

Beyond the Socket: NUMA-Aware GPUs

ABSTRACT

GPUs achieve high throughput and power efficiency by employing many small single instruction multiple thread (SIMT) cores. To minimize scheduling logic and performance variance they utilize a uniform memory system and leverage strong data parallelism exposed via the programming model. With Moore’s law slowing, for GPUs to continue scaling performance (which largely depends on SIMT core count) they are likely to embrace multi-socket designs where transistors are more readily available. However when moving to such designs, maintaining the illusion of a uniform memory system is increasingly difficult. In this work we investigate multi-socket non-uniform memory access (NUMA) GPU designs and show that significant changes are needed to both the GPU interconnect and cache architectures to achieve performance scalability. We show that application phase effects can be exploited allowing GPU sockets to dynamically optimize their individual interconnect and cache policies, minimizing the impact of NUMA effects. Our NUMA-aware GPU outperforms a single GPU by $1.5\times$, $2.3\times$, and $3.2\times$ while achieving 89%, 84%, and 76% of theoretical application scalability in 2, 4, and 8 sockets designs respectively. Implementable today, NUMA-aware multi-socket GPUs may be a promising candidate for scaling GPU performance beyond a single socket.

1. INTRODUCTION

In the last decade GPUs computing has transformed the high performance computing, machine learning, and data analytics fields that were previously dominated by CPU-based installations [1, 2, 3, 4]. Many systems now rely on a combination of GPUs and CPUs to leverage high throughput data parallel GPUs with latency critical execution occurring on the CPUs. In part, GPU-accelerated computing has been successful in these domains because of native support for data parallel programming languages [5, 6] that reduce programmer burden when trying to scale programs across ever growing data sets.

Nevertheless, with GPUs nearing the reticle limitation for maximum die size and the transistor density growth rate slowing down [7], developers looking to scale the performance of their single GPU programs are in a precarious position. Multi-GPU programming models support explicit programming of two or more GPUs, but it is challenging to leveraging mechanisms such as Peer-2-Peer access [8] or a combination of MPI and CUDA [9] to manage multiple

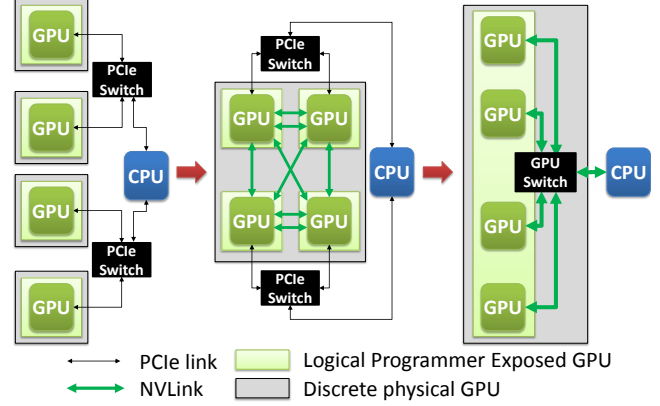


Figure 1: The evolution of GPUs from multiple discrete PCIe devices to single logical, multi-socket accelerators utilizing switched interconnects.

GPUs. These programming extensions enable programmers to employ more than one GPU for high throughput computation, but they require re-writing of traditional single GPU applications which has slowed their adoption rate.

Recently GPUs have started to expand beyond the traditional PCIe peripheral interface to enable a single protocol interconnection between both the GPUs and CPUs [10, 11, 12], as shown in Figure 1. As a result, some GPUs are being delivered on high pin-count socketed modules to provide high-speed links for interconnect bandwidth that is up to an order of magnitude higher than PCIe connections. The expansion of GPUs from single pluggable devices to closely coupled multi-socket designs is a natural progression as GPU–GPU and CPU–GPU link bandwidth becomes a performance critical system component.

The onset of multi-socket GPUs provides a pivot point for GPU and system vendors. On one hand, vendors can continue to expose multi-socket GPUs as individual GPUs and force developers of multi-GPU systems to use multiple programming paradigms to leverage multiple GPUs. On the other, vendors could expose multi-socket designs as a single non-uniform memory access (NUMA) GPU resource. By extending the single GPU programming model to multi-socket GPUs, applications can scale beyond the bounds of Moore’s law, while simultaneously retaining the programming interface that GPU developers have become accustomed.

Several groups have previously examined aggregating multiple GPUs together under a single programming model [13, 14]; however this work was done in an era where GPUs had limited memory addressability and relied on high latency, low bandwidth PCIe interconnects. As a result, prior work focused primarily on improving the multi-GPU programming experience rather than achieving highly scalable performance. Building upon this work, we propose a multi-socket *NUMA-aware* GPU architecture and runtime that aggregates multiple GPUs into a single programmer transparent logical GPU. We show that in the the era of unified virtual addressing [15], cache line addressable high bandwidth interconnects [16], and dedicated GPU and CPU socket PCB designs [11], scalable multi-GPU performance may be achievable under existing single GPU programming models. In this work, we make the following contributions:

- We show that traditional NUMA memory placement and scheduling policies are not sufficient for multi-socket GPUs to achieve performance scalability. We then demonstrate that inter-socket bandwidth will be the primary performance limiter in future NUMA GPUs.
- By exploiting program phase behavior we show that inter-socket links (and thus BW) should be dynamically and adaptively reconfigured at runtime to maximize link utilization. Moreover, we show that link policy must be determined on a per GPU basis, as global policies fail to capture per GPU phase behavior.
- We show that both the GPU L1 and L2 caches should be made NUMA-aware and dynamically adapt their caching policy to minimize NUMA effects. We demonstrate that in NUMA GPUs, extending existing GPU cache coherence protocols across multiple sockets is a good design choice, despite the overheads.
- We show that multi-socket NUMA-aware GPUs can allow traditional GPU programs to scale efficiently to as many as 8 GPU sockets, providing significant headroom before developers must re-architect applications to obtain additional performance.

2. MOTIVATION AND BACKGROUND

Over the last decade single GPU performance has scaled thanks to a significant growth in per-GPU transistor count and DRAM bandwidth. For example, in 2010 NVIDIA’s Fermi GPU integrated 1.95B transistors on a 529mm² die, with 180GB/s of DRAM bandwidth [17]. In 2016 NVIDIA’s Pascal GPU contained 12B transistors within a 610 mm² die, while relying on 720GB/s of memory bandwidth [18]. Unfortunately, transistor density is slowing significantly and many integrated circuit manufacturers are not providing roadmaps beyond 7nm. Moreover, GPU die sizes, which have been also slowly but steadily growing generationally, are expected to slow down due to limitations in lithography and manufacturing cost.

Without either larger or denser dies, GPU architects must turn to alternative solutions to significantly increase GPU performance. Recently 3D die-stacking has seen significant interest due to its successes with high bandwidth

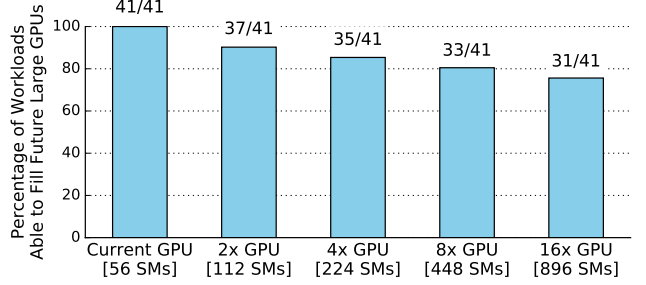


Figure 2: Percentage of workloads that are able to fill future larger GPUs (average number of concurrent thread blocks exceeds number of SMs in the system).

