Is Disaggregation possible for HPC Cognitive Simulation?

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Abstract—Cognitive simulation (CogSim) is an important and emerging workflow for HPC scientific exploration and scientific machine learning (SciML). One challenging workload for CogSim is the replacement of one component in a complex physical simulation with a fast, learned, surrogate model that is “inside” of the computational loop. The execution of this in-the-loop inference is particularly challenging because it requires frequent inference across multiple possible target models, can be on the simulation’s critical path (latency bound), is subject to requests from multiple MPI ranks, and typically contains a small number of samples per request. In this paper we explore the use of large, dedicated Deep Learning / AI accelerators that are disaggregated from compute nodes for this CogSim workload. We compare the trade-offs of using these accelerators versus the node-local GPU accelerators on leadership-class HPC systems.

Index Terms—deep learning, cognitive simulation, surrogate models, accelerator performance

I. INTRODUCTION

Recent advances in deep learning are leading to a resurgence of interest in the use of surrogate or reduced-order models to replace complex, first-principle physics packages in scientific simulations. Rather than expecting data-driven models to provide end-to-end predictions that replace entire multi-physics simulations, we explore new hybrid workflows that intertwine data-driven, learned models with traditional scientific simulation, a pattern in the community called Cognitive Simulation (CogSim). In this work, we explore the use of discrete, disaggregated accelerator hardware coupled with leadership class HPC resources to accelerate these workflows.

CogSim applications leverage AI for surrogate modeling of costly physics calculations [1]–[4], and are being developed to improve searching in multi-dimensional spaces and to automate manual processes. While the AI portion of the workflow is important and consumes significant compute resources, traditional scientific simulation CogSim workflows still require significant HPC compute capabilities – often at higher numerical precision.

Most of today’s leadership-class HPC systems use GPU accelerators for both the HPC and AI portions of CogSim workflows. While many research efforts are looking into other accelerators for HPC compute, no leading contenders have emerged. However, a large number of startup companies and traditional chip vendors are producing AI optimized accelerators, up to and including dedicated self-hosted machines. These machines show promising speedups for a variety of machine learning workloads [5].

With different processor architectures optimal for different pieces of a CogSim workload, an opportunity exists to build more cost efficient machines through the addition of another accelerator type. Accelerators could be integrated into every node or they can be treated as a disaggregated resource by being placed on the high-speed network of a supercomputer. Each approach has different cost and performance characteristics and the best approach may be workload dependent.

Using the newly installed SambaNova DataScale® system at Lawrence Livermore National Laboratory (LLNL) as motivation, we explore the question of whether the disaggregated design approach is viable for two test applications: a non-local thermodynamic equilibrium (NLTE) collisional-radiative atomic physics package [1] and a materials interface reconstruction task. We look at high-frequency machine learning inference of surrogate models due to the challenges this use case presents in time to solution and data transfer needs, and we aim to identify the most effective methods for performing it on each system. We aim to answer the question of should Deep Learning (DL) inference be performed on the GPUs, competing with the primary calculations, or could they be off-loaded to dedicated AI accelerators, where the DL calculation could occur simultaneously as the physics progresses on the GPU.

In this paper we make the following novel contributions:

- We describe a first of its kind Disaggregated CogSim system and how it was built. We also discuss how CogSim applications can use the disaggregated system and the relative challenges and costs.
- We show the trade-offs in terms of latency and throughput for processing inference samples on GPUs and AI accelerators for two independent surrogate models.

Our initial experimentation shows that disaggregated AI accelerators are viable from a technical perspective to speed-up surrogate inference calculations that occur in the Hydra multi-physics code used for simulating experiments on the National Ignition Facility at LLNL. Furthermore, our experimentation
with a material interface reconstruction surrogate demonstrates some of the challenges with both developing these models and mapping them to emerging accelerators.

II. Disaggregated Heterogeneous System Architectures for CogSim

CogSim is an emerging field that is generating new workflows and system requirements. Combining traditional HPC with ML to solve scientific problems, CogSim applications have unique requirements from traditional HPC applications. In this section, we describe one disaggregated system we have built for CogSim workloads and discuss how various CogSim use cases can benefit.

