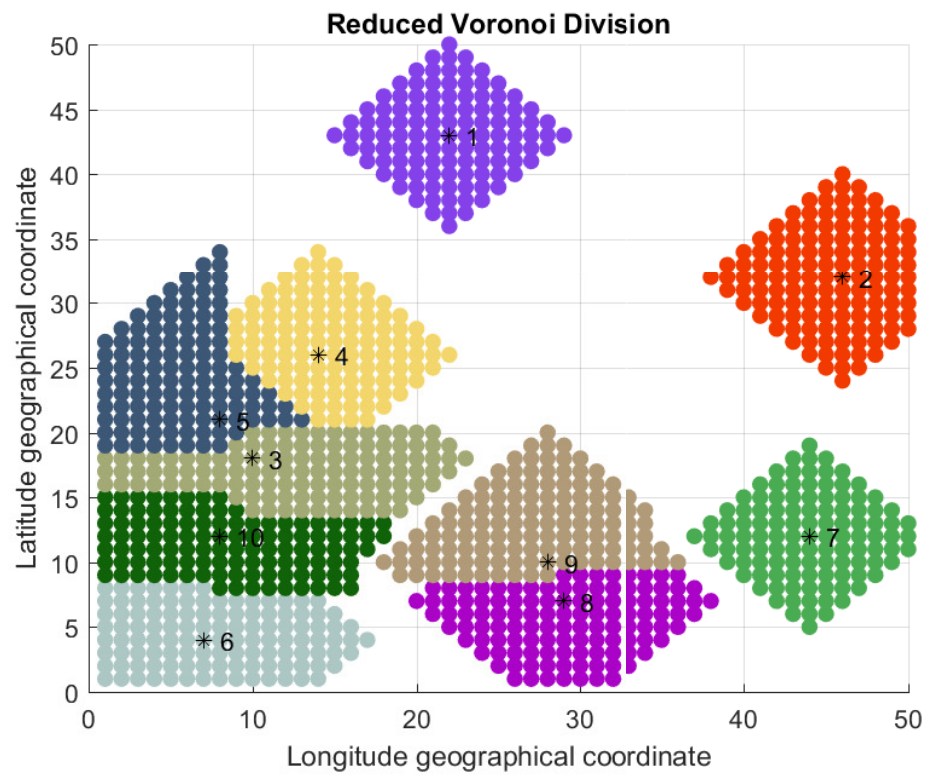


Figure 4.20: 10 reduced Manhattan distance modified store and forward switching Voronoi cells

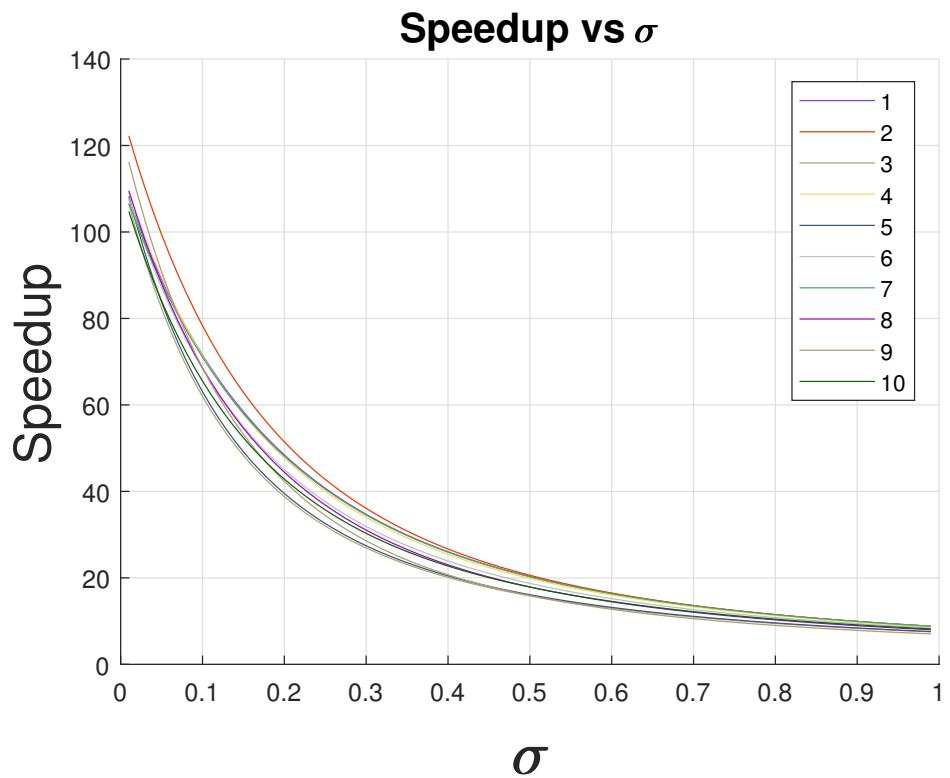


Figure 4.21: 10 reduced Voronoi cells' store and forward switching speedup curves

We save about 53.4% processors and finish the task within the same time.

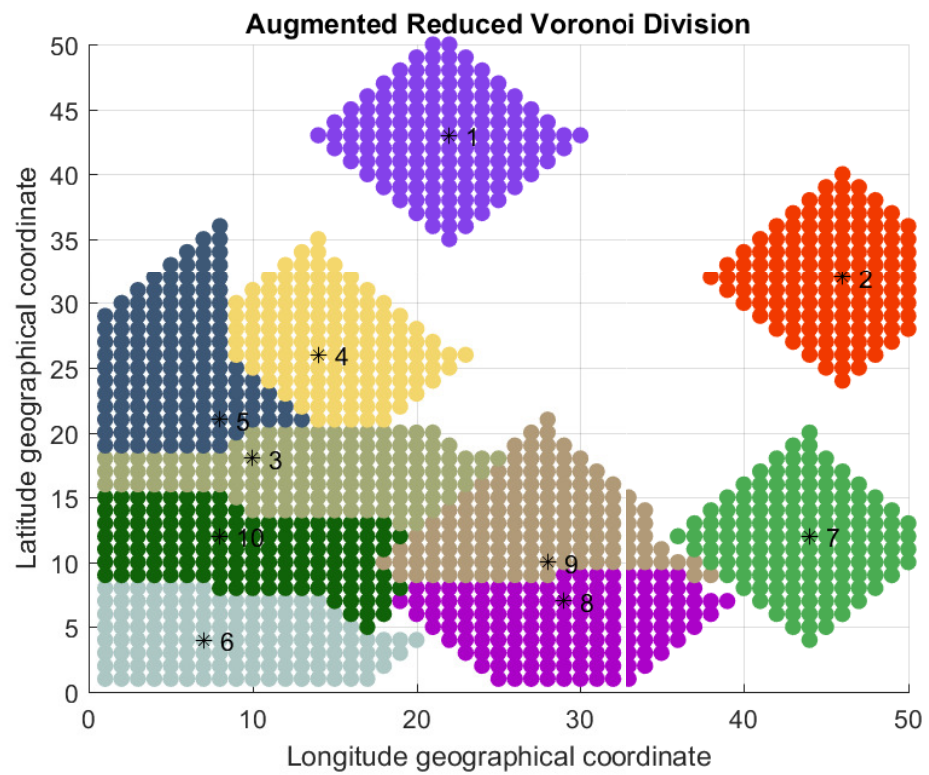


Figure 4.22: 10 reduced Manhattan distance modified store and forward switching Voronoi cells

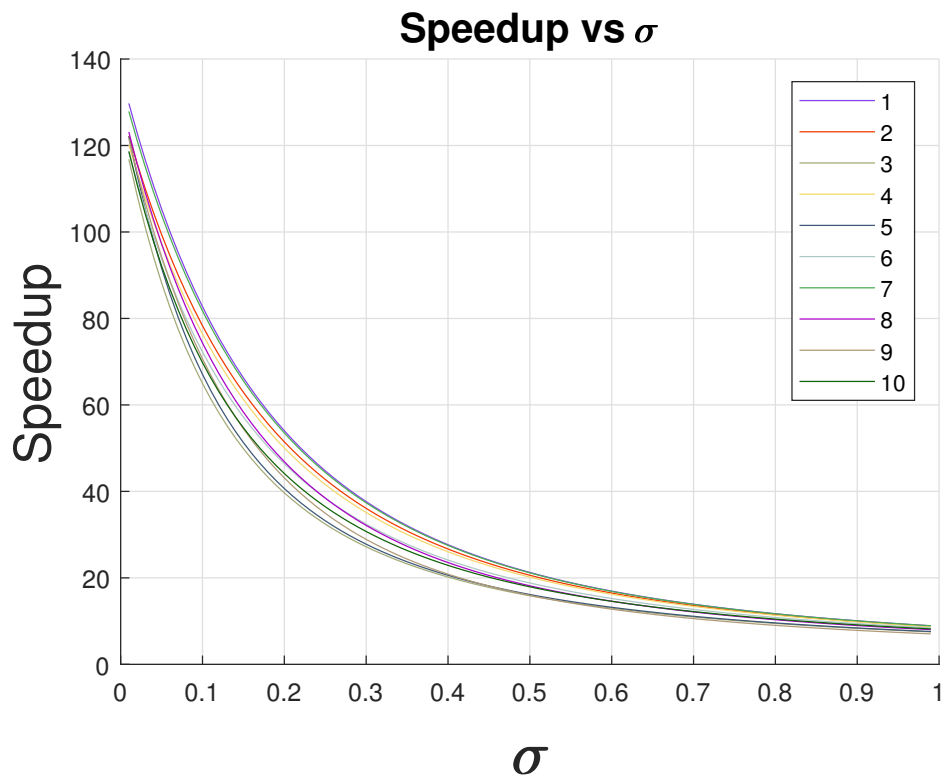


Figure 4.23: 10 reduced Voronoi cells' store and forward switching speedup curves

Fig. 4.22 and Fig. 4.23 show ARMDVDA simulation result and the new heuristic algorithm accelerates about 10% speedup.

Model III

We utilize the Reduced Constrained Manhattan Distance Voronoi Diagram Extended Algorithm to address this problem.

4.1.6 Multi-source Non-uniform Data Injection

We utilize the similar methodology and algorithms in virtual cut-through situation to deal with these problem.

Chapter 5

Toroidal Rectangle Network

5.1 Toroidal Rectangle Network Notions

The toroidal network is a regular complex in that each polygonal face has the same number of links and each node is connected to the same number of links. According to the paper [1], there are three different torus mesh networks. In this paper, our intent is not to propose one model to “fit all” problems but rather to indicate one normal case.

Considering the toroidal rectangle network Fig. 5.1 and Fig. 5.2.

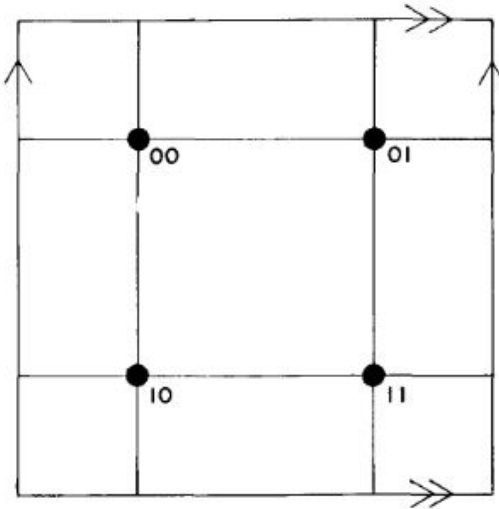


Figure 5.1: The rectangular toroidal network. Figure citing from [1]

- m : There are m processors on the longitude.

