Improving the Accessibility of Heterogeneous System Resources for Application Developers using Programming Abstractions

Verbesserung der Zugänglichkeit heterogener Systemressourcen für Anwendungsentwickler durch Programmierabstraktionen

Max Frederik Plauth

Dissertation zur Erlangung des Doktorgrades
Doktor der Ingenieurwissenschaften (Dr.-Ing.)
der Digital Engineering Fakultät
der Universität Potsdam
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Datum der Einreichung: 03. Mai 2022
Datum der Disputation: 05. Juli 2022

Published online on the Publication Server of the University of Potsdam:
https://doi.org/10.25932/publishup-55811
https://nbn-resolving.org/urn:nbn:de:kobv:517-opus4-558118
Abstract

The heterogeneity of today’s state-of-the-art computer architectures is confronting application developers with an immense degree of complexity which results from two major challenges. First, developers need to acquire profound knowledge about the programming models or the interaction models associated with each type of heterogeneous system resource to make efficient use thereof. Second, developers must take into account that heterogeneous system resources always need to exchange data with each other in order to work on a problem together. However, this data exchange is always associated with a certain amount of overhead, which is why the amounts of data exchanged should be kept as low as possible.

This thesis proposes three programming abstractions to lessen the burdens imposed by these major challenges with the goal of making heterogeneous system resources accessible to a wider range of application developers. The lib842 compression library provides the first method for accessing the compression and decompression facilities of the NX-842 on-chip compression accelerator available in IBM Power Central Processing Units (CPUs) from user space applications running on Linux. Addressing application development of scale-out Graphics Processing Unit (GPU) workloads, the CloudCL framework makes the resources of GPU clusters more accessible by hiding many aspects of distributed computing while enabling application developers to focus on the aspects of the data parallel programming model associated with GPUs. Furthermore, CloudCL is augmented with transparent data compression facilities based on the lib842 library in order to improve the efficiency of data transfers among cluster nodes. The improved data transfer efficiency provided by the integration of transparent data compression yields performance improvements ranging between 1.11× and 2.07× across four data-intensive scale-out GPU workloads. To investigate the impact of programming abstractions for data placement in Non-Uniform Memory Access (NUMA) systems, a comprehensive evaluation of the PGASUS framework for NUMA-aware C++ application development is conducted. On a wide range of test systems, the evaluation demonstrates that PGASUS does not only improve the developer experience across all workloads, but that it is also capable of outperforming NUMA-agnostic implementations with average performance improvements of 1.56×.

Based on these programming abstractions, this thesis demonstrates that by providing a sufficient degree of abstraction, the accessibility of heterogeneous system resources can be improved for application developers without occluding performance-critical properties of the underlying hardware.
Zusammenfassung

Die Heterogenität heutiger Rechnerarchitekturen konfrontiert Anwendungsentwickler mit einem immensen Maß an Komplexität, welches sich aus zwei großen Herausforderungen ergibt. Erstens müssen Entwickler fundierte Kenntnisse über die Programmiermodelle oder Interaktionsmodelle verfügen, welche eine Voraussetzung sind um die jeweiligen heterogenen Systemressourcen effizient nutzen zu können. Zweitens müssen Entwickler berücksichtigen, dass heterogene Systemressourcen immer auch Daten untereinander aus tauschen müssen, um ein Problem gemeinsam zu bearbeiten. Dieser Datenaustausch ist aber auch immer mit einem gewissen Mehraufwand verbunden, weshalb die ausgetauschten Datenmengen so gering wie möglich gehalten werden sollten.


Um die Auswirkungen von Programmierabstraktionen auf die Datenplatzierung in NUMA-Systemen zu untersuchen, wird eine umfassende Evaluierung des PGASUS-Frameworks für NUMA-gewährte C++-Anwendungsentwicklung durchgeführt. Unter Verwendung einer breiten Palette von Testsystemen zeigt die Evaluierung, dass PGASUS nicht nur die Entwicklung von NUMA-gewährten Anwendungen erleichtert, sondern auch in der Lage ist, die Leistung von NUMA-agnostischen Implementierungen im Mittel um 1,56× zu übertrifft.

Auf der Grundlage dieser Programmierabstraktionen zeigt diese Dissertation, dass heterogene Systemressourcen durch die Bereitstellung angemessener Abstraktionsmechanismen einfacher von Anwendungsentwicklern erschlossen werden können, ohne dass leistungsrelevante Eigenschaften der zugrunde liegenden Hardware verdeckt werden.
Acknowledgements

This thesis would have never been completed without the support I have received from many people whom I would like to thank. I would like to thank my advisor Andreas Polze, who has guided my work by sharing his advice for countless times. First as the advisor and later as my colleague and friend, Frank Feinbube has sparked my interest in returning to academia in order to pursue the endeavors of earning a doctoral degree. At the beginning of my undertakings, the SSICLOPS project, or rather the project partners involved in it, have provided me with a research direction when I was not sure which direction I wanted to pursue. Collaborating with the IBM Germany R&D Lab in Böblingen over the course of the Hybrid DB project was an incredible opportunity. Special thanks are due to Wolfgang Maier and his team, who did not only provide valuable feedback, but based on their support, I have been awarded with the IBM PhD Fellowship Award in 2017.

Over the years I have spent at the Operating Systems and Middleware Group led by Andreas Polze, I have enjoyed both the collaboration and the numerous coffee talks with the ‘inhabitants’ of the ‘hardware corner’, namely Felix Eberhardt, Andreas Grapentin, Sven Köhler, and Lukas Wenzel. I am especially grateful to Sven Köhler and Lukas Wenzel, who both have been a great source of inspiration not only professionally, but also on a personal level. Over time, Bernhard Rabe, Christian Neuhaus, Daniel Richter, Frank Feinbube, Lena Feinbube, Marcel Taeumel, Tobias Pape, and Robert Schmid have helped me advance in my endeavors. At the later stages of my undertakings, the collaboration with Timo Hönig has provided me with several new perspectives and ideas that I hope I will be able to pursue in my postdoctoral life.

Without the support of Sabine Wagner, many bureaucratic acts would have been a nightmare. Also, this work would not have been possible without the FutureSOC Lab. More important than the compute equipment provided by the lab was the relentless support of Bernhard Rabe, Tobias Pape, and Ayleen Oswald.

The biggest thanks are due to my wife Annabell and my children. Without their tremendous loving support, I could never have pulled this through.
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1 Introduction

Over the last few decades, the uninterrupted growth of data quantities accumulating in the age of digitization has been driving an ever-growing demand for compute capacity. Up until the early 2000s, this demand could be easily satisfied based on the performance gains provided by frequency scaling. With frequency scaling however having reached its limits around 2006, computer architectures had to resort to different approaches in order to continuously provide improved compute capacities. Even though the first response to the end of frequency scaling was to invest the steadily increasing transistor count into multicore CPUs, the pressure to innovate created by the end of frequency scaling has promoted the entry of heterogeneous system resources into mainstream computer architectures [146]. As such, most CPU vendors have adopted cache-coherent NUMA architectures to scale multiprocessor systems to dimensions that were not feasible with Uniform Memory Access (UMA) approaches [92,107]. Similarly, all major GPU microarchitectures have adopted unified shader architectures, paving the way for the utilization of GPUs as general purpose compute resources [39,5,116].

Figure 1.1: The development of single-precision peak performance of both CPUs and GPUs demonstrates that even though CPU performance has caught up based on the introduction of excessively wide Single Instruction Multiple Data (SIMD) extensions, heterogeneous compute resources such as GPUs have delivered sustained growth of compute capacity over the years. The plot has been generated based on the data provided by Karl Rupp [176].

Ever since, heterogeneous system resources have not only become indispensible in providing significant improvements in compute capacity as illustrated in Figure 1.1 but
the degree of heterogeneity in computer architectures has also been steadily increasing over the following years \cite{14}. Upon closer examination, today’s state-of-the-art systems ranging from mobile phones to high-end servers are brimmed with heterogeneous system resources such as dedicated inference engines, video compression engines, digital signal processing engines, and many more. A crucial factor driving this trend is the specialization of heterogeneous compute resources offering superior performance per watt ratings compared to the general purpose compute resources of CPUs.

1.1 Heterogeneous System Resources

In the context of computer architectures, heterogeneity is a largely overloaded term. For example, heterogeneous computing may refer to the use of different classes of compute resources (e.g., CPUs and GPUs), the use of CPUs with varying Instruction Set Architectures (ISAs), or the use of CPUs with the same ISA but differences in their microarchitectural properties such as the mixed use of performance-optimized cores and efficiency-optimized cores. Therefore, the goal of this section is to establish a mutual understanding about what types of heterogeneous system resource are considered in the context of this thesis.

This thesis considers the three types of heterogeneous system resources highlighted in Figure 1.2:

- **On-Chip Accelerators** Many modern CPUs and System on a Chip (SoC) designs are equipped with on-chip accelerators that unlike SIMD ISA extensions are not easily accessible from user-space. While these types of accelerators are tightly integrated into the on-chip communication fabric, interacting with these types of accelerators can be particularly challenging as the methods for invoking their resources may vary even among individual CPU models produced by the same vendor.

- **Off-Chip Accelerators** As the probably most common approach for heterogeneous computing, compute resources of the CPU are augmented with a different type of compute resource such as a GPU which is connected to the CPU using an off-chip
1.2 Problem Statement

While the ever-growing degree of heterogeneity facilitates continuously increasing compute capacities, it is also the source of two major challenges in dealing with heterogeneous system resources from the perspective of application developers:

1. Many types of heterogeneous system resources require application developers to adapt specific programming models or interaction models to make use of its capabilities. With the large variety of heterogeneous system resources available in state-of-the-art computer architectures, application developers are therefore confronted with an immense degree of complexity.

2. Heterogeneous system resources have to exchange data in order to process a workload collectively. Since moving data across heterogeneous system resources can be a performance bottleneck, application developers have to be wary about balancing the use of heterogeneous system resources against the overhead associated with data transfers. However, there are certain scenarios in which prevalent system abstractions may make it hard for application developers to gauge whether their actions may trigger unnecessary data transfers or not.

With the imminent advent of disruptive memory technologies the effects of these challenges are very likely to intensify [136]. It is therefore necessary to provide application developers with programming abstractions that improve the accessibility of heterogeneous system resources without obscuring performance-critical system properties and that help developers to reduce the amount of data that has to be exchanged among heterogeneous system resources.
Introduction

The goal of this thesis is to address these challenges by proposing programming abstraction approaches for each type of heterogeneous system resource considered in this work (cf. Section 1.1). In the context of this thesis, libraries and frameworks that hide certain complexities of the underlying hardware or the programming models therewith are considered as programming abstractions. For this endeavor, each approach has to factor in the peculiarities of the corresponding resource type:

On-Chip Compression Accelerators Originally intended for the use-case of transparent main-memory compression [17], one goal of this thesis is to enable the use of the NX-842 on-chip compression accelerator available in IBM POWER CPUs to improve the efficiency of data transfers across heterogeneous system resources. On POWER7+ and POWER8 CPUs, the NX-842 on-chip compression accelerator is only accessible from kernel-space using the privileged Initiate Coprocessor Store Word Indexed (icswx) instruction and is therefore inaccessible from user-space applications. Even though the Virtual Accelerator Switchboard (VAS) facilities have been introduced in the POWER9 microarchitecture with the goal of providing user-space applications access to on-chip accelerator resources such as the NX-GZIP compression accelerator, access to the NX-842 units via the VAS facilities is still restricted to kernel-space unless the skiboot firmware is modified correspondingly. Therefore, a programming abstraction is necessary that exposes kernel-space resources to the user-space in order to make the compression facilities of the NX-842 accessible for user-space applications. The two NX-842 units available per POWER CPU are tightly integrated and can process up to 36.8 GB/s, exceeding even the throughput of dedicated compression accelerators attached via PCIe 4.0.

Scale-Out GPU Clusters The demand for GPU compute resources has been steadily increasing over the last few years to the point where workloads such as deep learning applications require entire GPU clusters [84] to satisfy their demand for compute resources. Application development for scale-out GPU workloads is very challenging, as developers have to be adept using both data parallel programming models (e.g., Open Computing Language (OpenCL)) and distributed memory parallel programming models (e.g., Message Passing Interface (MPI)). To produce relief, a programming abstraction mechanism is required that enables developers to focus on a single parallel programming model. However, data transfers between CPUs and GPUs can already be a bottleneck in single node scenarios. The limited bandwidth available 10 Gbit/s, 25 Gbit/s, and even 40 Gbit/s Ethernet networks which are still the norm in the vast majority of data centers [19] necessitates that the volume of data transfers is kept minimal in order not to impede the scalability of scale-out GPU workloads.

Non-Uniform Memory Access Systems Even though CPUs have become popular in many data-intensive application domains, many workloads still rely on the flexibility and versatility of multicore CPUs [208]. While several of these CPU-based workloads can be adapted to scale-out across multiple systems to provide sufficient compute resources, certain workloads such as In-Memory Databases (IMDBs) [25] or de novo genome assembly [133] are inherently hard to scale out and therefore require as many resources as possible in a single scale-up system. For such workloads, state-of-the-art NUMA systems support
Research Question

The research question of this thesis is concerned with improving the accessibility of heterogeneous system resources for applications developers. A central assumption of the research question is that suitable programming abstractions can help to address two major challenges inherent to heterogeneous system resources: Foremost, this thesis hypothesizes that a certain degree of the complexity conditioned by the large variety of heterogeneous system resources considered in the context of this work can be encapsulated using programming abstractions without obscuring performance-critical system properties. Furthermore, this thesis hypothesizes that these abstractions can help to mitigate the performance penalty associated with data transfers across heterogeneous system resources by either improving data transfer efficiency or by avoiding unnecessary data transfers altogether. For these endeavors, it is of utmost importance to find the right balance between providing a sufficient degree of abstraction on one without burying the heterogeneous system resources under too many layers of abstraction on the other hand.

Contributions

The author of this thesis, provides several contributions to the field of software engineering. First, the author proposes a programming abstraction mechanism in the form of the lib842 compression library that facilitates user-space access to the resources of NX-842 on-chip compression accelerators. To enable interoperability of data compressed using the proprietary 842 compression algorithm with arbitrary CPUs or GPUs, the library also provides the first publicly available user-space facilities for software-based 842 compression and decompression on CPUs, as well as OpenCL-based decompression on GPUs. Both the hardware as well as the software-based compression facilities provide sufficient throughput to saturate 10 Gbit/s, 25 Gbit/s, and even 40 Gbit/s Ethernet networks which are still the norm in the vast majority of data centers. Second, the author introduces the CloudCL framework which provides a single-paradigm programming experience for scale-out GPU workloads. By abstracting away many aspects of the distributed memory parallel programming model commonly used in scale-out scenarios, the framework allows application developers to focus on the data parallel programming model associated with
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To warrant efficient data exchange across compute nodes of a GPU cluster, the CloudCL framework implements transparent on-the-fly data compression based on the lib842 compression library. Third, the author provides a comprehensive evaluation of the impact of the programming abstractions for NUMA-aware C++ application development provided by the PGASUS framework on both developer experience and application performance. To facilitate reproducibility, all software-artifacts presented or used in this thesis are curated in a freely accessible repository.

1.5 Overview

This thesis is structured as follows. After the introduction, Chapter 2 analyses the state of the art of heterogeneous system resources and presents related work from the field of software engineering. Chapter 3 presents the concept, implementation, as well as the evaluation of the lib842 compression library which provides user-space access to the hardware-based compression facilities of the NX-842 on-chip compression accelerator. Focusing on scale-out GPU workloads, Chapter 4 highlights the programming abstraction facilities provided by the CloudCL framework that enable application developers and domain experts to focus their efforts on the data parallel programming model associated with GPUs. The improved developer experience offered by CloudCL is demonstrated in Chapter 4 by showcasing CloudCL-based implementations of two exemplary workloads. To improve the efficiency of data-exchange across compute nodes of a GPU cluster, Chapter 4 also presents the concept, implementation and evaluation of transparent data compression based on the lib842 compression library. Chapter 5 provides an overview of the programming abstractions provided by the PGASUS framework for NUMA-aware C++ application development, which has been proposed in the master’s theses of Wieland Hagen [68] and Karsten Tausche [198]. The improved developer experience facilitated by the PGASUS framework is demonstrated in Chapter 5 by comparing PGASUS-based implementations of three exemplary workloads to NUMA-aware implementations based on the Open Multi-Processing (OpenMP) interface or the pthreads library. As another central aspect of Chapter 5, the performance impact of the PGASUS framework is evaluated. Chapter 6 discusses the contributions of this thesis and reflects on potential links for future research before the thesis concludes in Chapter 7.

1.6 Publications

I have already published partial results about many central aspects of the major contributions I am going to present in the following chapters of this thesis. With this section I would like to provide a list of the papers that have directly shaped the contributions I am presenting in this thesis and which have been published in peer-review workshops, conferences, and journals. In my research efforts leading to these publications, I have been supported by the work of many colleagues and master’s students, with their support being acknowledged in each publication’s respective list of authors.

https://github.com/plauth/thesis-artifacts

6
1.7 Context

Over the years that have led to this work, I had the honor and pleasure to work with many wonderful colleagues in several research projects, collaborations, or teaching activities. Therefore, the goal of this section is to provide a brief overview of the activities that have both accompanied and influenced my work.

1.7.1 Scalable and Secure Infrastructures for Cloud Operations (SSICLOPS)

Funded from the European Union’s Horizon 2020 research and innovation program 2014-2018 under grant agreement No. 644866, the SSICLOPS project lasted from February 2015 to January 2018. As outlined in Figure 1.3, the major goal of the project was to investigate resource management strategies in federated private cloud infrastructures. Operating under the guiding principle “resource management from core to cloud”, the work package dealing with workload and data placement strategies in private cloud infrastructures
Figure 1.3: The SSICLOPS project has investigated federated private cloud infrastructures. 

turned out to be a productive environment for my colleague Felix Eberhardt and me to investigate workload and data placement strategies on the scope of NUMA systems. In our joint research efforts centered around NUMA systems, we have co-supervised the master’s thesis by Wieland Hagen, which has yielded the initial version of the PGASUS framework. PGASUS provides the foundation for my contributions I am going to present in Chapter 5. Beyond the NUMA topic, I have supervised the seminar project by Karsten Tausche, in which he investigated the dOpenCL API forwarding library to evaluate whether GPU resources can be virtualized on the abstraction level of the OpenCL interface. The successful evaluation yielded by this seminar has served as the basis for the Dynamic OpenCL framework presented in the master’s thesis by Florian Rösler, which again has laid the groundwork for my work on the CloudCL framework, which I am going to present in Chapter 4. Together with my colleagues I have undertaken further research efforts in the work package on policy and security as well as the work package on scenario integration. Even though these research efforts do not align with the topics covered in my thesis, they have yielded several joint publications.

1.7.1 Project-Related Publications

The joint research efforts I have conducted together both with my colleagues Felix Eberhardt and Stefan Halfpap (Klauck) from HPI as well as our collaboration partners from the SSICLOPS project have yielded several peer-reviewed publications. Even though these publications are not necessarily related to my thesis or have only influenced my research efforts indirectly, they are listed hereinafter to highlight the research activities I have conducted alongside my work on this thesis:


Max Plauth, Christoph Sterz, Felix Eberhardt, Frank Feinbube, and Andreas Polze. “Assessing NUMA Performance Based on Hardware Event Counters”. In: Proceedings of the IEEE International Parallel and Distributed Processing Symposium Workshops (IPDPSW). IEEE. May 2017, pages 904–913. DOI: 10.1109/IPDPSW.2017.51


1 Introduction


### 1.7.1.2 Project-Related Master’s Theses

In the context of the SSICLOPS project, I have (co-)supervised the following master’s theses:


In the master’s programs at HPI, the master’s thesis is preceded by a larger project comparable to the scope of a final year project. Over the course of the SSICLOPS project, I have (co-)supervised the following master’s projects:


The Hybrid DB project has been conducted in cooperation with IBM Germany Research and Development from 2016 through 2021 as a follow-up to preceding research efforts centered around NUMA systems undertaken by my colleague Felix Eberhardt. Hence, a central aspect of the Hybrid DB project was the investigation of heterogeneous properties of main memory resources in large scale-up NUMA systems. In this continuation of NUMA-related research, I have supervised the master’s thesis by Karsten Tausche in which he has contributed improvements to the PGASUS framework. PGASUS serves as the
foundation for my contributions I am going to present in [Chapter 5]. As another central aspect of the HybridDB project was the investigation of new approaches for interacting with heterogeneous system resources, the project provided me with an ideal environment for focusing my research efforts on this topic. In this environment, I have co-supervised the master’s theses by Lukas Wenzel [209] and Robert Schmid [181], who have explored programming abstractions for leveraging coherently integrated Field-Programmable Gate Array (FPGA) accelerators for data-intensive workloads such as IMDBs leveraging the then unique Coherent Accelerator Interface Architecture (CAIA) facilities introduced with the POWER8 microarchitecture. Their work on data-intensive workloads has sparked my research interests centered around the NX-842 on-chip compression accelerators, which I am going to introduce in [Chapter 3]. Building up on top of the work of Karsten Tausche (cf. dOpenCL [199]) and Florian Rösler (cf. Dynamic OpenCL [175]) conducted within the framework of the SSICLOPS project (cf. Section 1.7.1), I have furthermore investigated strategies for virtualizing GPU compute resources in multi-tenant scenarios in the context of the Hybrid DB project. For this work, I have been awarded with the IBM PhD Fellowship Award in 2017. In these GPU-related research efforts, I have proposed the CloudCL framework which I am going to present in [Chapter 4].

**Figure 1.4:** Introduced with the POWER8 microarchitecture, the CAIA facilities are comprised of the Coherent Accelerator Processor Proxy (CAPP) on the side of the host CPU and the POWER Service Layer (PSL) on the side of the accelerator, with both components communicating via the Coherent Accelerator Processor Interface (CAPI) protocol. Image source: [209].

### 1.7.2.1 Project-Related Publications

The research efforts I have conducted together with my colleagues Felix Eberhardt, Robert Schmid, and Lukas Wenzel have yielded several peer-reviewed publications. Even though these publications are not necessarily related to my thesis or have only influenced my research efforts indirectly, they are listed hereinafter to highlight the research activities I have conducted alongside my work on this thesis:


1.7 Context

1.7.2.2 Project-Related Master’s Theses

In the context of the Hybrid DB project, I have (co-)supervised the following master’s theses:


1.7.2.3 Project-Related Master’s Projects

In the master’s programs at HPI, the master’s thesis is preceded by a larger project comparable to the scope of a final year project. Over the course of the Hybrid DB project, I have (co-)supervised the following master’s projects:


1.7.3 Memento: Energy-Efficient Memory Placement

Together with my colleagues Sven Köhler and Lukas Wenzel, I have performed preliminary research in collaboration with the Bochum Operating Systems and System Software (BOSS) research group at the Ruhr University Bochum (RUB) headed by Prof. Dr. Timo Hönig from 2019 through 2021, which has led to the submission of the joint Memento proposal to the Priority Program 2377 on Disruptive Memory Technologies by the German Research Foundation. Based on an ISA-agnostic workload representation, our initial goal was to improve energy-efficiency in data centers by placing workloads on the resources that are best suited for the respective workload as outlined in Figure 1.5

Figure 1.5: The goal of the proposed Memento project is to improve energy efficiency based on memory placement strategies that consider the varying characteristics of diverse memory resources available in state-of-the-art computer architectures. Image source: [79].

During the initial phase of the preliminary work, our research efforts were focused on establishing common APIs for measuring the energy demand of workloads across various hardware using the Pinpoint utility [104]. However, as the diversification of memory resources has culminated in the approval of the Priority Program 2377 on Disruptive Memory Technologies, we have refocused our research efforts on exploiting the heterogeneity of memory resources with the goal of improving energy efficiency. Even though a funding
decision for Memento proposal is still pending at the time of writing, we are hoping to commence the active work on the Memento project some time around summer 2022.

### 1.7.3.1 Project-Related Publications

The joint research efforts I have conducted together both with my colleagues Sven Köhler and Lukas Wenzel from HPI as well as our collaboration partners Timo Hönig and Benedict Herzog from RUB have yielded several peer-reviewed publications. Even though these publications are not necessarily related to my thesis or have only influenced my research efforts indirectly, they are listed hereinafter to highlight the research activities I have conducted alongside my work on this thesis:


### 1.7.3.2 Project-Related Master’s Theses

During the preparatory work leading to the submission of the Memento proposal, I have (co-)supervised the following master’s theses:


### 1.7.3.3 Project-Related Master’s Projects

In the master’s programs at HPI, the master’s thesis is preceded by a larger project comparable to the scope of a final year project. Over the preparation phase leading to the submission of the Memento proposal, I have (co-)supervised the following master’s projects:
1 Introduction


1.7.4 Teaching Activities

The research efforts leading to this work were accompanied by various teaching activities specified hereinafter. Especially the lectures on parallel programming and heterogeneous computing are tightly associated with the topics covered in this thesis, whereas the remaining courses are motivated by the research projects highlighted in the preceding sections.

| Winter 2015/16 | Parallel and Distributed Systems Project Seminar, Master |
| Summer 2016 | IBM Power Systems Block Lecture, Bachelor and Master |
| Winter 2017/18 | File System From Scratch Project Seminar, Master |
| Winter 2017/18 | Embedded Operating Systems Lecture, Master |
| Winter 2018/19 | IBM Power Systems Block Lecture, Bachelor and Master |
| Summer 2019 | Parallel Programming and Heterogeneous Computing Lecture, Master |
| Winter 2019/20 | Energy-Aware Computing in Heterogeneous Data Centers Project Seminar, Master |
| Summer 2020 | Parallel Programming and Heterogeneous Computing Lecture, Master |
| Summer 2021 | Parallel Programming and Heterogeneous Computing Lecture, Master |
2 State of the Art and Related Work

This chapter presents the state of the art and related work in the research field of heterogeneous computer architectures. To deepen the appreciation of the heterogeneous system resources considered in this thesis, this chapter starts by elaborating on the origins of using Graphics Processing Unit (GPU) resources for general purpose computations as well as the development of Non-Uniform Memory Access (NUMA) architectures. Afterwards, upcoming trends in heterogeneous computer architectures are discussed to point out upcoming challenges for the use of heterogeneous system resources. Finally, the canonical approaches for interacting with the heterogeneous system resources covered by this thesis are discussed alongside approaches that provide programming abstractions to deepen the understanding of the complexities application developers are confronted with by heterogeneous system resources.

2.1 The Origins of Heterogeneous System Resources

Revisiting the origins of heterogeneous system resources helps to deepen the understanding of today’s heterogeneous computer architectures. Therefore, this section begins with elaborating on how GPU resources have gained general purpose computing capabilities. Afterwards, the formation of modern NUMA architectures is revisited to stress the omnipresence of heterogeneous memory resources in most modern server systems.

2.1.1 Non-Uniform Memory Access Architectures

Of the various parallel architectures available in the 1980s, Symmetric Multiprocessing (SMP) and message-passing emerged as the major multiprocessor architectures [71]. Especially for smaller multiprocessor systems, shared memory SMP systems were prevalent, using a bus to interconnect all processors with main memory and I/O resources [71] as illustrated in Figure 2.1. The success of bus-based SMP systems lies in the circumstance, that smaller instances approach the properties of the Paracomputer model [106], in which “identical processors (each with a conventional order-code set) share a common memory which they can read simultaneously in a single cycle” [135]. Using a shared medium to attach the processors to main memory alleviated the introduction of coherent caches based on bus snooping in order to reduce memory access latencies in these Uniform Memory Access (UMA) systems in practice [71]. For a larger number of processors however, the traffic caused by the snooping-based coherence mechanisms results in bus contention, limiting the scalability of this approach to configurations ranging between 4 and 16 processors per system, depending on the system at hands [172][71].