DRAM [19]. Unfortunately 3D die-stacking still has a significant engineering challenges related to power delivery, energy density, and cooling [20] when employed in power hungry, maximal die-sized chips such as GPUs. Thus we propose GPU manufacturers are likely to re-examine a tried and true solution from CPU world, *multi-socket GPUs*, to scaling GPU performance while maintaining the current ratio of floating point operations per second (FLOPS) and DRAM bandwidth.

Multi-socket GPUs are enabled by the evolution of GPUs from external peripherals to central computing components, considered at system design time. GPU optimized systems now employ custom PCB designs that accommodate high pin count socketed GPU modules [10] with inter-GPU interconnects resembling QPI or HyperTransport [16, 21, 22]. Despite the rapid improvement in hardware capabilities, these systems have continued to expose the GPUs provided as individually addressable units. These multi-GPU systems can provide high aggregate throughput when running multiple concurrent GPU kernels, but to accelerate a single GPU workload they require layering additional software runtimes on top of native GPU programming interfaces such as CUDA or OpenCL [6, 23]. Unfortunately, by requiring application re-design many workloads are never ported to take advantage of multiple GPUs.

Extending single GPU workload performance significantly is a laudable goal, but we must first understand if these applications will be able to leverage larger GPUs. Today NVIDIA’s largest GPUs contain 56 SMs; across a benchmark set of 41 applications (later described in Section 3.2), Figure 2 shows that most single GPU optimized workloads already contain sufficient data parallelism to fill GPUs that are 2–8× larger than today’s biggest GPUs. While these applications are unlikely to ever scale to tens of thousands of GPUs across an entire data center, we believe that programmer transparent workload scaling (up to 8×) will be attractive to many GPU developers.

In this work, we examine the performance of a future 4-module NUMA GPU to understand the effects that NUMA will have when executing applications designed for UMA GPUs. We will show (not surprisingly), that when executing UMA-optimized GPU programs on a multi-socket NUMA GPU, performance does not scale. We draw on prior work and show that before optimizing GPU microarchitecture for NUMA-awareness, several software optimizations must be in place to preserve data locality. Along these SW improve-

ments are not sufficient to achieve scalable performance however and interconnect bandwidth is a significant hindrance on performance. To overcome this bottleneck we propose two classes of improvements to reduce the observed NUMA penalty.

First, in Section 4 we examine the ability of switch connected GPUs to dynamically repartition their ingress and egress links to provide asymmetric bandwidth provisioning when required. By using existing interconnects more efficiently the effective NUMA bandwidth ratio of remote memory to local memory decreases, improving performance. Second, to minimize traffic on oversubscribed interconnect links we propose GPU caches need to become NUMA-aware in Section 5. Traditional on-chip caches are optimized to maximize overall hitrate, thus minimizing off-chip bandwidth. However, in NUMA systems, not all cache misses have the same relative cost and thus should not be treated equally. Due to the NUMA penalty of accessing remote memory, we show that performance can be maximized by preferencing cache capacity (and thus improving hitrate) towards data that resides in slower remote NUMA zones, at the expense of data that resides in faster local NUMA zones. To this end, we propose a new NUMA-aware cache architecture that dynamically balances cache capacity based on memory system utilization. Before diving into microarchitectural details and results, we first describe the locality optimized GPU SW runtime that enables our proposed NUMA-aware architecture.

3. A NUMA-AWARE GPU RUNTIME

Current GPU software and hardware is co-designed together to optimize throughput of processors based on the assumption of uniform memory properties within the GPU. Fine grained interleaving of memory addresses across memory channels on the GPU provides implicit load balancing across memory but destroys memory locality. As a result, thread block (CTA) scheduling policies need not be sophisticated to capture locality, which has been destroyed by the memory system layout. For future NUMA GPUs to work well, both system software and hardware must be changed to achieve both functionality and performance. Before focusing on architectural changes to build a NUMA-aware GPU we describe the GPU runtime system we employ to enable multi-socket GPU execution.

Prior work has demonstrated it is possible to design a framework and a runtime system that transparently decomposes GPU kernels in sub-kernels and executes them on multiple PCIe attached GPUs in parallel [14]. For example, on NVIDIA GPUs this can be implemented by intercepting and remapping each kernel call, GPU memory allocation, memory copy, and GPU-wide synchronization issued by the CUDA driver. Special care needs to ensure that per-GPU memory fences are promoted to system level and seen by all GPUs as well as guaranteeing that sub-kernels CTA identifiers are properly managed to reflect those of the original kernel. In [14] these two problems were solved by introducing code annotations and an additional source-to-source compiler which was also responsible for statically partitioning data placement and computation.

In our work, we follow a similar strategy but without using

an source-to-source translation. Unlike prior work, we are able to rely on NVIDIA’s Unified Virtual Addressing [15] to allow dynamic placement of pages into memory at runtime rather than static memory placement. Similarly, technologies with cache line granularity interconnects like NVIDIA’s NVLink [16] allow transparent access to remote memory without the need to modify application source code to access local or remote memory addresses. Due to these advancements, we assume that through dynamic compilation of PTX to SASS at executions the GPU runtime will be able to statically identify and promote system wide memory fences as well as manage sub-kernel CTA identifiers.

Current GPUs perform fine grained memory interleaving at a sub-page granularity across memory channels. In a NUMA GPU this policy would destroy locality and result in 75% of all accesses to be to remote memory in a 4 GPU system, an undesirable effect in NUMA systems. Similarly, a round-robin page level interleaving could be utilized, similar to the Linux INTERLEAVE page allocation strategy, but despite the inherent memory load balancing, this still results in 75% of memory accesses occurring over low bandwidth NUMA links. Instead we leverage UVM page migration functionality to migrate pages on-demand from system memory to local GPU memory as soon as the first access (also called first-touch allocation) is performed as described by Arunkumar et. al [24].

On a single GPU fine grain dynamic assignment of CTAs to SMs is performed to achieve good load balancing. Extending this policy to a multi-socket GPU system is not possible due to the relatively high latency of passing sub-kernel launches from software to hardware. To overcome this penalty the GPU runtime must launch a block of CTAs to each GPU-socket at coarse granularity. To encourage load balancing, each sub-kernel could be comprised of an interleaving of CTAs using modulo arithmetic. Alternatively a single kernel can be decomposed into N sub-kernels, where N is the total number of GPU sockets in the system, assigning equal amount of contiguous CTAs to each GPU. This design choice potentially exposes workload unbalance across sub-kernels, but it has been shown to preserve data locality present in applications where contiguous CTAs also access contiguous memory regions [14, 24].

3.1 Performance Through Locality

Figure 3 shows the relative performance of a 4-socket NUMA GPU with respect to a single GPU under the two possible CTA scheduling and memory placement strategies explained above. The green bars show the relative performance of traditional single GPU scheduling and memory interleaving policies when adapted to a NUMA GPU. The light blue bars show the relative performance of using locality optimized GPU scheduling and memory placement, consisting of contiguous block CTA scheduling and first touch page migration. We can clearly see that the *Locality-Optimized* solution almost always outperforms the traditional GPU scheduling and memory interleaving. Without these runtime locality optimizations, in average a 4-socket NUMA GPU is not able to even match the performance of a single GPU despite the large increase in hardware resources. Thus, using variants of prior proposals [14, 24], we now