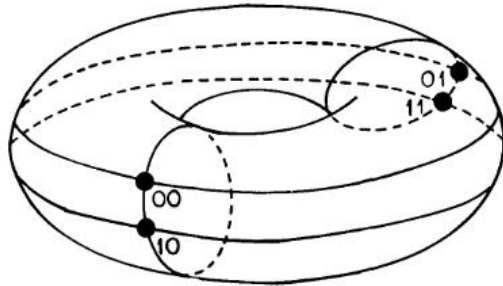


Figure 5.2: The rectangular toroidal network. Figure citing from [1]

- n : There are n processors on the latitude.
- L : The load injection site.
- L_x : The L 's longitude coordinate.
- L_y : The L 's latitude coordinate.
- D_k : P_k 's shortest Manhattan distance to L .
- $D_{k,x}$: P_k 's longitude shortest Manhattan distance to L .
- $D_{k,y}$: P_k 's latitude shortest Manhattan distance to L .

$$\left\{ \begin{array}{l} D_k = D_{k,x} + D_{k,y} \\ D_{k,x} = \min\{\|D_{k,x} - L_x\|, m - \|D_{k,x} - L_x\|\} \end{array} \right. \quad (5.1) \quad (5.2)$$

$$D_{k,y} = \min\{\|D_{k,y} - L_y\|, n - \|D_{k,y} - L_y\|\} \quad (5.3)$$

5.2 Virtual Cut-through Switching Scenario

5.2.1 Data Injection On The Grid Processor

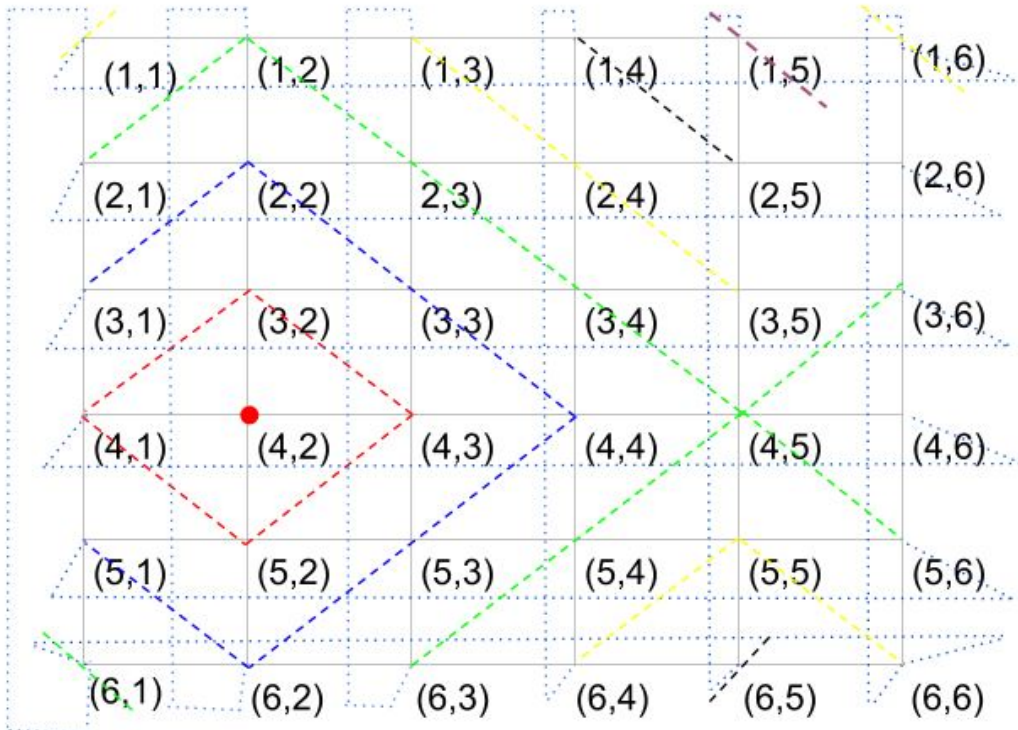


Figure 5.3: The $m*n$ toroidal rectangle network and the data injection is $P_{4,2}$

In $m * n$ ($m = 6, n = 6$) Fig. 5.3 toroidal rectangle network, L happens on grid position $(4, 2)$. We calculate the $D_{k,i}$ table Table 5.1 by breadth first search(**BFS**) algorithm.

D_i	Number
0	1
1	4
2	8
3	10
4	8
5	4
6	1

Table 5.1: D_i vs Number

The flow matrix closed-form is :

$$\begin{bmatrix} 1 & 4 & 8 & 10 & 8 & 4 & 1 \\ 1 & -1 & 0 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & 1 & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & \sigma & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_2 \\ \alpha_3 \\ \alpha_4 \\ \alpha_5 \\ \alpha_6 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (5.4)$$

The simulation result is :

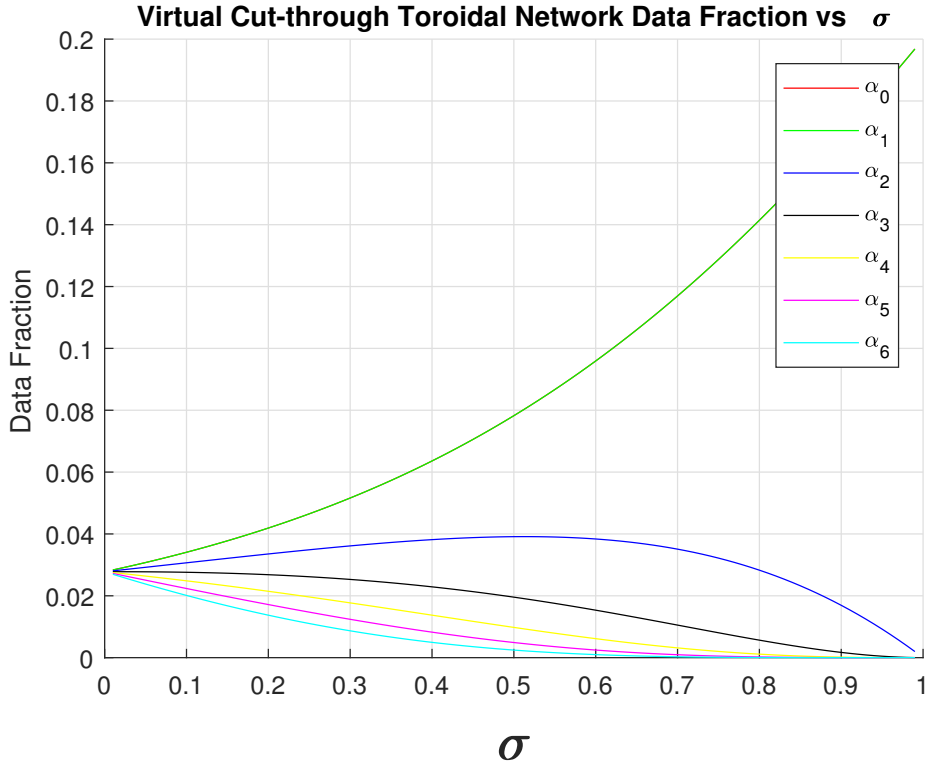


Figure 5.4: Data fraction curve of toroidal rectangle network Fig. 5.3

Here for fast communication ($\sigma \approx 0$), each of 36 processors gets $\frac{1}{36}$ of the load as all 36 processors receive a uniform amount of load. If communication is slow, each of P_0 and four P_0 neighbors receive $\frac{1}{5}$ of the load (i.e. 0.2 of the load). The load sharing by these five processors is economical.

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0}$$

D_i	Number
0	1
1	4
2	8
3	8
4	4

Table 5.2: D_i vs Number of processor on each D_i

5.2.2 Sensitivity Analysis of Toroidal Rectangle Network

Considering a $5 * 5$ toroidal rectangle network, the $level_i$ table shows Table 5.2.

So the simulation result illustrates in Fig. 5.5

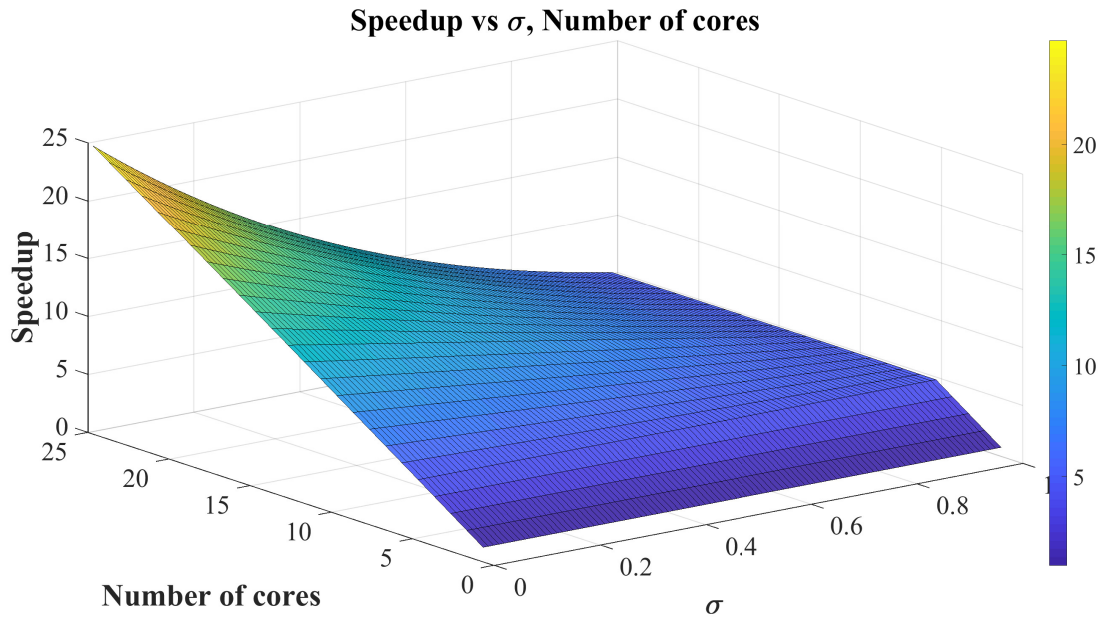


Figure 5.5: Sensitivity analysis of $5*n$ torus network (where $n = 1 \cdots 5$)

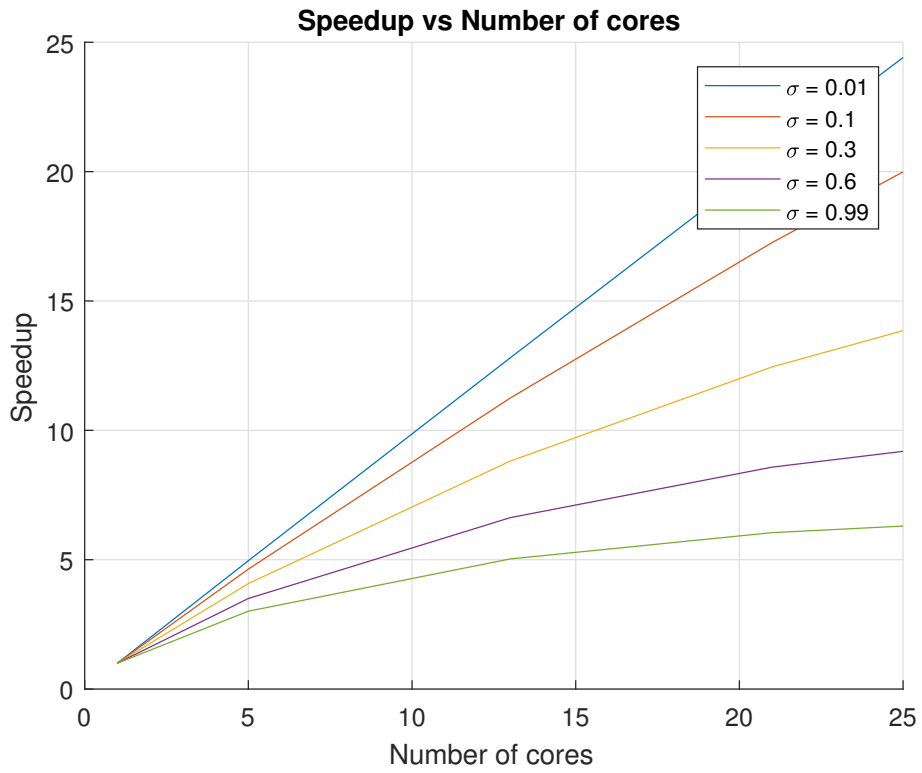


Figure 5.6: Data fraction curves of a $5 \times n$ torus network (where $n = 1 \dots 5$) vs σ

Fig. 5.6 illustrates that as the value σ decreases, the speedup efficiency increases. For fast communication ($\sigma = 0.01$), the speedup is almost 25, as all 25 processors are uniformly engaged in processing. For slow communication ($\sigma = 0.99$), the speedup is about 5 as there are P_0 and its four neighbors are engaged into the processing load.

5.2.3 Multi-source Uniform Data Fraction

We extend the single source assignment to multi-source assignment problem [55] [16]. According to each processor, we concentrate on the processors' geographical location P_i , data fraction assigned α_i .