A. CogSim Disaggregated System Architectures

Disaggregated system architectures with heterogeneous node types have multiple advantages. By offering disparate node types, different parts of a calculation can execute on the most efficient compute resource. In addition, these architectures allow each application to run on the right mix of node types for its compute needs. This solves the stranded resource problem that arises when a system is built with a single heterogeneous node type with fixed ratios and multiple accelerators, where accelerator or CPU resources may be wasted. Disaggregation comes with costs though as network attached resources are connected with lower bandwidth and higher latency than node-local accelerators.

CogSim is a natural place to test whether disaggregated heterogeneous system architectures could work in practice. New accelerators show significant performance gains for ML applications. In addition, neural network training and inference calculations have small input and output requirements relative to their compute needs. Therefore, the impact of reduced bandwidth and additional latency from network attached resources should be minimal with this workload.

Using the DataScale AI accelerator from SambaNova Systems purchased by LLNL, we tested its ability to serve as a disaggregated CogSim system. This accelerator was integrated into the high speed network of the Corona supercomputer, which has a theoretical peak of over 11 PF. The DataScale node is attached to a top of the rack (TOR) switch that connects into the Corona core switches. Communication between Corona compute nodes and the DataScale system happens across a Mellanox Infiniband ConnectX-6 with up to 100Gb/s bandwidth and less than 1μs latency.

The DataScale system houses 8 SambaNova Reconfigurable Dataflow Units (RDU)™ that utilize the SambaNova Reconfigurable Dataflow Architecture (RDA)™. DL models are compiled and run on the system using the SambaNova SambaFlow™ software stack. Scheduling on the DataScale system is done with the SLURM job manager.

B. CogSim Workloads on Disaggregated Machines

Within the field of CogSim there are multiple ways that AI is incorporated into scientific computations. Figure 1 shows examples of three “classes” of possible use cases and how they are embedded within the simulation: in-the-loop, on-the-loop, and around-the-loop. Starting from the center, in-the-loop represent some of the most challenging tasks for integration because AI models are part of the inner most loop and thus typically on the critical path. Tasks can range from inference calculations of surrogate models that replace physics computations that occur on every simulated zone for every timestep, to inference calculations that help reconstruct mesh zones on material boundaries. These in-the-loop problems typically work with small batch sizes and should be tuned to minimize inference latency. Additionally, it is quite possible to have multiple models for in-the-loop inference that are used in different physics regimes, and thus should support concurrent execution.

The around-the-loop training problem presents the easiest case for adapting to a disaggregated system. Outer loop training takes inputs from one or more simulations. Since all of the data needs to be aggregated in one place for training and training is more compute intensive than inference, the relative network bandwidth and latency requirements are relaxed compared to the other use cases that we present. The main challenge for outer loop training is balancing resource types and making sure models are retrained fast enough to improve the quality of the workflow as new data are produced.

Presenting more of a challenge is training/retraining during a simulation or inference that occurs on-the-loop every \( x \) timesteps. While more tightly coupled, updating these models is not urgent and retraining of the model will often include old data that is best stored elsewhere and not on the memory-constrained HPC compute nodes. On-the-loop inference problems often permit significant time for the computation to occur before an inference result is needed.

The two outermost use cases result in system balance and allocation challenges to avoid stranded resources due to bursty use. They also present bandwidth and latency requirements, but those are not the strictest requirements. Of the three use cases, the most tightly coupled physics surrogate models that run in-the-loop present the largest challenge to using a disaggregated system. This use case sends significant data per simulated zone to the accelerator, and it has low latency tolerance for a response. Due to these challenges we focus on such in-the-loop calculations in this paper. We believe that if performing these calculations are viable using a disaggregated accelerator, then it follows that the other two use cases will be as well.
III. RELATED WORK

In this paper, we focus on many new and emerging technologies and techniques, like CogSim and dataflow architectures. There are a small number of prior publications we can look to for context of our work. For example, in [5] the SambaNova Reconfigurable DataFlow is compared to an Nvidia V100 GPU. The authors focus on the ability to train larger models on the DataScale that would typically not fit on GPU. They demonstrate how larger networks and input data sizes on dataflow architectures generate more accurate models than the GPU in the same number of epochs. This work is largely focused on typical ML workloads, such as image classification and NLP tasks.