Aiming towards better scalability, Distributed Shared Memory (DSM) architectures emerged in the late 1970s and early 1980s, giving up cache coherence to overcome the
scaling limitations of bus-based 
SMP systems. Notable DSM multiprocessor approaches include the Carnegie Mellon Cm* [197], the IBM RP3 [147], the Honeywell Information Systems Italia (HSI) XPS-100 series [32], [200], and the Bolt, Baranek, and Newman (BBN) Butterfly [108]. As depicted in Figure 2.2 each processor in a DSM architecture is equipped with local main memory and I/O resources, and all processors are interconnected using a scalable interconnection network. Even though memory resources are physically distributed in this approach, the shared memory programming model still applies, as all memory resources are accessible through a single shared address space [71]. As a result of the shared address space, each processor can access remote memory resources attached to another processor through the scalable interconnection network using load and store semantics. Without a shared medium as an interconnection network, snooping-based coherency protocols were no longer feasible [71]. From an application developers perspective, DSM systems were much more challenging to develop for, as remote data access operations could not be cached due to the lack of cache coherency, resulting in remote access times that could take 10 times longer compared to local access operations [71]. With varying access times being a performance-critical property of DSM architectures, they are more commonly referred to as NUMA architectures. Due to the vast performance penalty of uncached remote access operations in such NUMA systems, developers had to carefully decide which data should be shared [187].

To combine the scalability of DSM architectures with the application developer productivity of bus-based SMP architectures, directory-based schemes for maintaining cache coherence in large multiprocessor systems have been proposed as a promising approach in 1988 [4]. By augmenting each processor with the local partition of a distributed directory as visualized in Figure 2.3, the coherence state of cache lines can be tracked without congesting the interconnection network. With the Stanford DASH multiprocessor, the first Cache Coherent Non-Uniform Memory Access (ccNUMA) multiprocessor architecture has been introduced in 1992 [112]. In the DASH prototype, Silicon Graphics Power Station

Figure 2.1: In SMP architectures, the bus used to interconnect all processors with main memory and I/O resources can be exploited to implement coherent caches based on bus snooping. However, the shared medium also represents a bottleneck, limiting the scalability of the approach. Figure adapted from [71].
2.1 The Origins of Heterogeneous System Resources

![Diagram of heterogeneous system resources](image)

**Figure 2.2:** DSM architectures avoid the bottleneck of SMP systems by equipping each processor with local main memory and I/O resources at the cost of giving up coherent caches, making remote memory access extremely costly. Figure adapted from [71].

4D/340 4-way SMP systems were used as base clusters. By augmenting the memory bus of a base cluster system with directory memory and an inter-cluster interface, up to 16 base clusters could be interconnected to form a ccNUMA system with up to 64 Central Processing Units (CPUs). Another important contribution of the DASH project is that it formalized weak memory consistency models, including release consistency as further means to improve the scalability of multiprocessor architectures [57].

![Diagram of ccNUMA system](image)

**Figure 2.3:** By augmenting the local memory resources with directories for tracking the location of cache-lines, ccNUMA systems manage to combine the ease of use of SMP systems with the scalability of DSM architectures. Figure adapted from [71].

Defining a directory-based, coherent high-speed interconnection standard, the approval of the Scalable Coherent Interface (SCI) (IEEE 1596-1992) paved the way for the first wave of commercially available ccNUMA systems. The Convex Exemplar SPP-1000 series intro-
duced in 1994 marks the first commercial ccNUMA system on the market. Based on SCI, the Exemplar SPP-1000 was available in configurations with up to 16 hypernodes comprised of 8 HP PA-7100 CPUs each \[20\]. In 1996, the Sequent NUMA-Q series, internally dubbed Sequent: The Next Generation (with Intel inside) (STiNG), has been introduced, which employed SCI to interconnect up to 16 quads comprised of 4 Intel Pentium Pro CPUs \[119\]. A similar approach has been taken with the Data General AViiON AV 2000 series introduced in 1997, which also relied on SCI to interconnect up to 8 quads comprised of 4 Intel Pentium Pro CPUs \[35\]. All three approaches have in common that like the DASH prototype, they used conventional, bus-based SMP configurations as building blocks and augmented them with glue logic to interconnect multiple blocks. In this glue-based approach, bus-based snooping to used to implement coherency on the building block level, whereas the directory-based coherence mechanisms of SCI are used to maintain coherence across building blocks.

While the glue-based approaches provided the flexibility that unmodified CPU designs could be used, they came at the cost of increased remote access latencies caused by the two-leveled design. With the Silicon Graphics Origin 2000 series, the first commercially available glueless ccNUMA architecture was introduced in 1997 \[110\]. Unlike the glue-based approaches, the two CPUs in each node do not form a bus-based SMP cluster, but are directly connected to the interconnection Hub. In theory, the employed architecture would have supported a maximum of 1024 CPUs, but customer configurations were only available with up to 128 CPUs and only one system with 512 CPUs was installed internally at Silicon Graphics.

With a certain delay, all major CPU vendors adopted ccNUMA architectures in their processor designs. IBM followed by introducing a ccNUMA design in their POWER 4 processors in 2002 \[202\]. In 2003, the Digital Equipment Corporation (DEC) adopted a ccNUMA architecture in their Alpha 21364 (EV7) processors \[57\]. With the introduction of the AMD Opteron series of processors also in 2003, a ccNUMA based design entered the mass market \[92\]. Finally, Intel caught up in 2009 by introducing ccNUMA based designs in their lineup of both x86_64 and Itanium processors with the Nehalem \[107\] and Tukwila \[190\] microarchitectures, respectively.

Since ccNUMA approaches have become the predominating manifestation of NUMA architectures since the 1990s, the prefix for indicating the cache coherent variant is commonly omitted and the term NUMA is used to refer to ccNUMA architectures.

### 2.1.2 GPU Computing

An initial level of programmability has made its entry into professional graphics hardware in the mid 1980s, as programmable graphics architectures have been introduced as an alternative to fixed rendering pipelines \[42\]. To achieve programmability, early examples such as the Pixar CHAP architecture \[113\] or the Ikonas platform \[42\] employed microcodable Single Instruction Multiple Data (SIMD) processors in order to process vertex and pixel data in parallel. In the late 1990s, programmable Multiple Instruction Multiple Data (MIMD) architectures such as PixelFlow \[44\] or the Silicon Graphics InfiniteReality \[132\] system became available in the high-end range of professional graphics workstations.

The notion of exploiting CPU hardware for general purpose computations was triggered by the introduction of programmable vertex and fragment shaders in consumer
2.1 The Origins of Heterogeneous System Resources

GPUs Quickly after the introduction of corresponding hardware such as the NVIDIA GeForce 3 or the ATI Radeon 8500 in 2001, an attempt at offloading matrix multiplications to GPUs has been presented [109]. Even though this first approach was unable to achieve performance improvements compared to CPU-based execution, it marks an important milestone as the first successful attempt to use GPUs for general purpose computations. Only one year later in 2002, another approach has managed to break even with CPU performance, achieving a $3.2 \times$ speed-up for GPU-based matrix multiplication using $1500 \times 1500$ matrices [205].

Despite the high potential of using GPUs for general purpose computations, widespread adoption of this approach was limited as it was very taxing even for application developers with in-depth knowledge of graphics Application Programming Interfaces (APIs) such as OpenGL to set up compute workflows [23]. To simplify the process, several third party compute libraries started to appear in the mid 2000s, with Brook [23] being one of the most influential examples. However, even with such compute libraries at hands, early generations of consumer GPUs with programmable shaders were still limited in various ways, such as the number of instructions that could be used per shader. On an architectural level, the most limiting factor was that fixed pipelines with separate vertex and pixel processors were used, which limited their flexibility for graphics workloads as well as for general purpose computations.

![Diagram of shader architecture]

**Figure 2.4:** Earlier generations of GPUs equipped with programmable shaders suffered from the limitation that each stage of the graphics pipeline required dedicated hardware, making it hard for vendors to decide how much of their chip area they want to spend on what types of resources to provide decent performance across a wide range of applications. Image source: [178].

As the first approach to replace fixed pipeline designs with unified shader architectures, the TerraScale microarchitecture by ATI was introduced in the form of the Xenos GPU employed in the Microsoft Xbox 360 gaming console [59]. The TerraScale microarchitecture employed a SIMD architecture based on Very Long Instruction Word (VLIW) characteristics with the goal of maximizing Instruction-Level Parallelism (ILP) whereas the Tesla microarchitecture presented by competitor NVIDIA in 2006 used a SIMD architecture based on Reduced Instruction Set Computer (RISC) characteristics, which exploits
Thread-Level Parallelism (TLP)\cite{116}. While graphics workloads were able to benefit from unified shader architectures, the novel architecture had an even larger impact on compute tasks and paved the way for a widespread adoption of employing GPUs as general purpose compute accelerators. In 2007, NVIDIA released the initial version of the Compute Unified Device Architecture (CUDA) framework\cite{98} to officially support and promote the use NVIDIA GPUs for general purpose computations. With similar capabilities, the Khronos group has released the initial version of the OpenCL specification\cite{137} as a vendor-independent alternative to CUDA, also including support for other compute resources such as CPUs, Digital Signal Processors (DSPs), Field-Programmable Gate Arrays (FPGAs), and others.

![Diagram](Image-Source-178)

**Figure 2.5:** In unified GPU architectures, the special-purpose resources employed in fixed-pipeline GPU designs are replaced with generic compute resources that can be used to process most stages of a graphics pipeline, greatly increasing the flexibility of GPUs. Image source: \cite{178}.

Over the course of the 2010s, all major GPU microarchitectures adopted SIMD compute resources with RISC characteristics. Not only dedicated GPUs improved continuously, but also integrated GPUs advanced significantly as GPU and CPU resources reached a much tighter level of integration, sharing the same memory hierarchy not only physically but also on a logical level\cite{165}. With the introduction of the IBM Power System AC922 in 2018, the combination of IBM POWER9 CPUs, NVIDIA V100 GPUs, and the cache-coherent NVLink 2.0 interconnection technology have yielded the first system setup to achieve cache-line level integration of dedicated GPUs into the main memory hierarchy of the host CPUs\cite{171}.
2.2 Trends in Heterogeneous System Resources

At the time of writing, innovations in the area of interconnection standards are driving extensive transformations in heterogeneous computer architectures. Most notably, the memory hierarchy is about to become more diverse and an increasing number of accelerators such as GPUs, FPGAs, or other domain-specific accelerators are gaining fine-grained, coherent integration into the host system. In this section, the major aspiring interconnection standards are reviewed in Section 2.2.1 and the importance of disruptive memory technologies are outlined in Section 2.2.2.

2.2.1 Coherent Interconnects

Introduced in 2003, the Peripheral Component Interconnect Express (PCIe) standard has served as the standard intra-node interconnect technology for almost two decades. As illustrated in Figure 2.6, frequent releases of new revisions of the PCIe standard roughly every 3.5 years have yielded continuous bandwidth improvements on par with the progression of memory bandwidth available per CPU socket throughout the first three revisions of the standard [63, 64, 65]. After the release of PCIe 3.0 however, the pace of development has slowed down significantly, as the first commercially available product to support the succeeding PCIe 4.0 standard was released seven years later in 2017 with the introduction of the IBM POWER9 CPU [177, 66]. Unfortunately, widespread adoption of the newer standard has only started with the introduction of the AMD Zen 2 CPU microarchitecture in 2019 [196], but even at the time of writing not all major CPU vendors have adopted the PCIe 4.0 standard.

Over the extensive lifespan of PCIe 3.0 as the dominating standard, continuous improvements in the performance of CPUs and accelerators have been accompanied by proportionate improvements of their respective memory subsystems. Due to the overdue progression of interconnect technology, the severe gap between the capabilities of interconnects and the performance of CPUs and accelerators as well as their respective memory subsystems have created a strong need for innovation in the field of interconnect technologies. Consequently, several consortia have formed to bring forward a new generation of interconnection standards such as Open Coherent Accelerator Processor Interface (OpenCAPI), Compute Express Link (CXL), Cache Coherent Interconnect for Accelerators (CCIX), and Gen-Z. Differing in technical details, all of these approaches have in common that they do not only facilitate improved bandwidth and latency, but they also introduce new features such as coherent integration of devices into the main memory hierarchy, as well as serial facilities for attaching memory. While OpenCAPI, CXL and CCIX are intended for short-reach, intra-node connectivity, the Gen-Z specification has been drafted with rack-scale connectivity in mind. In addition to vendor-independent technologies, proprietary approaches such as NVIDIA NVLink have been introduced.

2.2.2 Disruptive Memory Technologies

Over the last decades, most computer architectures have employed homogeneous memory resources on each layer of the memory hierarchy. In these systems, the properties of memory resources were sufficiently similar to conceal them behind the flat address
state of the art and related work

Figure 2.6: Over the years, a certain gap has emerged between the development of Dynamic Random Access Memory (DRAM) bandwidth compared to the bandwidth provided by the PCIe standard. The pressure to innovate created by this discrepancy has fueled a competition for next-generation interconnection technologies. The figure has been adapted from [78] and augmented with additional data from [176].

Figure 2.6

space of the virtual memory abstraction. However, the recent introduction of various novel memory technologies has initiated a fundamental shift in the design of computer architectures towards supporting heterogeneous memory resources on various layers of the memory hierarchy. The near end of the memory hierarchy is extended with large, potentially self-managed caches based on stacked Static Random-Access Memory (SRAM) or on-package High-Bandwidth Memory (HBM) memory [212 15]. One step further out in the memory hierarchy, volatile DRAM resources are augmented with Graphics Double Data Rate (GDDR) memory or HBM resources for memory-bound algorithms and non-volatile memory for persistent data, either exclusively [213] or in combination with conventional Double Data Rate (DDR) memory resources. Originating from the respective ecosystems of the OpenCAPI and CXL interconnection standards, the mixed used of different memory technologies is enabled based on the introduction of technology-agnostic memory interfaces [191]. At the far end of the memory hierarchy, the new interconnection technologies discussed in Section 2.2.1 also provide the foundation for disaggregated memory resources, which have the potential to offer enormous memory capacities [148]. The diversity of employable memory technologies implies a disruptive degree of heterogeneity regarding the characteristics of memory resources, which have to be considered.

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2.3 Programming Abstractions for Heterogeneous System Resources

The prevalence of heterogeneous system resources has implied a wide range of programming abstractions that have been presented over the last decade. To put the programming abstractions proposed in this thesis in perspective with the latest insights from the field, this section provides an overview of programming abstractions and related approaches for the heterogeneous system resources considered in this work.

2.3.1 Memory Compression and Compressed Data Transfers

Using compression as means for improving utilization of main memory has a well-established history. This is well reflected by the work of Mittal et al. [131], which provides a comprehensive survey of the widely researched field of data compression mechanisms for CPU-based main memory and cache resources. Regarding memory compression on GPUs, a number of approaches have been published as well. Targeting memory-bound applications on GPUs, Sathish et al. [179] have proposed using hardware-based compression to increase the efficiency of access to off-chip device memory, yielding up to 37% better performance compared to the uncompressed case. A similar approach has been published by Vijaykumar et al. [207], who are also employing memory and register compression to increase the utilization of all GPU resources, yielding up to 2.6× speed-up across a variety of memory-bound applications. Following the same goal, Lu et al. [120] have recently proposed a low-latency, hardware-based compression architecture optimized for floating point data that reduces bandwidth demand and energy consumption of GPUs by 44.46% and 44.34%, respectively. Focusing entirely on the register level, Lee et al. [111] have explored register compression on GPUs with the goal of reducing the energy consumption of graphics hardware. All approaches have in common that they are using custom, domain-specific compression algorithms instead of general-purpose compression algorithms and rely on custom hardware designs. With the introduction of their latest A100 GPUs, NVIDIA has introduced hardware support for device-sided memory compression with the Compute Data Compression feature [143], promising up to 4× improvements in effective DRAM and L2 bandwidth.

Both CPU-based and GPU-base approaches for memory compression can improve both effective bandwidth and capacity of memory resources significantly. Regrettably, all approaches presented thus far only consider the isolated scope of the memory resources attached to either a CPU or a GPU only and do not consider the scenario of exchanging data across devices in compressed form. Patel et al. [145] have explored the feasibility of on-the-fly data compression for data transfers between CPU and GPU using a generic compression algorithm. The authors conclude that on-the-fly data compression is not feasible using their software-based compression approach implemented on the CPU. Using a data-specific compression algorithm however, Kaczmarzski et al. [86] have successfully demonstrated that on-the-fly data compression can be used to speed-up transfers between main memory and GPU memory for CPU-based In-Memory Database (IMDB) use cases. Khavari Tavana et al. [93] have also investigated GPU-based compression approaches using compression algorithms tailored to the characteristics of floating point data. In
contrast to improving transfer efficiency between CPU and GPU, they are using on-the-fly data compression to improve data transfers among multiple GPUs.

### 2.3.2 General Purpose Computing on GPUs

When the concept of performing general-purpose computations on GPUs emerged, in-depth knowledge of graphics APIs were necessary to set up compute workflows based on fragment shaders. Considering these origins, fundamental frameworks for GPU computing Open Computing Language (OpenCL) and CUDA provide a decent level of abstraction. However, further approaches have been presented with the goal of abstracting the use of GPUs both on the single-node level and in scale-out deployments. The goal of this section is to provide a brief overview of abstractions for GPU compute resources ranging from the basics at the level of fundamental compute frameworks such as OpenCL to high-level abstractions concerned with scale-out GPUs resources.

#### 2.3.2.1 OpenCL

The OpenCL standard [204] defines a framework for executing parallel compute kernels on heterogeneous compute resources available in a single system such as CPUs, GPUs, FPGAs, DSPs, and possibly even further device types. At a conceptual level, the OpenCL standard is built around four models: a platform model, an execution model, a memory model, and a programming model.

The platform model describes the basic hierarchy of a host that is equipped with an arbitrary number of compute devices, which are comprised of compute units as an organizational structure for grouping individual processing elements. An OpenCL implementation must provide a platform in the form of an Installable Client Driver (ICD), which enables the host to control compute devices belonging to the platform. Multiple platforms may even coexist on a single host, as multiple ICDs may be available on a system (e.g., for using heterogeneous compute resources from diverse vendors).

The execution model distinguishes between the host application and device kernels. To provide an organizational structure for managing kernel instantiations, the execution model defines an index range that comprises a hierarchy of work-items and work-groups in order to specify what data individual kernel instances operate on. A work-item represents the work performed by a single kernel instance, whereas work-groups provide an organizational unit for grouping multiple work-items. In a context, the host application defines devices, kernels, program objects, and memory objects. Furthermore, the host application is responsible for managing queues, which are used to define a sequence of kernel executions, operations on memory objects (e.g., transfers between host and device memory), as well as synchronization points.

The memory model is tightly complemented with the peculiarities of the platform model and the execution model and defines different types of memory that vary in their scope of accessibility as well as their performance characteristics. Main memory resources of the host system are referred to as host memory. On the side of compute devices, the memory hierarchy is composed of global memory, constant memory, local memory, as well as private memory. The global memory and constant memory resources comprised by a context are typically served by off-chip GDDR memory or HBM and can be accessed by both the host...
and kernels. In contrast to that, local memory is often backed by on-chip SRAM and is only shared among the work-items belonging to a single work-group. Private memory refers to memory resources at the register level and is therefore only accessible from the work-item it belongs to.

With OpenCL supporting a wide range of heterogeneous compute resources, the employed programming model is flexible enough to support both a data parallel programming model and a task parallel programming model. For the data parallel case, a strict one-to-one mapping between a few data elements and work-items is used, whereas the task parallel case can be achieved by defining an index-range with a single work-item per task. OpenCL kernels are implemented using OpenCL C, which is a superset of C99, containing additional keywords that are used to specify the employed memory resources defined by the memory model as well as synchronization mechanisms. The host application is usually implemented in C or C++, however other language bindings exist as well.

Listing 2.1 presents a minimal OpenCL kernel implementing the addition of two vectors. Each parameter is a pointer to an array of float values residing in the global memory of the heterogeneous compute resource. As the kernel is instantiated for every work-item, each kernel instance identifies its work-item by querying its index using the in-built function get_global_id.

Listing 2.1: Exemplary OpenCL vector addition kernel.

```c
__kernel void vec_add(__global float *a, __global float *b, __global float *c) {
  int i = get_global_id(0);
  c[i] = a[i] + b[i];
}
```

While the simplistic vector kernel amounts to very few lines of kernel code, OpenCL is notorious for its verbosity regarding the host code necessary to choose an appropriate device, initiate data transfers, execute the kernel, and perform other auxiliary tasks. Even though the host code exemplified in Listing 2.2 uses the C++ bindings of OpenCL which are much less verbose than the native C API, roughly 30 lines of code are necessary to execute the vec_add kernel. Using the native C API, the same application requires roughly 50 lines of code.

Listing 2.2: Minimal C++ OpenCL host application necessary to execute the vec_add kernel.

```c
int main(){
  // Initialize arrays on host
  float array_a[10] = {0.0, 1.0, 2.0, 3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0};
  float array_b[10] = {0.0, 1.0, 2.0, 3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0};
  float* array_result = new float[10];
  size_t size = 10 * sizeof(float);

  // OpenCL initialization
  cl::Platform platform;
  cl::Platform::get(&platform);
  std::vector<cl::Device> devices;
  platform.getDevices(CL_DEVICE_TYPE_GPU, &devices);
  cl::Context context(devices);
```
cl::Program program(context, OPENCL_PROGRAM);
program.build(devices);
cl::Kernel kernel(program, "vec_add");
cl::CommandQueue queue(context, devices[0]);
cl::Buffer d_a(context, CL_MEM_READ_ONLY, size);
cl::Buffer d_b(context, CL_MEM_READ_ONLY, size);
cl::Buffer d_c(context, CL_MEM_WRITE_ONLY, size);

// data transfers & kernel invocation
queue.enqueueWriteBuffer(d_a, CL_FALSE, 0, size, array_a);
queue.enqueueWriteBuffer(d_b, CL_FALSE, 0, size, array_b);
queue.enqueueNDRangeKernel(kernel, cl::NullRange, size, cl::NullRange);
queue.enqueueReadBuffer(d_c, CL_TRUE, 0, size, array_result);
}

2.3.2.2 Single-Node Wrappers for OpenCL

For many application developers and domain experts not well versed in the C or C++ programming languages, the verbosity of OpenCL can be overwhelming. Hence, a wide range of OpenCL wrappers have been presented for various higher-level programming languages such as Java, C#, and Python. As one of the best known OpenCL wrappers, the Aparapi framework makes the resources of OpenCL compatible compute devices available in Java. Aparapi is not just a simple wrapper and aims for a transparent integration of heterogeneous compute resources, completely abstracting away device handling tasks and data transfers. Inspired by Aparapi, the Hybrid.Parallel project attempts to go even one step further by implementing an OpenCL backed drop-in replacement for the Parallel.For construct of the Task Parallel Library (TPL) in the .NET ecosystem \[50, 48\]. In contrast to these tightly integrated approaches, approaches such as PyOpenCL offer basic OpenCL wrappers \[101\].

2.3.2.3 Scale-Out Extensions for OpenCL

Scaling out GPU workloads across multiple compute nodes of a cluster is a recurring issue which has been investigated in a wide range of publications. The canonical approach for distributing GPU workloads relies on mixing GPU compute frameworks such as OpenCL or CUDA with implementations of the Message Passing Interface (MPI) standard. However, this approach requires application developers to deal with multiple levels of abstraction, using different programming models and synchronization semantics.

A commonly employed approach for achieving a uniform developer experience is to present devices scattered across compute nodes as if they were local devices. Relying on means of API forwarding, this strategy exploits the mechanisms available in OpenCL for coordinating work across multiple devices and has been applied by various approaches such as dOpenCL \[90\], SnuCL \[95\], or VirtualCL \[7\]. However, these approaches require developers to partition and schedule their workloads across multiple, potentially heterogeneous devices, increasing the complexity of applications significantly.

To address the problem of increased application complexity, DistCL \[38\] as well as another approach \[64\] have proposed to fuse multiple compute devices into a single
logical device. DistCL creates the illusion of a single, logical device by splitting kernels and the data they operate on it into multiple sub-ranges. To make this automatic splitting process possible, application developers have to supply a meta-function that specifies the memory access pattern. Based on the provided meta-functions, DistCL can identify data relevant for a sub-range and transfers it to the corresponding heterogeneous compute resource.

2.3.3 Data Placement in NUMA Architectures

The variety of approaches available for controlling data placement in NUMA systems ranges from approaches that assume a shared memory parallel programming model on one end to approaches that assume a distributed memory parallel programming model on the other end. Both strategies are valid, as the DSM architecture employed by NUMA systems exhibits certain characteristics of a distributed system, all while the cache coherent address space spanning across all NUMA nodes facilitates the system behavior of a shared memory system. Situated somewhere in between, several approaches employ the Partitioned Global Address Space (PGAS) model which enables the differentiation between local and remote memory resources. Hereinafter, popular approaches from all three categories are highlighted.

2.3.3.1 Shared Memory Model

Since many software developers are highly accustomed to the shared memory parallel programming model, the idea of extending this environment with means for controlling data placements on NUMA stands to reason. This section provides an overview of some of the most important approaches that have attempted to achieve that very same goal.

OpenMP

The Open Multi-Processing (OpenMP) standard specifies compiler-based extensions for the C/C++ and Fortran programming languages. Based on annotations, OpenMP enables application developers to instruct the compiler to define tasks that can be executed concurrently using a fork-join workflow. The OpenMP runtime library is responsible for mapping tasks onto the threading and synchronization primitives provided by the operating system. As OpenMP has no concept of memory locality, developers have to consider data placement themselves. Even though several approaches have been presented that extend OpenMP with task-to-data associations and a locality-aware scheduling policies, none of these approaches have gained traction. As a result thereof, the canonical methods for making OpenMP-based applications NUMA aware are to rely on the first touch memory allocation policy of the operating system or to use the libnuma API to manually manage the placement of memory resources. Both approaches are cumbersome to use as they result in cluttered code that harder to maintain.

Threading Building Blocks

The Threading Building Blocks (TBB) C++ template library provides a framework for task and data parallelism for shared memory systems. Based on a parallel tasking infrastructure, synchronization primitives, atomic operations and concurrent data-structures, the library enables application developers to implement
2 State of the Art and Related Work

parallel algorithms. Similar to OpenMP, the framework does not consider NUMA characteristics on its own. Building up on top of TBB, Majo et al. have presented TBB-NUMA, which extends the TBB scheduler with task affinities that can be specified manually or automatically by annotating parallel constructs with distribution templates. These task affinities are then considered by the scheduler.

Polymorphic Allocators Standardized in C++17, Polymorphic Allocators can be constructed using specific memory pools which can be used to represent different NUMA domains or different memory characteristics (e.g., volatility, latency, or bandwidth). By serving allocations from the memory pool they have been constructed upon, polymorphic allocators enable developers to enforce specific memory placement policies of the active scope. Unfortunately, polymorphic allocators cannot be used to modify data placement policies of allocations performed by nested data-structures transparently as polymorphic allocators have to be used explicitly. The explicit use of polymorphic allocators makes it necessary to modify nested data-structures in order to pass allocator objects to the nested allocations.