Assume the data fraction is uniformly distributed. For example, the workload is unit 1 and there are k different data injection options. So each data injection is assigned $\frac{1}{k}$ workload.

From the data injection position relationship we consider three different situations :

- Data injection positions consist of a subgraph G_L of G .
- Data injection processor doesn't connect with each other.
- Data injection positions consist of some connected subgraphs and individual injection.

Model I

We choose the Equivalence Processor Scheduling Algorithm to address the problem.

Torus connected subgraph data injection Fig. 5.7 Fig. 5.9 show a connected subgraph data injection situation.

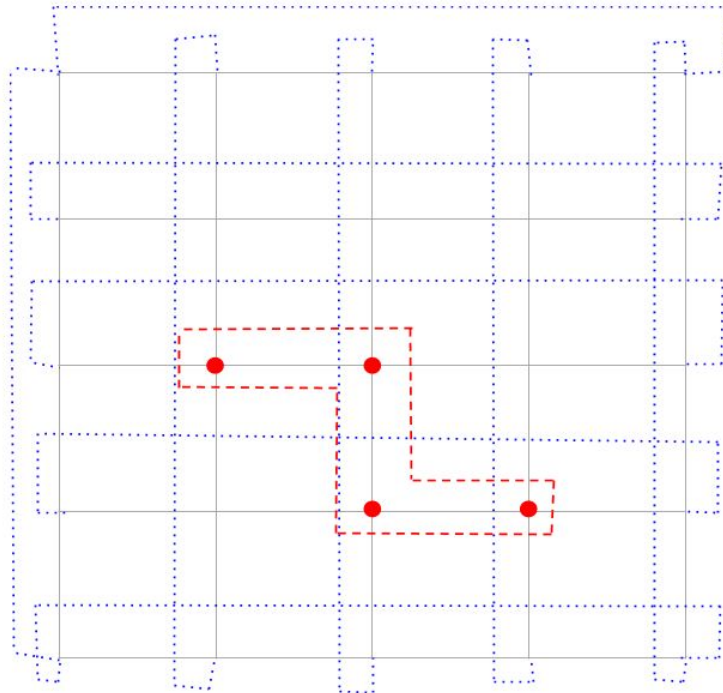


Figure 5.7: Toroidal mesh network subgraph data injection

A connected subgraph Fig. 5.7's flow matrix equation is

$$\begin{bmatrix} 4 & 8 & 9 & 4 \\ 1 & -1 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (5.5)$$

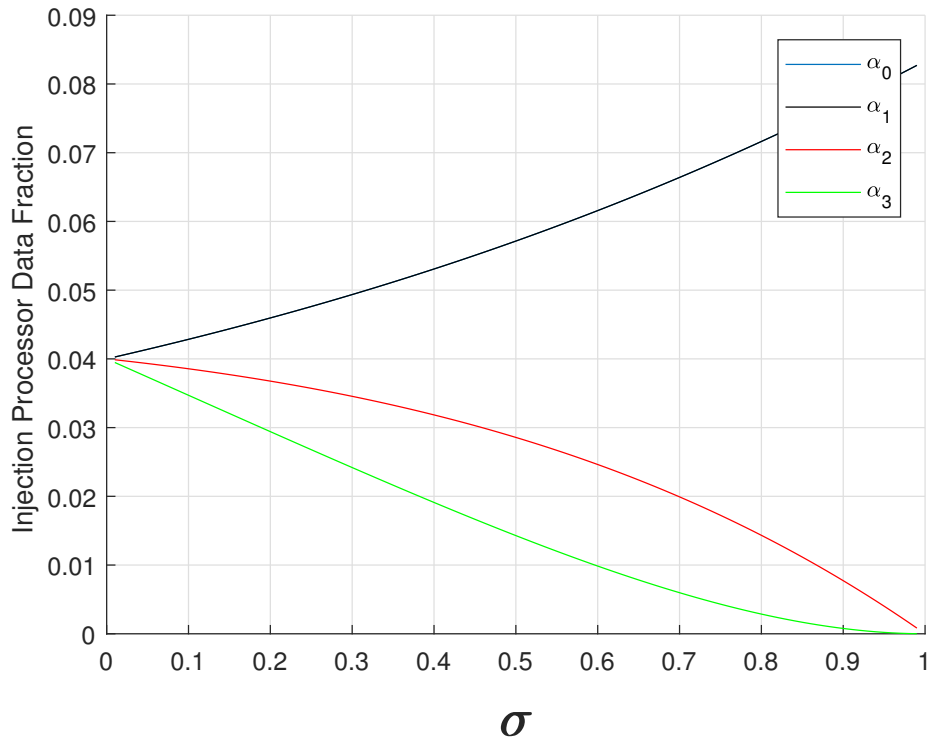


Figure 5.8: Data fraction curves of connected subgraph Fig. 5.7

The load fraction are 0.04 a small σ (fast communication) as all 25 processors are uniformly engaged processing. For slow communication, each of 12 processors (4 in the cluster and 8 in the adjacent neighbors) receives $\frac{1}{12}$ of the load as it is not economical to distribute load beyond those 12 processors.

Fig. 5.9 shows another connected subgraph data injection example.

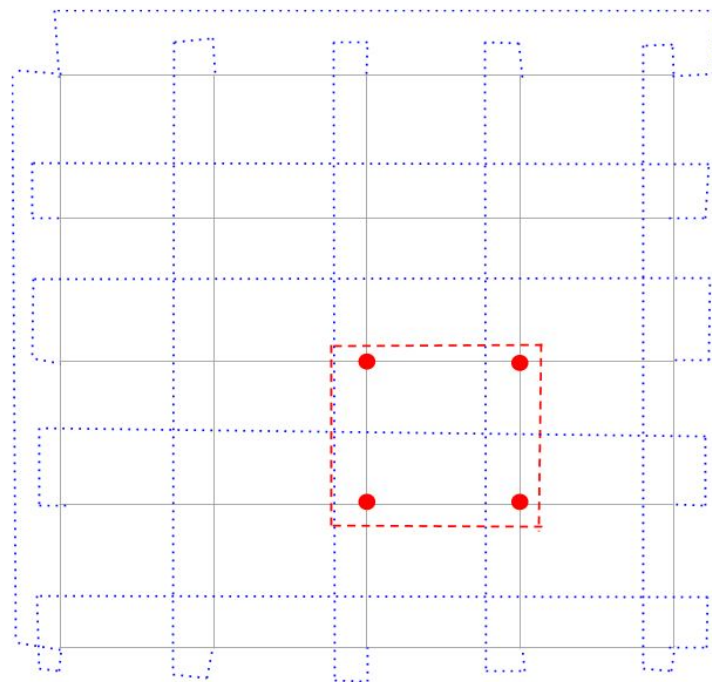


Figure 5.9: Toroidal mesh network subgraph data injection

Fig. 5.9's flow matrix equation is :

$$\begin{bmatrix} 4 & 8 & 9 & 4 \\ 1 & -1 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_2 \\ \alpha_3 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (5.6)$$

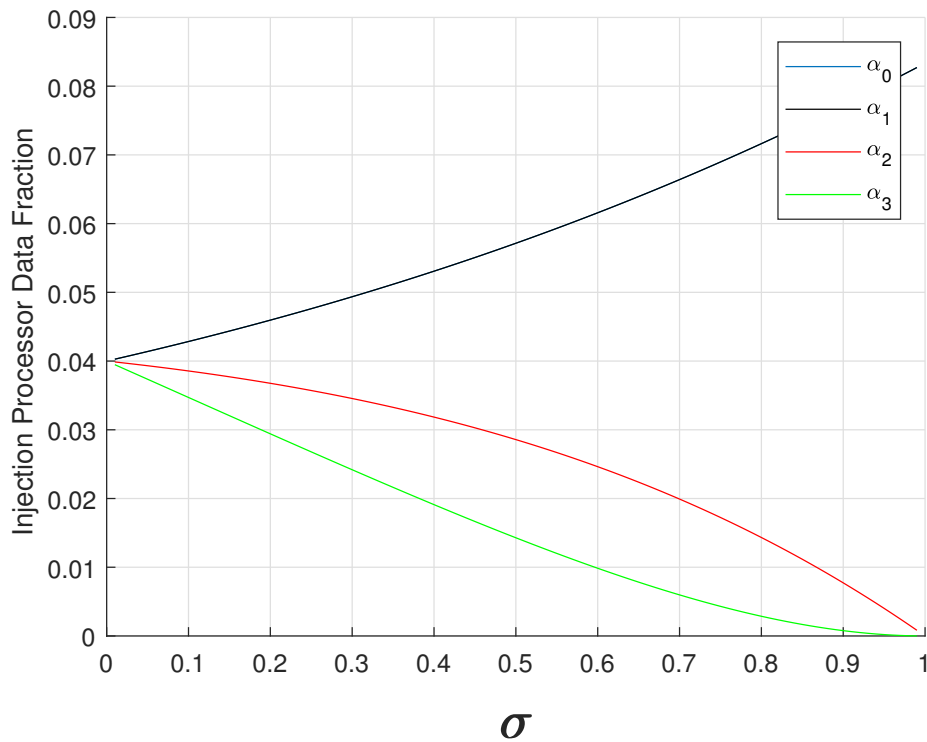


Figure 5.10: Data fraction on different type processors for Fig. 5.9

Since Fig. 5.10 also involves 25 total processors and 12 cluster processor adjacent to the source, it is identical to Fig. 5.8.

Model II

We choose the Reduced Voronoi Diagram Algorithm to address the problem.

Fig. 5.12 [63] provides a torus Voronoi method, which is used here. It extends the original domain to 8 copies and calculate the Voronoi Diagram as a planar algorithm [64]. Then the corner part is the torus Voronoi diagram.

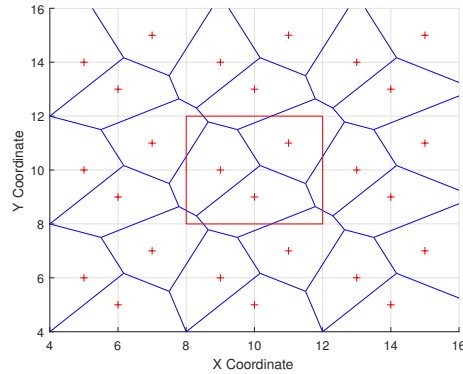


Figure 5.11: Copy the original domain to 8 copies

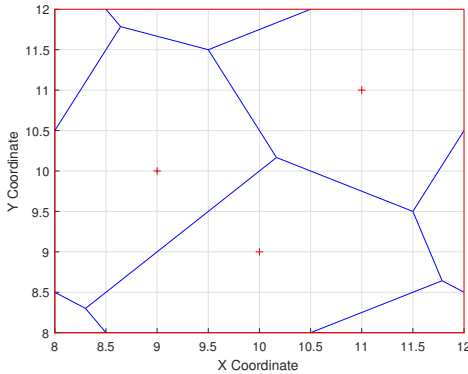


Figure 5.12: How to calculate torus Voronoi Diagram

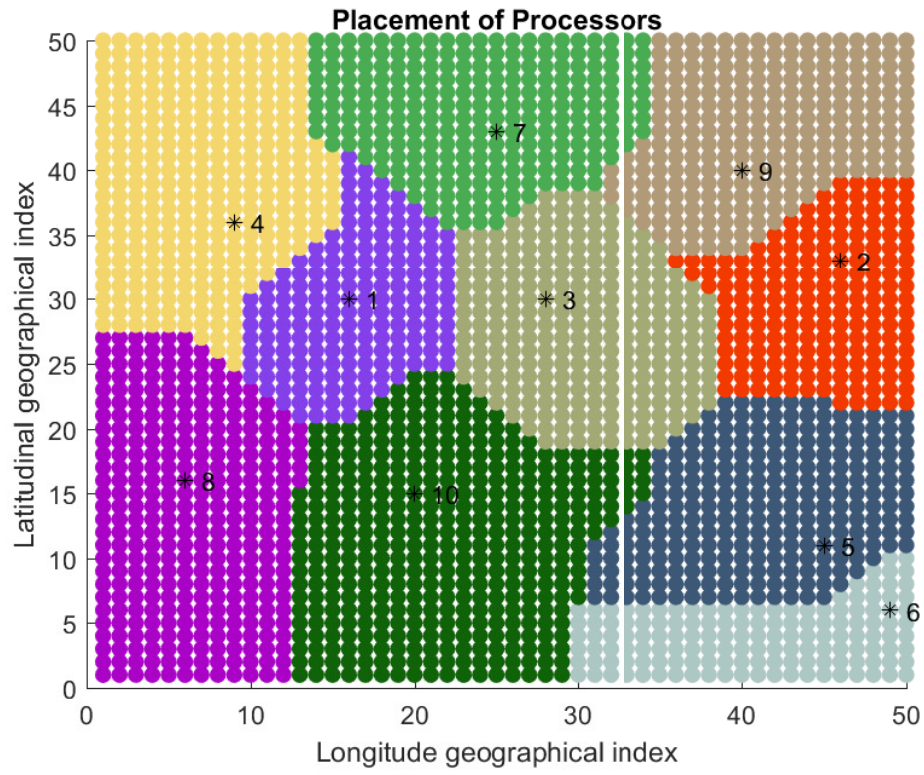


Figure 5.13: Initial Manhattan distance Voronoi Diagram

Fig. 5.13 says the initial Voronoi division of a 50×50 mesh network. The base is the data injection and the processor Voronoi cell tackle base's workload together. The user case contains 10 data injections and each cell's speedup curve shows in Fig. 5.16.