A number of publications demonstrate the use of surrogate modeling in physics applications [2–4]. These works introduce the use of surrogate models for in-the-loop inference with physics applications like inertial confinement fusion [2,3] and fluid simulation [4]. While these publications focus on how the physics applications can be performance optimized through CogSim, they do not explore how different architectures and configurations (e.g., remote vs local inference) can be adapted to the CogSim workload. In this paper, we extend the current scope of the literature by focusing on how traditional GPU architectures and newer dataflow architectures perform with CogSim workloads.

In extending the CogSim and dataflow architecture literature, we also look to many publications related to Deep Learning GPU performance. For example, in [6] the authors survey the scope of recent publications that demonstrate optimizing DL tasks on GPUs. Similar to publications outlined in [6], we test different APIs, toolkits, software, mini-batch sizes, and hardware to optimize the performance on both GPU and dataflow architectures. More specifically, we use PyTorch [7] and focus on inference performance [8–10]. We test and compare neural network inference performance on both Nvidia and AMD GPUs [11]. For each GPU architecture, we compare multiple generations of GPU [12]. We also test toolkits and APIs like TensorRT [13] for improving inference performance on Nvidia GPUs [10,14].

From these works, we gather best practices that we applied in our own performance experiments. For example, one way that we ensure a fair comparison between architectures in the context of CogSim workloads is considering data movement and if it should be included in latency and throughput measurements [15]. Unlike these previous publications, we extend the performance measurement work to dataflow architectures. In doing so, we establish novelty of our work in the neural network inference performance and CogSim space.

IV. IN-THE-LOOP INFEERENCE

As discussed in Section II-B, one of the most challenging tasks for a disaggregated AI accelerator is to support in-the-loop inference. The challenges are that the inference is typically on the critical path of the primary scientific calculation, may require multiple independent models to serve different inference tasks, will typically have small numbers of requests per time-step, and will have to serve requests from multiple compute nodes (and distributed MPI ranks). These system integration challenges are why this paper focuses on two data-driven surrogate models: Hermit and MIR. It is also important to note that neither these models are massively large, or utilize complex neural network architectures or layers, and as a result allow this study to focus on the system level issues.

A. Hermit: a surrogate model for NLTE collisional-radiative atomic physics

The Hermit neural network model described in [1] is diagrammed in Figure 2a. The model consists of 21 fully connected layers across 3 sub-structures: an encoder, Deep Jointly-Informed Neural Network (DJINN) layers, and a decoder. The total input size for each sample is just 42 values. The encoder has 4 layers with a maximum hidden layer width of 19 and the decoder has 6 layers with a maximum hidden layer width of 27 neurons. The bulk of the network size from the DJINN layers, which reach a maximum width of 2050 neurons. In total, there are 2.8M parameters in the Hermit model.

We use the description of the Hermit model integration with the Hydra physics simulation code in [16] to characterize the needs of our in-the-loop inference requirements and test our hypothesis that disaggregated systems are feasible for CogSim applications. Hermit is used to replace the Detailed Configuration Accounting (DCA) package and requires two or three inference calculations per zone in each timestep of the simulation. Typical simulations can be tens of thousands of timesteps. The relationship between DCA or Hermit and the rest if Hydra is shown in Figure 2b and 2c.

While problem and use case dependent, typical Hydra problems using DCA physics are run with only a few zones per node due to high memory capacity needs and/or time to solution requirements [17]. On Nvidia V100 GPUs, users typically run 100-1,000 zones per GPU when using DCA physics models. Swapping in the Hermit model both accelerates the calculation and reduces the memory footprint, enabling the user to run with more zones per GPU. In Hydra, two or three inference calculations are required per zone in each timestep of the simulation. With 10,000 zones per GPU, 20,000 - 30,000 inference calculations are needed per timestep. In addition, inference requests from each MPI rank are submitted to different Hermit models, where each model is trained to represent a particular material. An MPI rank might typically require results for 5-10 different materials. The low number of inference calculations needed and the fact that they are spread across multiple models means small batch size performance is key to Hermit performance.