AutoNUMA With memory management being a central aspect of operating systems, it makes sense to facilitate NUMA-awareness through means of operating system facilities. Since version 3.8, the Linux Kernel implements a transparent mechanism called AutoNUMA. The mechanism provides two strategies: memory-follow-cpu and cpu-follow-memory. The former approach unmaps the process pages in regular intervals and tracks the NUMA-node of the CPU which has triggered the page-fault. Based on that information, pages are migrated to the identified NUMA-node to facilitate data locality. The latter strategy uses fault statistics to migrate tasks to CPU cores residing on the same NUMA node where most of the memory resides. These approaches work well assuming that data-structures are page-aligned. However, issues arise when data-structures are scattered among pages or if they are placed alongside other data-structures that are accessed from CPU cores residing on different NUMA nodes.

2.3.3.2 Distributed Memory Model

As the de-facto standard for implementing scale-out applications based on the distributed memory parallel programming model, the MPI standard defines an extensive list of message-based communication patterns, including point-to-point and group communication as well as reductions. Based on the Single Program Multiple Data (SPMD) paradigm, the MPI execution model assumes one application such as the one exemplified in Listing 2.3 that is deployed on all compute nodes participating in the computation. In a NUMA-agnostic scenario, one process is instantiated per compute node. Each process is identified by a unique identifier, and processes exchange messages to coordinate their work.

An MPI application does not necessarily have to be executed on a set of distinct cluster nodes, but can also launch multiple MPI processes on a single UMA or NUMA system. Since MPI applications are typically designed with the goal of minimizing communication among processes, they scale very well when deployed on a NUMA system. However,
using MPI to implement NUMA awareness on a shared memory system does not allow application developers to exploit the strengths of large scale-up NUMA systems.

Listing 2.3: Simple message passing example implemented in C using MPI.

```c
#include <stdio.h>
#include <string.h>
#include <mpi.h>

int main(int argc, char **argv){
    char msgBuffer[64];
    int rank, processCount;

    MPI_Init(&argc, &argv);
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
    MPI_Comm_size(MPI_COMM_WORLD, &processCount);

    if(rank == 0) {
        for(int i = 1; i<processCount;i++) {
            sprintf(msgBuffer, "This is a message from Process 0.");
            MPI_Send(msgBuffer, sizeof(msgBuffer), MPI_CHAR, i, 0, MPI_COMM_WORLD);
        }
    } else {
        MPI_Recv(msgBuffer, sizeof(msgBuffer), MPI_CHAR, 0, 0, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
        printf("Process %d receives: %s\n", rank, msgBuffer);
    }

    MPI_Finalize();
    return 0;
}
```

2.3.3.3 Partitioned Global Address Space Model

The PGAS programming model has been designed for data-parallel workloads, maintaining a global address space regardless of the underlying system architecture. The PGAS model has been employed by a multitude of programming languages, libraries and extensions such as X10 [29], Chapel [28], Legion [9], Sequoia [46], Unified Parallel C (UPC) [41], or High Performance Fortran (HPF) [170]. A central aspect of the PGAS model is the distinction between local and remote memory resources, which would make it a perfect fit for NUMA systems. However, even though the PGAS model could be applied to both shared memory architectures and distributed memory architectures, the latter case is much more common and the shared memory level is usually only considered in order to improve the performance of node-local inter-process communication [16].

2.4 Summary

Each type of heterogeneous system resource considered in this thesis has widely varying characteristics, which is well reflected in the way the approaches for providing abstractions vary from one type of heterogeneous system resource to another. While main memory
2 State of the Art and Related Work

compression in itself has been researched extensively [131], the different implementations and prototypes available vary on a per-product basis, rendering generic abstractions unreasonable. In the field of GPU computing, many approaches exist for alleviating access to GPUs. However, many approaches offer a similar range of functionality, often times neglecting challenges of distributed computing such as the potential overhead of inter-node data transfers. For NUMA systems, the APIs necessary for controlling data placement are available but very difficult to use correctly. This has resulted in many application developers either hoping for automatic approaches such as AutoNUMA to yield decent performance, or to ignore the coherent shared memory view available by treating NUMA systems as distributed systems which is a common strategy in the High Performance Computing (HPC) community.
This chapter lays out the foundation for investigating the impact of on-the-fly data compression for data transfers across **Central Processing Units (CPUs)** and **Graphics Processing Units (GPUs)** in scale-out GPU clusters in Chapter 4. For this purpose, the current chapter is focused on presenting the *lib842* compression library, which provides user-space access to the high-throughput compression facilities of the NX-842 on-chip compression accelerator available in all IBM POWER processors introduced since the POWER7+ [17].

With the proprietary 842 algorithm [54] employed by the NX-842 accelerator lacking any software-based implementations available for user-space applications, the *lib842* compression library introduces software-based high-throughput implementations of the algorithm to enable interoperability with arbitrary **CPUs** and **GPUs**.

The following master’s theses were supervised alongside the research leading to this chapter, fostering scholarly exchange between this work and the supervised theses:


Furthermore, partial results of the work presented in this chapter have been published:


- Max Plauth and Andreas Polze. “GPU-Based Decompression for the 842 Algorithm”. In: *Proceedings of the Seventh International Symposium on Computing and Networking Workshops (CANDARW)*. IEEE. Nov. 2019, pages 97–102. DOI: [10.1109/CANDARW.2019.00025](https://doi.org/10.1109/CANDARW.2019.00025)

This chapter is structured as follows. Section 3.1 motivates the demand for hardware-accelerated and software-based high-throughput compression facilities. The $842$ compression algorithm employed by the $\text{NX-842}$ on-chip hardware compression accelerator is introduced in Section 3.2. Section 3.3 then introduces the $\text{lib842}$ compression library on a conceptual level. The implementation details of all major hardware-based and software-based compression facilities provided by the $\text{lib842}$ library are detailed in Section 3.4. Both the compression ratio and the throughput of all implementations are evaluated in Section 3.5 using a wide range of test systems. Finally, the chapter is summarized in Section 3.6.

### 3.1 Motivation and Problem Statement

Combining the strengths of diverse heterogeneous system resources such as CPUs and other accelerators in state of the art heterogeneous computer architectures is vital to keep up with the demand for compute capacity imposed by the abundance of data accumulating in the age of digitization. To solve a given task using the respective strengths of the different heterogeneous system resources at hands, transferring data back and forth among the local memories of the involved system resources is a critical aspect of heterogeneous systems. Except for highly integrated approaches where all major system resources can be accommodated on a single System on a Chip (SoC) or package, data transfers across the local memories of dedicated system resources in heterogeneous systems are usually entailed by a certain level of performance degradation and increased energy demand compared to operations that can be completed without accessing the memory of another compute resource. Unfortunately, the number of workloads exceeding the compute capacity of single nodes increases with growing data volumes, requiring computations to be scaled out across numerous systems. For example, the soaring popularity of deep learning applications has drastically increased the demand for both Cloud-based and private GPU clusters. As the penalties related to data transfers already affect certain workloads executed on a single system, scale-out workloads are more susceptible to the overhead of data transfers as data has to be conveyed across comparatively slow inter-node interconnects. While the forthcoming commoditization of coherent interconnection technologies has the potential to improve upon the situation, the penalties associated with data transfers will not disappear entirely.

To further improve the efficiency of data transfers, on-the-fly data compression has already been demonstrated as a viable approach both in academic research and in commercial products. On the level of single compute nodes, most of the generally applicable approaches are restricted to individual system resources. Belonging to this category, the compute data compression feature introduced by NVIDIA with their latest generation of Ampere GPUs serves as a notable example of a product-grade approach, yielding up to $4 \times$ effective bandwidth improvements for accessing data residing in the local GPU memory or in the L2 cache of the GPU. While all the previously mentioned on-the-fly data compression approaches rely on custom hardware-based compression facilities, a software-based lightweight compression technique has been used to improve data transfer efficiency among system resources on the intra-node level. Regrettably, the compression technique used in this approach is only applicable to workloads operating...
3.2 The 842 Compression Algorithm

The 842 algorithm \cite{54,17} is a generic compression algorithm that has been designed with the use case of transparent main memory compression in mind. As this use case requires compression and decompression facilities that offer high throughput and low latency, the algorithm has been designed accordingly to enable hardware-based implementations that can be placed directly on transmission channels \cite{17}. The first implementation of the algorithm is a hardware-based implementation that has been introduced with the NX-842 on-chip compression accelerator, which is available in all IBM POWER processors introduced since the POWER7+ \cite{17}. The 842 algorithm can be attributed to the family of Lempel-Ziv derivatives \cite{54}. The compression process deviates from the original Lempel-Ziv algorithm \cite{215} in several aspects. However, decompression works very similar compared to LZ’ \cite{77} \cite{54}.

\url{https://github.com/torvalds/linux/tree/master/drivers/crypto}
The 842 compression algorithm operates in units of 8 bytes, treating the input data as sub-phrases of 8, 4 and 2 bytes length. The algorithm uses a fixed set of template codes (see Table 3.1) to encode 8 bytes of raw data by specifying a permutation of offsets to past occurrences or literals of 8, 4 and 2 bytes length, as demonstrated in the example.

As illustrated in Figure 3.1, the 842 algorithm [14, 17] operates on units of 8 bytes, treating input data as sub-phrases of 8, 4 and 2 bytes length, respectively. For each phrase length, a hash table holds offsets to sub-phrases that have already appeared in the raw data stream within a certain window. Depending on the outcome of the lookup, a compression template is chosen from the fixed list of available templates (see Table 3.1), with each template encoding 8 bytes of raw data. Each 5-bit template encodes a permutation of offsets or literals of 8, 4 and 2 bytes length, followed by the actual offsets and literals. Offsets to 2 and 8-byte phrases are encoded using 8 bits, whereas offsets to 4-byte phrases are encoded using 9 bits, resulting in the parameter sizes specified for each template in Table 3.1. With a clock frequency of 2.3 GHz, and the ability to ingest 8 bytes per cycle, one NX-842 unit can achieve a maximum throughput of 18.4 GB/s [17]. With two NX-842 units per socket, the total compression throughput of a POWER processor can be as high as 36.8 GB/s [17].

The example provided in Figure 3.1 demonstrates how the 32-byte string PITTERPATTERPATTERPATTERLISTENTO is compressed using four templates. The template 0x00 is used to encode the raw literal PITTERPA, since no matching sub-phrases have appeared in the raw data stream beforehand. The second template 0x13 encodes TTERPITT by providing offsets to two 2-byte phrases in the uncompressed data stream at the positions 2 (TT) and 4 (ER), as well as an offset to a 4-byte phrase at the position 0 (PITT). The third template 0x18 encodes ERPATTER by providing offsets to two 4-byte phrases at the positions 4 (ERPA) and 8 (TTER). Finally, the last template 0x00 encodes LISTENTO as a raw literal.
### Table 3.1: A 5-bit template encodes precisely 8 bytes of raw data using four consecutive actions. Actions include raw data phrases D and index references I, both in variants yielding 8, 4 and 2 bytes respectively. No-op actions N0 are used to fill up unused action slots (not shown).

<table>
<thead>
<tr>
<th>Template Parameter</th>
<th>Actions and corresponding bytes encoded by the template.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0x00 64 bits</td>
<td>D8 D2 I2</td>
</tr>
<tr>
<td>0x01 56 bits</td>
<td>D4 I2 D2</td>
</tr>
<tr>
<td>0x02 56 bits</td>
<td>D4 I2 I2 D2</td>
</tr>
<tr>
<td>0x03 48 bits</td>
<td>D4 I2 I2</td>
</tr>
<tr>
<td>0x04 41 bits</td>
<td>D4 I2 I2</td>
</tr>
<tr>
<td>0x05 56 bits</td>
<td>D2 I2 D4</td>
</tr>
<tr>
<td>0x06 48 bits</td>
<td>D2 I2 D4</td>
</tr>
<tr>
<td>0x07 48 bits</td>
<td>D2 I2 I2 D2</td>
</tr>
<tr>
<td>0x08 40 bits</td>
<td>D2 I2 I2</td>
</tr>
<tr>
<td>0x09 33 bits</td>
<td>D2 I2 I4</td>
</tr>
<tr>
<td>0x0A 56 bits</td>
<td>I2 D2 I4</td>
</tr>
<tr>
<td>0x0B 48 bits</td>
<td>I2 D2 I4</td>
</tr>
<tr>
<td>0x0C 48 bits</td>
<td>I2 D2 I2 I2 D2</td>
</tr>
<tr>
<td>0x0D 40 bits</td>
<td>I2 D2 I2</td>
</tr>
<tr>
<td>0x0E 33 bits</td>
<td>I2 D2 I4</td>
</tr>
<tr>
<td>0x0F 48 bits</td>
<td>I2 I2 D4</td>
</tr>
<tr>
<td>0x10 40 bits</td>
<td>I2 I2 D4</td>
</tr>
<tr>
<td>0x11 40 bits</td>
<td>I2 I2 I2 D2</td>
</tr>
<tr>
<td>0x12 32 bits</td>
<td>I2 I2 I2</td>
</tr>
<tr>
<td>0x13 25 bits</td>
<td>I2 I2 I4</td>
</tr>
<tr>
<td>0x14 21 bits</td>
<td>I4 I2 I4</td>
</tr>
<tr>
<td>0x15 33 bits</td>
<td>I4 D2 I4</td>
</tr>
<tr>
<td>0x16 33 bits</td>
<td>I4 I2 D2</td>
</tr>
<tr>
<td>0x17 25 bits</td>
<td>I4 I2 I4</td>
</tr>
<tr>
<td>0x18 18 bits</td>
<td>I4 I8 I4</td>
</tr>
<tr>
<td>0x19 08 bits</td>
<td>Reserved / Unused</td>
</tr>
<tr>
<td>0x1A 00 bits</td>
<td>Repeat preceding 8 B for N times</td>
</tr>
<tr>
<td>0x1B 00 bits</td>
<td>Emit 8 B of zeros</td>
</tr>
<tr>
<td>0x1C 00 bits</td>
<td>Reserved / Unused</td>
</tr>
<tr>
<td>0x1D 00 bits</td>
<td>End of compressed bitstream</td>
</tr>
</tbody>
</table>
### 3.3 Lib842: A User-Space Library for 842 Compression

The availability of high-throughput and low-latency compression and decompression facilities accessible from user-space are stringently required in order to enable application developers to improve data transfer efficiency among heterogeneous system resources based on data compression. Even though the 842 compression algorithm has been designed with transparent main memory compression in mind (cf. Section 3.2), it should be well suited for compressing data transfers among system resources, too since the requirements of both use cases are virtually identical in that they require high-throughput and low-latency compression and decompression facilities. Prior to this work however, both hardware-based and software-based implementations of the 842 compression algorithm have been widely inaccessible for user-space applications. The only way for user-space applications to leverage the resources of the NX-842 on-chip compression accelerator was to use IBM’s proprietary [(Advanced Interactive eXecutive (AIX)](https://www.ibm.com/analytics/aix) operating system, where an in-built user-space API exposes the hardware-accelerated compression facilities. On Linux however, both the NX-842 on-chip compression accelerator and a rudimentary software-based fallback implementation were only accessible from kernel-space through the Linux Crypto API. To make high-throughput 842 compression and decompression facilities available to user-space applications on Linux, this chapter introduces the lib842 compression library.

As its most prominent contribution, the lib842 library introduces the first available approach for making the resources of the NX-842 on-chip compression accelerator available to user-space applications running on Linux. To provide high-throughput 842 compression and decompression facilities on compute resources that are not equipped with a corresponding compression accelerator, library contributes software-based compression and decompression facilities several execution targets. Hiding the implementation details of all hardware and software-based implementations from any users of the library, lib842 exposes all available implementations through an implementation-agnostic interface as illustrated in [Figure 3.2](#). Of course, the implementation-agnostic interface also facilitates extensibility, making it relatively easy to add further implementations (e.g., for other heterogeneous system resources such as [Field-Programmable Gate Arrays (FPGAs)](https://www.xilinx.com)). A brief overview of all implementations that are currently provided by the lib842 library is outlined hereinafter:

**Hardware-based On-Chip Compression Accelerator (NX-842)** Based on a custom kernel module, this implementation exposes the compression and decompression facilities of all available NX-842 on-chip compression accelerators to user-space.

**Software-based Compression and Decompression (CPU Baseline)** This version is a user-space port of the software-based fallback-implementation of 842 compression and decompression facilities provided by the Linux kernel in case an NX-842 accelerator is not available. Even though this version merely provides minor compression and decompression throughput, it served as a golden unit during the development of all other implementations.
Software-based Compression and Decompression (CPU Optimized) Building up on top of the CPU-based baseline implementation, this version uses several optimization techniques to yield significantly higher compression and decompression throughput.

Software-based Decompression using OpenCL (GPU) To provide high-throughput decompression facilities for 842 compressed data on a wide range of GPUs this version is implemented using the Open Computing Language (OpenCL) framework. Relying on many optimization-techniques used in the optimized CPU-based implementation, some additional GPU-specific optimizations are employed in this implementation.

3.4 Implementation

This section provides insights into all major implementations provided by the lib842 compression library as highlighted in Section 3.3. Section 3.4.1 presents the first available approach for leveraging hardware-accelerated 842 compression on Power CPUs from user-space applications running on Linux. Serving as a golden unit during development, Section 3.4.2 briefly outlines the challenges of porting Linux kernel code to user-space for the software-based baseline implementation. Section 3.4.3 elaborates on the optimization strategies applied to the optimized software-based compression facilities that can be executed on arbitrary CPUs. Finally, Section 3.4.4 documents the optimization techniques used in the GPU-based decompression component implemented in OpenCL.
3.4.1 Hardware-based On-Chip Accelerator (NX-842)

On all POWER microarchitectures available at the time of writing, the NX-842 on-chip compression accelerators can only be accessed directly from kernel-space. While the AIX operating system provides a corresponding user-space Application Programming Interface (API), comparable interfaces are not available in the Linux kernel even though the NX-842 accelerators have already been in use for some time to implement the zram memory compression feature analogue to the Active Memory Expansion (AME) feature in AIX. To the best of the author’s knowledge, the approach presented in this section is the first to make the resources of the NX-842 on-chip compression accelerators available to user-space applications running on Linux. The presented approach involves several layers of indirection, as visualized in Figure 3.3.

Figure 3.3: On the kernel level, a driver for the NX-842 accelerator, a wrapper for the Linux Kernel Crypto API, as well as a modified version of the cryptodev-linux out-of-tree kernel module are required to expose the resources of the NX-842 on-chip compression accelerator to user-space applications. These components are augmented by a user-space wrapper provided by lib842 for interacting with the /dev/crypto special file exposed by the cryptodev-linux kernel module through ioctl() system calls.

Closest to the hardware, the nx842 driver available in the Linux kernel interacts with the hardware accelerator. Various code paths are available in the driver to cover different hardware configurations (e.g., to differentiate between native hardware and the PowerVM hypervisor). Another aspect the driver needs to provide different code paths for is that the method for interacting with the NX-842 units differs depending on the employed generation of the POWER microarchitecture. On POWER7+, POWER8, and POWER8+ CPUs, all on-chip accelerators are accessed through the privileged Initiate Coprocessor Store Word Indexed (icswx) instruction and its associated communication protocol [17]. From POWER9 onwards, the Virtual Accelerator Switchboard (VAS) facilities have been introduced with the goal of making on-chip accelerator resources accessible from user-space [83]. For user-space access to be enabled, the firmware has to be modified to initialize accelerators and make them available in the device tree. While corresponding modifications are available for other on-chip accelerators such as the NX-GZIP [164], comparable firmware modifications are thus far unavailable for NX-842 accelerators. As a result thereof, the NX-842 remains to be accessible from kernel-space only even on VAS-enabled POWER CPUs.
A wrapper then hides the various code paths available in the nx842 driver as well as the software-based fallback implementation of the 842 algorithm available in the Linux kernel [118], utilizing the interface for compression algorithms in the Linux Kernel Crypto API. Unfortunately, the subset of the Linux Kernel Crypto API exposed to user-space through the AF_ALG socket type does not include compression facilities. Even though the cryptodev-linux out-of-tree kernel module [‡] provides user-space access to a bigger portion of the Linux Crypto API compared to the AF_ALG socket type, it too does not provide access to any compression facilities. Therefore, the cryptodev-linux kernel module was extended in the context of this thesis to expose the resources of the NX-842 as well as other hardware accelerated compression facilities available through the Linux Kernel Crypto API through the /dev/crypto special file.

On the user-space side, lib842 takes care of interacting with the special file through ioctl() system calls. Since each ioctl request on the /dev/crypto special file involves a system call, the interface was augmented with a batching method that enables lib842 to submit multiple chunks for compression or decompression using a single system call. To further optimize the interaction between lib842 and /dev/crypto, session caching was implemented in order to re-use sessions. With these optimizations in place, the proposed approach is able to achieve high throughput for compression or decompression from user-space applications with minimal load on the CPUs.

### 3.4.2 Software-based Compression and Decompression (CPU Baseline)

Prior to this work, the only software-based implementation of the algorithm available to the public prior has been the fall-back implementation in the Linux kernel [118], which is also only accessible from kernel-space. As a first step, this basic implementation was ported to user-space by replacing all kernel dependencies with corresponding equivalents. The majority of dependencies could be easily resolved by consolidating various preprocessor definitions of constants and simple functions spread across the kernel in a single header file. More complex however was the task of replacing the generic hash table facilities provided by the Linux kernel [¶] with the uthash§ C library. Finally, concatenating strings of bits on the sub-byte level required careful consideration of the execution targets endianness, as the NX-842 operates in big-endian byte order. Delivering meager performance for both compression and decompression, the biggest value of this version is its use as a golden unit for all other software-based implementations.

### 3.4.3 Software-based Compression and Decompression (CPU Optimized)

Building up on top of the baseline implementation discussed in Section 3.4.2, various optimization efforts brought forward software-based high-throughput compression and decompression facilities that can be executed on arbitrary CPUs. Recalling the basic workflow of 842 compression illustrated in Figure 3.1, all major operations were optimized as documented hereinafter.

[‡] https://github.com/cryptodev-linux/cryptodev-linux

[§] https://troydhanson.github.io/uthash/
3.4.3.1 Fast Hash Tables

With efficient hash table lookups being the major potential bottleneck of the compression process, the general-purpose hash tables used in the baseline version were replaced with a very simplistic hashing mechanism. First, the 8, 4, and 2-byte sub-phrases are stored in a vector of 64-bit unsigned integer values. The vector-based representation with a uniform data type enables compilers to perform auto-vectorization of most subsequent operations. Using a vector-scalar-multiplication, all fields of the vector are multiplied with the largest prime number that falls within the range of a 64-bit unsigned integer. A right shift operation truncates the result of the multiplication results to its \( n_{hash} \) most significant bits, yielding a vector of hashes. For each sub-phrase length, two buffers are used to form a basic hash table structure: an index array with \( 2^n_{hash} \) unsigned short integer values and a FIFO buffer of \( 2^8, 2^9, \) and \( 2^8 \) elements for 2, 4, and 8-byte sub-phrases, respectively. The latter exponents are fixed constants defined by the FIFO sizes employed by the hardware-based NX-842 implementation. The index array uses hash-based addressing to store the offsets of corresponding values in the FIFO buffer. To retrieve the best possible performance, the hash size \( n_{hash} \) has to be chosen carefully to yield acceptable collision rates at a memory footprint that still fits into the CPU caches. For \( n_{hash} = 10 \), the total memory footprint amounts to \( 3 \times 2^{10} \times \text{sizeof(uint16\_t)} + 2^8 \times \text{sizeof(uint16\_t)} + 2^9 \times \text{sizeof(uint32\_t)} + 2^8 \times \text{sizeof(uint64\_t)} = 10.5KiB \) and should fit into the L1 data cache of most recent CPU architectures. Several tests were performed to make sure the presented hashing mechanism has minimal effects on compression ratio.

3.4.3.2 Efficient Template Lookup

In the baseline implementation, the hash tables are queried for known occurrences of a phrase in a complex hierarchy of if-else blocks in order to determine the most suitable template code for the data at hand. This mechanism was replaced with a simple look-up mechanism, where the template key is computed as exemplified in Listing 3.1. The resulting template key is used to retrieve the template code (as specified in Table 3.1) at the i-th position of a fixed lookup table. Since the \( \text{isInHashTable} \) flags can be computed branchless, the entire look-up is free of any branches.

Listing 3.1: Prime numbers 13, 53, and 149 are used to encode matches of 2, 4 or 8 byte phrases, respectively. To encode the action slot of the match, prime numbers 3, 5, 7, and 11 are used to encode a matching phrase in the first, second, third, or forth action slot. When a known value is found in a hash table, the primes indicating phrase-length and position are multiplied. The prime numbers have been chosen so that higher template keys indicate more efficient template codes.

```c
// isInHashTable has been computed earlier on without branches
uint16_t templateKey_21 = (13 * 3) & isInHashTable_21;
uint16_t templateKey_22 = (13 * 5) & isInHashTable_22;
uint16_t templateKey_23 = (13 * 7) & isInHashTable_23;
uint16_t templateKey_24 = (13 * 11) & isInHashTable_24;
uint16_t templateKey_41 = (53 * 3) & isInHashTable_41;
uint16_t templateKey_42 = (53 * 5) & isInHashTable_42;
uint16_t templateKey_81 = (149 * 3) & isInHashTable_81;
```
3.4 Implementation

// prefer one 4-byte matches over two 2-byte matches
uint16_t high = max(templateKey_41, templateKey_21 + templateKey_22);
uint16_t low = max(templateKey_42, templateKey_23 + templateKey_24);

// prefer one 8-byte match over two 4-byte matches
uint16_t templateKey = max(templateKey_81, high+low);

3.4.3.3 Optimized Template Encoder

The unoptimized baseline version encodes the template code and the four action parameters by calling an append function on the output buffer for each data item independently. Calls to the append function require a certain degree of overhead due to bookkeeping tasks for the bitstream writer. To reduce the number of append calls, fused calls to the append function were implemented for each template code, as exemplified in Listing 3.2.

As an additional optimization, the append function was replaced with the buffered bitstream writer¶ employed by the zfp library [117]. It accumulates bitstrings until a full 64-bit data sequence can be written to the output buffer. The buffering technique significantly reduces the complexity of appending sub-byte bitstring to the output buffer.

Listing 3.2: For all templates except for 0x00, the template key and all action parameters are packed into a single value, which reduces the number of calls to the stream_write_bits() function from five invocations to a single invocation. Template 0x00 requires two invocations.

```
uint64_t out = 0;
switch(TEMPLATE_KEY) {
  case 0x00:  // [ D8, N0, N0, N0 ], 64 bits
    stream_write_bits(p->stream, TEMPLATE_KEY, OP_BITS);
    stream_write_bits(p->stream, rawPhrase_81, D8_BITS);
    break;
  case 0x01:  // [ D4, D2, I2, N0 ], 56 bits
    out = (((uint64_t) TEMPLATE_KEY) << (D4_BITS + D2_BITS + I2_BITS)) |
          (((uint64_t) rawPhrase_41) << (D2_BITS + I2_BITS)) |
          (((uint64_t) rawPhrase_23) << (I2_BITS)) |
          (((uint64_t) indexOffset_24));
    stream_write_bits(p->stream, out, OP_BITS + D4_BITS + D2_BITS + I2_BITS);
    break;
  ...
}
```

3.4.4 Software-based Decompression using OpenCL (GPU)

An important design goal for the GPU-based implementation [158] of 842 decompression is that it must remain fully compatible with the compressed data streams produced by the NX-842 hardware compression accelerator. With this limitation in mind, the compression format of the NX-842 unit (see Section 3.2) leaves no obvious venues for parallelism at the intra-chunk level of granularity. Due to the sliding window mechanism used to encode known phrases within the window as index offsets, there are no entry points that guarantee the absence of data dependencies within a chunk of compressed data.