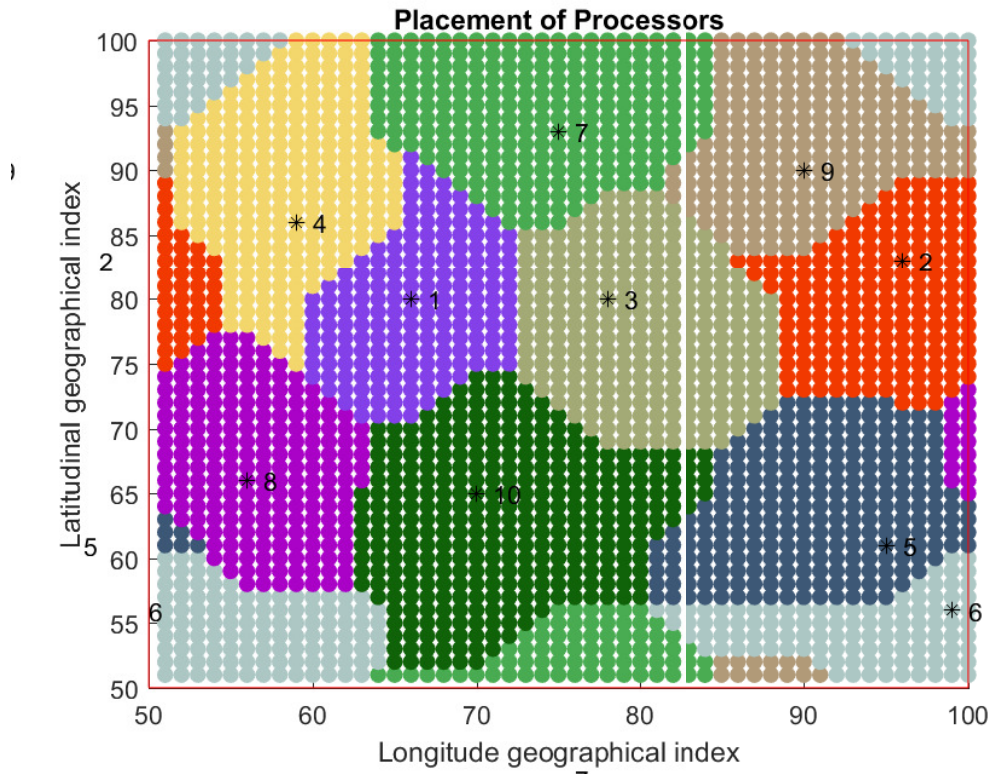


Figure 5.14: Torus Manhattan distance Voronoi diagram

Torus Manhattan distance Voronoi diagram Fig. 5.14 shows a toroidal Voronoi division result. The same color parts connect with each other and consist of a whole division in toroidal rectangle network. For example, the four gray corners consist of a whole area and the injection processor index is 6.

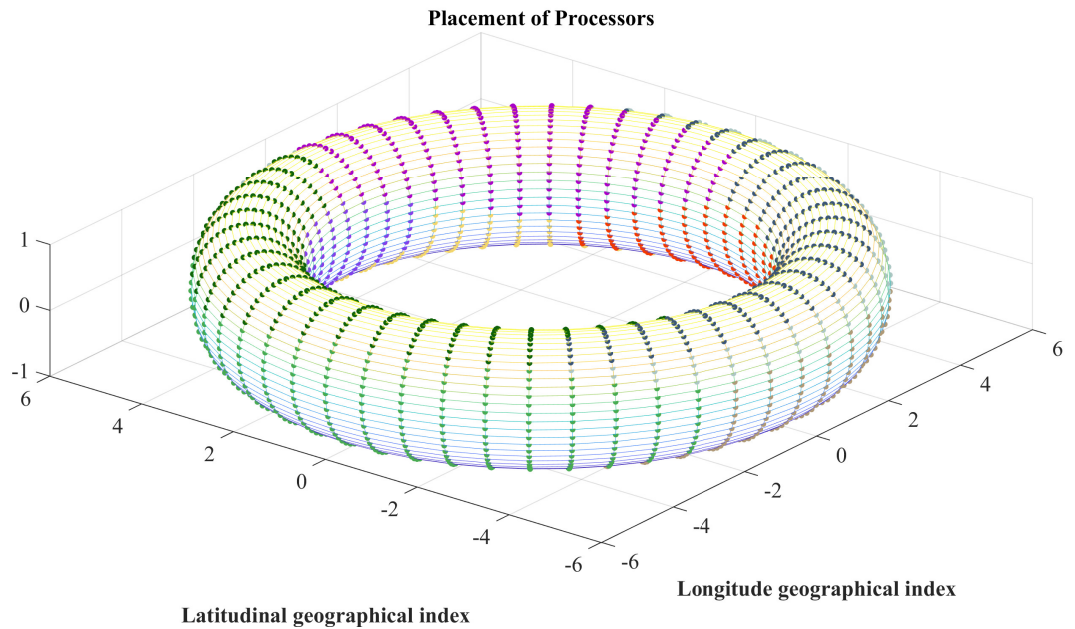


Figure 5.15: Voronoi diagram casting to the torus

Fig. 5.15 shows the Fig. 5.14 casts to the toroidal rectangle network result.

- The dot represents the processor.
- Different color represents the different cell in planar graph.
- Merge the left and right side of Fig. 5.14 to obtain a cylinder.
- Merge the top and bottom side of cylinder to obtain a torus.

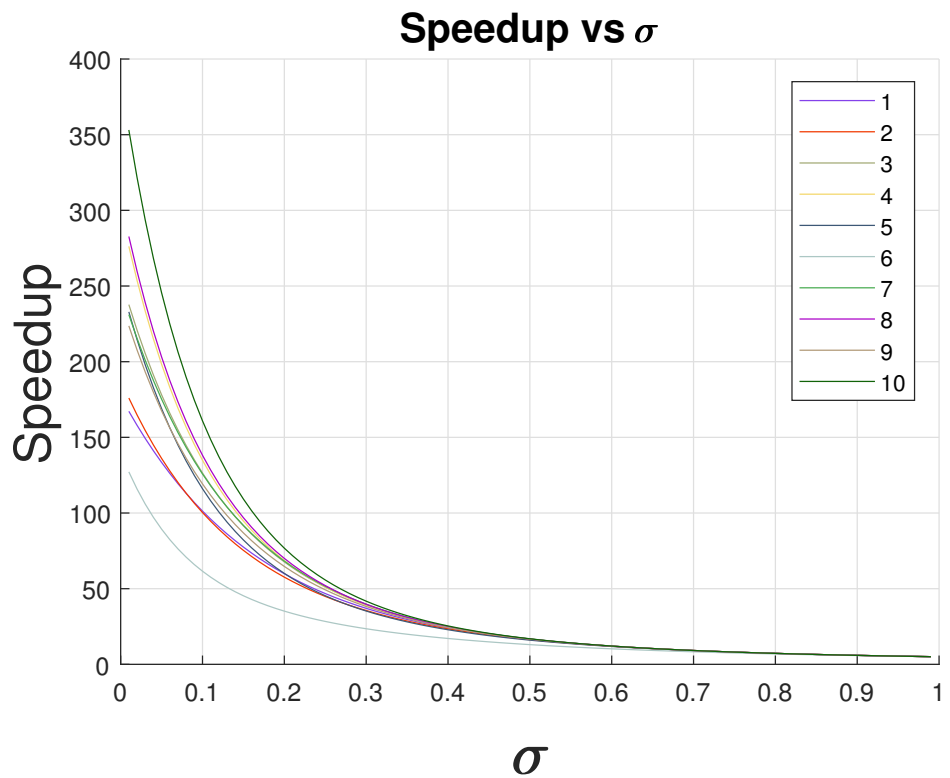


Figure 5.16: Toroidal Voronoi cells' speedup curves

- Different color represents the different Voronoi cell.
- Each curve represents the speedup efficiency of each Voronoi cell in different σ value.
- The maximum size of Voronoi cell is about 351 and the minimum size of Voronoi cell is 128.
- If $\sigma \approx 0$, the speedup ratio is similar with the number of processors in each cell.
- Even the processor number of Voronoi cell is different, $\sigma \geq 0.4$, the speedup ratio is about 1 in different cell.