B. Material Interface Reconstruction (MIR) Model

The MIR model is a neural network used for material interface reconstruction in physics simulations, in which boundaries are constructed between immiscible materials based on volume fractions for each zone in the environment. Current methods tradeoff continuity and conservation of material,
often to the detriment of the reconstruction accuracy. Figure 3a compares outputs for one current reconstruction method, PLIC, and the MIR model. PLIC conserves the volume of the material in each zone but leads to discontinuous boundaries. The MIR model is able to create continuous boundaries while conserving volumes and looks remarkably like the ground truth.

The MIR model is a convolutional autoencoder and is diagrammed in Figure 3b. It is composed of 4 convolution layers with pooling, layernorm after every convolution, 3 fully connected layers, two of which with 4608 neurons each, and transposed convolution layers to return the output of the fully connected layers to the size of the original image. The weights of the convolution and transposed convolution layers are tied as a form of regularization. In total, there are 700K parameters in the MIR model.

Mixed zones, or zones with more than one material in them, are processed each simulation timestep. The workflow of the simulations is shown in Figure 3c. The number of mixed zones per GPU for each simulation timestep ranges from the thousands to the hundreds of thousands depending on the application. The number of zones per timestep may vary throughout the simulation in some applications. To not impede the physics calculations going on around it, the target throughput of the model is 100,000 samples per second per MPI rank.

C. Optimizing the MIR Model for dataflow architectures

The original MIR model, which was much larger, did not translate well to a dataflow architecture. In order to optimize for dataflow, batchnorm layers were replaced with layernorm layers and work was done to shrink particularly large fully connected layers. The GPU model was subsequently changed to match the version run on the DataScale system.

V. Evaluation

To test the efficacy of these new accelerators for in-the-loop inference workloads we use the Hermit and MIR models, described in Section IV. We measure the inference performance of 3 accelerator architectures: 1) Nvidia GPUs, 2) AMD GPUs, and 3) SambaNova DataScale. We describe our experimental setup, demonstrate optimizing performance on GPU and dataflow architectures, and compare the architectures for in-the-loop CogSim workloads.

A. Experimental Setup

In this section, we describe the experimental setup for each of the tested architectures. We established a fair comparison across different architectures by working with the vendors to ensure optimal configurations and accurate measurements of the hardware performance. We implemented the models in PyTorch and used them as references to generate the models on all tested hardware. For the GPU tests, the models were run with the PyTorch API as well as used to generate compiled models for Nvidia’s TensorRT and CUDA Graphs APIs. On the DataScale, the SambaFlow software stack compiles a model from the PyTorch reference model.

In our experiments for all hardware, we measured half-precision inference latency and throughput bandwidth. We performed these measurements across a range of mini-batch sizes (i.e., 1, 4, 16, 64, 256, 1K, 2K, 4K, 8K, 16K, and 32K) to capture the performance landscape and record how it is modulated by the number of samples being pushed through a model. We adjusted the total number of samples run through a model, with respect to the mini-batch size, such that the total wall-clock time is greater than 10s for each performance run. Latency was measured in milliseconds (ms) as the average time across all mini-batches for running inference on a single mini-batch of samples. Throughput was measured in samples per second across all samples of a given mini-batch size. Before each performance measurement on the GPUs, we “warmed-up” the hardware by running inference on 10 mini-batches. The performance measurement occurred directly after the warm-up phase.