¶https://github.com/LLNL/zfp/blob/develop/src/inline/bitstream.c

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Therefore, naïve parallel decompression of chunks remains as the only viable venue for parallelization. However, decent decompression throughput can be achieved on various GPU hardware using the optimization strategies explained hereinafter.

### 3.4.4.1 Avoiding Divergent Execution

Most importantly, the amount of divergent execution among threads had to be reduced to a minimum. For the implementation at hands, divergent execution could be reduced by replacing a naïve case differentiation required to process each template code with a branch-free implementation. As outlined in Listing 3.3, the branch-free implementation strategy relies on a dictionary using the template code as a key for which it provides all parameters necessary to interpret the four actions encoded by a template code (e.g. type of action, parameter length in the compressed bitstream, and the length of the decompressed literal). Furthermore, the bitstream reader yielding an arbitrary number of bits from the compressed data stream has been reformulated to come by with very few case differentiation.

**Listing 3.3:** The array `dec_templates` serves as a dictionary, specifying the four actions associated with a template code. For each action, it holds the parameter size of the action (specified in bits), a tag specifying whether the action is an index action or not, and the number of raw bytes produced by the action. Based on this information, templates can be decoded without requiring a complex hierarchy of case differentiations.

```c
#define OP_DEC_D8 {(D8_BITS | NO_INDEX_OP), 8}
#define OP_DEC_D4 {(D4_BITS | NO_INDEX_OP), 4}
#define OP_DEC_D2 {(D2_BITS | NO_INDEX_OP), 2}
#define OP_DEC_N0 {(N0_BITS | NO_INDEX_OP), 0}
#define OP_DEC_I8 {(I8_BITS | IS_INDEX_OP), 8}
#define OP_DEC_I4 {(I4_BITS | IS_INDEX_OP), 4}
#define OP_DEC_I2 {(I2_BITS | IS_INDEX_OP), 2}

__constant uint8_t dec_templates[26][4][2] = {
	{OP_DEC_D8, OP_DEC_N0, OP_DEC_N0, OP_DEC_N0}, // template code 0x00
	{OP_DEC_D4, OP_DEC_D2, OP_DEC_I2, OP_DEC_N0}, // template code 0x01
	{OP_DEC_D4, OP_DEC_D2, OP_DEC_d2, OP_DEC_D2}, // template code 0x02
	...
	{OP_DEC_I8, OP_DEC_N0, OP_DEC_N0, OP_DEC_N0}, // template code 0x19
};

__kernel void decompress(__global uint64_t *in, __global uint64_t *out) {
    ...
    uint64_t template = read_bits(&buffer, OP_BITS);
    ...
    for (int i = 0; i < 4; i++) { // there are four actions to be decoded
        uint32_t dec_template = dec_templates[template][i][0];
        uint32_t is_index = (dec_template >> 7);
        uint32_t dst_size = dec_templates[template][i][1];
        uint64_t value = read_bits(&buffer, dec_template & 0x7F);
    }
}
```
3.4 Implementation

3.4.4.2 Avoid Template Lookups

Even though the approach outlined in Section 3.4.4.1 greatly reduces the occurrence of branching and improves decompression throughput on GPUs to a certain degree, the performance improvements yielded from this approach are much more distinct on CPUs where the lookup table easily fits the L1 data cache. On most GPUs however, the limited performance improvements gained by this approach are caused by frequently accessing the constant memory for each lookup, which is a far more expensive operation than accessing the L1 data cache on CPUs. However, the regular pattern of data and index actions used in templates 0x00 through 0x18 (cf. Table 3.1) can be used to compute parameter size, action type, as well as the number of bytes produced by each action as demonstrated in Listing 3.4. It should be noted that this technique is not applicable for template 0x19, however this is not a problem as it can be handled earlier on in the codebase alongside with the special templates 0x1A through 0x1E. Even though the computation of all necessary information involves expensive operations such as modulus and division operations, this approach provides higher throughput across all tested GPU hardware (cf. Table 3.2).

Listing 3.4: The regular patterns in the templates (cf. Table 3.1) can be exploited to compute parameter size, action type, as well as the number of bytes produced by each action. With this information available, the dec_templates dictionary used in Listing 3.3 can be removed in order to avoid costly memory access operations on each lookup in addition to avoiding divergent execution.

```cpp
__kernel void decompress(__global uint64_t *in, __global uint64_t *out) {
    uint64_t template;
    do {
        op = read_bits(&buffer, OP_BITS);
        ...
        opbits = 64 - ((op % 5) + 1) / 2 * 8 - ((op % 5) / 4) * 7
            - ((op / 5) + 1) / 2 * 8 - ((op / 5) / 4) * 7;
        uint64_t params = read_bits(p, opbits);
        for (int i = 0; i < 4; i++) {
            uint8_t opchunk = (i < 2) ? op / 5 : op % 5;
            uint32_t is_index = (i & 1) * (opchunk & 1)
                + ((i & 1) ^ 1) * (opchunk >= 2);
            uint32_t dst_size = 2 + (opchunk >= 4)
                * (1 - 2 * ((i % 2)) * 2);
            uint8_t num_bits = (i & 1) * (16 - (opchunk % 2) * 8
                - (opchunk >= 4) * 16) * ((i & 1) ^ 1)
                * (16 - (opchunk / 2) * 8 + (opchunk >= 4) * 9);
```
3.4.4.3 Optimized Memory Access Patterns

Another important optimization step was to reduce the number of global memory access operations by modifying the bitstream reader logic borrowed from the zfp library to cache data from global memory in registers, using the granularity of a native machine word. Based on this method, significant speed-up was achieved since not every read operation on the compressed input data results in a global memory access operation.

Transposing the compressed input data with the goal of achieving coalesced memory access operations was also evaluated as an optimization technique even though it would break compatibility with the data format generated by the NX-842 unit. However, this approach did not yield any measurable performance improvements. The reason for this is that each chunk is very likely to use a different series of template codes with differing parameter length each (cf. Table 3.1). Therefore, each thread ingests a differing amount of data in each step so that sooner rather than later threads request data from different offsets in their respective chunks, breaking the coalesced access pattern.

Finally, another attempt at improving memory access efficiency was undertaken by caching the output data in local memory. In theory, this would improve performance for index actions when phrases are copied from earlier positions of the output buffer. While this approach was able to deliver roughly 2× speed-up, it could only do so for very small chunk sizes (≤ 256 bytes). With each thread requiring the equivalent of one chunk of local memory, the overall consumption of local memory becomes too high for reasonable chunk sizes (≥ 4 KiB), resulting in poor occupancy and thus worse performance.

3.5 Evaluation

In this section, compression throughput, the energy demand, and compression ratio of all major implementations contributed by the lib842 compression library are evaluated. The list of evaluated implementations includes hardware-based compression and decompression using the NX-842 on-chip compression accelerator, optimized software-based compression and decompression on CPUs as well as GPU-based decompression implemented in OpenCL. Laying out the foundation for the evaluation, Section 3.5.1 documents the testing environment as well as the benchmark procedures used for the evaluation. Afterwards, the compression ratio delivered by both high-throughput compressors available in lib842 is investigated in Section 3.5.2. Finally, Section 3.5.3 determines the throughput of compression and decompression operations for all evaluated implementations using the wide range of test systems provided by the testing environment.
3.5 Evaluation

3.5.1 Testing Environment & Benchmark Procedure

To evaluate the throughput of the compression and decompression facilities provided by *lib842* across a wide range of CPU and GPU systems, a total number of six different test systems were employed. The detailed hardware configurations used for the evaluation are documented in Table 3.2, ranging from seasoned hardware configurations to state-of-the-art high-end hardware configurations.

All compression throughput measurements presented hereinafter were performed after a fresh reboot in order to ensure a clean system state. Furthermore, no other active users or background tasks were running on the involved servers. Both for the evaluation of compression rate and compression throughput, a chunk size of 64 KiB was used.

In order to retrieve a sufficiently meaningful dataset, each benchmark was executed 25 times. Error bars are used in all plots to report the standard deviation for each measurement. Furthermore, each benchmark was preceded by a warm-up run in order to eliminate any confounding factors. All measurements presented in this work are reported as average values including standard deviation (\(n = 25\)).

Throughput was determined by dividing the size of the uncompressed test data set through the isolated execution time of the compression and decompression functions. The measured execution times only include the execution of the compression or decompression function, respectively, excluding potential confounding variables such as the time required for setup, data transfers and teardown.

To compare the energy demand of NX-842-based and software-based compression facilities in *lib842*, the energy demand of a test application performing a compression operation immediately followed by a decompression operation on the contents of a given file was measured on the IBM Power System S824L test system using two Microchip MCP39F511N dual-channel power measurement devices \(^{129}\) and the PINPOINT \(^{104}\) utility. Since these measurements cover the entire execution of the test application, the compression and decompression cycle is repeated 30 times in the test application in order to reduce the impact of setup, data transfers, and teardown on the overall energy draw measurements. From these measurements, the idle power draw of the test system is deducted in order to only report the share of energy demand caused by the compression and decompression process itself. Similarly, the energy demand of the CPU-based decompression process was measured using the PINPOINT utility \(^{104}\) and the energy readings provided by the NVIDIA Management Library \(^{141}\).

3.5.2 Compression Ratio

As the performance optimization techniques for compression algorithms can often have an impact on the compression ratio \(r\), the compression ratios achieved by both the hardware-based NX-842 units (see Section 3.4.1) and the optimized CPU-based software implementation (see Section 3.4.3) is investigated hereinafter. The basic characteristics such as a brief description, size, and compression ratio of all employed datasets used for this evaluation are documented in Table 3.3. However, the reported compression ratios indicate that the differences in compression efficiency are negligible across all datasets. To facilitate replicability, this investigation employs well-disseminated, publicly available datasets whenever...
### Table 3.2: Specifications of the test systems used to evaluate the throughput of lib842.

<table>
<thead>
<tr>
<th>Model</th>
<th>S824L</th>
<th>m710p</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU</td>
<td>2×IBM POWER8 (Murano), 3.42 GHz, 10C/80T each</td>
<td>Intel Xeon E3-1284Lv4, 2.90 GHz, 4C/8T</td>
</tr>
<tr>
<td>Memory</td>
<td>1024 GB DDR3 ECC, 1600 MHz</td>
<td>32 GB DDR3, 1600 MHz</td>
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<tr>
<td>GPU</td>
<td>n/a</td>
<td>Iris Pro Graphics P6300</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 20.04.4</td>
<td>Ubuntu 18.04.4</td>
</tr>
<tr>
<td>Kernel</td>
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<td>4.15.0</td>
</tr>
<tr>
<td>Compiler</td>
<td>GCC 10.2.1 (AT 14.0)</td>
<td>GCC 7.4.0</td>
</tr>
<tr>
<td>OpenCL</td>
<td>n/a</td>
<td>OpenCL 2.1 NEO</td>
</tr>
<tr>
<td>GPU Driver</td>
<td>n/a</td>
<td>20.09.15980 (NEO)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Model</th>
<th>DL380 G9</th>
<th>Tyan</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU</td>
<td>2×Intel Xeon E5-2620v4, 2.20 GHz, 10C/20T each</td>
<td>2×AMD EPYC 7282, 2.80 GHz, 16C/32T each</td>
</tr>
<tr>
<td>Memory</td>
<td>256 GB DDR4 ECC, 2133 MHz</td>
<td>256 GB DDR4 ECC, 3200 MHz</td>
</tr>
<tr>
<td>GPU</td>
<td>8×NVIDIA Tesla K80</td>
<td>NVIDIA Tesla T4</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 20.04.4</td>
<td>Ubuntu 20.04.4</td>
</tr>
<tr>
<td>Kernel</td>
<td>5.4.0</td>
<td>5.4.0</td>
</tr>
<tr>
<td>Compiler</td>
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<td>GCC 9.4.0</td>
</tr>
<tr>
<td>OpenCL</td>
<td>OpenCL 1.2 CUDA</td>
<td>OpenCL 1.2 CUDA</td>
</tr>
<tr>
<td>GPU Driver</td>
<td>470.103.01</td>
<td>510.47.03</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Model</th>
<th>DGX-1</th>
<th>DGX A100</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU</td>
<td>2×Intel Xeon E5-2698v4, 2.20 GHz, 10C/20T each</td>
<td>2×AMD EPYC 7742, 2.25 GHz, 64C/128T each</td>
</tr>
<tr>
<td>Memory</td>
<td>512 GB DDR4 ECC, 2133 MHz</td>
<td>1024 GB DDR4 ECC, 3200 MHz</td>
</tr>
<tr>
<td>GPU</td>
<td>8×NVIDIA Tesla V100</td>
<td>8×NVIDIA Tesla A100</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 20.04.4</td>
<td>Ubuntu 20.04.4</td>
</tr>
<tr>
<td>Kernel</td>
<td>5.4.0</td>
<td>5.4.0</td>
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</tr>
<tr>
<td>GPU Driver</td>
<td>470.103.01</td>
<td>470.103.01</td>
</tr>
</tbody>
</table>
possible. For the remaining, artificial data sets, an additional description is provided hereinafter.

The artificial datasets periodic, zeros, and random are self-explanatory and are used to quantify the compression ratio $r$ for extreme cases ranging from the best case (periodic, zeros) to the worst case (random). Distinguishing periodic and zeros makes sense because zeros triggers a special template in the 842 algorithm, whereas the periodic dataset has to be encoded using the regular templates described in Section 3.2.

### 3.5.3 Compression Throughput and Energy Demand Benchmark

To gauge the compression and decompression performance characteristics of all major implementations available in the lib842 library, throughput measurements were performed on all test systems presented in Table 3.2. The throughput measurements retrieved for hardware-based compression and decompression using the NX-842 on-chip compression accelerator, optimized software-based compression and decompression on CPUs and GPUs, and CPU-based decompression are illustrated in Figure 3.4. As a compression payload, the enwik9 dataset (see Table 3.3) was employed.

Using a total number of four NX-842 accelerators available in the dual-socket test-system, the throughput measured for the hardware-accelerated implementation approaches the theoretical throughput offered by two NX-842 accelerators available per Power CPU, yielding a utilization efficiency of roughly 38%. One might come to expect a certain level of performance loss caused by the many abstraction layers involved in the implementation (cf. Section 3.4.1). However, it appears as if the utilization efficiency of 43.48% achieved for accessing the NX-GZIP accelerator from user-space based on the VAS facilities [3] does only provide slightly improved utilization efficiency. For the software-based compression operation, even dual-socket systems equipped with high-end state-of-the-art CPU models can only provide roughly one-third of the compression throughput provided by the NX-842 units available in our dual-socket Power test-system. While there still may be some room for minor optimizations in the software-based compression process, the general picture is unlikely to change fundamentally even with further optimizations in place. On the side of the decompression operation however, software-based implementations on CPUs and GPUs are able to achieve throughput levels on the high-end test systems comparable to the NX-842 accelerators available in our dual-socket Power test-system.

Additional tests using other datasets did not reveal a significant impact of the dataset on the compression throughput, except for the zeros dataset. There, the compression throughput for CPU-based compression roughly doubled across all node test systems. This effect is likely caused by the special compression template used for encoding an eight-byte sequence of zeros as well as the special template for encoding a repeated occurrence of eight-byte sequences. When a special template is encoded, the entire hash-and-lookup operations during the compression process is skipped, likely yielding the observed speed-up. Since both special templates address corner cases, this effect rarely occurs and compression throughput should remain stable across many datasets.
<table>
<thead>
<tr>
<th>Dataset Description</th>
<th>Source</th>
<th>Size (Bytes)</th>
<th>Compression Ratio (NX-842)</th>
<th>Compression Ratio (SW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Periodic pattern of 256 bytes (00 01 02 ... FD FE FF)</td>
<td>Space Telescope</td>
<td>4,050,000</td>
<td>0.390</td>
<td>0.701</td>
</tr>
<tr>
<td>Periodic pattern of the Curiosity Mars Rover</td>
<td>Curation</td>
<td>8,135,406</td>
<td>1.04</td>
<td>1.000</td>
</tr>
<tr>
<td>Zeros</td>
<td>Dump of the Open Library works catalog</td>
<td>10,565,490</td>
<td>0.54</td>
<td>0.410</td>
</tr>
<tr>
<td>Zeros</td>
<td>Wikipedia</td>
<td>76,134,477,181</td>
<td>0.57</td>
<td>0.681</td>
</tr>
<tr>
<td>Random</td>
<td>Books</td>
<td>76,134,477,181</td>
<td>0.84</td>
<td>0.705</td>
</tr>
<tr>
<td>Random</td>
<td>Book reviews from Amazon.com</td>
<td>22,361,816</td>
<td>0.69</td>
<td>0.692</td>
</tr>
<tr>
<td>Random</td>
<td>The full 2006-03-03 Wikipedia dump.</td>
<td>121</td>
<td>0.701</td>
<td>0.705</td>
</tr>
<tr>
<td>OLW dump of the Open Library works catalog</td>
<td>OLW</td>
<td>10,565,840,859</td>
<td>0.54</td>
<td>0.410</td>
</tr>
<tr>
<td>OLW dump of the Open Library works catalog</td>
<td>OLW</td>
<td>10,565,840,859</td>
<td>0.54</td>
<td>0.410</td>
</tr>
<tr>
<td>Curiosity Stitched panorama from the Curiosity Mars Rover</td>
<td>OLW</td>
<td>8,135,406</td>
<td>1.04</td>
<td>1.000</td>
</tr>
<tr>
<td>OLW dump of the Open Library works catalog</td>
<td>OLW</td>
<td>8,135,406</td>
<td>1.04</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Table 3.2: Characteristics of the evaluated data sets and the respective compression ratios achieved using hardware and software-based compression.
Even though analyzing the energy-efficiency is not a central concern of this work, the energy demand of all major implementations provided by lib842 was measured (cf. Figure 3.5) to investigate the energy-efficiency of different implementations and heterogeneous compute resources. The energy draw measurements for a compression/decompression cycle were only performed on the IBM Power System S824L test system with the goal of enabling commensurability among hardware-based and software-based implementations. The optimized software-based implementation requires almost four times the energy required by the NX-842 on-chip compression accelerator to perform the same compression/decompression cycle on the employed test dataset. While the superior energy-efficiency of the NX-842 should not come as a surprise, the clear difference in energy consumption underlines the importance of being able to leverage the resources of the dormant on-chip compression accelerators in place of software-based equivalents whenever possible. On the side of CPU-based decompression, the energy required to perform the decompression process was measured on all employed NVIDIA GPUs to compare the energy efficiency across various generations of GPU microarchitectures. The measurements demonstrate unequivocally how the energy-efficiency of GPUs improves with each microarchitecture. With the lower-end T4 GPU almost approaching the energy-efficiency of the NX-842 unit, there seems to be a huge potential for improving the energy-efficiency of certain workloads when end-users are willing to relinquish a certain level of throughput in favor of improving the energy-efficiency of their application.

3.6 Summary

With the introduction of the lib842 compression library, this chapter presented the first user-space approach for providing compression and decompression facilities based on the proprietary 842 compression algorithm. Relying on a modified version of the cryptodev-linux out-of-tree kernel module, the implementation details for making the high-throughput and low-latency compression and decompression facilities of NX-842
Figure 3.5: Panel (a) compares the energy required to compress and decompress the enwik9 test dataset using the software-based implementation and the NX-842-accelerated approach on the IBM Power System S824L test system. In panel (b), the energy required to decompress the compressed enwik9 test dataset on various generations of NVIDIA GPUs microarchitectures is compared.

On-chip compression accelerators accessible to user-space applications through lib842 were discussed. To enable compressed data exchange across heterogeneous system resources, the hardware-accelerated approach was complemented with the introduction of highly optimized software-based compression and decompression routines for arbitrary CPUs as well as OpenCL-based decompression facilities for arbitrary GPUs. The detailed evaluation of all major implementations available in lib842 has revealed that while hardware-based compression and decompression clearly outperforms software-based approaches both in terms of throughput and energy-efficiency, the latter still provide decent throughput across the wide range of employed test systems. On higher-end systems, both CPU-based and GPU-based decompression is almost approaching the decompression throughput provided by the hardware-accelerated implementation. By making these high-throughput compression and decompression facilities available in the form of the lib842 compression library, this chapter lays out the foundation for investigating the impact of On-the-Fly I/O Link Compression for data transfers among CPUs and GPUs in scale-out GPU clusters in Chapter 4.
This chapter introduces the CloudCL framework, which attempts to make scale-out Graphics Processing Unit (GPU) resources more accessible to a wider audience by providing abstractions that hide many aspects of the distributed memory parallel programming model associated with scale-out workloads. Based on these abstractions, the framework enables application developers and domain experts to focus on the data parallel programming model associated with GPUs. By implementing a naïve form of GPU resource disaggregation based on the dOpenCL Application Programming Interface (API) forwarding library for the Open Computing Language (OpenCL) ecosystem, the framework also helps operators to improve the utilization of their GPU clusters. To improve the data transfer efficiency of the API forwarding approach in commodity 10 Gbit/s Ethernet networks, the dOpenCL library is augmented with transparent on-the-fly data compression for inter-node data transfers based on the lib842 compression library presented in Chapter 3.

The following master’s thesis were supervised alongside the research leading to this chapter, fostering scholarly exchange between this work and the supervised thesis:


Furthermore, partial results of the work presented in this chapter have been published:

This chapter is structured as follows. Section 4.1 motivates the need for programming abstractions in the context of scale-out GPU workloads. Providing such abstractions, the CloudCL framework is introduced in Section 4.2. To demonstrate the developer experience of CloudCL, Section 4.3 exemplifies the implementation of two data-intensive GPU scale-out workloads using CloudCL. Section 4.4 then proposes a strategy for transparently integrating on-the-fly data compression into the CloudCL framework to improve the efficiency of inter-node data transfers. The pipelined approach for implementing the previously proposed integration strategy is elaborated in Section 4.5. To test the impact of transparently compressed data transfers both on the effective data transfer throughput between nodes and on the overall performance of scale-out GPU workloads, a thorough evaluation is conducted in Section 4.6. Finally, the chapter is summarized in Section 4.7.

4.1 Motivation and Problem Statement

Over the last decade, the use of GPUs as a general purpose compute resource has become prevalent across arbitrary domains [24, 87, 188, 174]. Consequently, the demand for GPU compute resources has been steadily increasing over the last few years to the point where many use cases even require multiple GPUs to satisfy their resource demands. The soaring popularity of deep learning applications for example has drastically increased the demand for both Cloud-based and private GPU clusters [84].

As a result of scale-out GPU workloads becoming increasingly common, the following issues arise:

1. Application development for scale-out GPU workloads is becoming very challenging, as developers have to be adept using both data parallel programming models (e.g., OpenCL cf. Section 2.3.2.1) and distributed memory parallel programming models (e.g., Message Passing Interface (MPI) cf. Section 2.3.3.2) in addition to considering the dynamic aspects of Infrastructure as a Service (IaaS) based resources.

2. To provide dynamic GPU resources based on public or private GPU clusters, their operators are often faced with the problem of resource fragmentation [123] as illustrated in Figure 4.1.

Numerous programming abstractions are available for alleviating access to GPU compute resources in single-node scenarios [6, 101, 50]. To tackle the first issue however, a larger number of GPUs spread out across multiple nodes have to be considered. Even though several approaches exist that make GPUs scattered across compute nodes appear as if they were local resources based on API forwarding techniques [90, 95, 7], programming abstractions targeting this larger scale however have to provide the means for splitting workloads into partitions that can be processed mostly independent of each other, without requiring fine-grained communication between GPUs [195, 13].
To resolve the second issue, resource disaggregation is considered a promising approach to improve the efficiency of data centers [31, 125, 80], as resource disaggregation eliminates many resource allocation issues such as fragmentation. With no implementation of resources disaggregation being ready for production yet, simple forms of resources disaggregation can be implemented in software already today. Using API forwarding techniques for example to scale out multi-GPU applications across multiple compute nodes can be considered a naïve form of resource disaggregation [123]. Usually, the lessened data transfer performance available between the host Central Processing Unit (CPU) and remote GPUs in such a naively disaggregated setup has to be compensated by either employing expensive high-end inter-node network technology or by restricting the range of employed workloads to compute-bound workloads operating on small datasets.

With the introduction of the CloudCL framework, this chapter presents an approach that unites both solution strategies by extending an existing programming abstraction framework for GPUs with the scale-out capabilities offered by an API forwarding library for the OpenCL ecosystem. Furthermore, the programming abstraction mechanisms are extended with the means necessary for defining workload partitions and for managing the resources of an ad-hoc GPU cluster. Building up on top of the efforts presented in Chapter 3, the potential of mitigating the limitations of such a naïvely disaggregated setup based on on-the-fly data compression is investigated.

4.2 CloudCL: Single-Paradigm Scale-Out GPU Computing

In addition to hiding the complexity of the distributed memory programming model for scale-out GPU workloads, the CloudCL framework presented in this chapter provides ad-hoc GPU clusters tailored specifically to the resource requirements of each workload. By enabling developers and domain experts to focus on the data parallel programming model, one goal of CloudCL is to make scale-out GPU resources accessible to a wider audience using a single-paradigm approach. Simultaneously, CloudCL attempts to improve the utilization of public or private GPU clusters by disaggregating GPU resources. As illustrated in Figure 4.2, CloudCL heavily relies on the dOpenCL library and the Aparapi framework as underlying technologies to achieve these goals. Building up on top of these technologies, CloudCL provides enhancements to Aparapi with the goal of optimizing the framework for scale-out GPU workloads. Both the underlying technologies and the enhancements are detailed hereinafter.
Programming Abstractions for Scale-Out Graphics Processing Unit Clusters

Figure 4.2: The CloudCL framework builds up on top of the dOpenCL API forwarding library for OpenCL and the Aparapi framework for executing native Java code on OpenCL-enabled GPUs. The underlying technologies are extended with a job infrastructure to hide most aspects of the distributed memory parallel programming model during the development of scale-out GPU workloads.

4.2.1 Underlying Technologies

CloudCL heavily relies on the dOpenCL library and the Aparapi framework as underlying technologies. Therefore, both technologies are introduced in greater detail hereinafter.

4.2.1.1 dOpenCL

Serving as the foundation of CloudCL, the dOpenCL API forwarding library for OpenCL enables applications to transparently utilize OpenCL devices installed in remote machines [90]. The library provides its own Installable Client Driver (ICD), which forwards the API calls to specified remote machines in the network, which run a dOpenCL daemon. The calls are received by the daemon and are executed using the available native OpenCL ICDs on the remote compute node with the results being returned via network. With this approach, OpenCL kernels do not require changes to run remotely as dOpenCL hides network transfers behind the standard OpenCL API. An overview of the architecture of an exemplary compute cluster based on dOpenCL is shown in Figure 4.3.

4.2.1.2 Aparapi

Since the verbose nature of OpenCL can still be overwhelming for many domain experts, CloudCL employs Aparapi as an abstraction layer on top of OpenCL. Aparapi is a framework that drastically simplifies the usage of the OpenCL API and that minimizes development efforts of OpenCL kernels [6]. For CloudCL, the most important feature of Aparapi is that it enables developers to implement kernels in a subset of Java which is then translated to valid OpenCL kernels. Of similar importance to CloudCL is that Aparapi takes care of tedious setup-tasks and releases developers from the task of moving data back and
forth between host and accelerator. Based on these features, the framework enables developers to express the same functionality of native OpenCL with much fewer lines of code, reducing code complexity significantly as demonstrated in Listing 4.1. The reduced complexity enables developers to implement algorithms considerably faster and offers beginners and domain experts easy access to OpenCL features without profound knowledge of low-level mechanisms.