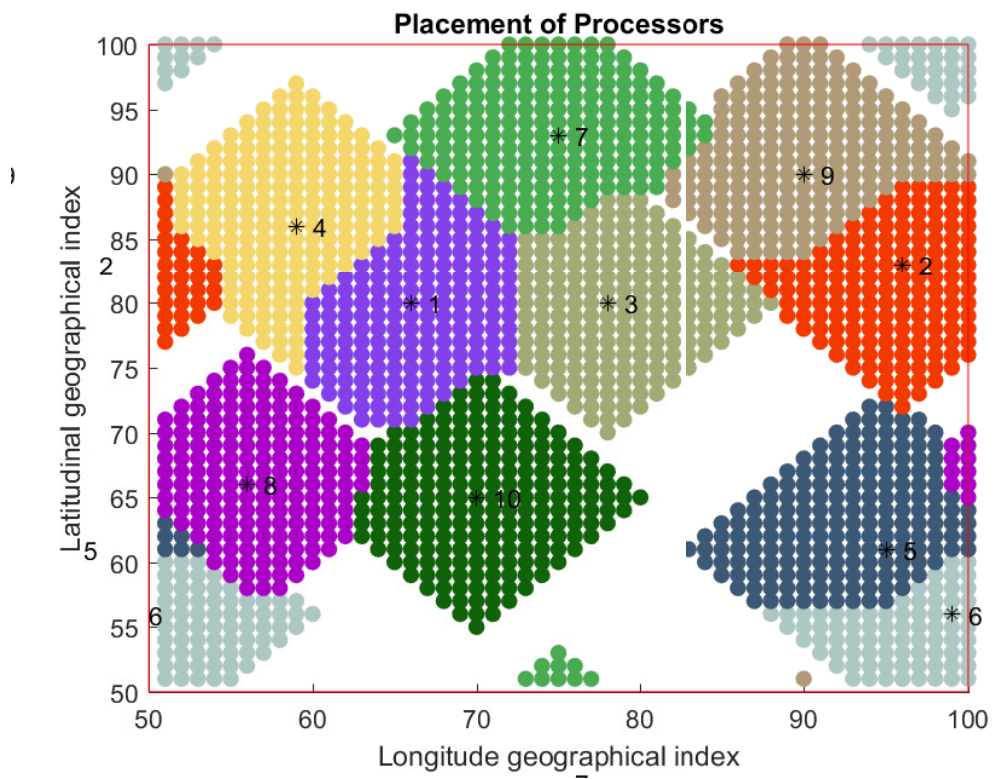


Figure 5.17: Torus reduced Voronoi diagram

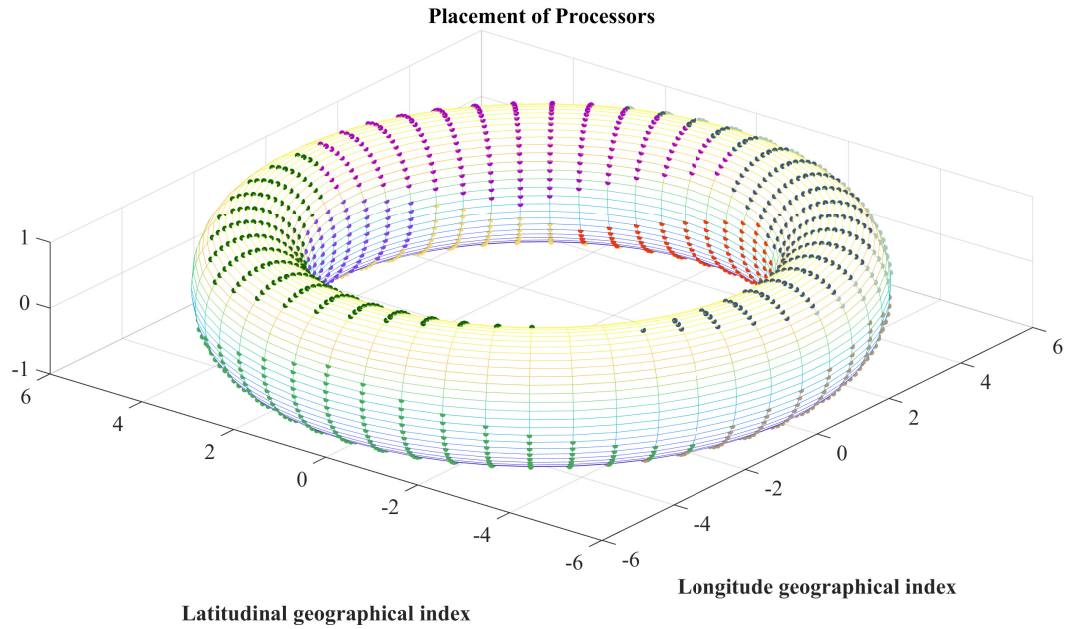


Figure 5.18: Torus reduced Voronoi diagram casting to a toroidal rectangle network

Fig. 5.18 shows the Fig. 5.17 casts to the toroidal rectangle network result.

- The dot represents the processor.
- The mesh grid without dot represents there is empty without processor.
- Different color represents the different cell in planar graph.
- Merge the left and right side of Fig. 5.14 to obtain a cylinder.
- Merge the top and bottom side of cylinder to obtain a torus.

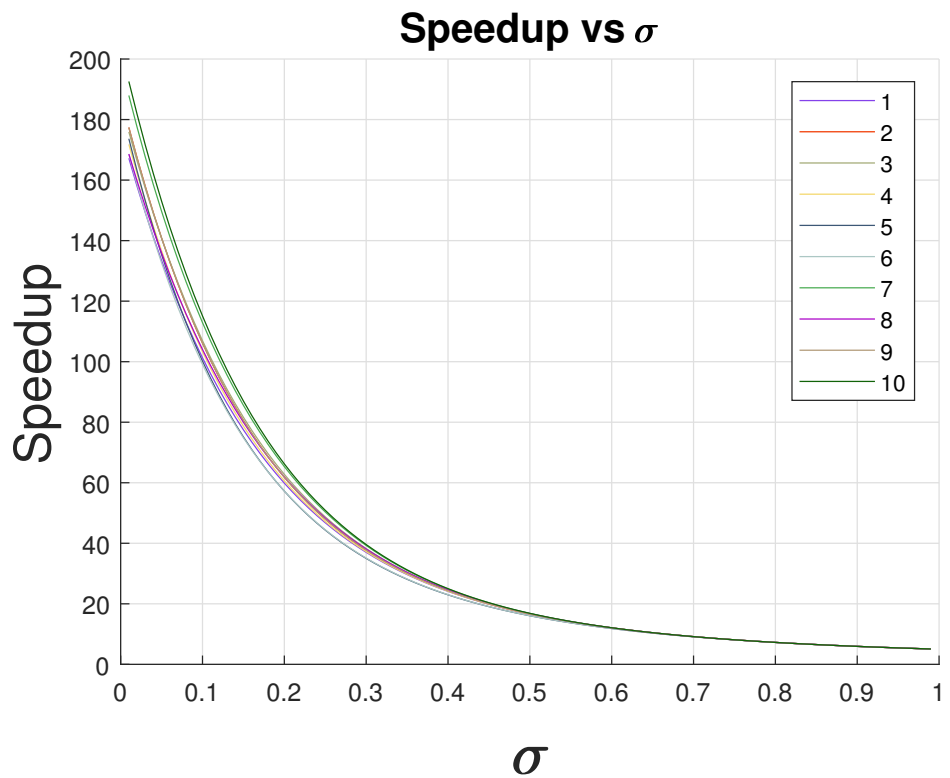


Figure 5.19: Torus reduced Voronoi diagram speedup curves

- Different color represents the different Voronoi cell.
- Each curve represents the speedup efficiency of each Voronoi cell in different σ value.
- The maximum size of Voronoi cell is about 191 and the minimum size of Voronoi cell is 168.
- Even the processor number of Voronoi cell is different, $1 > \sigma > 0$, the speedup ratio is about 1 in different Voronoi cell.

In the reduced model Fig. 5.18 and Fig. 5.19, reduced toroidal Voronoi diagram save 27% processors for the same processing capacity.

The ratio speedup is about $\frac{Sp_{max}}{Sp_{min}} = \frac{351}{125} \approx 2.816$. After the reduced stage, the ratio is $\frac{191}{168} \approx 1.137$. That is the reduced heuristic algorithm obtaining more balanced computation capacity distribution.

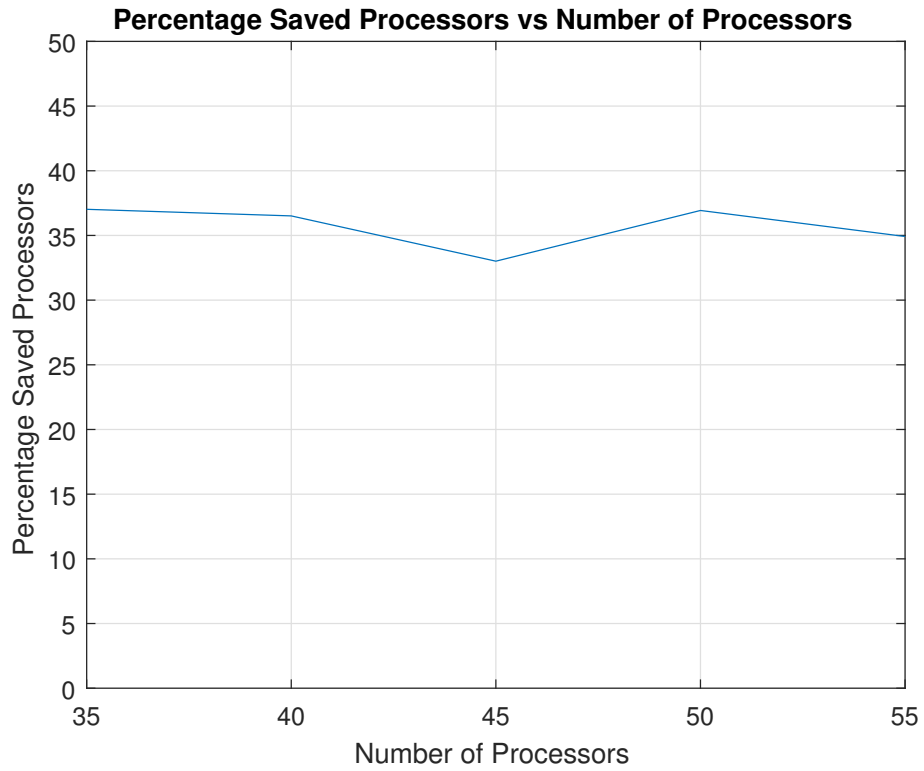


Figure 5.20: Torus reduced Voronoi diagram average saved processors

Model III

We utilize the Reduced Constrained Voronoi Diagram Algorithm to address this kind of question.

Table 5.3: Constrained Voronoi Diagram and RCVD

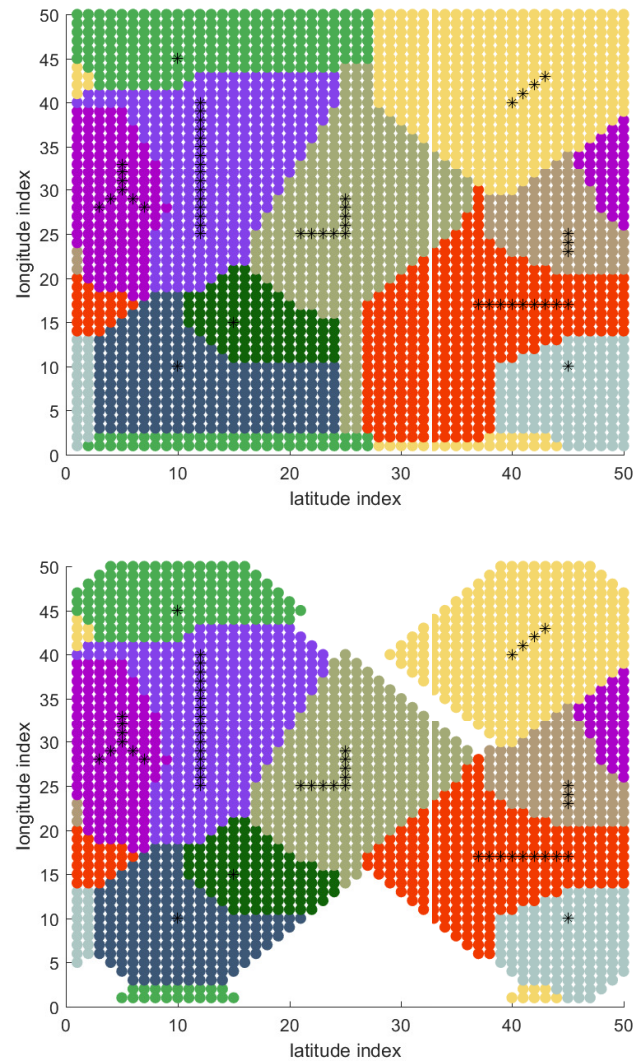
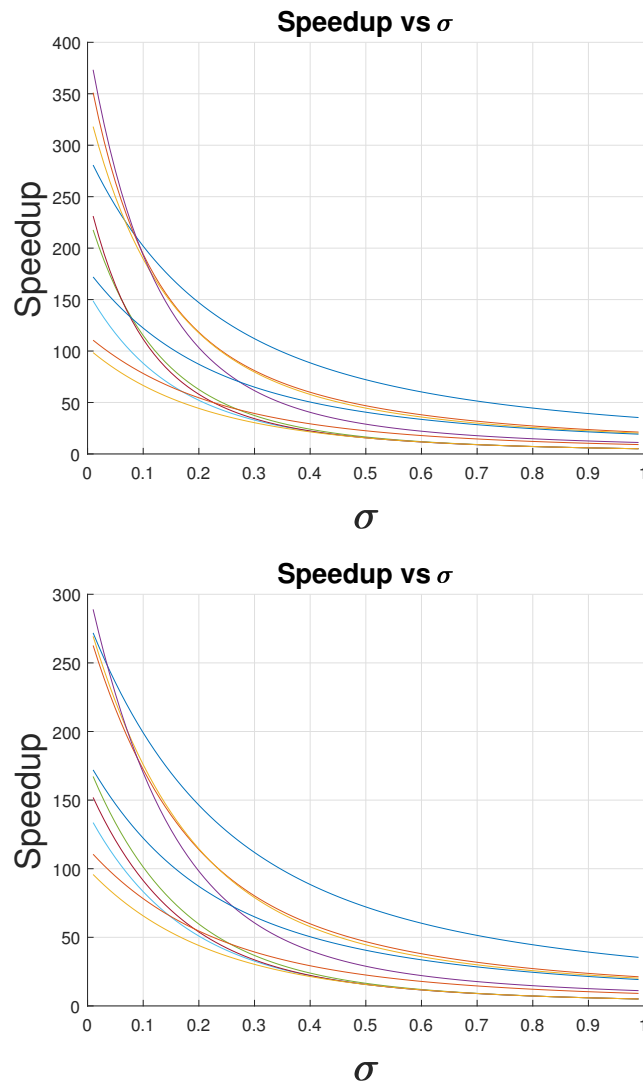


Table 5.4: Speedup curves Vs RCVDA speedup curves



Save about $436/2500 \approx 17\%$ processors.