We performed two types of experiments: node-local inference and remote inference. As discussed in Section II, surrogate models can reside on the same GPU as the CogSim simulation. Therefore we measured the node-local inference for GPUs, where the input data is generated and model inference is executed on the same GPU. Also discussed in Section II new dataflow architecture accelerators, such as the SambaNova DataScale, can reside on separate nodes and be made available to compute nodes via a high-speed InfiniBand
network. We first measure node-local inference on the DataScale to optimize model performance, sans network latency, to obtain an upper bound on performance. We then measured remote inference with the DataScale, where input data is generated on a compute node, sent across the network to the DataScale node, and inference results are sent back across the network to the originating compute node. Communication was done via a prototype C++ API and library, but these tests mimic the expected use case where multiple MPI ranks would issue queries to the DataScale node. Additionally, the GPU measurements include no data movement between system memory and the accelerator to mimic the real-world CogSim application, where the simulation and models are both resident on the GPU. Measurements on the DataScale do include movement to and from system memory for both node-local and remote inference.

We tested several generations of GPU architecture from Nvidia (i.e., P100, V100, and A100) and AMD (i.e., MI50 and MI100). For each of these hardware types, we use PyTorch 1.9.0. Our Nvidia setup uses CUDA 11.4, cuDNN 8.2.2.26, and TensorRT 8.0.1.6. Our AMD setup uses ROCm 4.2 and MIOpen 2.11.0. Models were run in half-precision with FP16 on the GPUs. In addition to our baseline PyTorch implementation, we also worked with the vendor to optimize performance for an A100 GPU. We tested TensorRT and CUDA Graphs APIs in Python and C++. TensorRT, with the Torch2TRT library, compiles our PyTorch model to optimize performance through kernel selection and by combining layers. CUDA Graphs allows the model to be called from a single CPU operation and reduces kernel launch overhead from the PyTorch API. While TensorRT, CUDA Graphs, and C++ add complexity to running the model, the benefit-cost ratio is high for a frequently used inference model (e.g., a CogSim surrogate model).

For the dataflow architecture, we tested the DataScale AI accelerator from SambaNova Systems. Our surrogate models are run in half-precision with BF16 on the DataScale. The SambaFlow 1.8 library generated a compiled model from the PyTorch model. Given the relative novelty of this hardware, we performed a more extensive survey of the performance landscape compared to GPU-related experiments. The DataScale system is composed of 8 SN10 RDUs. Each RDU contains 4 tiles, which are discrete compute and memory units of the RDU. A single model can be deployed in various configurations, ranging from $\frac{1}{4}$ of an RDU (i.e., 1 tile) up to a complete RDU (i.e., 4 tiles). The RDU uses an additional micro-batch size parameter that the GPUs do not have. This parameter determines the size of accumulated data sent across the RDU tiles during inference. The micro-batch size must always be equal to or less than the mini-batch size. In our exploration of DataScale performance, we tested how inference performance scales with RDU tiles using different combinations of mini-batch and micro-batch sizes. From these results we established a baseline for PyTorch node-local performance of the DataScale and then worked with the vendor to optimize the model using C++ APIs and hand-tuned model placement on the hardware.

Remote inference performance tests on the DataScale were run with the configurations (i.e., RDU tile count, mini-batch size, and micro-batch size, optimizations) that provided the best performance. In remote inference experiments, data is generated on a remote compute node, sent across a high-speed network, inference is performed on the DataScale, and the results are returned to the remote client. The high-speed network in our tests was the Mellanox Infiniband ConnectX-6 with a bandwidth of 100 Gb/s. Latency measurements include the additional round-trip of data transfer. Throughput was maximized in these tests by allowing asynchronous communication between the client and SN10-8 server. The client sends mini-batch $n+1$ to the server before inference results for mini-batch $n$ are returned to the client.

All experiment measurements were replicated 5 times. The figures in the remainder of this section plot the mean of the 5 measurements with error bars indicating the 95% confidence interval.