Listing 4.1: Example of a vector addition kernel and the corresponding host code using Aparapi.

```java
final double[] a = new double[]{0, 1, 2, 3, 4, 5, 6, 7, 8, 9};
final double[] b = new double[]{0, 1, 2, 3, 4, 5, 6, 7, 8, 9};
final double[] c = new double[10];

Kernel kernel = new Kernel() {
    @Override
    public void run() {
        int i = getgId();
        c[i] = a[i] + b[i];
    }
};
kernel.execute(10);
System.out.println(Arrays.toString(c));
```
4.2.2 Enhancements

CloudCL provides several enhancements to Aparapi with the goal of optimizing the framework for scale-out GPU workloads. These enhancements are explicated hereinafter.

4.2.2.1 Jobs

CloudCL introduces the concept of Jobs not only to serve as a unit of scheduling, but Job classes are also the main venue for specifying the strategy how workloads are partitioned into multiple kernel instances operating on independent sub-ranges of the input data as exemplified in [Listing 4.2] Providing numerous kernel instances for a workload that can be processed independently is crucial in order for CloudCL to scale with the number of available GPUs. Even though semi-automatic approaches for identifying sensible sub-ranges were investigated during the development of CloudCL [94][114], a manual approach where developers have to specify compartmentalization strategies themselves was favored over the immense complexity associated with semi-automatic classification mechanisms. While the manual approach requires the developer’s attention, partitioning strategies tailored to a workload are likely to outperform generic approaches.

Listing 4.2: The VecAddJob class is responsible for dividing the input data into partitions that can be processed independently by multiple VecAddKernel instances.

```java
1 public class VecAddKernel extends CloudCLKernel{
2    int[] a, b, result;
3
4    public VecAddKernel(CloudCLJob job, Range range, int[] a, int[] b) {
5        super(job, range);
6        this.a = a;
7        this.b = b;
8        this.result = new int[a.length];
9    }
10
11    @Override
12    public void run() {
13        result[getgId()] = a[getgId()] + b[getgId()];
14    }
15 }
16
17 public class VecAddJob extends CloudCLJob {
18    public VecAddJob(int vectorSize, int partCount, DevicePreference pref, ThreadFinishedNotifyable notify) {
19        super("VecAdd", notify);
20        int[] a = new int[vectorSize];
21        int[] b = new int[vectorSize];
22        int[] result = new int[vectorSize];
23
24        int partWidth = vectorSize / partCount;
25        for(int i = 0; i < partCount; i++){
26            int[] aPart = Arrays.copyOfRange(a, i * partWidth, (i + 1) * partWidth);
27            int[] bPart = Arrays.copyOfRange(b, i * partWidth, (i + 1) * partWidth);
28            Range range = Range.create(partWidth);
29        }
30    }
31 }
```
Unlike the C API of OpenCL, all commands such as kernel invocations in Aparapi are synchronous, requiring developers to use threading in order to launch kernels on multiple OpenCL devices simultaneously. To unburden developers, the job infrastructure takes care of launching all independent kernels concurrently as well as of monitoring their execution status. Furthermore, performance metrics such as data volume, transfer time and execution time are collected for all kernels to provide a solid basis for scheduling decisions.

4.2.2.2 Job Scheduler

CloudCL employs a pluggable two-tiered architecture as outlined in Figure 4.4. The first tier (Job Scheduler) operates on the job level and does not consider individual kernel instances and the corresponding data partitions. Operating at this high level of abstraction, the first tier is mainly concerned with fairness and can be configured to use either a First In, First Out (FIFO) or a Round Robin scheduling strategy to decide which jobs are becoming eligible for being scheduled by the second tier. The first tier then hands over the kernel instances belonging to a job to the second tier (Device Scheduler) which is responsible for assigning individual kernels to available compute devices.

**Figure 4.4:** The two-tier scheduling mechanism employed by CloudCL considers jobs on the first-tier to make sure jobs are processed in a fair order. The second tier assigns the individual kernels encapsulated by jobs to compute devices whilst taking into consideration statistics regarding data volume, transfer time, and execution time of preceding runs.

Especially in the context of CloudCL, the net performance of a compute-device heavily depends on the speed of the network connection used to tie in the compute device to the CloudCL master node. Therefore, the second tier assigns kernels to devices based on the performance metrics collected on the job level. It also considers the fact that the performance of OpenCL kernels can vary significantly depending on what kind of heterogeneous compute resource they are executed on. For example, kernels that may
perform exceptionally well on a CPU may run poorly on a CPU because of varying microarchitectural properties. Therefore, CloudCL enables developers to express preferences which device type their kernels should be executed on using the Device Preference attribute demonstrated in Listing 4.2. Valid values for the attribute include: None, CPU only, GPU preferred, GPU only and GPU preferred.

4.2.2.3 Dynamic Scaling Capabilities

To exploit the potential of private or public IaaS-based compute resources, one important goal of CloudCL is that resources can be dynamically added to or removed from the pool of compute resources utilized by the framework. However, both OpenCL and Aparapi are built to run on a single machine and as such assume a fixed topology of compute devices during operation. Fortunately, dOpenCL supports adding and removing devices virtually by adding or removing compute nodes at runtime using the custom methods clCreateComputeNodeWWU and clReleaseComputeNodeWWU. To further provide information about the relation between compute nodes and devices, dOpenCL introduces another method called clGetDeviceIDsFromComputeNodeWWU. Especially the latter feature is crucial for removing resources dynamically, where it has to ensured that no kernels are running on a device that is about to be removed.

Since Aparapi is bound to the standard interface specified by the OpenCL specification, the framework cannot make use of the offered dOpenCL extensions. To let CloudCL make use the dynamic resource scaling capabilities of dOpenCL, Aparapi was extended in the context of this work by implementing the two new Java Native Interface (JNI) methods addNode and removeNode. Both call the respective dOpenCL functions, with addNode also reporting back the available devices of the added node.

4.3 Developer Experience of CloudCL

One major goal of CloudCL is to make resources of GPU clusters more accessible to developers and domain experts. To demonstrate the developer experience of the CloudCL framework, this section showcases the implementations of two workloads that are also used to evaluate the performance impact of employing on-the-fly data compression for data transfers in CloudCL (cf. Section 4.6). A Semi-Sparse Matrix Multiplication workload and an Analytical Database Query workload are employed as data-intensive, memory-bound workloads to evaluate the performance of CloudCL outside its comfort zone. Except for the Java syntax, CloudCL kernels themselves remain fairly similar to kernels implemented directly in OpenCL. The aspect that changes significantly however is the host code surrounding the kernel. Therefore, the Job classes of both workloads are presented hereinafter to demonstrate the brevity and simplicity of CloudCL.

4.3.1 Semi-Sparse Matrix Multiplication

The SemiSparseMatMulJob class exemplified in Listing 4.3 performs a dense matrix multiplication kernel to of matrix \( A \) \((N \times M)\) and matrix \( B \) \((M \times P)\), yielding matrix \( C \) \((N \times P)\) as a result. Matrices \( A \) and \( B \) are filled with random data except for a configurable fraction of cells that are zeros. This fraction of zeroed cells is controlled by the sparsity argument,
where 0 means all cells are filled with random data and 1 means all cells contain zeros. To focus on the aspect of partitioning the workload across multiple kernel instances, the process of filling the matrices was excluded from the code example for reasons of brevity. The semi-sparse scenario is intended to represent the varying degrees of compressibility encountered across the various use cases and application domains that rely on efficient matrix multiplication. Even though sparse matrix representations such as Compressed Sparse Row (CSR) are employed in use cases with a high-degree of sparsity, there is a certain gray area where the space gains provided by sparse representations are not sufficient to justify the additional complexity introduced by sparse representations. A tiling strategy is employed that splits up matrix A horizontally and distributes it independent workload partitions, whereas the entire second matrix B is used by all workload partitions. The dense matrix multiplication kernel itself is implemented using a naïve implementation strategy.

**Listing 4.3**: Using CloudCL, the SemiSparseMatMulJob class initializes all relevant data structures on the host and specifies the strategy for partitioning the Semi-Sparse Matrix Multiplication workload into multiple independent kernel instances.

```java
public class SemiSparseMatMulJob extends CloudCLJob{
    public SemiSparseMatMulJob(int sizeN, int sizeM, int sizeP, float sparsity, int tiles, DevicePreference pref, ThreadFinishedNotifyable notify)
    {
    super("SemiSparseMatMul", notify);
    final float[] a = new float[sizeN*sizeM];
    final float[] b = new float[sizeM*sizeP];

    // fill matrices with random data and sparsity
    int tileHeight = sizeN/tiles;
    for (int tile=0; tile<tiles; tile++)
    {
        float[] aPart = Arrays.copyOfRange(a, tile*tileHeight*sizeM, (tile+1)*tileHeight*sizeM);
        Range range = Range.create2D(tileHeight, sizeP, 100, 1);
        DenseMatMulKernel kernel = new DenseMatMulKernel(this, range, aPart, b, sizeM);
        kernel.setDevicePreference(pref);
        addKernel(kernel);
    }
}
```

### 4.3.2 Analytical Database Query

The DatabaseQueryJob class presented in **Listing 4.4** orchestrates an analytical database query modeled after Query 1 of the TPC-H benchmark [206]. This query involves filtering and grouping operations to perform an aggregation. To focus on the aspect of partitioning the workload across multiple kernel instances, the process of data generation was excluded from the code example for reasons of brevity. At this point, the author wishes to stress that for reasons of simplicity, neither the query nor the data generator fully complies with the very complex TPC-H specification. As such, the Analytical Database Query workload must not be mistaken for a subset TPC-H benchmark. To facilitate efficient execution
in scale-out deployments, the table entries are split up horizontally in order to yield partitions that can be processed independently.

**Listing 4.4:** Using CloudCL, the DatabaseQueryJob class initializes all table columns on the host and specifies the strategy for partitioning the Analytical Database Query workload horizontally to yield multiple independent kernel instances.

```java
public class DatabaseQueryJob extends CloudCLJob{
    public DatabaseQueryJob(Integer size, int tiles, DevicePreference pref,
                        ThreadFinishedNotifyable notify) {
        super("DatabaseQueryJob", notify);
        int[] colQuantity = new int[size];
        int[] colExtPrice = new int[size];
        int[] colDiscount = new int[size];
        int[] colTax = new int[size];
        int[] colReturnFlag = new int[size];
        int[] colLineStatus = new int[size];
        int[] colShippingDate = new int[size];

        // generate line items
        int tileHeight = size/tiles;
        for(int tile=0; tile<tiles; tile++){
            int start = tile*tileHeight, end = (tile+1)*tileHeight;
            int[] colQuantitySpl = Arrays.copyOfRange(colQuantity, start, end);
            int[] colExtPriceSpl = Arrays.copyOfRange(colExtPrice, start, end);
            int[] colDiscountSpl = Arrays.copyOfRange(colDiscount, start, end);
            int[] colTaxSpl = Arrays.copyOfRange(colTax, start, end);
            int[] colReturnFlagSpl = Arrays.copyOfRange(colReturnFlag, start, end);
            int[] colLineStatusSpl = Arrays.copyOfRange(colLineStatus, start, end);
            int[] colShippingDateSpl = Arrays.copyOfRange(colShippingDate, start, end);

            Range range = Range.create(end - start, 256);
            DatabaseQueryKernel kernel = new DatabaseQueryKernel(this, range,
                        colQuantitySpl, colExtPriceSpl, colDiscountSpl, colTaxSpl,
                        colReturnFlagSpl, colLineStatusSpl, colShippingDateSpl);
            kernel.setDevicePreference(pref);
            addKernel(kernel);
        }
    }
}
```

### 4.3.3 Summary

As a central construct of CloudCL, the job classes take care of initialization, splitting up the workload into independent partitions, and launching kernel instances. The examples demonstrated in Section 4.3.1 and Section 4.3.2 show that only few lines of code are necessary to accomplish these tasks. Considering the verbosity of the native OpenCL API this level of abstraction unburdens developers while still giving them full control over the data partitioning strategies as a performance-critical aspect.
4.4 Augmenting CloudCL with Data Transfer Compression

The overhead caused by inter-node data transfers in CloudCL is the biggest limitation of the framework, restricting its utility to compute-bound workloads operating on small datasets \([161]\). Preceding efforts of the research community have identified compression as a viable method for improving data transfer efficiency for certain application domains \([189, 55]\). To work around the issue of insufficient compression throughput, preceding investigations have proposed the use of offline data compression, where the payload for data transfers is available in a pre-compressed form. In Chapter 3, this thesis has demonstrated that both hardware-accelerated and purely software-based compression facilities can deliver throughput levels sufficient to saturate 10 Gbit/s, 25 Gbit/s, and even 40 Gbit/s Ethernet networks which are still the norm in the vast majority of data centers \([19]\). Based on this observation, this thesis hypothesizes that on-the-fly data compression can be used to improve data transfer efficiency and consequently overall performance of data-intensive scale-out GPU workloads, as illustrated in Figure 4.5.

![Diagram](image-url)

Figure 4.5: Compared to uncompressed data transfers (left), on-the-fly data compression may increase the effective bandwidth (right).

To test this hypothesis and to open up CloudCL to a wider range of workloads, an approach is presented hereinafter for augmenting CloudCL with on-the-fly data compression with the goal of improving the efficiency of data transfer across the master node and compute nodes. Building up on top of the work conducted in Chapter 3, Section 4.4.1 motivates the reasons for using the 842 compression algorithm to implement on-the-fly data compression in CloudCL. Since the concept of CloudCL detailed in Section 4.2 envisions a very arbitrary cluster model, the integration of on-the-fly data compression mandates that the assumed cluster model is defined more precisely as outlined in Section 4.4.2. Finally, the strategy for implementing on-the-fly data compression based on the integration of lib842 into CloudCL is elaborated in Section 4.4.3.

4.4.1 Choice of Compression Algorithm

To improve data transfer efficiency in CloudCL based on on-the-fly data compression, the 842 algorithm is very well suited as it has been designed for the purpose of transparent main memory compression as discussed in Section 3.2. Unlike lightweight compression approaches that can achieve high throughput by exploiting specific characteristics of datasets such as the employed data type or a restricted range of values, the 842 algorithm is a generic compression algorithm that can be used to compress arbitrary data. As it was demonstrated in Section 3.5.2, this generic approach yields sufficient compression ratios across various data sets, including floating point data.

Another important reason for using 842 compression in this work is the availability of NX-842 on-chip compression accelerators, which are part of all IBM Power CPUs.
introduced since the POWER7+ microarchitecture [17]. The lib842 compression library presented in Chapter 3 makes the resources of the NX-842 accelerators accessible from user-space (cf. Section 3.4), providing very high compression throughput without having to spend excessive amounts of CPU cycles on the task. The software-based implementations provided by lib842 are capable of providing compression throughput high enough to saturate common 10 Gbit/s, 25 Gbit/s, and even 40 Gbit/s Ethernet network links using high-end CPUs (cf. Section 3.5.3). However, the software-based approach is only used as a fall-back option in situations where no NX-842 on-chip compression accelerators may be available.

4.4.2 Assumed Cluster Model

To saturate fast commodity networks such as 10 Gbit/s, 25 Gbit/s, and even 40 Gbit/s Ethernet or faster, on-the-fly data compression requires sufficiently high compression throughput both on the side of the master node and on the side of compute nodes. As depicted in Figure 4.6, the master node must be able to saturate the network interfaces of all compute nodes, therefore having to deal with much larger traffic volumes than each compute node. Therefore, the cluster model assumed in this work employs a master node equipped with NX-842 on-chip compression accelerators, which are available in IBM Power CPUs. On the side of compute nodes, arbitrary CPU types can be used as decompression is handled by the GPU-based 842 decompression facilities provided by lib842. Based on the assumption that the results computed by each compute node are usually smaller in volume compared to the input data, CPU-based software compression based on the lib842 compression library is sufficient to transfer results back to the master node.

Figure 4.6: The cluster model assumed by this work includes a master node equipped with NX-842 on-chip compression accelerators available in IBM POWER CPUs to accelerate compression, and compute nodes equipped with GPUs that use lib842 (cf. Chapter 3) for OpenCL-based decompression on the CPU. All nodes are interconnected using a 10 Gbit/s Ethernet network.

4.4.3 Integration Strategy

The CloudCL architecture outlined in Section 4.2 offers various potential venues for integrating transparent on-the-fly data compression into the CloudCL software stack. A total number of four integration strategies were identified: On-the-fly data compression could be integrated at the level of CloudCL, by introducing extensions at the level of the OpenCL
4.5 Implementation

This section is focused on documenting the cornerstones of implementing on-the-fly compression transparently behind the curtains of the dOpenCL. From the rules and requirements mandated by the OpenCL specification for moving data, three categories of data transfers can be derived: Data transfers from the host to a device, data transfers from a device to the host, and data transfers between devices. While the former two categories can be handled by the OpenCL API, the latter requires a different approach.

For the integration of on-the-fly data compression, dOpenCL uses the lib842 compression library introduced in Chapter 3 to provide access to the hardware-based compression and decompression facilities of the NX-842 on-chip compression accelerators, if available. As a fallback option, the library uses the optimized, software-based implementation for both compression and decompression. On the side of compute nodes, dOpenCL is also responsible for coordinating the workflow of compressed data transfers, using the lib842 compression library to decompress data in GPU memory based on an OpenCL-based decompression kernel. The decompressed buffers are left in the GPU device memory, so that the actual application kernel can work on them without any additional overhead. After the execution of the application kernel has completed, buffers that should be transferred back to the master node are first copied back to the main memory of the compute node. There, the CPU-based software compressor available as part of the lib842 compression library is used to compress data prior to being sent back to the master node.
Figure 4.7: On-the-fly data compression is transparently integrated into CloudCL by modifying dOpenCL to incorporate the compression facilities of lib842.

Operating under the assumption that many workloads are ingesting more data than they egress and considering that the master node in dOpenCL has to supply data to a potentially larger number of compute nodes, the most important type of transfers in dOpenCL that have to be augmented with on-the-fly data compression are data transfers from the master node to compute nodes, as elaborated in Section 4.5.1. Section 4.5.2 then outlines the workflow for realizing compressing data transfers from compute nodes back to the master node in dOpenCL. Even though CloudCL explicitly targets workloads that can be partitioned into independent tasks that do not have to exchange data, the makeshift strategy for implementing compressed data transfers across compute nodes in dOpenCL is documented in Section 4.5.3 to comply with the OpenCL specification.
4.5 Implementation

Figure 4.8: Visualization of the workflows for uncompressed data transfers (a), naïvely compressed data transfers (b), and pipelined compressed data transfers (c).

4.5.1 Master Node to Compute Node Data Transfers

In OpenCL, data transfers from the host to a device triggered explicitly by a call to clEnqueueWriteBuffer, a call to clEnqueueMapBuffer with the CL_MAP_WRITE or CL_MAP_WRITE_INVALIDATE_REGION bits set in the map_flags argument, or a call to clEnqueueUnmapMemObject. The same kind of data transfer can be triggered implicitly by the first use of a buffer created by calling clCreateBuffer with the CL_MEM_USE_HOST_PTR or CL_MEM_COPY_HOST_PTR bits set in the flags argument. To realize host to device transfers in dOpenCL, the corresponding workflow for transparently compressed data transfers from the master node to a compute node are visualized in Figure 4.9. On both the side of the master node and the compute node, all opportunities for interleaving the operations are exhausted. The approach for interleaving the compression process with network transfers on the side of the master node as well as the strategy for pipelining network transfers, device upload, and decompression on the side of compute nodes are documented hereinafter.

Figure 4.9: The workflow for data transfers from the master node to compute nodes leverages hardware-based compression if available to transparently compress buffers prior to sending them out to compute nodes, where they are decompressed on the GPU.
4.5.1.1 Workflow on the Master Node

To interleave the compression and send stages of the workflow on the master node, an OpenCL buffer is partitioned into smaller units, so-called micro-batches, that can be processed independently as illustrated in Figure 4.10. In this approach, a micro-batch is composed of 16 chunks of 64 KiB, each, resulting in a payload of 1 MiB. To fully utilize the resources of the available NX-842 on-chip compression accelerators, a pool of compression threads is used to compress multiple micro-batches concurrently. In the send stage, a network thread then takes care of sending out all micro-batches that have already cleared the compression stage to the compute node. Especially for the send stage, pooling multiple chunks into micro-batches greatly increases the efficiency of network transfers significantly.

Figure 4.10: On the master node, the compression stage and the send stage are interleaved by sending out compressed micro-batches as soon as they are compressed using a pool of compression threads.

4.5.1.2 Workflow on the Compute Node

Similar to the send stage on the master node, pooling the chunks into micro-batches greatly increases the efficiency of the reception stage. On the side of the compute node, a double buffering mechanism using a pair of receive buffers allocated on the OpenCL device is used to interleave the reception stage with the device upload stage as well as the device upload stage with the decompression stage. At any single moment of time, only one buffer is mapped to the host, allowing the network thread to write received micro-batches directly into device memory. To allow for efficient device-based decompression, both buffers are dimensioned large enough to accommodate up to 512 micro-batches each, yielding a macro-batch of 512 MiB. Once the host-mapped receive buffer is filled up, it is unmapped and its sibling buffer is mapped to the host before the decompression kernel is launched on the former buffer. As demonstrated in Figure 4.11, this process is repeated in alternating order until the entire, decompressed OpenCL buffer is available in device memory.
4.5 Implementation

4.5.2 Compute Node to Master Node Data Transfers

The OpenCL standard specifies that data transfers from a device to the host can be triggered explicitly by a call to `clEnqueueReadBuffer`, a call to `clEnqueueMapBuffer` with the `CL_MAP_READ` bits set in the `map_flags` argument, or a call to `clEnqueueUnmapMemObject`. To implement device to host transfers in dOpenCL, the corresponding workflow for transparently compressed data transfers from a compute node to the master node are demonstrated in Figure 4.12.

Both on the side of the compute node and on the side of the master node, the strategies for interleaving stages work analogous to the approaches outlined in Section 4.5.1. Due to the lack of an OpenCL-based compression kernel at the time of writing, the major exception here is that the compression stage is performed on the CPU of the compute node. Since the cluster model assumed in this work (cf. Section 4.4.2) does not assume the availability of on-chip compression accelerators in compute nodes, the software-based compression routine provided by `lib842` are utilized. The software-based compression facilities provided by `lib842` provide high compression throughput on decent CPUs (cf. Section 3.5.3), this means that workloads yielding large result sets are more susceptible to experiencing performance hits for transferring the results from the compute nodes back to the master node.
4.5.3 Compute Node to Compute Node Data Transfers

According to the OpenCL specification, data transfers from one device to another device can be triggered explicitly by calling `clEnqueueCopyBuffer`. Alternatively, device-to-device transfers can be triggered implicitly by any command that depends on an event generated by another command that involves the manipulation of a buffer. Even though CloudCL specifically targets workloads that can be partitioned into independent tasks that do not require any device-to-device interaction, this case has to be addressed to comply with the OpenCL specification. As such, the resulting rudimentary strategy for enabling compressed data transfers across devices in dOpenCL is pointed out in Figure 4.13.

![Data Transfer Diagram](image)

**Figure 4.13:** Due to the lack of peer-to-peer communication among compute nodes in dOpenCL, the workflow for data transfers among compute nodes requires buffers to be transmitted to the master node from which they are forwarded to the respective compute node.

Compressed data transfers among compute nodes re-use the workflows employed on the side of compute nodes for sending (cf. Section 4.5.2) and receiving (cf. Section 4.5.1.2) compressed data. With dOpenCL employing a host-centric architecture, peer-to-peer communication is not available and hence both workflows are stitched together using the master node, which merely forwards the compressed micro-batches received from the sending compute node to the receiving compute node. While this approach certainly provides subpar performance for device-to-device transfers, this scenario is merely covered for the sake of completeness.

### 4.6 Evaluation

The evaluation presented in this section focuses on investigating the performance impact of applying on-the-fly data compression to scale-out GPU workloads implemented using either CloudCL or dOpenCL. Laying out the foundation for the evaluation, Section 4.6.1 documents the testing environment as well as the benchmark procedures used for the evaluation. The effects of on-the-fly data compression on the effective data transfer performance are investigated in Section 4.6.2, Section 4.6.3 then evaluates the impact of applying data transfer compression on the total execution time of four different scale-out GPU workloads. Finally, Section 4.6.4 summarizes the major findings brought forward by the evaluation.
4.6 Evaluation

4.6.1 Testing Environment & Benchmark Procedure

To evaluate the effect of compressed data transfers on the execution time of scale-out GPU workloads across hardware configurations with varying levels of performance, three different classes of compute nodes are employed in addition to the master node to represent potential low, medium, and high-performance configurations of compute nodes. The detailed hardware configurations of each node type are documented in Table 4.1. The medium and high performance compute nodes are equipped with eight GPUs each, and are connected to the same 10 Gbit/s Ethernet switch as the master node. To simulate scale-out behavior, up to eight Docker containers with one GPU attached to each container were employed as depicted in Figure 4.14. By instantiating a varying number of containers, a varying node count could be emulated.

![Diagram of the testing environment and benchmark procedure](image)

Figure 4.14: To simulate a varying number of compute nodes, the employed GPU servers were partitioned into eight compute nodes with one GPU each using Docker containers.

For the low power compute nodes however, up to eight individual bare-metal micro-servers were used instead of the container approach. All micro-servers are attached to the same 10 Gbit/s Ethernet switch as the master node. Across all tests, the same master node was used to warrant a certain degree of commensurability among the different compute node classes.

All performance measurements presented hereinafter were performed after a fresh reboot in order to ensure a clean system state. Furthermore, no other active users or background tasks were running on the involved servers and the network switch was idle. As discussed in Section 4.5, a chunk size of 64 KiB, a micro-batch size of 1 MiB (16 chunks), and a macro-batch size of 512 MiB (512 micro-batches) were employed.

In order to retrieve a sufficiently meaningful dataset, each benchmark was executed 25 times. Error bars are used in all plots to report the standard deviation for each measurement. Furthermore, each benchmark was preceded by a warm-up run in order to eliminate any confounding factors. All measurements presented in this chapter are reported as average values including standard deviation (n = 25).

Execution time was measured from the point where the application is started until it terminated. Therefore, all execution time measurements include the entire execution of a program, including setup, data transfers, computation phases, as well as teardown.
### Table 4.1: Specifications of the test systems used to evaluate the performance impact of applying compressed data transfers in scale-out GPU workloads based on CloudCL and dOpenCL.