5.2.4 Multi-source Non-uniform Data Injection

We utilize the same methodology and algorithm used in the mesh network.

5.3 Modified Store and Forward Switching Scenario

5.3.1 Data Injection On The Grid Processor

We implement a modified store and forward policy where each node must receive its entire load before it starts processing. However, it is assumed once a node receives its entire load it can forward load to successive nodes with zero delay in a virtual cut-through switching like mode. Certainly other policies can be modeled and solved which is not done here for space reasons.

We utilize the σ^* to present $-(\sigma + 1)$. The flow matrix closed-form of Fig. 5.3 is:

$$\begin{bmatrix} 1 & 4 & 8 & 10 & 8 & 4 & 1 \\ 1 & \sigma^* & 0 & 0 & 0 & 0 & 0 \\ 1 & -\sigma & \sigma^* & 0 & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma^* & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -\sigma & -\sigma & \sigma^* \end{bmatrix} \times \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \alpha_2 \\ \alpha_3 \\ \alpha_4 \\ \alpha_5 \\ \alpha_6 \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (5.7)$$

The fraction curve result is :

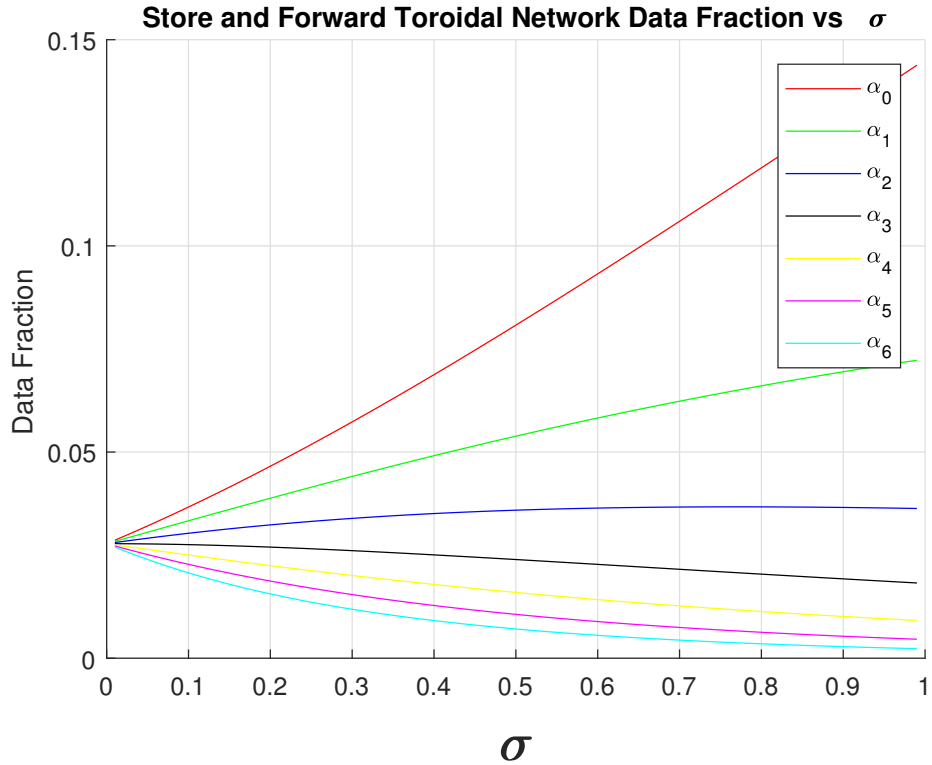


Figure 5.21: The data fraction curve of 6×6 torus network Fig. 5.3

From Fig. 5.21, we see that as the value σ grows, more and more workload is assigned to the $P_{4,2}$ and its one hop neighbors. Since there are 36 processors for $\sigma \approx 0$ (fast communication), $\frac{1}{36}$ of the load is assigned uniformly to each processor. That is, as the communication ability decreases, the economical method is to locally process the job.

D_i	Number
0	1
1	4
2	8
3	8
4	4

Table 5.5: D_i vs Number of processor on D_i

5.3.2 Sensitivity Analysis of Toroidal Rectangle Network

Considering a $5 * n$ ($n = 1 \dots 5$) toroidal rectangle network, the $level_i$ table shows Table 5.5:

So the simulation result illustrates in Fig. 5.22

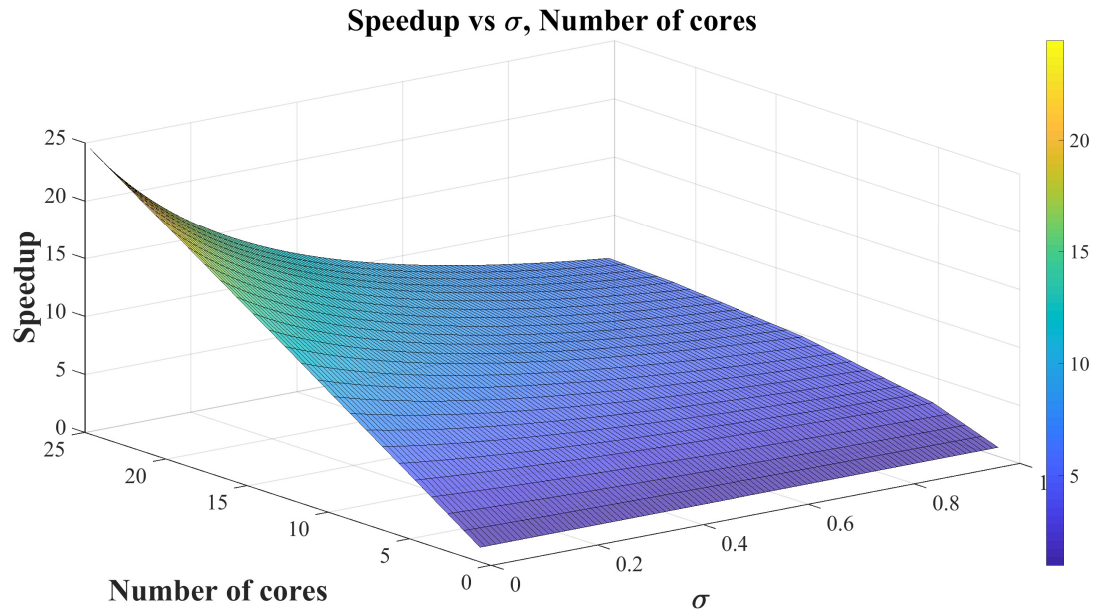


Figure 5.22: Sensitivity analysis result of $5 * n$ torus network ($n = 1 \dots 5$)

5.3.3 Multi-source Uniform Data Injection

We choose the same algorithm in virtual cut-through scenario to deal with this problem.

5.3.4 Multi-source Non-uniform Data Injection

We choose the same algorithm in virtual cut-through scenario to deal with this problem.

5.4 Comparison Result

5.4.1 Comparison Result Between Mesh Network and Toroidal With Same Number of Processors

Considering a $5 * n$ ($n = 1 \cdots 5$) mesh network Fig. 2.1, the best position for data injection is P_{12} . Other positions, for example P_8 , P_{13} , they don't have the same speedup efficiency. Yet, for a toroidal $5 * 5$ regular mesh network, each position's efficiency is equal. Fig. 5.23 explores the comparison result between the toroidal and corner scenario difference.

5.4.2 Comparison Result With Corner Processor and Inner Grid Processor

For a $5 * n$ ($n = 1 \cdots 5$) mesh network, the inner grid position is P_{12} and the corner data injection position is P_0 .

The comparing result is Fig. 5.23.

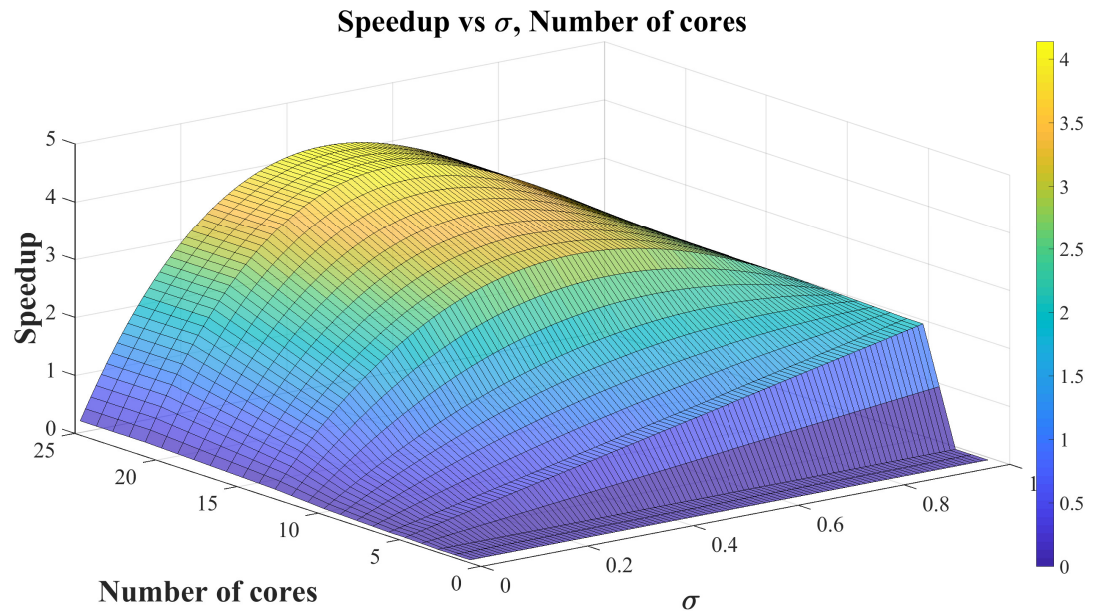


Figure 5.23: Speedup difference between inner grid injection and corner injection in $5 * n$ network ($n = 1 \cdots 5$)

Generally speaking, Fig. 5.23 says the inner grid position scenario has better performance than the corner injection option. If the grid node is 25 and $\sigma = 0.5$, the speedup difference is largest, which is 4.

5.4.3 Comparison Result Between Virtual Cut-through Switching Scenario and Modified Store and Forward Scenario

In the legend of figures, we use

- VCT presents virtual cut-through switching.
- SF presents store and forward switching.
- $VCT\alpha_0$ means the data fraction of P_0 in virtual cut-through switching scenario.
- $SF\alpha_0$ means the data fraction of P_0 in store and forward switching scenario.

Data Injection On the Corner Processor

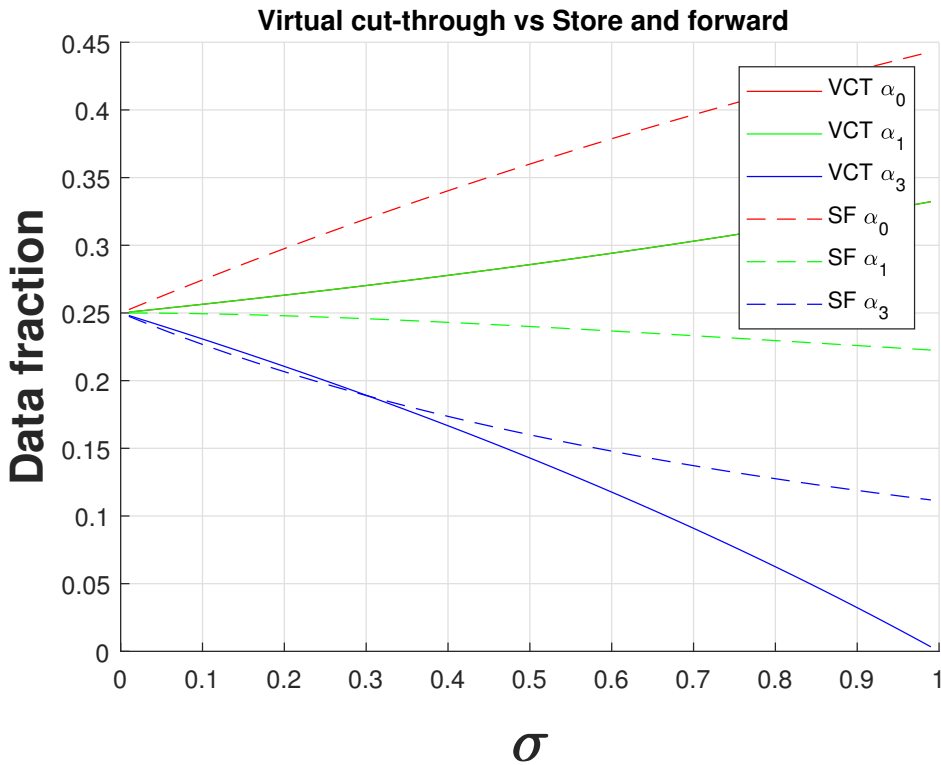


Figure 5.24: Comparing result between virtual cut-through switching and store and forward switching in $2 * 2$ mesh network

Fig. 5.24 says that P_0 takes more of the assigned load in store and forward switching scenario than virtual cut-through switching situation. As the value of σ increases, the fractions that are deployed to the deeper layers decreases in both cases. In the limit condition, for example, $\sigma = 1$, there is no data transmitted to P_3 in the virtual cut-through switching assumption, yet in the store and forward switching situation, there is still about 10% data fraction are communicated to P_3 . For fast communication ($\sigma \approx 0$), each of the four processors receive 25% of the load.