B. Tuning for the GPU architectures

We measured performance of 2 surrogate models, Hermit and MIR, across 3 generations of Nvidia GPUs: P100, V100, and A100. Figure 4 shows the inference latency of the Nvidia GPUs across different mini-batch sizes for the Hermit model. The left panel of the figure shows a nearly constant latency for each of the GPUs at mini-batch sizes smaller than 256. The A100 has the lowest single sample latency of 0.65ms. Perhaps unexpectedly, the V100 latency is larger than the P100 at these small mini-batch sizes. The Hermit model is a
small network, so with the “naïve PyTorch” implementation, performance at small mini-batch sizes is CPU-bound. This is due to the overhead of PyTorch CPU logic that determines which GPU kernels to launch. In context of the measured GPU performance, both the P100 and A100 systems use x86 architecture, while the V100 system uses Power9. The CPU-bound nature of the naïve PyTorch implementation at small mini-batch sizes is the cause of larger latency with the V100 compared to the P100 in Figure 4.

On the right side of Figure 4, we see that the latency increases more rapidly for the P100 than either the V100 or A100. This result suggests that at these larger mini-batch sizes for the Hermit model the P100 hardware becomes saturated. The P100 latency is more than 8x that of the A100 at the largest mini-batch size of 32K. The A100 has a latency of 3.92ms at this mini-batch size.

Measured throughput of the three Nvidia GPUs with the Hermit model is shown in Figure 5. We see a similar pattern as the latency measurements, with the V100 being slower than the P100 at the smallest mini-batch sizes. At larger sizes the additional transistor and memory hardware on the V100 and A100 are apparent as they achieve inference throughputs in excess of 5 Million samples/s. The throughput measurements of the A100 were largest for all mini-batch sizes, with 1 and 32k mini-batch throughputs of 1,534 and 8.35M samples/s, respectively.

In Figure 6 we show the measured latency for Hermit on two AMD GPUs, the MI50 and MI100. We observe near constant latency with the MI100 for mini-batch sizes at and below 1K. Single sample latency of the MI100 is measured at 0.96ms. MI50 performance was similar to P100 performance in Figure 4 as we see a marked increase in latency as the mini-batch size increases beyond 1K. Our measurements also show an unexpected drop in performance for the MI100 at a mini-batch size of 4K. At the maximum mini-batch size of 32k, we recorded a latency of 5.59ms on the MI100, corresponding to a maximum throughput of 5.85M samples/s.

A comparison of the fastest Nvidia GPU (i.e., A100) and fastest AMD GPU (i.e., MI100) is shown in Figure 6. We observe that the measured throughput of the A100 is larger than the MI100 at all tested mini-batch sizes. At the 32K mini-batch size, the A100 can process more than 2M additional samples per second than the MI100. We also note that the A100 has a lower TDP at 250W than the MI100 at 290W. We normalize throughput of the MI100 based on TDP and show these values in Figure 6 as well. At the smallest mini-batch sizes the A100 is superior to the MI100 with respect to latency, with measured single sample latencies of 0.65ms and...
Throughput (samples/s)

Latency (ms)

Fig. 7: Inference latency of the Hermit model on the Nvidia A100 and AMD MI100.

0.96ms. The unexpected plateau in performance of the MI100 between mini-batch sizes of 1K and 4K shown in Figure 7 may be explained by the beta support for AMD GPUs of PyTorch 1.9.0.

Fig. 8: Inference latency of the Hermit model on a Nvidia A100 GPU using combinations of Python and C++ APIs for PyTorch, CUDA Graphs, and TensorRT.

Given the baseline performance measurements that we obtained, we next optimized the performance of the Hermit and MIR models for the Nvidia A100 GPU. We worked closely with the vendor to identify and implement optimizations that would improve latency and throughput in our experimental measurements. We tested five configurations: (1) naïve PyTorch implementation, (2) PyTorch with TensorRT using the torch2trt library, (3) PyTorch with CUDA Graphs, (4) PyTorch with TensorRT and CUDA Graphs, and (5) C++ TensorRT API.

Figure 8 shows the measured inference latency with these five configurations on an A100 GPU with the Hermit model. The left panel of Figure 8 shows that all configurations are more than twice as fast as the initial naïve PyTorch implementation for single sample latency. PyTorch with TensorRT and CUDA Graphs provides the lowest inference latency for all mini-batch sizes, with a single sample latency of 0.12ms and a 32k samples latency of 1.52ms.