<table>
<thead>
<tr>
<th>Master Node</th>
<th>Low Performance Node</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>IBM Power System S824L</td>
</tr>
<tr>
<td>CPU</td>
<td>2×IBM POWER8 (Murano), 3.42 GHz, 10C/80T each</td>
</tr>
<tr>
<td>Memory</td>
<td>1024 GB DDR3 ECC, 1600 MHz</td>
</tr>
<tr>
<td>iGPU</td>
<td>n/a</td>
</tr>
<tr>
<td>dGPU</td>
<td>n/a</td>
</tr>
<tr>
<td>NIC</td>
<td>10 Gbit/s</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 20.04.4</td>
</tr>
<tr>
<td>Kernel</td>
<td>5.4.0</td>
</tr>
<tr>
<td>Compiler</td>
<td>GCC 10.2.1 (AT 14.0)</td>
</tr>
<tr>
<td>OpenCL</td>
<td>n/a</td>
</tr>
<tr>
<td>GPU Driver</td>
<td>n/a</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Medium Performance Node</th>
<th>High Performance Node</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>HPE ProLiant DL380 Gen9</td>
</tr>
<tr>
<td>CPU</td>
<td>2×Intel Xeon E5-2620v4, 2.20 GHz, 10C/20T each</td>
</tr>
<tr>
<td>Memory</td>
<td>256 GB DDR4 ECC, 2133 MHz</td>
</tr>
<tr>
<td>iGPU</td>
<td>n/a</td>
</tr>
<tr>
<td>dGPU</td>
<td>8×NVIDIA Tesla K80</td>
</tr>
<tr>
<td>NIC</td>
<td>10 Gbit/s</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 20.04.4</td>
</tr>
<tr>
<td>Kernel</td>
<td>5.4.0</td>
</tr>
<tr>
<td>Compiler</td>
<td>GCC 9.4.0</td>
</tr>
<tr>
<td>OpenCL</td>
<td>OpenCL 1.2 CUDA</td>
</tr>
<tr>
<td>GPU Driver</td>
<td>470.103.01</td>
</tr>
</tbody>
</table>
4.6.2 Effective Data Transfer Performance

To compare the effective transfer throughput between the master node and a single compute node with and without on-the-fly data compression, a modified version of the oclBandwidthTest sample application from the NVIDIA OpenCL SDK [142] was used. For this test, the synthetic periodic, zeros, and random datasets discussed in Section 3.5.2 were used as compression payloads. These artificial payloads are intended to test effective data transfer bandwidth for worst case and best case edge cases. To include more representative payloads, the enwik9, the OLW, as well as the Curiosity dataset (cf. Section 3.5.2) were included as well.

To evaluate effective transfer throughput in a scale-out scenario, the test application was modified in order to perform data transfers to eight nodes, simultaneously. This test uses the same data sets and only increases the data volume in proportion to the larger number of nodes. The effective transfer throughput is aggregated across all nodes.

![Figure 4.15](image)

**Figure 4.15**: The effective transfer throughput with and without on-the-fly data compression is documented in panel (a). Here, measurements were performed between the master node and a single compute node of each class. The aggregated effective bandwidth for simultaneous transfers to eight compute nodes is illustrated in panel (b).

The measurements for the effective single-node transfer throughput and the effective scale-out transfer throughput are presented in the panels (a) and (b) of Figure 4.15 respectively. The transfer throughput tests demonstrate that using the enwik9, OLW, and Curiosity datasets, compressed data transfers improve effective transfer throughput between 1.29× and 1.81× using medium or high performance compute nodes in the single node scenario, and between 1.40× and 1.91× using any compute node configuration in the scale-out scenario. Random data as the worst-case payload has no negative impact on the throughput, whereas the benevolent periodic and zeros datasets can yield drastic performance improvements. A closer look at the scale-out results for real-world payloads reveals that the limited bandwidth, especially on the master node network interface, remains as a major bottleneck, as the aggregated effective bandwidth of the scale-out tests only slightly exceeds the single node test bandwidth.
4.6.3 Workload Benchmarks

To evaluate whether on-the-fly data compression can be used to mitigate the performance overhead caused by scaling out GPU workloads across multiple compute nodes, the best strategy is to use established benchmark suites. To compensate for the limited choice of multi-GPU benchmarks implemented in OpenCL, four custom benchmarks were implemented in the course of this work using either Java and CloudCL or the C++ bindings of OpenCL. In the custom benchmarks, independent kernel instances are used to process partitions of the input data in order to avoid inter-GPU communication. For each benchmark, the total execution time is measured using regular, uncompressed data transfers for inter-node communication as a performance baseline. By performing the same measurements with compressed data transfers enabled, the baseline performance measurements can be used to quantify the performance improvements introduced by the transparent integration of on-the-fly data compression discussed in Section 4.4 and Section 4.5.

The list of custom benchmarks includes a Semi-Sparse Matrix Multiplication workload, an Analytical Database Query, a Text Search, as well as an Image Downscaler workload. Implemented in Java using CloudCL, the Semi-Sparse Matrix Multiplication workload and the Analytical Database Query were already introduced in Section 4.3. The remaining workloads are implemented in C++, using the C++ bindings of OpenCL. Each benchmark is introduced with a brief description before the results are presented and discussed.

4.6.3.1 Semi-Sparse Matrix Multiplication

As demonstrated in Section 4.3.1, this workload benchmark is implemented in Java using the CloudCL framework. It assumes a matrix multiplication workload where a certain fraction of cells can be assumed to hold zero values, but where this fraction is hard to determine or where it is not large enough to justify the use of a sparse matrix representation such as Compressed Sparse Rows (CSR). The benchmark performs a dense multiplication of matrix $A (N \times M)$ and matrix $B (M \times P)$, yielding matrix $C (N \times P)$ as a result. The dimensions used for $A$ and $B$ in this benchmark are $N = 13125 \times n_{nodes}$, $M = 20000$, and $P = 25$. For node counts larger one, a tiling strategy is employed that partitions matrix $A$ horizontally and distributes it across compute nodes, whereas the second matrix $B$ is sent out to all compute nodes. The matrix multiplication kernel itself is implemented using a naïve implementation strategy. The amount of data to be transferred to compute nodes roughly amounts to $4 \times (N \times M + M \times P \times n_{nodes})$ bytes, and the computation requires roughly $N \times M \times P$ flops.

To regulate the sparsity of the input matrices, both $A$ and $B$ are generated with cells holding either random numbers or a zero value according to the sparsity parameter $S$. For a value of $S = 0$, all cells hold randomly generated values, whereas $S = 1$ results in a fully zeroed matrix. Therefore, the matrices used in this benchmark can be compressed with a ratio $r$ of roughly $r = 1 - S$. The impact of compressed data transfers was tested for the sparsity parameters $S = 0.33$, $S = 0.50$, and $S = 0.67$. 
4.6 Evaluation

Figure 4.16: Panels (a), (b), and (c) present the execution time measurements for the Semi-Sparse Matrix Multiplication workload using low, medium and high power compute nodes, respectively. For each node type, the uncompressed baseline performance is compared to the performance achieved with compression enabled for the sparsity parameters $S = 0.33$, $S = 0.50$, and $S = 0.67$. 
The measurements presented in Figure 4.16 demonstrate that compression can provide performance improvements across the entire range of tested sparsity parameters. On the low performance compute nodes, compression only pays off for larger node counts \((n \geq 4)\) with performance improvements ranging between \(1.11 \times\) for \(S = 0.33\) and \(1.54 \times\) for \(S = 0.66\).

In contrast to the low performance compute nodes however, the medium and high performance compute node types show slight but consistent performance improvements even for low node counts \((1 \leq n \leq 2)\). For larger node counts \((n \geq 4)\), compression yields performance improvements between \(1.23 \times\) for \(S = 0.33\) and \(1.87 \times\) for \(S = 0.67\). However, it should be noted that this benchmark is dominated by transfer time and only a fraction of the execution time is spent on computation.

### 4.6.3.2 Analytical Database Query

For this benchmark, an Analytical Database Query was implemented in Java using the CloudCL framework as discussed in Section 4.3.2. The implemented workload resembles the characteristics of a column-oriented in-memory database executing an analytical query that resembles Query 1 of the TPC-H Benchmark \[206\]. Test data for this benchmark is generated loosely following the principles of the DBGEN data generator \[206\], with the minor modification that the Java pseudo random number generator is used instead of the custom pseudo random number generator specified by the DBGEN data generator.

At this point, the author wishes to reiterate that for reasons of simplicity, neither the query nor the data generator fully complies with the very complex TPC-H specification. As such, the Analytical Database Query benchmark must not be mistaken for a subset TPC-H benchmark. The implemented query is a relatively simple, join-free aggregation query that involves a simple filter statement. Only the relevant data columns are transferred to the GPUs using a columnar layout. The data volume to be transferred and processed amounts to \(28 \times 100000000\) bytes per node, and \(28 \times 100000000 \times n_{\text{nodes}}\) bytes in total. The GPUs perform the aggregation, to the extent possible, in parallel.

The benchmark results depicted in Figure 4.17 demonstrate that compression allows the query execution to scale almost perfectly on up to four nodes, as the total execution time barely increases compared to a single node. For two to four nodes, on-the-fly data compression facilitates almost perfect scaling behavior across all node types, as the multi-node execution times are barely higher compared to the single-node configurations. For the \(n = 8\) nodes, performance improvements of \(1.85 \times\), \(1.9 \times\), and \(2.07 \times\) are achieved for low, medium and high performance compute nodes, respectively. Based on these observations, it seems safe to assume that the network interface on the master node is the major bottleneck. With a wider network interface available on the master node, compressed data transfers have the potential to provide perfect scale-out behavior for even larger node counts.

### 4.6.3.3 Text Search

Here, a simplistic Text Search kernel was implemented that checks for a match at each position of a large text file. Unlike the preceding benchmarks, this test was implemented in C++ using the OpenCL C++ bindings, and therefore runs directly on top of the dOpenCL
4.6 Evaluation

Figure 4.17: The execution time measurements for the Analytical Database Query workload are reported in panels (a), (b), and (c) for low, medium and high power compute nodes, respectively. Each node has to process the same volume of $28 \times 10^6$ bytes, meaning that a constant execution time across node counts expresses perfect scale-out behavior. Up to a node count of four, this ideal scale-out behavior is approached across all compute node types using on-the-fly data compression.
library. The benchmark is performed using the Books, Wikipedia, and OLW datasets as large text corpora (cf. Section 3.5.2). A simple, computationally expensive but yet powerful implementation strategy is employed that can match any pattern, even non-regular ones. Using this naïve approach also yields a workload that is dominated by compute time instead of data transfer time. Depending on the number of nodes, the first $1000000000 \times n_{\text{nodes}}$ bytes of an employed data set are transmitted to compute nodes, with each node having to process $1000000000$ bytes.

Figure 4.18: Panels (a), (b), and (c) report the execution times measured for the Text Search workload on low, medium, and high performance compute nodes, respectively. The black portion of each bar represents the performance baseline of using uncompressed data transfers. The data supports the assumption that this benchmark is dominated by compute time, as the different performance levels of compute nodes can be easily identified. Even though the benefit of on-the-fly data compression for data transfers is not as distinct as in other workloads, compression becomes more beneficial for higher node counts.

Considering the benchmark results provided in Figure 4.18, the first impression might be that compression does not help too much in this use case, especially for faster compute node configurations. However, as mentioned before, this benchmark is more compute-intensive, which can be seen based on the larger performance differences between the different compute node classes. With the tests being less sensitive to data transfer volumes, it is still notable to see that compression yields $1.14 \times$, $1.25 \times$, and $1.43 \times$ performance improvements using $n = 8$ low, medium and high performance compute nodes, respectively.
4.6 Evaluation

4.6.3.4 Image Downscaler

Last but not least, a simple Image Downscaler workload was implemented in C++ using the OpenCL C++ bindings. In this benchmark, a Tag Image File Format (TIFF) image is read and transferred to all available GPUs in the form of an RGBA pixel buffer. To utilize multiple GPUs, the workload is partitioned by segmenting the image horizontally. As reference payloads, the Curiosity and Telescope datasets are used (cf. Section 3.5.2). In contrast to the other workloads, this test does not clip the datasets proportionally to the number of available nodes $n_{\text{nodes}}$, but the entire image is processed regardless of the employed node count.

![Graphs](image)

**Figure 4.19:** Panels (a), (b), and (c) report the execution times measured for the Image Downscaler workload on low, medium, and high performance compute nodes, respectively. The black portion of each bar represents the performance baseline of using uncompressed data transfers. The measurements clearly illustrate that the workload is dominated by data transfers, as the execution time does not vary significantly across compute node classes and varying node counts. Here, the use of on-the-fly data compression yields significant performance improvements across most conditions.

The results presented in **Figure 4.19** illustrate that this workload is largely dominated by transfer time, as the baseline execution time remains almost constant across varying node counts. With the employed datasets being well compressible, this benchmark makes it easy to gauge the impact of on-the-fly data compression, which improves execution time considerably by up to $1.67 \times$, $1.71 \times$, and $1.89 \times$ on low, medium, and high performance compute nodes, respectively. Nevertheless, the test also demonstrates that scalability is ultimately limited by width of the network, even when data is compressed.
4.6.4 Summary

The preceding evaluation has successfully tested the hypothesis that on-the-fly data compression can improve the overall performance of scale-out GPU workloads using various compute node configurations by increasing the effective bandwidth between the master node and compute nodes. Ranging between $1.11 \times$ and $2.07 \times$, the performance improvements observed across various workloads may not appear drastic on first sight. However, it should be noted that this speed-up was achieved without assuming any workload-specific knowledge in the compression scheme, without necessitating any modifications in the workloads themselves, or without introducing any other kind of overhead. Considering that the presented approach is capable of introducing even modest speed-up to a very wide range of GPU-based scale-out workloads, performance improvements up to $2.07 \times$ appear much more attractive on second sight, especially in the context that the number of applications that require multiple GPUs to satisfy their resource demand is increasing by the day.

4.7 Summary

This chapter has presented two major contributions:

Building up on top of the dOpenCL API forwarding library for OpenCL and the Aparapi framework for executing native Java code on GPUs, the CloudCL framework was introduced. By extending the underlying technologies with a job infrastructure including a job scheduler, as well as dynamic scaling capabilities for dynamically available resources, the CloudCL framework hides several aspects of the distributed memory parallel programming model during the development of scale-out GPU workloads. These abstractions enable application developers and domain experts to focus on the data parallel programming model associated with GPUs, yielding a single-paradigm development experience which makes scale-out GPU resources more accessible to a wider audience. From an operations point of view, CloudCL can also improve resource utilization by disaggregating GPU resources. The improved developer experience provided by CloudCL was demonstrated by presenting the job class implemented using the CloudCL framework for two exemplary workloads.

Serving as the foundation of the CloudCL framework, the dOpenCL library was augmented with transparent on-the-fly data compression for inter-node data transfers based on the lib842 compression library presented in Chapter 3. Using a highly pipelined approach to interleave all stages of the workflow for transferring transparently compressed OpenCL buffers from the master node to compute nodes or vice versa, it was possible to improve the effective throughput across nodes. From a workload perspective, the improved data transfer efficiency provided by the integration of transparent compression has yielded performance improvements ranging between $1.11 \times$ and $2.07 \times$ across various data-intensive scale-out GPU workloads implemented using either the CloudCL framework or the OpenCL API directly.
5 Programming Abstractions for Scale-Up Non-Uniform Memory Access Architectures

In this chapter, the PGASUS C++ framework is introduced with the goal of alleviating application development for scale-up Non-Uniform Memory Access (NUMA) architectures by providing easy-to-use facilities for memory placement and NUMA-aware task-parallelism. The PGASUS framework was originally proposed and extended in the master’s theses by Wieland Hagen [68] and Karsten Tausche [199], respectively. Building up on top of the concept and the implementation of the PGASUS framework brought forward by these master’s theses, the contributions of this work in the context of NUMA architectures are focused on investigating the impact of the programming abstractions provided by PGASUS on both the developer experience and performance. The developer experience of the framework is investigated exemplarily based on three different workloads, including a data compression workload that builds up on the lib842 compression library presented in [Chapter 3]. Furthermore, a comprehensive evaluation is conducted to investigate the performance-impact of the PGASUS framework.

The following master’s theses were supervised alongside the research leading to this chapter, fostering scholarly exchange between this work and the supervised theses:

- Christoph Sterz. “Analyzing NUMA Performance Based on Hardware Event Counters”. Master’s thesis. Potsdam, Germany: Hasso Plattner Institute, University of Potsdam, July 2016. URL: https://osm.hpi.de/bookshelf/Details/530

Furthermore, partial results of the work presented in this chapter have been published:

- Wieland Hagen, Max Plauth, Felix Eberhardt, Frank Feinbube, and Andreas Polze. “PGASUS: A Framework for C++ Application Development on NUMA architec-
5 Programming Abstractions for Scale-Up Non-Uniform Memory Access Architectures

- Max Plauth, Felix Eberhardt, Andreas Grapentin, and Andreas Polze. “Improving the Accessibility of NUMA-Aware C++ Application Development Based on the PGASUS Framework”. In: Concurrency and Computation: Practice and Experience (Feb. 2022), e6887. doi: 10.1002/cpe.6887

This chapter is structured as follows. Section 5.1 motivates the demand for programming abstractions that make it easier for developers to exploit data locality in scale-up NUMA systems without disregarding their advantage of providing cache coherency across NUMA domains. After that, Section 5.2 explains why neither the C++ standard library nor operating system Application Programming Interfaces (APIs) such as libnuma do not provide suitable means for controlling data placement for C++ objects on NUMA systems. Section 5.3 then introduces the PGASUS framework and the facilities it introduces to alleviate NUMA-aware application development in C++. To demonstrate the developer experience of PGASUS, Section 5.4 compares PGASUS-based implementations of three different workloads with NUMA-aware implementations based on the Open Multi-Processing (OpenMP) interface or a combination of POSIX pthreads and the libnuma library. The performance impact of the abstractions introduced by PGASUS are investigated in a comprehensive evaluation in Section 5.5. Finally, the major insights from this chapter are summarized in Section 5.6.

5.1 Motivation and Problem Statement

Even though Graphics Processing Units (GPUs) have become popular in many data-intensive application domains, many workloads still rely on the flexibility and versatility of multicore Central Processing Units (CPUs) [208]. While several of these CPU-based workloads can be adapted to scale-out across multiple systems to provide sufficient compute resources, certain workloads such as in-memory databases [25] or de Novo genome assembly [133] are inherently hard to scale out and therefore require as many resources as possible in a single scale-up system.

As elaborated in Section 2.1.1 Uniform Memory Access (UMA) architectures have dominated multiprocessor systems for a long time. From the perspective of an application developer, UMA architectures align conveniently with the shared memory programming model. Unfortunately, sharing the memory subsystem with all other multicore CPUs severely limits the scalability of multiprocessor systems, both in the number of multicore CPUs and in the amount of memory that can be accommodated in a single system.

NUMA architectures avoid this bottleneck, as each multicore CPU is equipped with dedicated memory controllers. Memory attached to other multicore CPUs can still be accessed transparently through inter-CPU interconnects such as Ultra Path Interconnect (UPI), Infinity Fabric (IF) and Power with A-bus, X-bus, OpenCAPI, and NVLink (PowerAXON). However, remote memory access operations incur increased latencies and reduced bandwidth, especially on systems with more than four multicore CPUs, where fully meshed connectivity among CPUs is no longer feasible. State-of-the-art NUMA systems support
Data Placement in NUMA Systems

5.2 Data Placement in NUMA Systems

For developers intending to factor in the properties of NUMA systems in their applications, several challenges regarding data placement have to be considered during the development of C++ applications. Hereinafter, the implications of object placement and object migration on NUMA-aware application development are identified, where neither...
the C++ standard library nor operating system mechanisms provide sufficient means to express data locality on NUMA systems.

### 5.2.1 Object Placement

The C++ programming language and its standard library do not have any concept for considering data locality in NUMA topologies. As a consequence of the virtual memory abstraction, a flat, homogeneous address space is presented to applications, where regions can be made available through operating system APIs such as `libnuma`\(^\text{[100]}\). These regions are identified only by location and length, and are otherwise indistinguishable, as application developers are supposed to be indifferent about any details of the underlying hardware. As such, no mechanisms are provided to group data, prevent intra-page fragmentation or otherwise deal with the specific challenges of NUMA topologies, as illustrated in Figure 5.1.

![Figure 5.1: Example of how a `std::vector<std::string>` instance and its underlying data structures can be spread across many distinct pages.](image)

To control memory placement decisions at runtime, an application has to either provide a custom implementation of the `new` operator that overrides the default behavior, or memory has to be allocated manually in advance. However, neither method considers that every class used in a context sensitive to object placement needs to be allocated using the modified `new` operation. This cannot be guaranteed for classes that are defined outside of the program such as libraries that may implement custom allocation schemes. Also, many template-based container data-structures rely on the default behavior of placement `new` and implement their own memory management based on `malloc`. Lastly, overwriting `new` has no effect on data-structures implemented in C libraries that use `malloc`.

### 5.2.2 Object Migration

The issues faced during object migration are very similar to those described for object placement. The page migration mechanisms provided by the operating system for moving data to a different NUMA node does not consider the internal, potentially nested structure of objects. To move an instance of `std::vector<std::string>` to another node for example, not only the pages containing the `std::string` instances have to be moved to the other node, but also the pages holding the string data that are allocated separately on
the heap have to be considered for each `std::string` instance as illustrated in Figure 5.1. In the described scenario, the lack of control over the placement of the `std::vector` and `std::string` data storage is a problem. When more complex objects containing nested object hierarchies have to be migrated using the page migration mechanisms, all associated objects have to be allocated in contiguous memory and occupy a private set of pages that is not shared with other objects.

5.3 PGASUS: NUMA-Aware C++ Application Development

This section provides an overview of the PGASUS framework, which has been proposed in the master’s thesis by Wieland Hagen [68] and which has been further extended in the course of the master’s thesis by Karsten Tausche [198]. PGASUS is a C++ framework that employs PGAS semantics on NUMA systems in order to provide developers with the means to specify data placement policies based on the RAII idiom. To furthermore alleviate the co-location of data and tasks, PGASUS also provides a simple NUMA-aware tasking infrastructure. The PGASUS framework provides five major facilities to alleviate the development of NUMA-aware C++ applications: MemSources are used to represent logical memory regions that are bound to a specific NUMA node and provide the foundation for the concept of PlaceGuards, which configure an underlying memory allocator to serve allocations from a specific MemSource using the RAII idiom. An interface for discovering the NUMA topology of a system provides the means for developers to respond to the characteristics of a NUMA system at runtime, whereas NUMA-aware Task Parallelism is used to situate tasks based on the location of the data they are operating on. Finally, PGASUS provides a NUMA-aware hash-table to investigate the potential of NUMA-aware drop-in replacements of common data-structures. All five facilities are further detailed hereinafter.

5.3.1 MemSources

The `libnuma` API [100] in Linux provides two methods for influencing the placement of data: Using calls to `numa_alloc_onnode`, memory can be allocated explicitly on the specified node. Memory allocated using this method is always page aligned, which may lead to internal fragmentation for small allocations, leaving large parts of the memory in a page potentially unused. For existing code bases to incorporate this method, all allocations have to be replaced with calls to `numa_alloc_onnode` and the size of allocations has to be tracked to replace deallocations with the corresponding `numa_free` call.

Alternatively, using the `numa_set_membind` call implies a contextual approach. This method can be used to specify which NUMA nodes should be used to serve subsequent allocations. Like the other method, `numa_set_membind` also operates on the granularity of pages, which means that the specified placement policy only applies to pages that are faulted into the heap after the `numa_set_membind` call. This behavior can yield false sharing effects in situations where an allocator serves small allocations from a page that has been faulted into the heap prior to the call to `numa_set_membind` and therefore may be placed on a different NUMA.
To avoid these pitfalls, *PGASUS* introduces *MemSources* as the central means for representing logical memory regions that are bound to a specific NUMA node. As exemplified in Listing 5.1, *MemSources* can be used to serve allocations for objects of arbitrary size and type. Furthermore, *MemSources* provide a mechanism for conveniently migrating all of its pages to another node, and also making sure that allocations are served from that node thereafter.

**Listing 5.3:** *MemSources* represent a logical memory region that is bound to a specific NUMA node.

The *MemSource* interface can be used to group memory allocations, to control allocation placement, and to migrate groups of objects between NUMA nodes.

```cpp
1.  int initialSize = 1 << 24;        // 16 MiB
2.  MemSource msource = MemSource::create(targetNode, initialSize);
3.  Foo *foo = msource->construct<Foo>(); // create object
4.  void *buffer = msource->alloc(1024); // allocate memory
5.  msource->migrate(newHomeNode);    // migrate pages
```

A *MemSource* maintains a pre-allocated buffer bound to its home node using *libnuma* from which smaller allocations are served using an optimized memory allocator. At first sight, well-known high-concurrency allocators such as *jemalloc* and *TCMalloc* may seem like a good choice from a performance perspective. Considering the segregated allocation strategy employed by *jemalloc*, *TCMalloc*, as well as many other high-concurrency allocators, they cannot be applied directly to a single, externally allocated memory block such as the pre-allocated buffer provided by a *MemSource* as segregated allocators employ many distinct memory blocks to serve allocations of different size classes. As a result, *MemSources* are implemented using the more conservative *ptmalloc* allocator employing a best-fit allocation strategy. The allocator supports the *mspace_t* construct, which can be created within the pre-allocated buffers provided by the *MemSources*. These pre-allocated buffers and the *mspace_t* constructs therein will be referred to as * Arenas*. Large allocations are allocated directly via *mmap* and are bound to the home node of the *MemSources* using *libnuma*.

For object location querying and, more importantly, object de-allocation, it is imperative that each object allocated from a *MemSource* can be identified. Since the *free* function used for memory de-allocation in C only takes one pointer to the memory chunk, it has to be possible to query the *MemSource* belonging to a memory chunk by just using this pointer. For this purpose a data item called *Footer* is stored before each allocated chunk. This *Footer* contains a pointer to the *MemSource* and the *Arena* that the allocation was made from. For large chunks allocated using *mmap* directly, the arena pointer is set to NULL and the footer is extended to also include the block size and links to other *mmap*-allocated chunks, thus implementing a linked list.

### 5.3.2 Place Guards

Following the RAII idiom, *PGASUS* introduces the *PlaceGuard* construct to control the behavior of the *new* operator and all common memory allocation functions to allocate memory from a *Node* (cf. Listing 5.2) or a *MemSource* (cf. Listing 5.3) instead of using the NUMA-agnostic *malloc* implementation of the C library. After a *PlaceGuard* is created,
its effects are active until the PlaceGuard goes out of scope. Their effect can also be nested by specifying additional PlaceGuards within an already guarded scope, allowing for fine-grained control over the allocations of the application, and even of third-party code. When a PlaceGuard instance goes out of scope and is deallocated, its effect will end and subsequent allocations will be served by the previously active parent PlaceGuard, if any, and otherwise will return to the malloc implementation of the C library.

**Listing 5.2:** PlaceGuards enable developers to easily specify object placement by configuring the behavior of the underlying stacked malloc allocator. Both the allocated string object and its underlying string data buffer will be allocated on the specified targetNode.

```cpp
1 std::string* createStringOnTargetNone(Node targetNode) {
2     PlaceGuard guard(targetNode);
3     return new std::string("foo");
4     // PlaceGuard leaves scope and loses effect
5 }
```

**Listing 5.3:** PlaceGuards can also be specified in relation to a previously created MemSource instead of a NUMA node. In this case, the data buffer inside the string implementation will be allocated using the given MemSource.

```cpp
1 std::string* createStringInMemSource(MemSource source) {
2     PlaceGuard guard(source);
3     return new std::string("foo");
4     // PlaceGuard leaves scope and loses effect
5 }
```

To achieve this behavior, the PlaceGuard construct is backed by the stacked malloc allocator, which replaces all memory allocation functions defined by the C++ standard including inherited C interfaces as well as specific POSIX interfaces with the behavior described hereinafter. In stacked malloc, every thread maintains a stack of places, which can be either references to MemSources or NUMA nodes. When the PlaceGuard construct is invoked, the specified MemSource or NUMA node is pushed onto the stack. Upon leaving scope, the PlaceGuard construct removes the corresponding place from the stack. To serve allocations, stacked malloc consults the top element of the stack. In case a MemSource resides on the top of the stack, allocations are served thereof. When a node is the top element, the default MemSource residing on the specified node is used to serve the allocation.

Upon deallocation of a memory chunk, it has to be known from which MemSource the memory chunk was allocated. Thus the allocated memory chunks are annotated with information that facilitates a link to the MemSource it stems from. It is then possible to ask that MemSource to deallocate the given chunk. The basic algorithms for memory allocation and deallocations are shown in **Listing 5.4**.