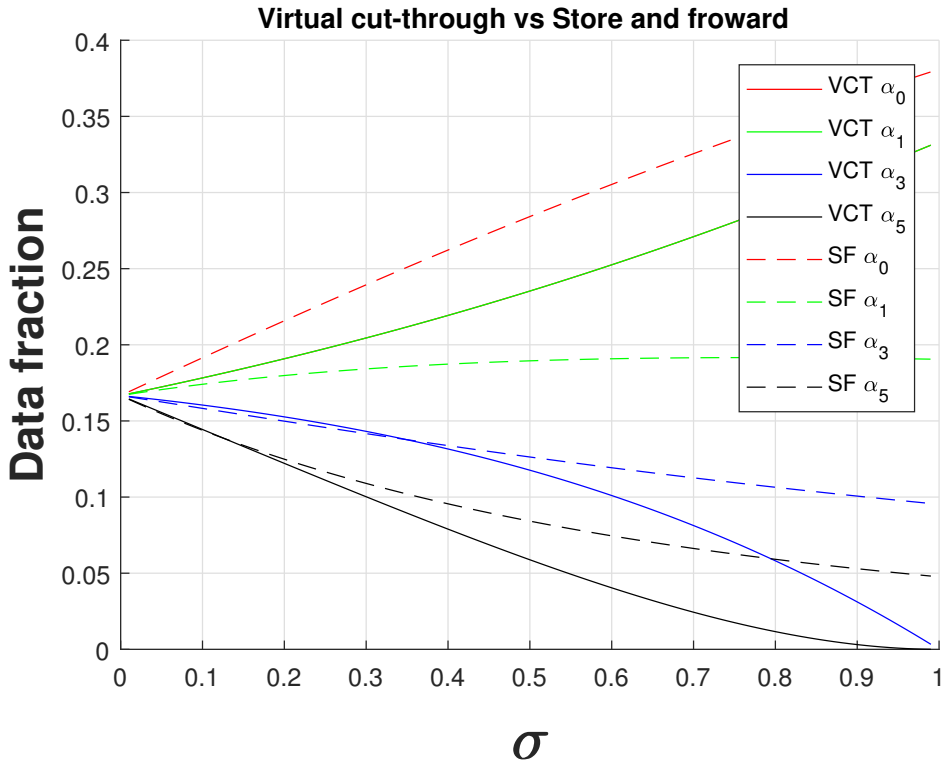


Figure 5.25: Comparing result between virtual cut-through switching and store and forward switching in 2×3 mesh network

Fig. 5.25 says that P_0 takes more assigned task in store and forward switching scenario than virtual cut-through switching situation. As the value of σ increases, the fractions that are deployed to the deeper levels decreases. In the limit condition, for example, $\sigma = 1$, there is no data transmitted to $level_3$, that is, P_5 in the virtual cut-through switching assumption. Yet in the store and forward switching situation, there is still about 5% data fraction is communicated to P_5 . For fast communication ($\sigma \approx 0$), each of the six processors receive $\frac{1}{6}$ of the load.

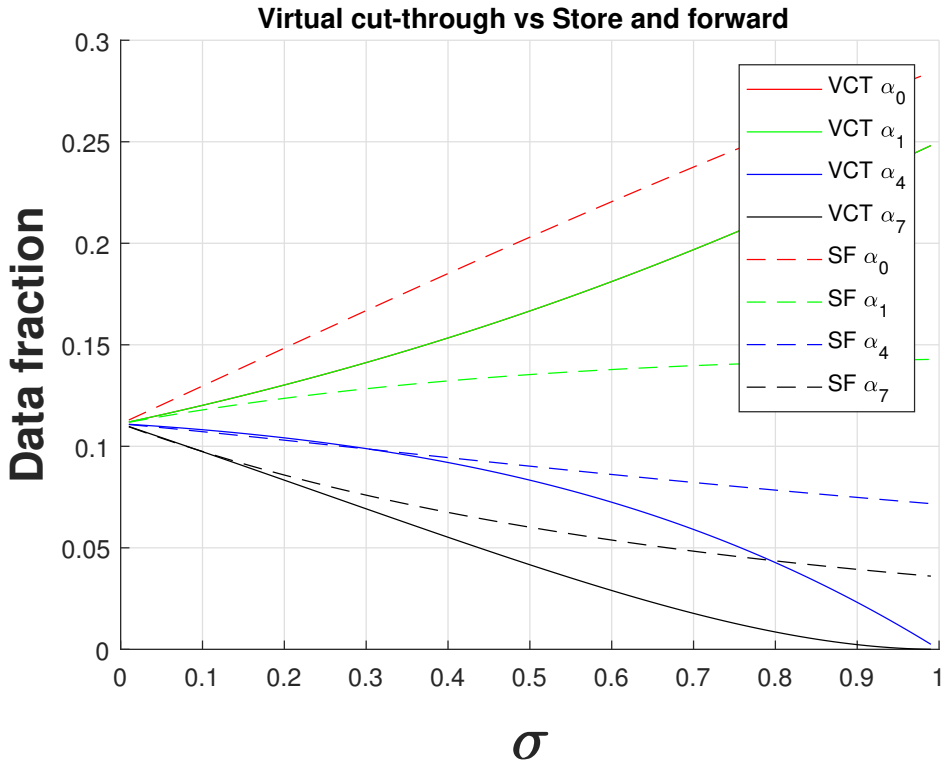


Figure 5.26: Comparing result between virtual cut-through switching scenario and store and forward switching scenario. The data injection is boundary processor of a 3×3 mesh network.

Fig. 5.26 says that P_0 takes more assigned task in store and forward switching scenario than virtual cut-through switching situation. As the value of σ increases, the fractions that are deployed to the deeper levels decreases. In the limit condition, for example, $\sigma = 1$, there is no data transmitted to $level_3$, that is, P_7 and P_8 in the virtual cut-through switching assumption. Yet in the store and forward switching situation, there is still about 5% data fraction is communicated to P_7 and P_8 . For fast communication ($\sigma \approx 0$), each of the nine processors receive $\frac{1}{9}$ of the load.

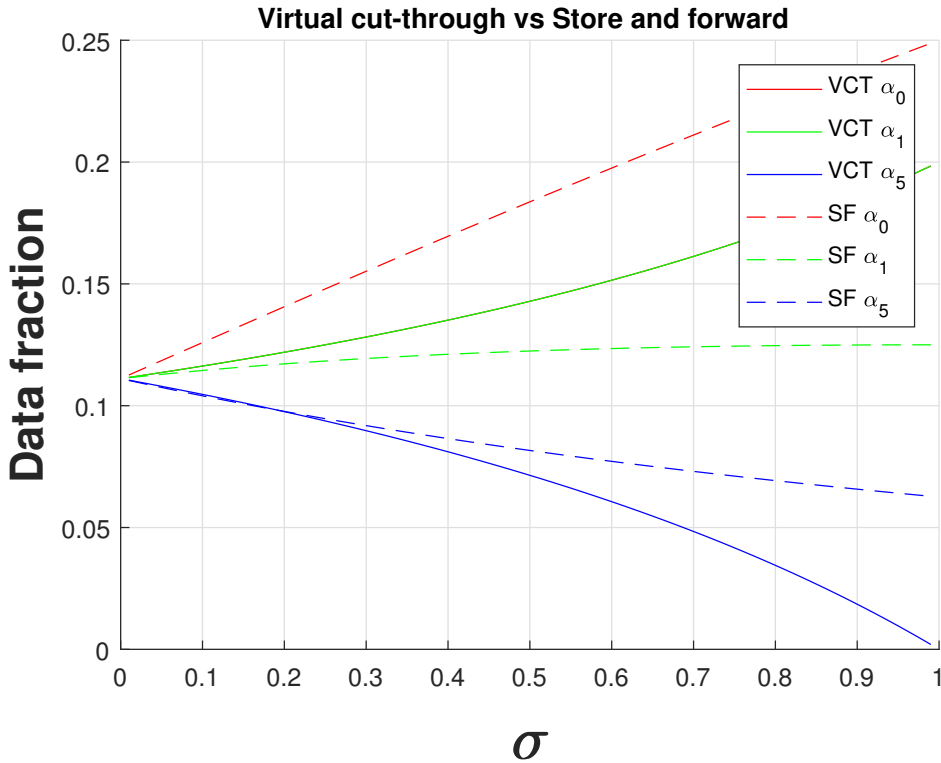


Figure 5.27: The comparing result between virtual cut-through switching processor with without virtual cut-through switching processor in 3×3 mesh network injection on inner grid processor

Fig. 5.27 says that P_0 takes more assigned task in store and forward switching scenario than virtual cut-through switching situation. As the value of σ increases, the fractions that are deployed to the deeper levels dropping down. In the limit condition, for example, $\sigma = 1$, there is no data transmitted to $level_2$, that is, P_5 , P_6 , P_7 and P_8 in the virtual cut-through switching assumption. Yet in the store and forward switching situation, there is still about 5% data fraction is communicated to P_5 , P_6 , P_7 and P_8 .

Comparing with Fig. 5.26, P_0 takes less workload in inner grid position than boundary data injection. The reason is there are 4 neighbor processors on the $level_1$, yet there is only three processors on $level_1$ on the boundary.

Fig. 5.28 shows the speedup difference between the virtual cut-through switching situation and modified store and forward switching scenario.

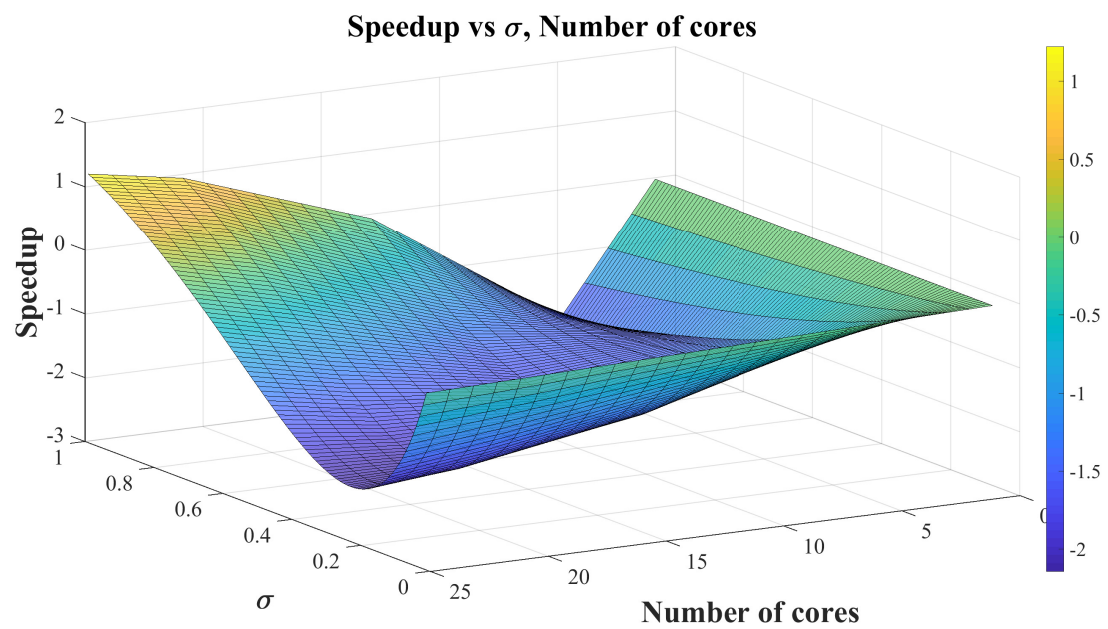


Figure 5.28: Speedup difference between modified store and forward switching and virtual cut-through switching in $5 * n$ mesh network ($n = 1 \cdots 5$)

Chapter 6

Hypercube Network and General Network

6.1 Hypercube Network Virtual Cut-through Switching Scenario

The hypercube topology has two nodes along each dimension and $\log_2 n$ dimensions. The construction of a hypercube goes as follows, in general a d -dimensional hypercube is constructed by connecting corresponding nodes of two $(d - 1)$ dimensional hypercubes Fig. 6.1.