Fig. 9: Inference throughput of the Hermit model on an Nvidia A100 GPU using combinations of Python and C++ APIs for PyTorch, CUDA Graphs, and TensorRT.

Fig. 10: Throughput of the MIR Model using combinations of PyTorch, CUDA Graphs, and TensorRT APIs.

Throughput measurements for different configurations on the A100 with the Hermit model are presented in Figure 9. PyTorch with TensorRT and CUDA Graphs had the largest bandwidth for all mini-batch sizes. We measured a single sample and 32K sample throughput of 8,240 samples/s and 21.6M samples/s. We observe that all the configurations using TensorRT provide very similar bandwidth performance across the tested mini-batch sizes.
We also measured performance of the MIR model on an A100 with 4 of the described configurations. Figure 10 shows that for this larger model, CUDA Graphs gives the greatest increase in throughput bandwidth. This figure also shows that configurations using TRT have measurably worse performance than the standard PyTorch implementation at mini-batch sizes larger than 64. We note that this is caused by the Torch2TRT library that we use to convert our PyTorch model to a TensorRT model. The library has unoptimized implementations of layernorm and unary functions that cause a performance bottleneck in the configurations using TensorRT. The vendor has indicated that the upcoming TensorRT release will address this issue. Figure 10 shows, that unlike with the Hermit model, the MIR model performance on the A100 with different configurations converge at the largest mini-batch size to nearly equal throughput bandwidth. This may indicate that at the largest tested mini-batch size, the compute capability of the A100 is saturated, regardless of the configuration.

### C. Tuning for a data flow architecture

As discussed in Section V-A, the DataScale has several tunable parameters. We measured the node-local performance of the Hermit model on a different amount of RDU compute resources (i.e., RDU tiles) and across a range of micro-batch and mini-batch sizes. Figure 11 shows the latency of inference requests with mini-batch sizes and micro-batch sizes ranging from 1 to 32K on a single RDU tile (i.e., 1/4 RDU). In Figure 11 we see that both mini-batch size and micro-batch size modulate the latency of the network on the DataScale. Each mini-batch size has a micro-batch size that provides optimal performance in terms of latency. We highlight the minimum latency values in purple for each mini-batch size. We also note that tuning the micro-batch and mini-batch size to be multiples of 6 offers additional performance in terms of latency and throughput by exploiting hardware properties of the DataScale. We show this below in Figures 13 and 14. However, for the purposes of these tests, we focus on common mini-batch sizes that are also optimized for GPU architectures.

![Fig. 11: Latency of the Hermit model on 1/4 RDU for a range of mini-batch and micro-batch sizes using the Python API. The lowest latency (mini-batch, micro-batch) pair for each mini-batch size is highlighted.](image)

![Fig. 12: Latency of the Hermit model on 1 RDU for a range of mini-batch and micro-batch sizes using the Python API. The lowest latency (mini-batch, micro-batch) pair for each mini-batch size is highlighted.](image)

Figure 12 shows the latency of inference requests with the same range of mini-batch size and micro-batch size as Figure 11 but on 4 RDU tiles (i.e., one RDU). Again, we highlight the combination of mini-batch size and micro-batch size that give optimal performance. We observe that providing more RDU tiles for model inference changes which mini-batch and micro-batch size combinations give optimal performance. In both Figure 11 and Figure 12 the white squares indicate configurations that are not valid (e.g., micro-batch size larger than mini-batch size) or failed to run on the hardware. From both figures, we can see that at low mini-batch sizes, the micro-batch size has benign effects on performance. As the mini-batch size increases, choosing the proper micro-batch size is important for optimizing performance. For example, in Figure 12 at a mini-batch size of 32K, the difference between the slowest and fastest micro-batch size is 10-fold. For this reason, we performed parameter sweeps of the (mini-batch, micro-batch) landscape for each tested configuration and report the maximum throughput and minimum latency for each mini-batch size in the remainder of this section.