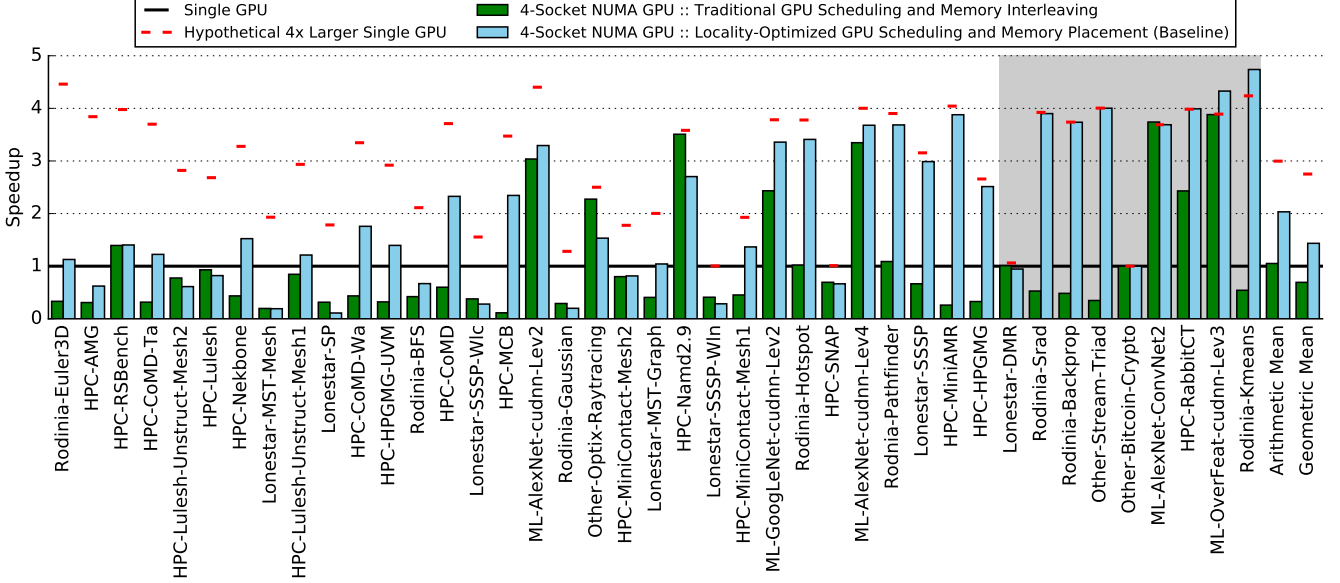


Figure 3: Performance of a 4-socket NUMA GPU relative to a single GPU and a hypothetical $4\times$ larger (all resources scaled) single GPU. Applications shown in grey achieve greater than 99% of performance scaling with SW-only locality optimization.

only consider this locality optimized GPU runtime for the remainder of the paper.

Despite the performance improvements that can come via locality optimized software runtimes, many applications are not scaling well on our proposed NUMA GPU system. To illustrate this, Figure 3 shows the speedup achievable by a hypothetical (unbuildable) $4\times$ larger GPU with a red dash. This red dash represent an approximation of the maximum theoretical performance we could expect from a perfectly architected (both HW and SW) NUMA GPU system. Figure 3 sorts the applications by the gap between relative performance of the Locality-Optimized NUMA GPU and hypothetical $4\times$ larger GPU. We observe that on the right side of the graph some workloads (shown in the grey box) can achieve or surpass the maximum theoretical performance. In particular for the two far-most benchmarks on the right, the locality optimized solutions can outperform the hypothetical $4\times$ larger GPU due higher cache hit rates because contiguous block scheduling is more cache friendly than traditional GPU scheduling.

However, for the applications on the left side there is a large gap between the Locality-Optimized NUMA design and theoretical performance. These are workloads in which either locality does not exist or the Locality-Optimized GPU runtime is not effective, resulting in large amount of remote data accesses still occurring. Because our goal is to provide scalable performance for single GPU optimized applications, in the rest of the paper we aim to close this performance gap through microarchitectural innovation. To simplify later discussion, we choose to exclude benchmarks that achieve $\geq 99\%$ of the theoretical performance with SW-only locality optimizations. However, we include all benchmarks in our final results to show the overall performance scalability achievable with NUMA-aware multi-socket GPUs.

3.2 Simulation Methodology

To evaluate the performance of future NUMA-aware multi-socket GPUs we use a proprietary, cycle-level, trace-driven simulator for single and multi-GPU systems. Our baseline GPU in both single GPU and multi-socket GPU configurations, approximates the latest NVIDIA Pascal architecture [25]. Each streaming multiprocessors (SM) is modeled as an in-order processor with multiple levels of cache hierarchy containing private, per-SM, L1 caches and multi-banked, shared, L2 cache. Each GPU is backed by local on-package high bandwidth memory [19]. Our multi-socket GPU systems contains two to eight of these GPUs interconnected through a full bandwidth GPU switch as shown on Figure 1. Table 1 provides a more detailed overview of the simulation parameters.

We study the scalability of multi-socket NUMA GPUs using 41 workloads taken from a range of production codes based on the HPC CORAL benchmarks [26], graph applications from Lonestar [27], HPC applications from Rodinia [28], in addition to several other in-house CUDA benchmarks. This set of workloads covers a wide spectrum of GPU applications used in machine learning, fluid dynamic, image manipulation, graph traversal, and scientific computing. Each of our benchmarks is hand selected to identify the region of interest deemed representative for each workload, which may be as small as a single kernel containing a tight inner loop or several thousand kernel invocations. We run each benchmark to completion for the determine region of interest. Table 2 provides both the memory footprint of these workloads as well as the average number of active CTAs in the workload (weighted by the time spent on each kernel) to provide a representation of how many parallel thread blocks (CTAs) are generally available during workload execution.

Parameter	Value(s)
Num of GPU sockets	4
Total number of SMs	64 per GPU socket
GPU Frequency	1GHz
Max number of Warps	64 per SM
Warp Scheduler	Greedy then Round Robin
L1 Cache	Private, 128 KB per SM, 128B lines, 4-way, GPU-side SW-based coherency
L2 Cache	Shared, 4MB per socket, 128B lines, 16-way, Memory-side non-coherent
GPU-GPU Interconnect	128GB/s per socket (64GB/s each direction) 8 lanes 8B wide each per direction 128-cycle latency
DRAM Bandwidth	768GB/s per GPU socket
DRAM Latency	100ns

Table 1: Simulation parameters for evaluation of single and multi-socket GPU systems.

4. ASYMMETRIC INTERCONNECTS

Figure 4(a) shows a switch connected GPU with symmetric and static link bandwidth assignment. Each link is comprised of equal number of uni-directional high-speed lanes in both directions, collectively comprising a symmetric bi-directional link. Traditional static design time link capacity assignment is very common and has several advantages. For example, only one type of I/O circuitry (egress drivers or ingress receivers) along with only one type of control logic need to be implemented at each on-chip link interface. Moreover, the multi-socket switches result in simpler designs that can easily support a statically provisioned bandwidth requirements. On the other hand, multi-socket link bandwidth utilization can have a large impact on overall system performance. Static partitioning of bandwidth, when application needs are dynamic, can leave performance on the table. Because I/O bandwidth is a limited and expensive system resource, NUMA-aware interconnects designs must look for innovations that can keep wire and I/O utilization high.

In multi-socket NUMA GPU systems, we observe that many applications have different utilization of egress and ingress channels on both a per GPU-socket basis and during different phases of execution. For example, Figure 5 shows a link utilization snapshot over time for HPC-HPGMG-UVM running on a SW locality optimized 4-socket NUMA GPU. Vertical dotted black lines represent kernel invocations that are split across the 4 GPU-sockets. We can see that several small kernels have negligible interconnect utilization. However, for the later larger kernels, GPU0 and GPU2 fully saturate their ingress links, while GPU1 and GPU3 fully saturate their egress links. At the same time GPU0 and GPU2, and GPU1 and GPU3 are underutilizing their egress and ingress links respectively.