**Listing 5.4:** Simplified operating principle employed by the stacked malloc allocator.

```cpp
1 MemSource *getMemSource() {
2     tls = getThreadLocalStorage();
3     places = tls.placeStack;
4     if (!places.empty() && places.top.isMemSource())
5         return places.top.memsoure;
```
Programming Abstractions for Scale-Up Non-Uniform Memory Access Architectures

```cpp
int node = places.empty() ? localNode : places.top.node;
if (node == localNode)
    return tls.localMemSource;
return tls.nodeData.msources[node];
```

```cpp
static const int offset = sizeof(MemSource::Footer);

void malloc(size_t sz) {
    MemSource *ms = getMemSource();
    return ms->alloc(sz);
}

void free(void *p) {
    void *chunk = p - offset;
    MemSource *ms = *((MemSource**) chunk);
    ms->free(p);
}
```

5.3.3 Topology Discovery

PGASUS provides means for retrieving the topology of a system based on the Node and NodeList classes outlined in Listing 5.5. The framework creates a model of the machine topology by using the information provided by hwloc and /sys/devices/system/node/node[X]/distance. Certain hardware configurations can result in NUMA node ids which may not be consecutive. Such situations include hypervisors such as PowerVM, systems with coherently attached accelerator memory, or disaggregated memory setups.

To better deal with a non-linear id space, the topology interface employs a logical node mapping using consecutive node ids. Similarly, there are configurations where NUMA nodes may only contain memory resources and no CPU resources. To address the possibility of compute-less NUMA nodes, the helper methods in the NodeList class provide additional variants considering only NUMA nodes equipped with CPU resources.

Listing 5.5: PGASUS exposes topology information using the Node and NodeList classes.
5.3 PGASUS: NUMA-Aware C++ Application Development

```cpp
static NodeList& logicalNodes(); // as detected at runtime
static size_t logicalNodesCount();

static const NodeList& logicalNodesWithCPUs();
static size_t logicalNodesWithCPUsCount();

static int physicalToLogicalId(int physicalId);
static size_t physicalNodesCount();
```

5.3.4 NUMA-aware Task-Parallelism

Even though a plethora of very mature parallel tasking libraries are readily available, they are too complex to prototypically incorporate the notion of MemSources and the partitions of the global address space they represent. To investigate the developer experience as well as the performance impact of a parallel tasking mechanism that respects the locality aspects of a partitioned global address space, PGASUS provides a simple parallel tasking infrastructure. As demonstrated in Listing 5.6 the tasking interface proposed for PGASUS follows the general concepts of the interfaces for threading and asynchronous calls in C++11 and onwards. Task functions are defined as a `std::function<T>` where T is the return value type which may also be void. For the specification of a task, a priority level as well as a target NUMA node can be specified that the task should be bound to. In addition to using named functions to define tasks functions, lambda expressions introduced with C++11 may be used to define anonymous task functions. Spawning a task returns a Future object that can be used to wait for the task to finish or to obtain the result value. These simple but versatile tasking facilities enable developers to easily co-locate tasks and data by specifying a target node a task will be bound to, which is typically the home node of a MemSource the task should operate on. If no target node is specified, the task is executed on an arbitrary node when no tasks bound to that specific node are available.

Listing 5.6: Spawning tasks using C++ lambdas yields a future object that can be used to wait for the completion of the task. This behavior enables asynchronous and synchronous task parallelism.

```cpp
auto task = numa::async(targetNode, []() {
    std::cout << "Executed on node "
              << Node::current() << std::endl;
    return 42;
});
// do something else in the meantime ...
auto result = task.wait(); // result = 42
```

5.3.5 NUMA-aware Hash Table

To investigate the potential of implementing NUMA aware drop-in replacement of common data-structures based on PGASUS, a NUMA-aware Hash Table is provided as a part of the PGASUS framework. The custom Hash Table was implemented using PGASUS and allows an arbitrary amount of concurrent insert, update, read and delete operations.
Developers only have to make sure that they only perform delete operations on objects that are not currently read, modified, or iterated upon.

The Hash Table is divided into $2^N$ buckets, each of which is resides on a specific NUMA node and is responsible for a part of the index space. The last $N$ bits of a key's hash value are used to identify the bucket responsible for storing that key. Each bucket is furthermore subdivided into $2^M$ bins. Bins are linked lists that store an arbitrary, but usually a very small amount of key-value pairs.

Synchronization is applied at a very fine-grained level via Reader-Writer locks. Bin entries are reference-counted to relax synchronization constraints. The Hash Table features a number of hierarchical iterators, allowing the iteration space to be divided into sub-iterations for each node. Automatic parallel iteration over the data-structure is implemented by collecting all bucket iterators for each node. Each worker thread then iterates over iterators from that node. Whenever there are iterators left for remote nodes after all local iterators have been processed, workers start stealing from this remote work pool.

### 5.4 Developer Experience

A central aspect of this chapter is to demonstrate the developer experience of the PGASUS framework. Therefore, this section introduces three carefully selected workloads that are used to demonstrate the capabilities and limitations of the framework: A Text Histogram application and a Data Compression workload are employed as embarrassingly data parallel workloads using a fine-grained and a coarse-grained task granularity, respectively. Furthermore, a Database Table Scan workload is used to represent a more challenging, irregular workload. For the Text Histogram and Data Compression workloads, three implementations are compared against each other: an implementation entirely based on PGASUS, a NUMA-agnostic implementation based on the OpenMP interface, and a NUMA-aware implementation that combines the mature task parallel computing capabilities of OpenMP with the data placement capabilities of PGASUS. The Database Table Scan workload is presented to compare a C-based implementation based on pthreads and libnuma with a C++-based implementation based on PGASUS.

### 5.4.1 Text Histogram

Using the massive corpus of public domain text books provided by Project Gutenberg [166], an embarrassingly parallel text histogram workload is used as a stress test for the PGASUS tasking component for fine-grained tasks. Counting the occurrence of each word, text histograms are computed on a per-book granularity, with each book representing in the order of hundreds of kilobytes of data that needs to be processed. As each book is stored in an individual text file, the proposed interface for each text file is outlined in [Listing 5.7].

**Listing 5.7:** For the Text Histogram workload, each book of the Project Gutenberg [166] corpus is represented using a TextFile object.

```cpp
class TextFile {
  std::string fileName;
  std::string fileContent;
```

Project Gutenberg [166]
5.4 Developer Experience

```cpp
std::map<std::string, int> wordHistogram; // occurrences of each word

void computeHistogram();
};
```

To compare the implementation effort for the simplistic parallelization strategy of the 
Text Histogram workload, a NUMA-agnostic OpenMP-based implementation, a NUMA-aware 
OpenMP-based implementation, and a PGASUS implementation are explicated 
hereinafter. The NUMA-agnostic OpenMP implementation presented in Listing 5.8 serves 
as a baseline that the NUMA-aware implementations can be compared to.

Listing 5.8: OpenMP-based implementation of the loadFiles and computeHistograms methods of the Text Histogram workload.

```cpp
void loadFilesOMP(const std::vector<std::string> &fileNames) {
    #pragma omp parallel for
    for (size_t i = 0; i < fileNames.size(); i++) {
        auto f = std::unique_ptr<TextFile>(new TextFile(fileNames[i]));
        #pragma omp critical(fileaccess)
        files[fileNames[i]] = std::move(f);
    }
}

void computeHistogramsOMP(const std::vector<TextFile*> &files) {
    #pragma omp parallel for
    for (size_t i = 0; i < files.size(); ++i) {
        files[i]->computeHistogram();
    }
}
```

During the initialization phase of the Text Histogram workload, the books of the Project Gutenberg corpus are loaded into main memory. All NUMA-aware implementations employ a simple round robin scheme to evenly distribute the resulting TextFile objects across NUMA nodes. The first NUMA-aware implementation outlined in Listing 5.9 combines the mature parallel tasking facilities of OpenMP with the data placement capabilities of PGASUS. In the loadFiles method, PGASUS is used to bind the nodeStorage elements to the respective nodes. The PGASUS topology interface is used extensively both in the loadFiles method as well as the computeHistograms method to avoid the complexity of performing topology discovery manually. Assuming the places policy being set to sockets upon launch, the NUMA-aware OpenMP-based implementation relies on two nested parallel statements in the computeHistograms method, with the outer statement scheduling one master thread per NUMA node. On the level of the nested parallel for statement, the location of each nodes master thread is inherited for the proc_bind statement in order to schedule threads for all logical cores on the current node.

Listing 5.9: NUMA-aware OpenMP-based implementation of the loadFiles and computeHistograms methods of the Text Histogram workload.

```cpp
void loadFilesPGASOMP(const std::vector<std::string> &fileNames) {
    const auto& numaNodes = numa::NodeList::logicalNodesWithCPUs();
    const size_t totalCPUCount = std::accumulate(...)
```
std::vector<std::vector<std::string>> perNodeFileNames(numaNodes.size());

// Distribute files/jobs to NUMA nodes according to local number of CPU cores
const float distFactor = float(fileNames.size()) / totalCPUCount;
size_t nextFileName = 0u;
for (size_t node = 0; node < numaNodes.size(); ++node) {
    const size_t localCount = std::ceil(numaNodes[node].cpuCount() * distFactor);
    for (size_t l = 0; l < localCount && nextFileName < fileNames.size(); ++l, ++nextFileName) {
        perNodeFileNames[node].push_back(fileNames[nextFileName]);
    }
}

nodeStorages.resize(numa::NodeList::logicalNodesCount());
#pragma omp parallel proc_bind(spread) num_threads(numaNodes.size())
{
    const auto node = numa::Node::curr();
    const numa::PlaceGuard placeGuard{ node };
    const auto nodeId = node.logicalId();
    nodeStorages[nodeId] = std::unique_ptr<NodeStorage>(new NodeStorage);
    NodeStorage &nodeStorage = *nodeStorages[nodeId];
    const auto& localFileNames = perNodeFileNames[nodeId];
    #pragma omp parallel for proc_bind(master) num_threads(node.threadCount())
    for (size_t i = 0; i < localFileNames.size(); ++i) {
        const std::string& fileName = localFileNames[i];
        auto f = std::unique_ptr<TextFile>(new TextFile(fileName));
        auto fPtr = f.get();
        #pragma omp critical(fileaccess)
        {
            nodeStorage.files.push_back(std::move(f));
            nodeStorage.filesMap.emplace(fileName, fPtr);
        }
    }
}

void computeHistogramsPGASOMP() {
    omp_set_nested(1);
    const auto& numaNodes = numa::NodeList::logicalNodesWithCPUs();
    #pragma omp parallel proc_bind(spread) num_threads(numaNodes.size())
    {
        const auto node = numa::Node::curr();
        std::vector<const TextFile*> &localFiles
            = nodeStorages[node.logicalId()]->files;
        #pragma omp parallel for proc_bind(master) num_threads(node.threadCount())
        for (size_t i = 0; i < localFiles.size(); ++i) {
            localFiles[i]->computeHistogram();
        }
    }
}
The PGASUS-based implementation exemplified in Listing 5.10 uses the NUMA-aware hash table introduced in Section 5.3.5 to keep the TextFile objects balanced across all nodes in the loadFiles method. By leveraging the NUMA-aware hash table implementation provided by PGASUS, a simple lambda expression in the computeHistograms method is sufficient to define tasks which are scheduled for execution on the NUMA node on which the respective TextFile object resides. In terms of code complexity, NUMA-awareness based on PGASUS can be achieved with minimal effort, whereas the nested parallel statements of the NUMA-aware OpenMP-based implementation are more complex.

Listing 5.10: PGASUS-based parallelization of the text histogram workload.

```cpp
t numa::HashTable<std::string, std::unique_ptr<TextFile>, 6> files;

virtual void loadFilesPGASUS(const std::vector<std::string> &fileNames) {
    std::list<TriggerableRef> waitList;
    for (const std::string &file : fileNames) {
        waitList.push_back(files.insertAsync(file, [file] { return std::unique_ptr<TextFile>(new TextFile(file)); }));
    }
    numa::wait(waitList);
}

void computeHistogramsPGASUS(const std::vector<TextFile*> &files) {
    std::list<TriggerableRef> waitList;
    for (const TextFile* : files) {
        waitList.push_back(numa::async<void>([this, file] { files[file]->wordHistogram(); }, 0, files.where(file).getNode()));
    }
    numa::wait(waitList);
}
```

5.4.2 Data Compression

To complement the fine-grained quality of the Text Histogram workload, a Data Compression workload is used to provide a coarse-grained task profile. The first 10⁹ bytes from the 2006-03-03 Wikipedia dump (enwik9)[121] are compressed and decompressed using the 842 compression algorithm [54]. Also belonging to the category of embarrassingly data parallel problems, the Data Compression workload employs a coarse-grained task profile with each task compressing or decompressing tens of megabytes of raw data using the highly optimized lib842 library presented in Chapter 3.

To compare the implementation effort of the parallelized Data Compression workload, a NUMA-agnostic OpenMP-based implementation, a NUMA-aware OpenMP-based implementation, and a PGASUS implementation are discussed hereinafter. The NUMA-agnostic OpenMP-based implementation of the chunk-wise compression routine of the lib842 li-
brary outlined in Listing 5.11 serves as a baseline that the NUMA-aware implementations can be compared to.

Listing 5.11: OpenMP-based parallelization of the Data Compression workload.

```c
#pragma omp parallel for
for (size_t chunkNum = 0; chunkNum < num_chunks; chunkNum++) {
  const uint8_t *chunk_in = input_buffer + (chunkNum * CHUNK_SIZE);
  uint8_t *chunk_out = compressed_buffer + (chunkNum * (CHUNK_SIZE * 2));
  ...
  compress(chunk_in, CHUNK_SIZE, chunk_out, ...);
}
```

During the initialization phase of the Data Compression workload, the enwik9 data set is loaded into main memory. Instead of loading the file into a single buffer, all NUMA-aware implementations split up the file contents into sub-buffers bound to each NUMA node. The [NUMA-aware OpenMP-based implementation outlined in Listing 5.12] again combines the mature parallel tasking facilities of OpenMP with the data placement capabilities of PGASUS to bind the sub-buffers residing in input_buffers and compressed_buffers to the respective NUMA nodes. The PGASUS topology interface is used in the NUMA-aware OpenMP-based implementation to avoid the complexity of performing topology discovery manually. Like the Text Histogram workloads, the places policy has to be set to sockets upon launch. This policy is used to schedule one master thread per NUMA node in the outer parallel section. Each nodes master thread then executes a nested parallel for section where the location of each nodes master thread is inherited for the proc_bind statement in order to schedule threads for all logical cores on the current node. To make sure that each thread operates on the correct sub-buffers and the respective chunks therein, the NUMA-aware OpenMP-based implementation has to compute the identifiers nodeId and localThreadId for each thread.

Listing 5.12: NUMA-aware OpenMP-based parallelization of the Data Compression workload.

```c
omp_set_nested(1);
const auto& numaNodes = numa::NodeList::logicalNodesWithCPUs();

#pragma omp parallel proc_bind(spread) num_threads(numaNodes.size())
{
  #pragma omp parallel proc_bind(master) num_threads(currentNode.threadCount())
  {
    size_t nodeId = numa::Node::curr().logicalId();
    size_t localThreadId = omp_get_thread_num();
    size_t chunkStart = localThreadId * chunks_per_cpu;
    size_t chunkEnd = chunkStart + chunks_per_cpu - 1;
    ...
    for(size_t chunkNum = chunkStart; chunkNum <= chunkEnd; chunkNum++) {
      // buffers are divided into local partitions per node
      const uint8_t *chunk_in = input_buffers[nodeId] + (chunkNum * CHUNK_SIZE);
      uint8_t *chunk_out = compressed_buffers[nodeId] + (chunkNum * (CHUNK_SIZE * 2));
      ...
      compress(chunk_in, CHUNK_SIZE, chunk_out, compressed_chunk_size);
    }
  }
```
Similar to the NUMA-aware OpenMP-based implementation, the PGASUS-based implementation of the Data Compression workload outlined in Listing 5.13 requires some additional effort to compute the identifiers nodeId and localThreadId each task is operating on. This additional degree of complexity is necessary in order to make sure that each thread operates on the correct sub-buffer elements of inputBuffers and compressedBuffers and the respective chunks therein. Even though these operations require additional complexity compared to a NUMA-agnostic OpenMP-based implementation, they are basic boilerplate operations that can be easily transferred to other workloads. In direct comparison with the NUMA-aware OpenMP-based implementation, PGASUS obviates the need for the tedious setup of the nested parallel sections. Considering that a manageable amount of boilerplate code is sufficient to facilitate NUMA-awareness, the slightly increased code complexity should be acceptable. Incorporating the prototypical nature of PGASUS, future iterations of the framework may further alleviate the use of the framework by providing thread and node indices through built-in helper functions.

Listing 5.13: PGASUS-based parallelization of the Data Compression workload.

```cpp
std::atomic<size_t> threadIds[numa::NodeList::logicalNodesCount()] = {}; 
numa::wait(numa::forEachThread(numa::NodeList::logicalNodesWithCPUs(), [&](){
size_t nodeId = numa::Node::curr().logicalId();
size_t localThreadId = threadIds[nodeId].fetch_add(1);
size_t chunkStart = localThreadId * chunks_per_cpu;
size_t chunkEnd = chunkStart + chunks_per_cpu - 1;
... 
for(size_t chunkNum = chunkStart; chunkNum < chunkEnd; chunkNum++) {
  // buffers are divided into local partitions per node 
  const uint8_t *chunk_in = inputBuffers[nodeId] + (chunkNum * CHUNK_SIZE);
  uint8_t *chunk_out = compressedBuffers[nodeId] + (chunkNum * CHUNK_SIZE * 2)
    ;
  ... 
  compress(chunk_in, CHUNK_SIZE, chunk_out, ...);
}
});
```

5.4.3 Database Table Scan

The PRESLEY benchmark by Felix Eberhardt and Andreas Grapentin [2] implements a Database Table Scan workload optionally using an index structure to test the impact of different types of indices on the throughput characteristics of a given workload. Currently implemented are the B-Plus Tree [1] index which is commonly found in conventional relational databases, as well as the Group Key index used in emerging in-memory database systems [47]. The original implementation of the PRESLEY benchmark utilizes pthreads for parallel execution and libnuma for data placement, whereas a new version of the benchmark was implemented in the context of a joint publication [152] to investigate the developer experience of PGASUS, using the framework for both parallel execution and
data placement. In a setup phase, PRESLEY creates a main data table of configurable size and uses either libnuma or PGASUS to place the data on a configurable primary NUMA node. The data is then shuffled using the fisher-yates algorithm \[52\] to ensure randomness of the data accesses in order to maximize the rate of cache misses. For the configurations relevant in the context of this work, the data is then indexed using one of the implemented index types. The index is either placed on the primary node or is replicated across all NUMA nodes in the system. After the setup phase, the benchmark performs lookups on the data table either by accessing the non-replicated or the replicated index in parallel on all CPUs available in the system.

The version of the benchmark implemented using pthreads and libnuma outlined in Listing 5.14 contains topology detection functionality to determine how many NUMA nodes and cores are available in the system to place the threads and data accordingly. Both the functionality of pthreads and libnuma is used to assign affinities to threads and to bind memory allocations on the desired NUMA nodes in order to implement the data placement and index replication. This manual approach proved to be more difficult to implement correctly and has several other drawbacks compared to the implementation of the benchmark based on the PGASUS framework as well.

**Listing 5.14:** Sequential generation of replicated index data across NUMA nodes based on libnuma.

```c
for (size_t i = 0; i < topology.nodes.n; ++i) {
    // wrapper to numa_membind_to_node
topology_membind_to_node(topology.nodes.nodes[i].num);

    // explicitly allocate on correct node to avoid reusing existing heap pages
    struct index_t *index =
        numa_alloc_onnode(sizeof(*index), topology.nodes.nodes[i].num);

    memset(index, 0, sizeof(*index));

    populate_data_index(index);

    data_index[topology.nodes.nodes[i].num] = index;
}

// wrapper to numa_membind_to_node
topology_release_membind();
```

In the PGASUS-based version of the benchmark demonstrated in Listing 5.15, topology information is already provided by the PGASUS framework and the manual topology detection is no longer needed. The PGASUS tasking functionality is used to execute the enclosed lambda function once on each NUMA node. PGASUS partitions the heap into separately managed spaces per NUMA node, thus operating on an allocation granularity as opposed to the page granularity implemented by libnuma. This means that using the PlaceGuard construct, the application has fine control over the allocations of third party code without modifications, while also avoiding the internal fragmentation or false sharing problems outlined in Section 5.3.1. The PlaceGuard construct ensures that all allocations in the current scope are placed on correct NUMA node. Since the PlaceGuard construct loses its effect when it goes out of scope, there is no need to manually undo the memory binding as it is the case in the pthreads and libnuma implementation. After making the
placement decision using the PlaceGuard construct, the remainder of the code uses regular calls to malloc and the new operator to setup the data-structures.

Listing 5.15: Parallel generation of replicated index data across NUMA nodes based on the PlaceGuard and tasking functionality of PGASUS.

```c
numa::wait(numa::onceForEachNode(numa::NodeList::logicalNodesWithCPUs(), [&]() {
    numa::PlaceGuard mguard(numa::Node::curr());

    struct index_t *index = new struct index_t;
    memset(index, 0, sizeof(*index));

    populate_data_index(index);
    data_index[current.logicalId()] = index;

    return 0;
}, 0));
```

5.4.4 Summary

Across the various workloads demonstrated in the preceding section, the PGASUS-based implementations were always less complex compared to the NUMA-aware implementations based on OpenMP or the pthreads library. With the programming abstractions provided by PGASUS reducing the complexity of NUMA-aware application development, the framework accomplishes its goal of unburdening developers. In terms of simplicity of code, only the NUMA-agnostic implementations can surpass the PGASUS-based implementations at the cost of completely ignoring the heterogeneity of the memory resources in NUMA systems.

5.5 Performance Evaluation

The goal of the evaluation presented in this section is to investigate the performance impact of the PGASUS framework using the workloads discussed in Section 5.4. To achieve this goal, Section 5.5.1 specifies all relevant details of the testing environment and the basic benchmark procedures to make the evaluation more repeatable. As the performance of memory allocations can have a big impact on the overall performance of workloads, a synthetic benchmark is presented in Section 5.5.2 which compares the memory allocation performance of the stacked malloc allocator provided by PGASUS to ptmalloc3, jemalloc, and TCMalloc. The performance measurements yielded for the workloads discussed in Section 5.4 are presented in Section 5.5.3. Finally, Section 5.5.5 summarizes central findings of the evaluation.

5.5.1 Testing Environment & Benchmark Procedure

All hardware configurations used for the evaluation of PGASUS are documented in Table 5.1. The set of employed machines covers the range from common two socket configurations up to high-end eight socket configurations. To analyze the behavior of different
processor designs and Instruction Set Architectures (ISAs) under different workloads, configurations using x86_64-based AMD EPYC and Intel Xeon CPUs and ppc64le-based IBM POWER8 CPUs are included.

Table 5.1: Specifications of the test systems used to evaluate the performance impact of PGASUS.

<table>
<thead>
<tr>
<th></th>
<th>Tyian</th>
<th>S824L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>Tyan TN83-B8251 [130]</td>
<td>IBM Power System S824L [81]</td>
</tr>
<tr>
<td>CPU</td>
<td>2 × AMD EPYC 7282, 2.80 GHz, 16C</td>
<td>2 × IBM POWER8 (Murano), 3.42 GHz, 10C/80T each</td>
</tr>
<tr>
<td>Memory</td>
<td>256 GB DDR4 ECC, 3200 MHz</td>
<td>1024 GB DDR3 ECC, 1600 MHz</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 18.04.5</td>
<td>Ubuntu 20.04.1</td>
</tr>
<tr>
<td>Kernel</td>
<td>5.4.0</td>
<td>5.4.0</td>
</tr>
<tr>
<td>Compiler</td>
<td>GCC 7.5.0</td>
<td>GCC 10.2.1 (AT 14.0)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>DL56o</th>
<th>E880</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>HPE ProLiant DL560 Gen10 [74]</td>
<td>IBM Power System E880 [82]</td>
</tr>
<tr>
<td>CPU</td>
<td>4 × Intel Xeon Gold 6148, 2.40 GHz, 20C each</td>
<td>8 × IBM POWER8 SCM (Turismo), 4.00 GHz, 12C each</td>
</tr>
<tr>
<td>Memory</td>
<td>1536 GB DDR4 ECC, 2666 MHz</td>
<td>6144 GB DDR3 ECC, 3200 MHz</td>
</tr>
<tr>
<td>OS</td>
<td>Ubuntu 18.04.5</td>
<td>Ubuntu 18.04.4</td>
</tr>
<tr>
<td>Kernel</td>
<td>4.15.0</td>
<td>5.3.0</td>
</tr>
<tr>
<td>Compiler</td>
<td>GCC 7.5.0</td>
<td>GCC 10.2.1 (AT 14.0)</td>
</tr>
</tbody>
</table>

All performance measurements presented hereinafter were performed after a fresh reboot in order to ensure a clean system state. Furthermore, no other active users or background tasks were running on the involved servers. In order to retrieve a sufficiently meaningful dataset, each benchmark was executed 30 times. Error bars are used in all plots to report the standard deviation for each measurement. As an additional measure, simultaneous multithreading was disabled on all systems to reduce the variance of the measurements. Furthermore, each benchmark was preceded by a warm-up run in order to eliminate any confounding factors.

5.5.2 Memory Allocation Performance

In this section, a synthetic benchmark is used to investigate the performance of the stacked malloc allocator provided by PGASUS. The performance of stacked malloc is compared to ptmalloc3, jemalloc, as well as TCMalloc. The ptmalloc3 allocator is used by PGASUS internally to serve small allocations from the arenas. Therefore, ptmalloc3 serves as measure for the overhead introduced by PGASUS itself. Finally, jemalloc and TCMalloc are included as state-of-the-art high-concurrency memory allocators that are currently used by major software companies.

Measuring the exact time spent in a single malloc or free call is generally not possible, as the measurement would introduce too much overhead compared to the duration of the
5.5 Performance Evaluation

call itself. Therefore, the duration of a large number of malloc/free cycles is measured as demonstrated in Listing 5.16 on the IBM Power System S824L [81].

Listing 5.16: Conceptual code of the malloc/free benchmark. The volatile pointer variable prevents the compiler from optimizing out the inner loop.

```c
for (size_t size = 512; size <= 1024*1024*1024; size *= 2) {
    const auto start = std::chrono::high_resolution_clock::now();
    for (size_t i = 0; i < repetitions; ++i) {
        volatile ptr = malloc();
        free(ptr);
    }
    const auto end = std::chrono::high_resolution_clock::now();
    timings[size] = duration(start, end) / repetitions;
}
```

Due to the repeated, equally sized allocations directly followed by deallocations, the benchmarking setup advantages caching allocators. Such allocators will serve most allocations with reused memory blocks, without requiring an actual system memory allocation. This setup seemed reasonable, as most real-work applications generate characteristic memory access patterns on limited ranges of block sizes [36]. Therefore, it is safe to assume that a well designed caching memory allocator will cover access patterns of realistic applications in most cases.

Furthermore, this benchmark does not access allocated memory. Thus, as far as not accessed by the allocator itself, memory pages are only virtually, but not physically allocated. This, however, is not true for PGASUS, which uses numa to bind every allocated memory page to a specific NUMA node. In Linux, memory bindings and policies are only applied once a page is physically allocated.