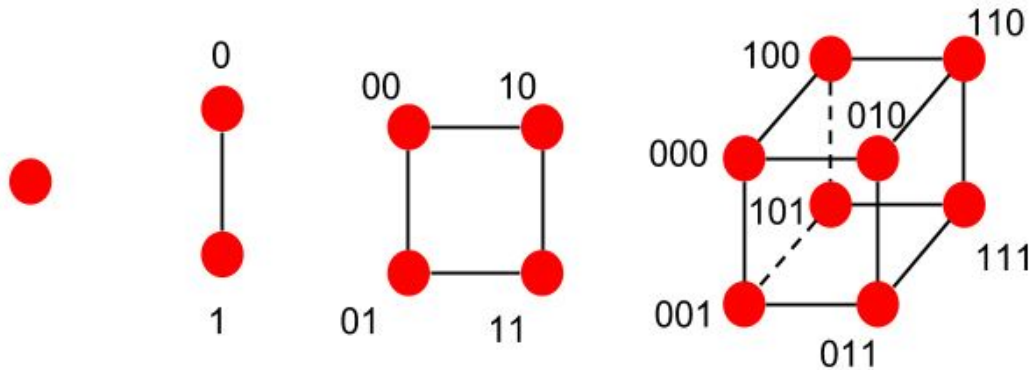


Figure 6.1: Hypercube in 0, 1, 2, 3 dimension. [2]

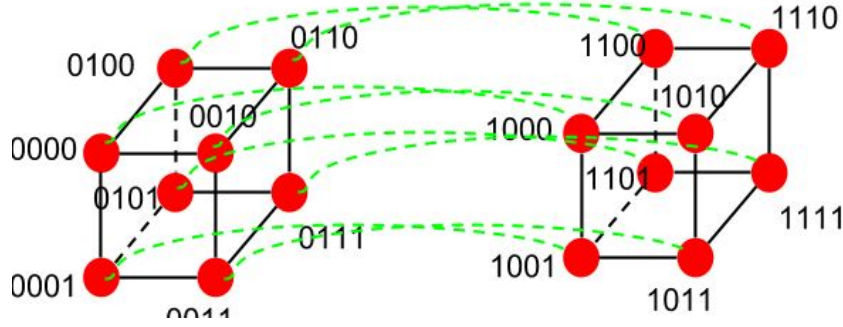


Figure 6.2: A hypercube network

6.1.1 Data Injection On The Grid Processor

For the hypercube of dimension d there are 2^d processor in the system. Each of the processors has direct links to d neighbors. A method of naming the processors is to use label consisting of a binary string d – position long. Further, the label of a processor is a binary number from the interval $[0, 2^d - 1]$

To address the qualitative model of computation, the critical problem is to calculate the number of processor on each D_i . Each node is connected by link and the hamming distance of their's label is 1. According to the lemma of [43],

Lemma 6.1.1. *In each layer i of D dimensional hypercube, there are $\binom{n}{i}$ processors each of which can be accessed through i communications links and is capable of transmitting to $d - i$ still idle processors.*

According to a $2 - D$ hypercube Fig. 3.1, the flow matrix is

$$A = \begin{bmatrix} \binom{2}{0} & \binom{2}{1} & \binom{2}{2} \\ 1 & -1 & 0 \\ 0 & \sigma - 1 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 2 & 1 \\ 1 & -1 & 0 \\ 0 & \sigma - 1 & 1 \end{bmatrix} \quad (6.1)$$

, which is investigated in Regular Network Chapter.

According to a $3 - D$ hypercube, the flow matrix is

$$A = \begin{bmatrix} (3)_0 & (3)_1 & (3)_2 & (3)_3 \\ 1 & -1 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} = \begin{bmatrix} 1 & 2 & 2 & 1 \\ 1 & -1 & 0 & 0 \\ 0 & \sigma - 1 & 1 & 0 \\ 0 & \sigma - 1 & \sigma & 1 \end{bmatrix} \quad (6.2)$$

The equivalence inverse processing speed :

$$\hat{T}_f = 1 * w_{eq} * T_{cp}$$

$$w_{eq} = \alpha_0 * w$$

The speedup is :

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = |-\det A|$$

A general case, D -dimension network, the flow matrix is :

$$A = \begin{bmatrix} \binom{n}{0} & \binom{n}{1} & \binom{n}{2} & \cdots & \binom{n}{n-2} & \binom{n}{n-1} & \binom{n}{n} \\ 1 & -1 & 0 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{bmatrix}$$

6.2 General Case Virtual Cut-through Switching Scenario

$$\begin{bmatrix} 1 & m_1 & m_2 & \cdots & m_{n-2} & m_{n-1} & m_n \\ 1 & -1 & 0 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & 1 & \cdots & 0 & 0 & 0 \\ 0 & \sigma - 1 & \sigma & 1 & 0 & \cdots & 0 \\ 0 & \sigma - 1 & \sigma & \sigma & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 0 & \sigma - 1 & \sigma & \cdots & \sigma & \sigma & 1 \end{bmatrix} \times \begin{bmatrix} \alpha_{l_0} \\ \alpha_{l_1} \\ \alpha_{l_2} \\ \alpha_{l_3} \\ \vdots \\ \alpha_{l_{n-1}} \\ \alpha_{l_n} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ \vdots \\ 0 \end{bmatrix} \quad (6.3)$$

The m_1, m_2, \dots, m_n are the number of processors on the $level_1, level_2, \dots, level_n$.
Also, the $\alpha_{l_0}, \alpha_{l_1}, \dots, \alpha_{l_n}$ are corresponding workload fraction.

Finally, the speedup is:

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = |-\det A|$$

6.3 Hypercube Network Modified Store and Forward Switching Scenario

According to a $2 - D$ hypercube Fig. 3.1, the flow matrix is

$$A = \begin{bmatrix} \binom{2}{0} & \binom{2}{1} & \binom{2}{2} \\ 1 & -(\sigma + 1) & 0 \\ 1 & -\sigma & -(\sigma + 1) \end{bmatrix} = \begin{bmatrix} 1 & 2 & 1 \\ 1 & -(\sigma + 1) & 0 \\ 1 & -\sigma & -(\sigma + 1) \end{bmatrix} \quad (6.4)$$

, which is investigated in Regular Network Chapter.

According to a $3 - D$ hypercube, the flow matrix is

$$A = \begin{bmatrix} \binom{3}{0} & \binom{3}{1} & \binom{3}{2} & \binom{3}{3} \\ 1 & -(\sigma + 1) & 0 & 0 \\ 1 & -\sigma & -(\sigma + 1) & 0 \\ 1 & -\sigma & -\sigma & -(\sigma + 1) \end{bmatrix} = \begin{bmatrix} 1 & 2 & 2 & 1 \\ 1 & -(\sigma + 1) & 0 & 0 \\ 1 & -\sigma & -(\sigma + 1) & 0 \\ 1 & -\sigma & -\sigma & -(\sigma + 1) \end{bmatrix} \quad (6.5)$$

The speedup is $|- \det A|$.

A general case, D -dimension network, the flow matrix is : We use σ^* to represents $-(\sigma + 1)$.

$$A = \begin{bmatrix} \binom{n}{0} & \binom{n}{1} & \binom{n}{2} & \cdots & \binom{n}{n-2} & \binom{n}{n-1} & \binom{n}{n} \\ 1 & \sigma^* & 0 & \cdots & 0 & 0 & 0 \\ 1 & -\sigma & \sigma^* & \cdots & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & \sigma^* & 0 & \cdots & 0 \\ 1 & -\sigma & -\sigma & -\sigma & \sigma^* & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 1 & -\sigma & -\sigma & \cdots & -\sigma & -\sigma & \sigma^* \end{bmatrix}$$

6.4 General Case Store and Forward Switching Scenario

$$\begin{bmatrix} 1 & m_1 & m_2 & \cdots & m_{n-2} & m_{n-1} & m_n \\ 1 & -(\sigma + 1) & 0 & \cdots & 0 & 0 & 0 \\ 1 & -\sigma & -(\sigma + 1) & \cdots & 0 & 0 & 0 \\ 1 & -\sigma & -\sigma & -(\sigma + 1) & 0 & \cdots & 0 \\ 1 & -\sigma & -\sigma & -\sigma & -(\sigma + 1) & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \ddots & \vdots \\ 1 & -\sigma & -\sigma & \cdots & -\sigma & -\sigma & -(\sigma + 1) \end{bmatrix} \times \begin{bmatrix} \alpha_{l_0} \\ \alpha_{l_1} \\ \alpha_{l_2} \\ \alpha_{l_3} \\ \vdots \\ \alpha_{l_{n-1}} \\ \alpha_{l_n} \end{bmatrix} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ \vdots \\ 0 \\ 0 \end{bmatrix} \quad (6.6)$$

The m_1, m_2, \dots, m_n are the number of processors on the $level_1, level_2, \dots, level_n$. Also, the $\alpha_{l_0}, \alpha_{l_1}, \dots, \alpha_{l_n}$ are corresponding workload fraction.

The speedup is

$$Speedup = \frac{T_f}{\hat{T}_f} = \frac{\omega T_{cp}}{\alpha_0 \omega T_{cp}} = \frac{1}{\alpha_0} = \frac{\det A}{\det A^*} = \left| \frac{\det A}{(\sigma^*)^{n-1}} \right|$$

Chapter 7

Conclusion and Future Work

7.1 Conclusion

In this work we investigate two problems. One problem is closed-form solutions for equivalence computation of divisible workload in mesh and toroidal networks and the other problem is scheduling divisible workloads from multiple sources in mesh and toroidal networks of processors.

We propose a flow matrix closed-form equation to determine the equivalence, which allows a characterization of the nature of minimal time solution and a simple method to determine when and how much load to distribute for processors.

Also, we propose the use of a reduced Manhattan distance Voronoi diagram algorithm (RMDVDA) to minimize the overall processing time of these workloads by taking advantage of the processor equivalence technique. Three case studies with 10 sources of workloads are presented to illustrate the general approach for multiple sources of workloads.

In the first phase, a Manhattan distance Voronoi diagram is used to obtain a network cluster division. In the second phase, we propose an efficient algorithm to obtain near-optimal load distribution among processors represented by equivalent processors. The algorithm minimizes the number of processors utilized.

The influence of Various factors, for example data injection position, data injection fraction, virtual cut-through switching and modified store and forward switching, the ratio of computation time over communication time, that influence the load distribution are shown in the numerical experiments. Experimental evaluation through simulations demonstrates that a task can be finished in the same suboptimal time and yet save over 40% of processor resources in mesh network and 11% – 17% in toroidal network.

7.2 Future Work

There are future projects that need attention.

- Given virtual cut-through and modified store and forward scenario, the hypercube network multi-source suboptimal algorithm to minimize the makespan.
- Given data injection workload, how to account for dynamic workload queues and how to find the suboptimal algorithm to minimize the makespan.
- Given the workload fractions distribution, how to choose the appropriate data injection positions. The objective to minimize the cost function, which has relationship with Earth Mover Distance [61] and Wasserstein Distance [65].
- Other topology networks, for example connected cubic network graph [66], heterogeneous networks [67], Gaussian network [68] .
- Other constraint assumptions, for example the front-end processor with finite-size buffer.

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