In many workloads we observe one common scenario, in which all CTAs writing to the same memory range at the end of a kernel (i.e. parallel reductions, data gathering). For CTAs running on one of the sockets, GPU0 for example, these memory references are local and do not produce any traffic on the inter-socket interconnections. However

Benchmark	Time-weighted Average CTAs	Memory Footprint (MB)
ML-GoogLeNet-cudnn-Lev2	6272	1205
ML-AlexNet-cudnn-Lev2	1250	832
ML-OverFeat-cudann-Lev3	1800	388
ML-AlexNet-cudnn-Lev4	1014	32
ML-AlexNet-ConvNet2	6075	97
Rodinia-Backprop	4096	160
Rodinia-Euler3D	1008	25
Rodinia-BFS	1954	38
Rodinia-Gaussian	2599	78
Rodinia-Hotspot	7396	64
Rodinia-Kmeans	3249	221
Rodinia-Pathfinder	4630	1570
Rodinia-Srad	16384	98
HPC-SNAP	200	744
HPC-Nekbone-Large	5583	294
HPC-MiniAMR	76033	2752
HPC-MiniContact-Mesh1	250	21
HPC-MiniContact-Mesh2	15423	257
HPC-Lulesh-Unstruct-Mesh1	435	19
HPC-Lulesh-Unstruct-Mesh2	4940	208
HPC-AMG	241549	3744
HPC-RSBench	7813	19
HPC-MCB	5001	162
HPC-NAMD2.9	3888	88
HPC-RabbitCT	131072	524
HPC-Lulesh	12202	578
HPC-CoMD	3588	319
HPC-CoMD-Wa	13691	393
HPC-CoMD-Ta	5724	394
HPC-HPGMG-UVM	10436	1975
HPC-HPGMG	10506	1571
Lonestar-SP	75	8
Lonestar-MST-Graph	770	86
Lonestar-MST-Mesh	895	75
Lonestar-SSSP-WIn	60	21
Lonestar-DMR	82	248
Lonestar-SSSP-Wlc	163	21
Lonestar-SSSP	1046	38
Other-Stream-Triad	699051	3146
Other-Optix-Raytracing	3072	87
Other-Bitcoin-Crypto	60	5898

Table 2: Application footprint and time weighted average number of thread blocks available during execution.

CTAs dispatched to other GPUs must issue remote memory writes, saturating their egress links while ingress links remain underutilized, but causing ingress traffic on GPU0. Such communication patterns typically utilize only 50% of available interconnect bandwidth. In these cases, dynamically increasing the number of ingress lanes for GPU0 (by turning around direction of egress lanes) and switching the direction of ingress lanes for GPUs 1–3, can substantially improve the achievable interconnect bandwidth. Motivated by these findings, we propose to dynamically control multi-socket link bandwidth assignments on a per-GPU basis resulting in dynamic asymmetric link capacity assignments as shown in Figure 4(b).

To evaluate this proposal we model point-to-point links containing multiple lanes, similarly to NVLink [25]. In these links, 8 lanes with 8GB/s capacity per lane yield an aggregate bandwidth of 64GB/s in each direction. We propose replacing uni-directional lanes with bi-directional lanes to which we apply an adaptive link bandwidth allocation mechanism that works as following. For each link in the system, at kernel launch the links are always reconfigured to contain

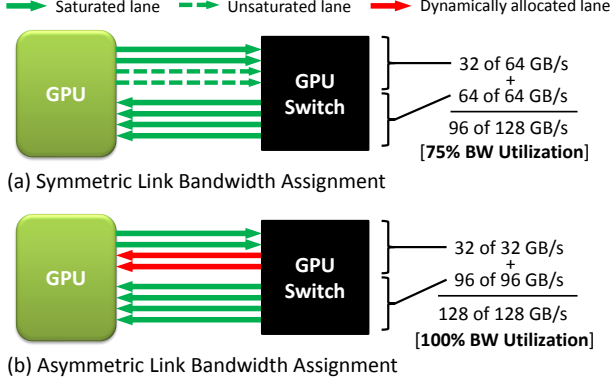


Figure 4: Example of dynamic link assignment to improve interconnect efficiency.

symmetric link bandwidth with 4 lanes per direction. During kernel execution the link load balancer periodically samples the saturation status of each link. If the lanes in one direction are not saturated, while the lanes in the opposite direction are 99% saturated, the link load balancer reconfigures and reverses the direction of one of the unsaturated lanes after quiescing all packets on that lane.

This sample and reconfigure process stops only when directional utilization is not oversubscribed or all but one lane is configured in a single direction. If both ingress and egress links are found to be saturated and in an asymmetric configuration, links are then reconfigured back towards a symmetric configuration to encourage global bandwidth equalization. While this process may sound complex, the circuitry for dynamically turning high speed single ended links around in a short number of cycles already is in use by modern high bandwidth memory interfaces such as GDDR; where the same set of wires is used for both memory reads and writes [29].

4.1 Results and Discussion

There are two important parameters that will affect the performance of our proposed mechanism (i) *SampleTime*: The frequency at which the scheme samples for a possible reconfiguration and (ii) *SwitchTime*: The cost of turning the direction of an individual lane. Figure 6 shows the performance improvement, with respect to our SW locality optimized GPU by exploring different values of the *SampleTime* indicated by green bars and assuming a *SwitchTime* of 100 cycles. The red bars in Figure 6 provide an upper-bound of performance speedups when doubling the available interconnect bandwidth to 256GB/s. For workloads on the right of the figure, doubling the link bandwidth has little effect, thus dynamic link policy will also show little improvement due to low GPU-GPU interconnect bandwidth needs. On the left side, we can see that for some applications, where improved interconnect bandwidth has a large effect, dynamic lane switching can improve application performance by as much as 80%. For some benchmarks like Rodinia-Euler-3D, HPC-AMG, and HPC-Lulesh, doubling the link bandwidth provides $2\times$ speedup, while our proposed dynamic link assignment mechanism is not able to significantly improve performance. Those are the workloads that

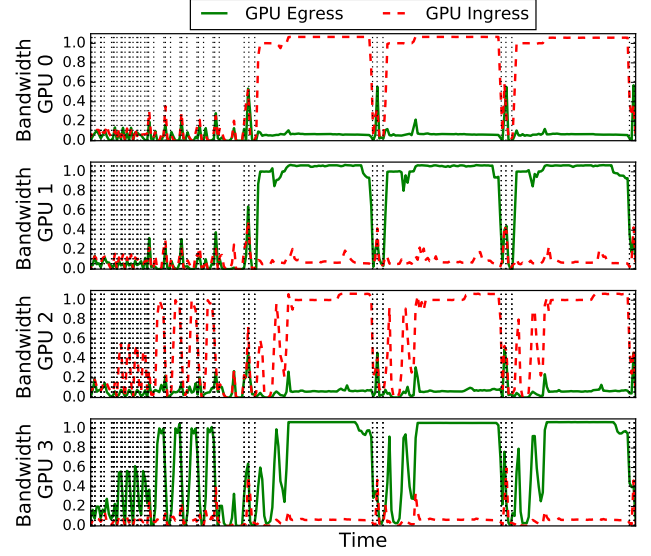


Figure 5: Normalized link bandwidth profile for HPC-HPGMG-UVM showing asymmetric link utilization between GPUs and within a GPU. Vertical black dotted lines indicate kernel launch events.

saturate both link directions, so there is no opportunity to provide additional bandwidth by turning links around.

Using a moderate 5K cycle sample time, dynamic link policy can improve performance by 14% on average over static bandwidth partitioning. If the link load balancer samples too infrequently application dynamics can be missed and performance improvement is reduced. However if the link is turned around too frequently, bandwidth is lost due to the overhead of turning the link. While we have assumed a pessimistic link turn time of 100 cycles, we performed sensitivity studies that show even if link turn time were increased to 500 cycles, our dynamic policy loses less than 2% of performance. At the same time, using a faster lane switch (10 cycles) does not significantly improve the performance over a 100 cycle link turn time. We note that the link turnaround times of modern high-speed on-board links such as GDDR5 [29] are about 8ns including both link and internal DRAM turn-around latency (which is less than 10 cycles at 1GHz).