An overview of the timings retrieved for all allocators is presented in Figure 5.2. At the border between small and large allocations, the duration of a malloc/free cycle of ptmalloc3, and PGASUS increase by 81× and 99×, respectively. Compared to that, jemalloc
and TCMalloc have a substantially higher performance and much smaller variations across different allocation sizes. At the border between small and large allocations, the duration of a malloc/free cycle increases by 15× and 8×, respectively.

For small allocations, PGASUS relies on ptmalloc3 for serving allocations from its arenas. As the performance of PGASUS is widely similar to that of ptmalloc3 for small allocations, it appears as if the overhead introduced by the stacked malloc allocator of PGASUS is minimal. Regarding large allocations, PGASUS is the slowest allocator in comparison. However, it should be noted that PGASUS handles large allocations itself in lists of mmap-allocated chunks, so that the internally used ptmalloc3 is not involved at this point.

As mentioned before, each large allocation request in PGASUS also results in physical allocation and a call to libnuma in order to bind the allocated pages to the requested NUMA node. ptmalloc3 serves large allocations approximately 40% faster than PGASUS jemalloc and TCMalloc implement caching even for large allocations. They use a central page heap implemented based on red-black trees [43] and free lists [58], respectively.

5.5.3 Workload Benchmarks

The main goal of this section is to demonstrate that the use of PGASUS can yield performance improvements across a wide range of workloads, including a Text Histogram workload, a Data Compression workload, and a Database Table Scan use case. All workloads benchmarked throughout this section are available in the implementations discussed in Section 5.4. The throughput measurements presented in this section are reported as average values including standard deviation (n = 30). For a statistically meaningful evaluation of the collected throughput data, t-tests are performed to assess statistical significance. To further verify that changes in throughput are caused by improved data locality, the performance counters PM_DATA_FROM_LMEM (data cache loaded from local memory), PM_DATA_FROM_RMEM (data cache loaded from remote memory), and PM_DATA_FROM_DMEM (data cache loaded from distant memory) are recorded for 10 repeated executions of each workload on the IBM Power System S824L [81]. Based on this data, the ratio between remote memory access and local memory access (RMA/LMA) is computed.

5.5.3.1 Text Histogram

The Text Histogram workload employs a very fine-grained task profile, as a task computes the word frequency histograms for one of the 64192 books curated by Project Gutenberg [166] by the end of 2020. Each book is stored in a dedicated .txt file, with an average file size of 360833 bytes. The total data volume processed by this workload amounts to 21.57 GiB. All three implementations discussed in Section 5.4.1 are used for the evaluation.

As illustrated in Figure 5.3, PGASUS achieves between 1.09× and 4.7× performance improvements for the fine-grained per-file task profile compared to the OMP baseline implementation. With performance improvements between 0.2× and 5.9×, the OpenMP+libnuma implementation yields mixed results, surpassing the performance improvements of PGASUS on x86_64-based systems and providing similar or worse performance POWER8-based systems. For each hardware configuration, a t-test has confirmed statistically significant (p < 0.000001) performance impact of the respective implementations compared to the OMP baseline. RMA/LMA ratios of 0.164, 0.024, and 0.005 were
5.5 Performance Evaluation

![Performance Evaluation Graph](image)

Figure 5.3: The throughput measurements (higher is better) yielded by the Text Histogram workload exhibit increased throughput using PGASUS compared to the OpenMP baseline across all hardware configurations. However, the superior performance of the OpenMP+libnuma implementation demonstrates what speed-up might be possible if the simple tasking facilities of PGASUS handled the fine-grained task profile more efficiently.

determined for the OpenMP, the OpenMP+libnuma, and the PGASUS implementations, respectively, confirming considerably improved locality for the NUMA-aware implementations. Additional performance profiling sessions have revealed that the simplistic NUMA-aware parallel tasking component of PGASUS is overwhelmed by the very fine-grained task profile of the text histogram workload. Furthermore, the huge drop of performance of the OpenMP+libnuma implementation on the IBM Power System E880 is caused by an excessive amount of time spent in the OpenMP implementation libgomp.so. However, the reason for this behavior could not be identified.

5.5.3.2 Data Compression

Unlike the preceding workload, the Data Compression workload exhibits a much more coarse-grained task profile. Each task processes multiple megabytes of data (1 GB / number of CPU cores), performing complex operations. The measurements are performed using the large text compression benchmark \[121\] as a payload, which is comprised of the first 10\(^9\) bytes from the 2006-03-03 Wikipedia dump. All three implementations discussed in Section 5.4.2 are used for the evaluation. Separate measurements are performed for the compression operation and the decompression operation.

The benchmark results documented in Figure 5.4 demonstrate that PGASUS provides performance improvements across all hardware configurations. For the compression operation, the PGASUS-based implementation achieves performance improvements between 1.08× and 1.75×. On the side of the decompression operation, the framework yields between 1.02× and 1.54× performance improvements compared to the baseline implementation. The OpenMP+libnuma implementation yields improvements in compression throughput ranging between 0.33× and 1.18×. Surprisingly, the same implementation results in consistent slowdowns for the decompression operation, delivering throughput ranging between 0.27× and 0.935× of the OMP baseline performance. While additional profiling sessions have not identified the source of the consistent slowdown in decompression performance, the huge drop of performance of the OpenMP+libnuma implementation on the IBM Power System E880 could be traced back to an excessive amount of time spent...
Figure 5.4: The throughput measurements (higher is better) for compression and decompression are reported in panels (a) and (b), respectively. For both operations, employing PGASUS provides performance improvements across all hardware configurations, with the compression operation experiencing slightly higher speed-up factors compared to the decompression operation. The OpenMP-libnuma version also demonstrates performance improvements for the compression operation, although not as marked as the PGASUS implementation. Surprisingly, the OpenMP-libnuma implementation fails to exceed the OpenMP baseline performance for decompression.

in the OpenMP implementation libgomp.so. For each hardware configuration, a t-test has confirmed statistically significant ($p < 0.000001$) performance improvements of the PGASUS-based implementation as well as the OpenMP-libnuma implementation compared to the OMP baseline. Finally, RMA/LMA ratios of 8.476, 2.680, and 2.771 were determined for the OMP, the OpenMP-libnuma, and the PGASUS implementations, respectively, confirming considerably improved locality for the NUMA-aware implementations.

5.5.3.3 Database Table Scan

The PRESLEY benchmark [2] implements a Database Table Scan workload looking for a value in the primary key column of a database table. Scan-threads corresponding to the number of logical cores in the employed machines are used, and the table has a single column with 1,000,000,000 unique integer values.

In the experiments, the throughput of the original implementation based on pthreads and libnuma is compared to a PGASUS-based implementation. Furthermore, two different configurations are compared for each implementation, with the first configuration performing an index-based scan using a B+ tree residing on a single NUMA node and the second configuration replicating the B+ tree across each NUMA node of the employed system. The lookup of the search-values in the tree resembles a pointer-chasing-based workload. This type of workload is latency-bound since every cacheline is only touched briefly and another cacheline, possibly in a distant location is accessed next. Therefore, varying memory latencies in a NUMA system are relevant.

Using both implementations discussed in Section 5.4.3, the replicated configuration eliminates almost all remote memory access operations, yielding RMA/LMA ratios of 0.0041 and 0.0001 for the pthreads-libnuma and PGASUS implementations, respectively. In comparison, the non-replicated configuration yields RMA/LMA ratios of 12.6515 and 4.6707 for the pthreads-libnuma and PGASUS implementations, respectively. The low RMA/LMA ratios of the replicated configurations are well reflected by the throughput measurements.

Database Table Scan

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5.5 Performance Evaluation

Figure 5.5: The throughput measurements (higher is better) yielded by the PRESLEY benchmark indicate improved performance across all hardware configurations when the B+ index is replicated across all NUMA nodes. In all configurations, the PGASUS-based implementation outperforms the pthreads+libnuma baseline implementation.

presented in Figure 5.5 yielding performance improvements between $1.293 \times$ and $3.23 \times$ for the replicated configurations in comparison to the non-replicated case. These improvements come at the price of the increased memory footprint caused by the replicated index copies. However, for irregular random access patterns this leads to a significant performance improvement and could very well be viable in situation where memory is abundant. Regarding the performance impact of the respective implementations, the PGASUS-based implementation yields performance improvements ranging between $1.19 \times$ and $1.83 \times$ compared to the pthreads+libnuma implementation. For each hardware configuration, a t-test has confirmed statistically significant ($p < 0.000001$) performance improvements.

5.5.4 Energy Demand Analysis

Even though analyzing the energy-efficiency is not a central concern of this work, the impact of NUMA-aware application development on the energy demand was briefly investigated using the Data Compression workload. The energy draw measurements for a compression/decompression-cycle of the enwik9 data set were performed using all three implementations on the IBM Power System S824L test system using two Microchip MCP39F511N dual-channel power measurement devices and the PINPOINT utility. Since these measurements cover the entire execution of the test application, the compression and decompression cycle was repeated 30 times in the test application in order to reduce the impact of setup, data transfers, and teardown on the overall energy draw measurements. From these measurements, the idle power draw of the test system is deducted in order to only report the share of energy demand caused by the compression and decompression process itself. The results illustrated in Figure 5.6 demonstrate that both NUMA-aware implementations provide considerably improved energy efficiency, using less than half of the energy required by the OpenMP-based, NUMA-agnostic implementation of the Data Compression workload.
Programming Abstractions for Scale-Up Non-Uniform Memory Access Architectures

Figure 5.6: For a compression/decompression-cycle of the enwik9 test dataset, both NUMA-aware implementations consume less than half of the energy required by the NUMA-agnostic OpenMP implementation.

5.5.5 Summary

The comprehensive evaluation conducted in this section has successfully demonstrated that PGASUS offers performance improvements compared to the OpenMP or pthreads+libnuma baseline implementations across all evaluated workloads. For the fine-grained task profile of the Text Histogram workload, the simplistic tasking facilities of PGASUS are overstrained and cannot provide the same level of performance improvements compared to the OpenMP+libnuma implementation on x86_64 based systems. In additional profiling sessions, the busy waiting locks in the PGASUS task scheduler were identified as a potential bottleneck. Therefore, it should be possible to improve the performance of PGASUS for fine-grained task profiles with some additional optimization work. Both for the Data Compression workload and the Database Table Scan workload, the use of PGASUS yielded notable performance improvements compared to the OpenMP or pthreads+libnuma baseline implementations. With average performance improvements of 1.56× and peak performance improvements of up to 4.67×, the evaluation demonstrated that PGASUS does not only improve the developer experience across all workloads, but that it also capable of outperforming the baseline implementations. Even though energy measurements were only performed for the Data Compression workload, it can be assumed that NUMA-aware implementations of the remaining workloads consume less energy as well. In addition to the improved performance provided by the PGASUS framework, the reduced energy demand provides one more reason to make data-intensive applications NUMA-aware.

5.6 Summary

The PGASUS framework used in this chapter applies the concept of providing explicit means for distinguishing between different memory partitions from the PGAS model and makes it available to C++ application developers targeting shared memory systems based on the RAII idiom. Even though the PGASUS framework has been presented in the master’s theses by Wieland Hagen [68] and Karsten Tausche [198], both theses have focused their evaluation efforts on micro-benchmarking aspects of the framework itself. To fill this gap, this chapter contributed a comprehensive evaluation based on three
5.6 Summary

exemplary workloads. First, the improved developer experience offered by the framework was demonstrated by comparing PGASUS-based implementations of all three workloads to NUMA-agnostic as well as NUMA-aware implementations based on OpenMP or the pthreads library. For the evaluation of performance aspects, test systems based on both the x86_64 and ppc64le ISAs and ranging from 2 to 8 socket configurations were employed. On these systems, the evaluation demonstrated that PGASUS does not only improve the developer experience across all workloads, but that it is also capable of outperforming NUMA-agnostic implementations with average performance improvements of $1.56 \times$ and peak performance improvements of up to $4.67 \times$. 
6 Discussion and Outlook

In this chapter, the achievements of this work are summarized, and the individual contributions presented in this thesis are reviewed. The chapter reviews how the contributions presented in this thesis address the problem statement formulated in Section 1.2. This chapter also discusses the limitations of the individual approaches and outlines ideas for future research based on the contributions of this thesis.

6.1 Overview

This thesis contributes mitigations to the challenges formulated in Section 1.2 by investigating programming abstractions for on-chip accelerators, off-chip accelerators, and non-uniform memory resources. For each type of heterogeneous resource, one programming abstraction mechanism is presented and evaluated. Chapter 3 introduces the lib842 compression library. The library does not only make the resources of the NX-842 compression accelerator accessible from user space, but also introduces the first freely accessible user-space implementations of software-based compression and decompression facilities for Central Processing Units (CPUs) as well as Graphics Processing Unit (GPU)-based decompression facilities. Chapter 4 introduces the CloudCL framework with the goal of hiding many aspects of distributed computing during the development of scale-out GPU workloads. To improve the scalability of the framework, the compression facilities of the lib842 compression library are used to implement transparent compression for data transfers. Targeting Non-Uniform Memory Access (NUMA) systems, Chapter 5 builds up on top of the PGASUS framework for NUMA-aware C++ application development brought forward by Wieland Hagen and Karsten Tausche. The chapter contributes a comprehensive evaluation of the impact of the programming abstractions provided by PGASUS on both developer experience and application performance.

6.2 Contributions and Future Research

In this section, the contributions presented in this thesis are reviewed and put in perspective with related abstraction mechanisms outlined in Section 2.3. Furthermore, potential starting points for future research efforts are identified, ranging from undertakings that build up on top of the contributions presented in this work to aspects that have not been investigated in the scope this work.
Lib842 Compression Library  The lib842 compression library presented in this thesis is the first user-space approach for providing compression and decompression facilities based on the proprietary 842 compression algorithm. Relying on a modified version of the cryptodev-linux out-of-tree kernel module, the implementation details for making the high-throughput and low-latency compression and decompression facilities of NX-842 on-chip compression accelerators accessible to user-space applications through lib842 are discussed. To enable compressed data exchange across heterogeneous system resources, the hardware-accelerated approach is complemented with the introduction of highly optimized software-based compression and decompression routines for CPUs as well as Open Computing Language (OpenCL)-based decompression facilities for arbitrary GPUs. In contrast to other approaches that only employ memory compression techniques on the isolated scope of the memory resources attached to either a CPU or a GPU, the lib842 compression library lays out the groundwork for exchanging data across heterogeneous system resources in compressed form. Another distinctive feature of the employed 842 algorithm is that it provides decent compression ratios across a wide range of workloads (cf. Section 3.5.2) and does not rely on characteristics of specific use cases [86, 93].

To further improve the interoperability of the lib842 library across heterogeneous system resources, an obvious choice would be to extend the library with GPU-based compression. Furthermore, implementations for additional resource types such as Field-Programmable Gate Arrays (FPGAs) should be included in future revisions of the library. Finally, assuming future optimization efforts can manage to improve the compression and decompression throughput of accelerator-based implementations to levels comparable to the NX-842, it might even be possible to improve the efficiency of intra-node data transfers based on compression.

CloudCL Framework for Single-Paradigm Scale-Out GPU Computing  The CloudCL framework presented in this thesis joins together the dOpenCL Application Programming Interface (API) forwarding library [90] for OpenCL and the Aparapi framework [6] for executing native Java code on GPUs. By extending the underlying technologies with a job infrastructure including a job scheduler, as well as dynamic scaling capabilities for dynamically available resources, the CloudCL framework hides several aspects of the distributed memory parallel programming model during the development of scale-out GPU workloads. These abstractions enable application developers to focus on the data parallel programming model associated with GPUs yielding a single-paradigm development experience which makes scale-out GPU resources more accessible to a wider audience. With the uniform developer experience enabled based on the job infrastructure and the support for adding or removing cluster resources dynamically at runtime, the CloudCL framework provides several distinguishing features compared to the use of plain API forwarding approaches [90, 95, 7] that create the illusion of local resources.

To further foster this illusion of local resources, the dOpenCL library is augmented with transparent on-the-fly data compression for data transfers based on the lib842 compression library in order to improve the efficiency of data transfers between the master node and compute nodes. Using a highly pipelined approach to interleave all stages of the workflow for transferring transparently compressed OpenCL buffers from the master node to a compute node or vice versa, it is possible to improve the effective throughput
Contributions and Future Research

across nodes. From a workload perspective, the improved data transfer efficiency provided by the integration of transparent compression yielded performance improvements ranging between $1.11 \times$ and $2.07 \times$ across various data-intensive scale-out GPU workloads implemented using either the CloudCL framework or the OpenCL API directly.

In its current form, the manual definition of independent workload partitions can be considered as one of the biggest limitations of the CloudCL. Therefore, future revisions of the framework shall investigate semi-automatic approaches such as the concept of meta-functions employed by the DistCL library [38]. Finally, another limitation of dOpenCL and therefore also the CloudCL framework is the lack of peer-to-peer communication among compute nodes, making device-to-device data transfers prohibitively expensive. Therefore, extending dOpenCL with support for peer-to-peer communication might open up CloudCL for workloads that require inter-device communication.

PGASUS Framework for NUMA-aware data-placement in C++

To investigate the impact of using non-uniform memory resources to its fullest potential, this thesis builds up on top of the PGASUS framework which been originally presented in the master’s theses by Wieland Hagen [68] and Karsten Tausche [198]. The PGASUS framework applies the concept of providing explicit means for distinguishing between different memory partitions from the Partitioned Global Address Space (PGAS) model (cf. Section 2.3.3) and makes it available to C++ application developers targeting shared memory systems based on the Resource Acquisition is Initialization (RAII) idiom [194]. The major contribution of this thesis to PGASUS is that it provides a comprehensive evaluation of the framework based on three exemplary workloads. First, the improved developer experience offered by the framework is demonstrated by comparing PGASUS-based implementations of all three workloads to NUMA-agnostic as well as NUMA-aware implementations based on the Open Multi-Processing (OpenMP) API or the pthreads library. For the evaluation of performance aspects, test systems based on both the x86_64 and ppc64le Instruction Set Architectures (ISAs) and ranging from 2 to 8 socket configurations were employed. On these systems, the results of the evaluation suggest that PGASUS does not only improve the developer experience across all workloads, but that it is also capable of outperforming NUMA-agnostic implementations with average performance improvements of $1.56 \times$ and peak performance improvements of up to $4.67 \times$.

PGASUS provides a rewarding alternative to the approaches for enabling NUMA-aware memory placement discussed in Section 2.3.3. Compared to implementing NUMA-aware applications based on OpenMP, PGASUS provides significant improvements in terms of developer experience, enabling them to specify data placement policies with distinctly fewer lines of code. In contrast to polymorphic allocators [72], PGASUS can transparently influence memory placement of nested data structures without having to modify them in order to make use of polymorphic allocators. Finally, PGASUS eliminates issues such as unintended inter-page fragmentation and false sharing, which can easily occur when operating system facilities such as libnuma [100] or AutoNUMA [34] are not used correctly.

In its current implementation, the simplistic tasking facilities of PGASUS leave room for improvements. With further optimizations, the combination of NUMA-aware data placement and task scheduling offers a lot of potential for NUMA-aware application development. Even though PGASUS already provides certain advantages on today’s
Discussion and Outlook

systems for coarse-grained task profiles, the author speculates that abstractions for data placement such as PGASUS will become vital to deal with the increasing diversity of memory resources in upcoming state-of-the-art computer architectures, as outlined in Section 2.2.

6.3 Review of Research Question

This section reviews the research question of this thesis (cf. Section 1.3), which seeks for programming abstractions that improve the accessibility of heterogeneous system resources for application developers. For this goal, two hypotheses are constructed: First, it is assumed that a certain degree of the complexity conditioned by the large variety of heterogeneous system resources considered in the context of this thesis can be encapsulated using programming abstractions without obscuring performance-critical system properties. Second, it is presumed that programming abstractions can help to mitigate the performance penalty associated with data transfers across heterogeneous system resources.

The implementations of the contributions presented in this thesis as well as their respective evaluation results demonstrate that programming abstractions can be used to make various heterogeneous system resources more accessible. For the NX-842 on-chip compression accelerators, exposing their resources to user space through the means of a software library makes them usable for applications in the first place. To test the hypothesis for the CloudCL framework as well as the PGASUS framework, the showcase of the developer experience of both frameworks demonstrates that they manage to reduce the code complexity necessary to make use of scale-out GPU resources and NUMA systems, respectively.

Similarly, the contributions presented in this thesis show how programming abstractions can contribute to mitigating the performance penalty associated with data transfers. With the tightly integrated on-chip connectivity of the NX-842 on-chip compression accelerator, data transfers are hardly a bottleneck for this type of heterogeneous system resource. However, the efficient use of the hardware-based compression facilities lays out the groundwork for improving the efficiency of data transfers that cannot be avoided. As such, the transparent integration of on-the-fly compression for inter-node data transfers in CloudCL confirms that programming abstractions can help to improve the efficiency of data transfers across heterogeneous system resources. Even more distinctive, the evaluation of the PGASUS framework demonstrates that programming abstractions for data placement can avoid unnecessary data transfers, delivering considerable performance improvements.

In summary, the results of this thesis show that programming abstractions can indeed be used to improve the accessibility of heterogeneous system resources for application developers. However, to make efficient use of these abstractions, developers have to provide a decent understanding of the underlying hardware characteristics.
7 Conclusion

Application developers bear a certain responsibility of leveraging the heterogeneous system resources available in state-of-the-art computer architectures. The proper use of heterogeneous resources does not only facilitate sustained performance improvements over the years, but it is also vital to improve the energy-efficiency of workloads across all application domains. Unfortunately, the heterogeneity of today’s state-of-the-art computer architectures is confronting application developers with an immense degree of complexity which can be ascribed to two major challenges. First, developers need to acquire profound knowledge about the programming models or the interaction models associated with each type of heterogeneous system resource to make efficient use thereof. Second, developers must take into account that heterogeneous system resources always need to exchange data with each other in order to work on a problem together. However, this data exchange is always associated with a certain amount of overhead, which is why the amounts of data exchanged should be kept as low as possible. To respond to these challenges, application developers cannot and should not expect tools like their compilers to take over the responsibility of making efficient use of heterogeneous system resources. However, application developers should also not be overwhelmed with the immense complexities that are implied by state-of-the-art computer architectures.

Standing on the shoulders of giants, this thesis has contributed to the state of the art in heterogeneous computing by presenting programming abstractions that lessen these burdens for three types of heterogeneous system resource. The lib842 compression library provides the first method for accessing the compression and decompression facilities of the NX-842 on-chip compression accelerator available in IBM Power CPUs from user space applications running on Linux. Addressing application development of scale-out GPU workloads, the CloudCL framework makes the resources of GPU clusters more accessible by hiding many aspects of distributed computing while enabling application developers to focus on the aspects of the data parallel programming model associated with GPUs. Furthermore, CloudCL is augmented with transparent data compression facilities based on the lib842 library in order to improve the efficiency of data transfers among cluster nodes. The improved data transfer efficiency provided by the integration of transparent data compression yields performance improvements ranging between 1.11× and 2.07× across four data-intensive scale-out GPU workloads. To investigate the impact of programming abstractions for data placement in NUMA systems, a comprehensive evaluation of the PGASUS framework for NUMA-aware C++ application development is conducted. On a wide range of test systems, the evaluation demonstrates that PGASUS does not only improve the developer experience across all workloads, but that it is also capable of outperforming agnostic implementations with average performance improvements of 1.56×. For the contributed programming abstractions, this thesis has demonstrated that they can indeed improve the accessibility of heterogeneous system resources by reducing the code complexity in terms of lines of code necessary to make
use of the respective resources without obscuring performance-critical system properties. Furthermore, the presented abstractions also help developers to reduce the amount of data that has to be exchanged among heterogeneous system resources, improving both the effective throughput and the energy efficiency of data transfers.

Fueled by the competition of coherent next-generation interconnection standards, the performance of both inter-node and intra-node interconnection technologies is finally catching up in upcoming computer architectures. In the light of the diversifying memory resources enabled by these novel interconnection standards, programming abstractions for data placement probably may have the brightest perspective for gaining traction. As such, the characteristics of diversifying memory resources might be exploited with the goal of improving energy efficiency based on memory placement decisions.
Bibliography


Bibliography


Bibliography


Bibliography


Glossary

icswx Initiate Coprocessor Store Word Indexed. 4 40

AIX Advanced Interactive eXecutive. 38 40
AME Active Memory Expansion. 40
API Application Programming Interface. 5 8 14 21 26 29 32 40 53 57 59 62 65 80 82 84 85 108 109

CAIA Coherent Accelerator Interface Architecture. 12
CAPI Coherent Accelerator Processor Interface. 12
CAPP Coherent Accelerator Processor Proxy. 12
CCIX Cache Coherent Interconnect for Accelerators. 23
ccNUMA Cache Coherent Non-Uniform Memory Access. 18 20
COTS Commercial Off-the-Shelf. 35
CPU Central Processing Unit. 1 3 12 19 23 26 30 33 35 38 41 43 47 49 51 54
      55 60 63 65 69 83 88 90 96 98 107 108 111
CUDA Compute Unified Device Architecture. 22 26 28
CXL Compute Express Link. 23 24

DDR Double Data Rate. 24
DRAM Dynamic Random Access Memory. 24
DSM Distributed Shared Memory. 17 19 29
DSP Digital Signal Processor. 22 26

FPGA Field-Programmable Gate Array. 12 22 23 26 38 108

GDDR Graphics Double Data Rate. 24 26
GPU Graphics Processing Unit. 11 17 19 20 23 25 26 28 32 35 39 44 47 49
      51 56 60 63 65 67 70 72 74 76 79 80 82 107 111
HBM High-Bandwidth Memory. 1 24 26
HPC High-Performance Computing. 32 35 83
IaaS Infrastructure as a Service. 54 60
ICD Installable Client Driver. 26 56 57
IF  Infinity Fabric.  82
ILP  Instruction-Level Parallelism.  21
IMDB  In-Memory Database.  4 12 25
ISA  Instruction Set Architecture.  2 14 98 105 109
JNI  Java Native Interface.  60
MIMD  Multiple Instruction Multiple Data.  20
MPI  Message Passing Interface.  4 28 30 31 54
NUMA  Non-Uniform Memory Access.  iii 11 12 8 11 12 18 20 20 12 81 91 93 97 99 105 107 109 111
OpenCAPI  Open Coherent Accelerator Processor Interface.  23 24
OpenCL  Open Computing Language.  4 5 8 26 28 39 46 52 57 59 60 62 64 70 74 76 79 80 108 109
OpenMP  Open Multi-Processing.  5 29 30 82 90 91 93 95 97 100 105 109
PCIe  Peripheral Component Interconnect Express.  3 4 23 24
PGAS  Partitioned Global Address Space.  29 31 83 85 104 109
PowerAXON  Power with A-bus, X-bus, OpenCAPI, and NVLink.  82
PSL  POWER Service Layer.  12
RAII  Resource Acquisition is Initialization.  83 85 86 104 109
RISC  Reduced Instruction Set Computer.  21 22
SCI  Scalable Coherent Interface.  19 20
SDRAM  Synchronous Dynamic Random-Access Memory.  3
SIMD  Single Instruction Multiple Data.  1 2 20 22
SMP  Symmetric Multiprocessing.  17 20
SoC  System on a Chip.  2 34
SPMD  Single Program Multiple Data.  30
SRAM  Static Random-Access Memory.  24 27
TBB  Threading Building Blocks.  29 30
TIFF  Tag Image File Format.  79
TLP  Thread-Level Parallelism.  22
UMA  Uniform Memory Access.  1 17 30 82
UPI  Ultra Path Interconnect.  82
VAS  Virtual Accelerator Switchboard.  4 10 40
VLIW  Very Long Instruction Word.  24
Eidesstattliche Erklärung


Potsdam, den 2. August 2022,

(Max Frederik Plauth)