Our results demonstrate that asymmetric link bandwidth allocation can be very attractive when inter-socket interconnect bandwidth is constrained by the number of on-PCB wires (and thus total link bandwidth). The primary drawback of this solution is that both types of interface circuitry (TX and RX) and logic need to be implemented for each lane in both the GPU and switch interfaces. We conducted an analysis of the potential cost of doubling the amount of I/O circuitry and logic based on a proprietary state of the art GPU I/O implementation. Our results show that doubling this interface area increases total GPU area by less than 1% while yielding a 12% improvement in average interconnect bandwidth which results in a 14% application performance improvement. One additional caveat worth noting is that the proposed asymmetric link mechanism optimizes link bandwidth in a given direction for each individual link, while the total switch bandwidth remains constant.

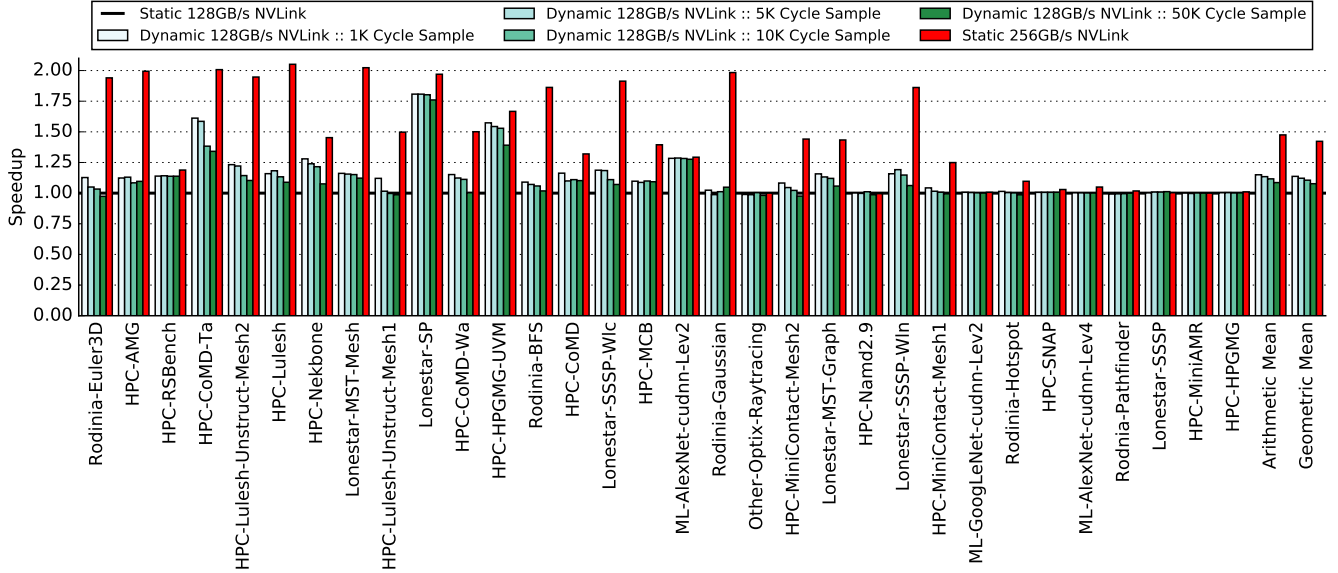


Figure 6: Relative speedup of the dynamic link adaptivity with respect to the baseline architecture by varying sample time and assuming switch time of 100 cycles. In red, speedup achievable by doubling link bandwidth.

5. NUMA-AWARE CACHE MANAGEMENT

In Section 4 we have shown that inter-socket bandwidth is an important factor in achieving scalable NUMA GPU performance. Unfortunately, because either the outgoing or incoming links must be underutilized for us to reallocate that bandwidth to the saturated link, if both incoming and outgoing links are saturated, dynamic link rebalancing yields minimal gains. To improve performance in situations where dynamic link balancing is ineffective, system designers can either increase link bandwidth, which is very expensive, or try and decrease the amount of traffic that crosses the low bandwidth communication channels. To decrease off-chip memory traffic, architects typically turn to caches to capture locality.

GPU cache hierarchies differ from traditional CPU hierarchies where in they are not supported by strong hardware coherence protocols [30]. They also differ from CPU protocols in that caches may be both processor side (where some form of coherence is typically necessary) or they may be memory side (where coherence is not necessary). As described in Table 1 and Figure 7(a), a GPU today is typically composed of relatively large SW managed coherent L1 caches located close to the SMs, while a relatively small, distributed, non-coherent memory side L2 cache resides close to the memory controllers. This organization works well for GPUs because their SIMT processor designs often allow for significant coalescing of requests to the same cache line, so having large L1 caches reduces the need for global crossbar bandwidth. By then placing the L2 caches memory-side they do not need to participate in the coherence protocol, reducing complexity.

5.1 Design Considerations

In NUMA designs remote memory references occurring across low bandwidth NUMA interconnections results in poor performance, as shown in Figure 3. Similarly, in NUMA GPUs utilizing traditional memory side L2 caches (that depend on fine grained memory interleaving for load

balancing) is a bad decision. Because memory side caches only able to cache accesses that originate in their local memory-side, they cannot cache memory from other NUMA zones and thus can not reduce NUMA interconnect traffic. Previous work has proposed that GPU L2 cache capacity should be split between memory-side caches and a new processor-side L1.5 cache that is an extension of the GPU L1 caches [24] to enable caching of remote data, shown in Figure 7(b). By balancing L2 capacity between memory side and remote caches (R\$), this design limits the need for extending expensive coherence operations (invalidations) into the entire L2 cache while still minimizing crossbar or interconnect bandwidth.

Flexibility: Designs that statically allocate cache capacity to local memory and remote memory, in any balance, may achieve reasonable performance in specific instances but they lack flexibility. Much like application phasing was shown to affect NUMA bandwidth consumption the ability to dynamically share cache capacity between local and remote memory has the potential to improve performance under several situations. First, when application phasing results in some GPU-sockets primarily accessing data locally while others are accessing data remotely, a fix partitioning of cache capacity is guaranteed to be sub-optimal. Second, while we show that most applications will be able to completely fill large NUMA GPUs, this may not always be the case. GPUs within the data center are being virtualized and there is on-going work to support concurrent execution of multiple kernels within a single GPU [31, 32]. If a large NUMA GPU is sub-partitioned, it is intuitive that system software attempt to partition it along the NUMA boundaries (even within a single GPU-socket) to improve the locality of small GPU kernels. To effectively capture locality in these situation, NUMA-aware GPUs need to be able to dynamically re-purpose cache capacity at runtime, rather than be statically partitioned at design time.

Coherence: To-date, single socket GPUs have not moved

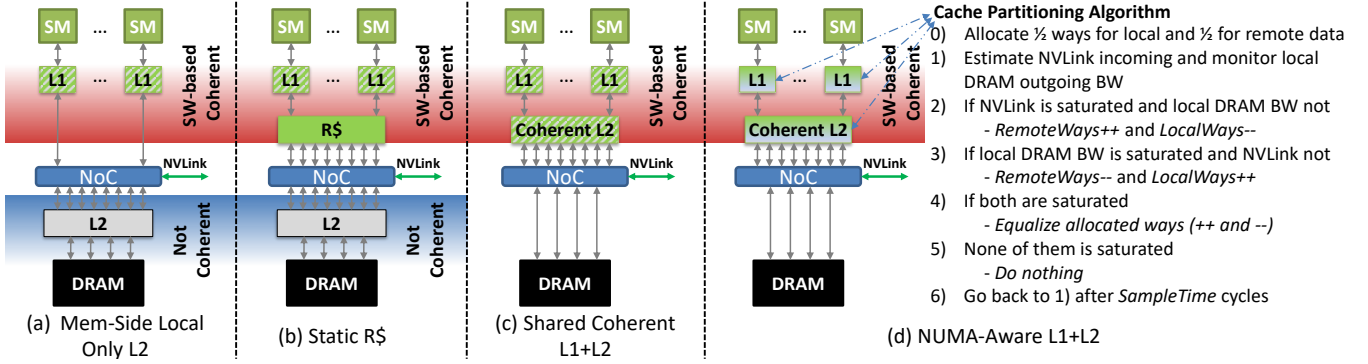


Figure 7: Potential L2 cache organizations to balance capacity between remote and local NUMA memory systems.

their memory-side caches to processor side because the overhead of cache invalidation (due to coherence) is an unnecessary performance penalty. Within a single socket GPU with a uniform memory system, there is little performance advantage to implementing L2 caches as processor side caches. However in a multi-socket NUMA design, the performance tax of extending coherence into L2 caches is offset by the fact that remote memory accesses can now be cached locally and may be justified; Figure 7(c) shows a configuration with a coherent L2 cache where remote and local data contend for L2 capacity as extensions of the L1 caches, implementing identical coherence policy.

Dynamic Partitioning: Building upon coherent GPU L2 caches, we posit that while conceptually simple, allowing both remote and local memory accesses to contend for cache capacity (in both the L1 and L2 caches) in a NUMA system is flawed. In UMA systems it is well known that performance is maximized by optimizing for cache hit rate, thus minimizing off-chip memory system bandwidth. However in NUMA systems, *not all cache misses have the same relative cost performance impact*. A cache miss to a local memory address has a smaller cost (in both terms of latency and bandwidth) than a cache miss to a remote memory address. Thus, it should be beneficial to dynamically skew cache allocation to preference caching remote memory over local data when it is determined the system is bottle-necked on NUMA bandwidth.

To minimize inter-GPU bandwidth in multi-socket GPU systems we propose a NUMA-aware cache partitioning algorithm, with cache organization and brief summary shown in Figure 7(d). Similar to our interconnect balancing algorithm, at initial kernel launch (after GPU caches have been flushed for coherence purposes) we allocate one half of the cache ways for local memory and the remaining ways for remote data (Step ①). After executing for a 5K cycles period, we sample the average bandwidth utilization on local memory and estimate the GPU-socket’s incoming read request rate by looking at the outgoing request rate multiplied by the response packet size. By using the outgoing request rate to estimate the incoming bandwidth, we avoid situations where incoming writes may saturate our link bandwidth falsely indicating we should preference remote data caching. Projected link utilization above 99% is considered to be bandwidth saturated (Step ①). In cases where the interconnect bandwidth is saturated but local memory bandwidth is not,

the partitioning algorithm attempts to reduce remote memory traffic by re-assigning one way from the group of local ways to the remote ways grouping (Step ②). Similarly, if the local memory BW is saturated and NVLink is not, the policy re-allocates one way from the remote group, and allocates it to the group of local ways (Step ③). To minimize the impact on cache design, all ways are consulted on look up, allowing lazy eviction of data when the way partitioning changes. In case where both the interconnect and local memory bandwidth are saturated, our policy gradually equalizes the number of ways assigned for remote and local cache lines (Step ④). Finally, if neither of the links are currently saturated, the policy takes no action (Step ⑤). To prevent cache starvation of either local or remote memory (which causes memory latency dramatically increase and a subsequent drop in performance), we always require at least one way in all caches to be allocated to either remote or local memory.

5.2 Results

Figure 8 compares the performance of 4 different cache configurations in our 4-socket NUMA GPU. Our baseline is a traditional GPU with memory side local-only L2 caches. To compare against prior work [24] we provide a 50–50 static partitioning where the L2 cache budget is split between the GPU-side coherent remote cache which contains only remote data, and the memory side L2 which contains only local data. In our 4-socket NUMA GPU static partitioning improves performance by 54% on average, although for some benchmarks, it hurts the performance by as much as 10% for workloads that have negligible inter-socket memory traffic. We also show the results for GPU-side coherent L1 and L2 caches where both local and remote data contend capacity. On average, this solution outperforms static cache partitioning significantly despite incurring additional flushing overhead due to cache coherence.

Finally, our proposed NUMA-aware cache partitioning policy is shown in dark grey. Due to its ability to dynamically adapt the capacity of both L2 and L1 to optimize performance when backed by NUMA memory, it is the highest performing cache configuration. By examining simulation results we find that for workloads on the left side of Figure 8 which fully saturate the NVLink bandwidth, NUMA-aware dynamic policy configures the L1 and L2 caches to be primarily used as remote caches. However, workloads on

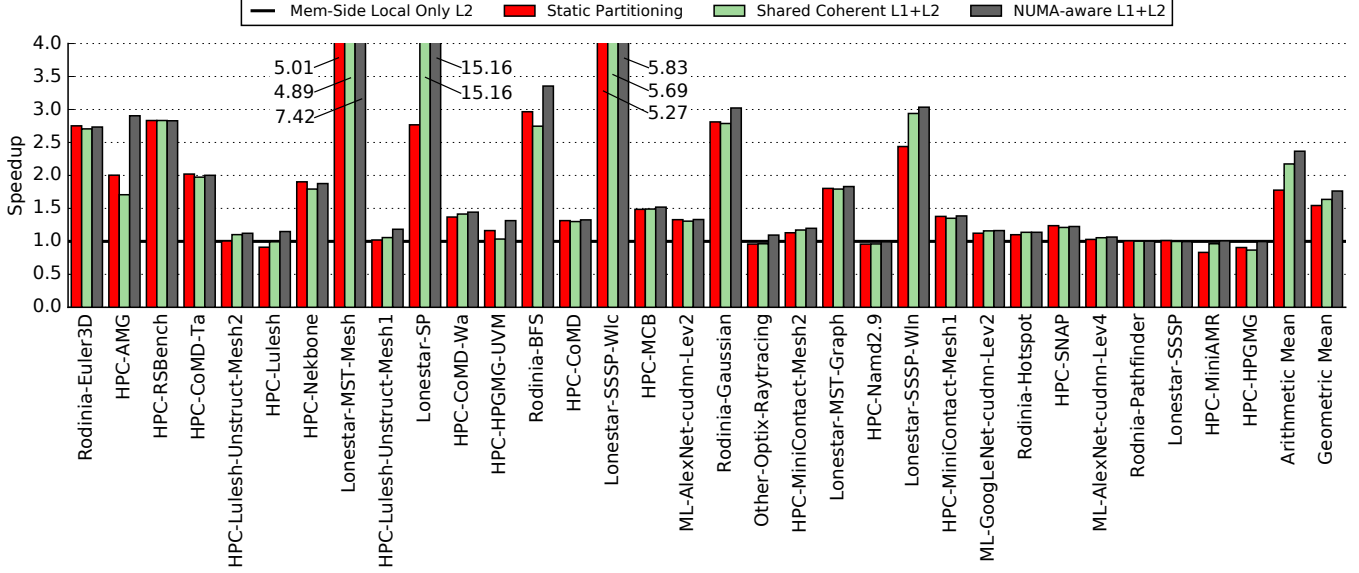


Figure 8: Performance of NUMA-aware dynamic cache partitioning in a 4-socket GPU compared to memory-side L2 and previously proposed static partitioning.

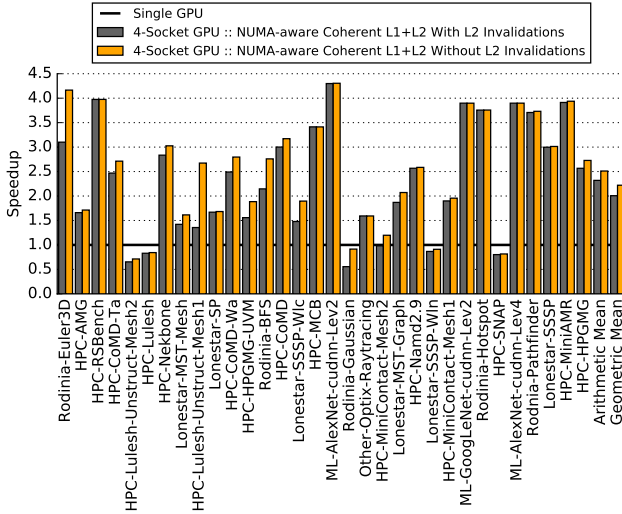


Figure 9: Performance overhead of extending current GPU software based coherence into the GPU L2 caches.

the right side of the figure tend to have good GPU-socket memory locality, and thus prefer L1 and L2 caches store primarily local data. NUMA-aware cache partitioning is able to flexibly adapt to varying memory access profiles and can improve average NUMA GPU performance 76% compared to traditional memory side L2 caches, and 22% compared to previously proposed static cache partitioning despite incurring additional coherence overhead.

When extending the software controlled GPU coherence protocol into the GPU L2 caches, L1 coherence operations (flushes) must also be extended into the GPU L2 caches. To further understand the impact these coherence operations have on our NUMA-aware cache performance we evaluated a hypothetical L2 cache which need not perform these operations. Figure 9 shows the impact that coherence operations have on application performance in our 4-socket NUMA

GPU. While significant for some applications, on average SW based GPU coherence overheads are only 10% even when extended into all GPU-socket L2 caches; we conclude that despite the coherence overheads the benefit of NUMA-aware coherent L2 caches on multi-socket GPUs is a worthy trade-off.

6. DISCUSSION

Combined Improvement: Sections 4 and 5 provide two techniques aimed at more efficiently utilizing scarce NUMA bandwidth within future NUMA GPU systems. The proposed methods for dynamic interconnect balancing and NUMA-aware caching are orthogonal and can be applied in isolation or combination. Dynamic interconnect balancing has an implementation simplicity advantage in that the system level changes to enable this feature are isolated from the larger GPU design. Conversely, enabling NUMA-aware GPU caching based on interconnect utilization requires changes to both the physical cache architecture and the GPU coherence protocol.

Because these two features target the same problem, when employed together their effects are not strictly additive. Figure 10 shows the overall improvement NUMA-aware GPUs can achieve when applying both techniques in parallel. For benchmarks such as CoMD, these features contribute nearly equally to the overall improvement, but for others such as ML-AlexNet-cudnn-Lev2 or HPC-MST-Mesh1, interconnect improvements or caching are the primary contributor respectively. On average, we observe that when combined we see $2.1\times$ improvement over a single GPU and 80% over the baseline software locality optimized 4-socket NUMA GPU using memory side L2 caches; best performance is clearly obtained when applying both features in unison.

Scalability: Ultimately, for vendors to produce multi-socket NUMA GPUs they must achieve high enough parallel efficiency to justify their design. To understand the scalability of our approach Figure 11 shows the perfor-

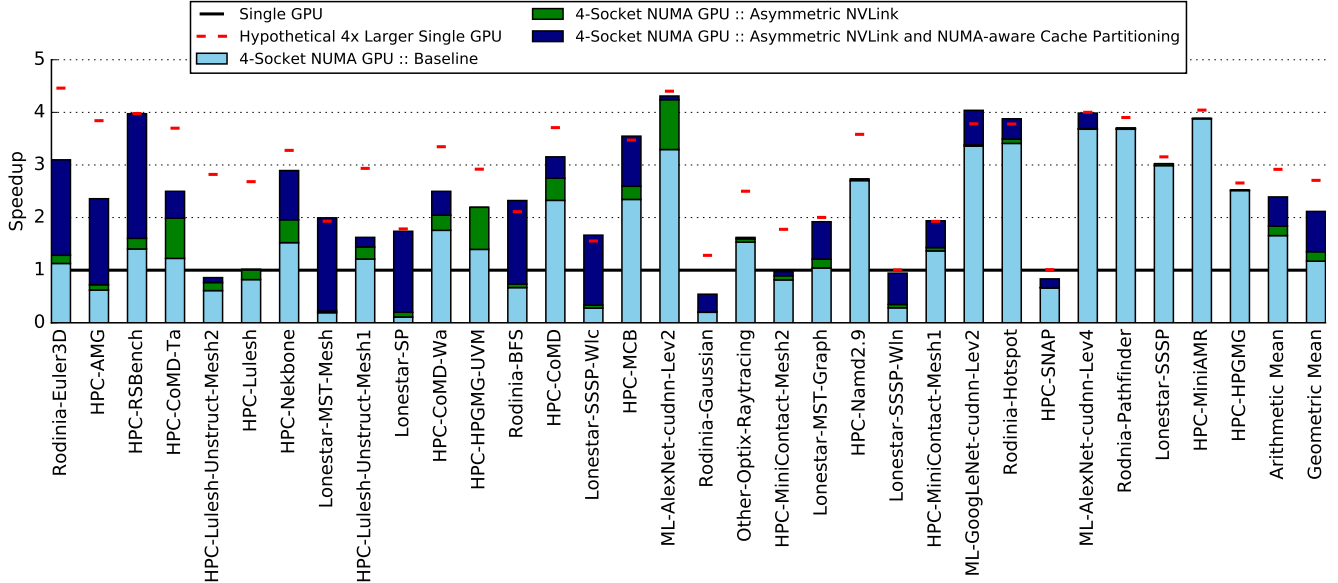


Figure 10: Final NUMA-aware GPU performance compared to a single GPU and $4\times$ larger single GPU with scaled resources.

mance of a NUMA-aware multi-socket GPU compared to a single GPU, when scaled across 2, 4, and 8 sockets respectively. On average a 2 socket NUMA GPU achieves $1.5\times$ speedup, while 4 sockets and 8 sockets achieve $2.3\times$ and $3.2\times$ speedups respectively. Depending on perspective these speedups may look attractive or lackluster; particularly when per-benchmark variance is included. However, the scalability of NUMA GPUs is not solely dependent on just NUMA GPU microarchitecture. We observe that for some applications, even if the application was run on larger hypothetical single GPUs, performance would scale similarly. This may be due to a variety of reasons beyond NUMA effects, including number of CTAs available, frequency of global synchronization, and other factors. Comparing our NUMA-aware GPU implementation to the scaling that applications could achieve on a hypothetical large single GPU, we see that NUMA-GPUs can achieve 89%, 84%, and 76% the efficiency of a hypothetical single large GPU in 2, 4, and 8 socket configurations respectively. This high efficiency factor indicates that our design is able to largely eliminate the NUMA penalty in future multi-socket GPU designs.

Multi-Tenancy on Large GPUs: In this work we have shown that many workloads today have the ability to saturate (with sufficient parallel work) a GPU that is at least $8\times$ larger than today’s GPUs. With deep-data becoming commonplace across many computing paradigms, we believe that the trend of having enough parallel thread blocks to saturate large single GPUs will continue into the foreseeable future. However when GPUs become larger at the expense of having multiple addressable GPUs within the system, questions related to GPU provisioning arise. Applications that cannot saturate large GPUs will leave resources underutilized and concurrently will have to multiplex across the GPU cooperatively in time, both undesirable outcomes.

While not the focus of this work, there is significant effort in both industry and academia to support finer grain sharing of GPUs through either shared SM execution [33], spatial

multiplexing of a GPU [31], or through improved time division multiplexing with GPU pre-emptability [32]. To support large GPU utilization any of these solutions could be applied to a multi-socket GPU in the cases where applications may not completely fill a larger GPU. Alternatively, with additional GPU runtime work multi-socket GPU designs could also be dynamically partitioned with a granularity of 1–N logical GPUs being exposed to the programmer, providing yet another level of flexibility to improve utilization.

Power Implications: As discussed earlier, arbitrarily large monolithic single GPUs are unfeasible, so multi-GPU systems connected with onboard high-speed links and switches are becoming an attractive solution for continuing GPU performance scaling. However, these onboard high-speed links and switches require additional power. We estimated the link overhead by assuming $10pJ/b$ of on board interconnect energy for combined links and switch (extrapolated from publicly available information for cabinet level Mellanox switches and links [34, 35]). Using this estimate we calculate an average (Geo-Mean) 30W of communication power for the baseline architecture composed of 4 GPUs, and 14W after our NUMA-aware optimizations are applied. Some applications such as Rodinia-Euler3D, HPC-Lulesh, HPC-AMG, HPC-Lulesh-Unstruct-Mesh2 are communication intensive, resulting in $\approx 130W$ of power consumption after our optimizations are considered. Assuming a typical TDP of 250W per GPU module, in a 4 GPUs system, the extra power due to the communication represents a 5% overhead across the full range of 41 evaluated benchmarks. While this power tax is not trivial, without alternative methods for building scalable large GPUs, interconnect power will likely become a large portion of the overall GPU power budget.

7. RELATED WORK

Multi-GPU programming is commonly used for scaling GPU performance via integration of multiple GPUs at the

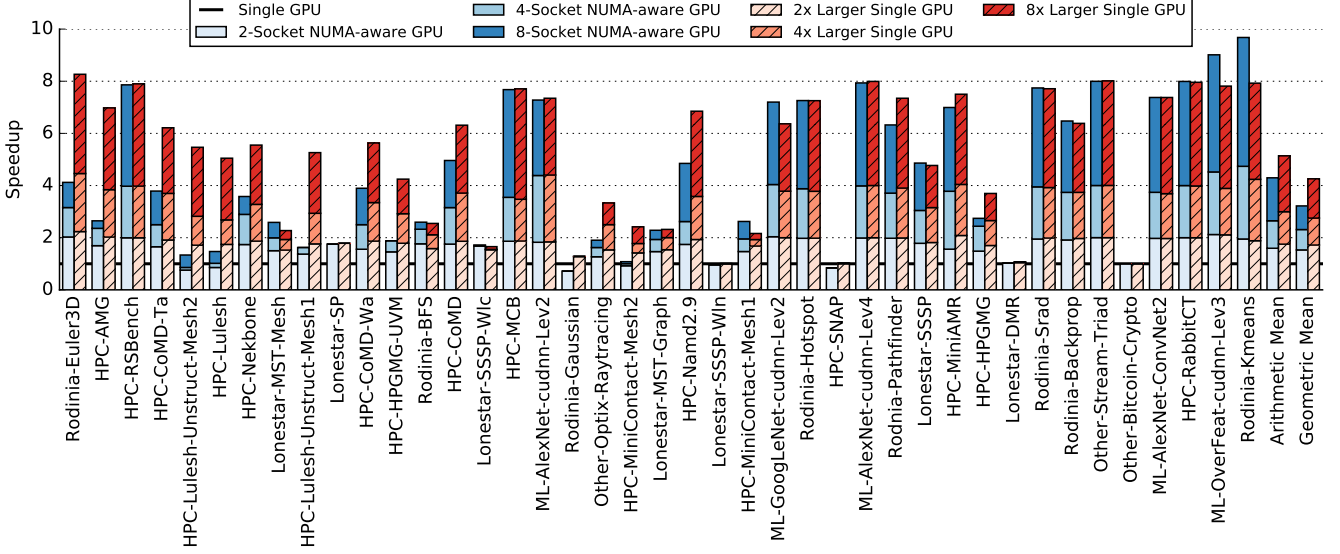


Figure 11: 1–8 socket NUMA-aware GPU scalability compared to hypothetical larger single GPU with scaled resources.

system level [1, 10, 25, 36] for a rapidly growing pool of applications [2, 3, 4, 26]. Similarly, multi-socket and multi-node CPU installations have been employed and studied in context of high performance computing and datacenter applications [37, 38, 39, 40]. Multi-GPU programming requires explicit design for multiple GPUs using SW APIs such as Peer-2-Peer access [8] or a combination of MPI and CUDA [9]. These extensions require unique programming experience and significant SW effort while adapting a traditional GPU application to a multi-GPU system. In this paper we execute single GPU applications on a NUMA multi-GPU system as if it was a single larger GPU via hardware innovations and extensions to the driver software stack; providing programmer and OS transparent execution, similarly to approaches proposed in the past [13, 14, 41].

Modern multi-socket CPU and GPU systems leverage advanced interconnect technologies such as NVLink, QPI and Infinity [10, 12, 21]. These modern fabrics utilize high speed serial signalling technologies over unidirectional lanes collectively comprising full-duplex links. Link capacity is statically allocated at design time and usually is symmetric in nature. In this paper we propose to dynamically re-allocate available link bandwidth resources by using same system wiring resources and on-chip I/O interfaces, while implementing both receiver and transmitter driver circuitry on each lane. This approach resembles previously proposed tristate bi-directional bus technologies [42] or former technologies such as the Intel front-side bus [43], albeit with just two bus clients. However our proposal leverages fast singled ended signalling while allowing a dynamically controlled asymmetric bandwidth allocation via on-the-fly reconfiguration of the individual lane direction within a link.

Static and dynamic cache partitioning techniques were widely explored in the context of CPU caches and QoS [44, 45, 46, 47, 48]. For example, Rafique et. al [46] proposed architectural support for shared cache management with quota-based approach. Qureshi et. al [47] proposed to partition cache space between applications. Jaleel et. al [48]

improved on this by proposing adaptive insertion policies. Recently, cache monitoring and allocation technologies were added to Intel Xeon processors, targeted for QoS enforcement via dynamic repartitioning of on-chip CPU cache resources [45] between applications. Efficient cache partitioning in the GPU has been explored in context of L1 caches [49] to improve application throughput. While dynamic cache partitioning has been widely used for QoS and L1 utilization, to the best of our knowledge it has never been used to try to optimize performance when caches are backed by NUMA memory systems.

8. CONCLUSIONS

With transistors growth slowing and multi-GPU programming requiring re-architecting of GPU applications, the future of scalable single GPU performance is in question. We propose that much like CPU designs have done in the past, the natural progression for continuous performance scalability of traditional GPU workloads is to move from a single to multi-socket NUMA design. In this work we show that applying NUMA scheduling and memory placement policies inherited from the CPU world is not sufficient to achieve good performance scalability. We show that future GPU designs will need to become NUMA-aware both in their interconnect management and within their caching subsystems to overcome the inherent performance penalty that NUMA memory systems introduce. By leveraging software policies that preserve data locality and hardware policies that can dynamically adapt to application phases, our proposed NUMA-aware multi-socket GPU is able to outperform current GPU designs by $1.5\times$, $2.3\times$, and $3.2\times$, while achieving 89%, 84%, and 76% of theoretical application scalability in 2, 4, and 8 GPU sockets respectively. Our results indicate that the challenges of designing a multi-socket NUMA GPU can be solved through a combination of runtime and architectural optimization, making NUMA-aware GPUs a promising technology for scaling GPU performance beyond a single socket.

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