Similar to the Nvidia A100 GPU, the DataScale has several configuration options that can optimize the model for latency and throughput. We tested the Python and C++ APIs and hand-optimized model placement for the DataScale to optimize node-local inference. Figure 13 shows the latency of the Hermit model on 1 RDU with several configurations that optimize for performance. We show the base performance of the Python API (i.e., “naïve”) and show how different modifications improve overall performance. Hand-optimized model placement (i.e., “optimized”) on the hardware provides benefits to the latency, especially at larger mini-batch sizes.
Throughput (samples/s)

Latency (ms)

0.03

2.0M

0.10

3.0M

0.12

4.0M

0.15

5.0M

0.18

2.0M

4.0M

6.0M

8.0M

10M

Fig. 13: Inference latency of the Hermit model on the DataScale with 1 RDU and 3 optimization methods.

We also show that switching to a C++ API (with hand-optimized model placement) provides additional benefits to latency. These benefits are significant for the smallest mini-batch sizes, where inference latency is more than halved compared to the Python API.

An additional optimization we show in Figure 13 is from making small adjustments to the mini-batch and micro-batch sizes. In Figure 12 we showed the performance landscape for mini-batch and micro-batch sizes that are powers of 2 (i.e., sizes \( \subset 2^n \)). However, the DataScale hardware design is such that multiples of 6 for micro-batch sizes and mini-batch sizes that are multiples of a given micro-batch size can provide better performance. We show the effect of using “preferred MB” in conjunction with the other optimizations in Figure 13. The lowest latency values are observed with the C++ API and hand-optimized model placement, with the exception of the 2 largest mini-batch sizes, where the Python API provides slightly lower latency. At the smallest mini-batch sizes we observe a minimum latency of 0.04ms. The “preferred MB” optimization provides additional reduction in latency.

We also measured the throughput bandwidth of the different configuration of the DataScale. Figure 14 shows this data. On the right panel of the figure, we see that the C++ API with hand-optimized model placement provides a maximum throughput bandwidth of 8.14 M samples/s at a mini-batch size of 16K. While the “preferred MB” optimization shows better latency and throughput, to maintain a fair comparison with GPU architectures we show only the powers of 2 mini-batch sizes in subsequent figures. Figure 13 and Figure 14 show that the C++ API with hand-optimized model placement provided the best overall performance on a single RDU in node-local performance tests. We use this configuration for the remote inference experiments. As such, the node-local performance will serve as an upper limit for the performance we can expect from remote inference.

Because the DataScale System will be disaggregated from other system compute nodes, the remote inference experiments provide information about expected performance with CogSim applications that do in-the-loop inference. Figure 15 shows the remote inference latency measurements for the Hermit model compared to node-local latency measurements. All models were run with the hand-optimized model placement, the remote inference is through the C++ API, and both Python and C++ APIs are shown for node-local. In Figure 15 we observe that remote inference adds additional latency overhead compared to both Python and C++ node-local inference. However, at the smallest batch sizes on the left panel of Figure 15 we see that C++ remote inference can be as fast or faster than Python node-local inference, with an average four sample latency of 0.05ms. At a mini-batch size of 16K, we observe the largest difference in performance.
between the node-local and remote inference with the C++ API at 1.14ms.

Remote inference throughput measurements for the DataScale are shown in Figure 16. In the left panel of the figure, we see that the C++ remote inference throughput was between node-local Python and node-local C++ inference measurements for small mini-batch. At mini-batch sizes greater than 1K, both node-local configurations exceeded the remote inference throughput. At a mini-batch size of 16K, a maximum remote inference throughput of 6.4M samples/s was recorded. Given the competitive performance of the DataScale remote inference compared to node-local performance shown in Figure 15 and Figure 16, we find a compelling reason to support disaggregated systems for CogSim workloads. In the following sections, we directly compare the performance of the dataflow architecture with the more traditional GPU accelerators.

D. Latency Comparison

The latency of inference is an important measurement for CogSim applications. With in-the-loop inference, the time to return an inference result needs to be short enough to avoid bottlenecking and degrading performance of the application. In Section V-B we measured the performance of GPU node-local inference in the context of a CogSim application running a surrogate model. And in Section V-C we measured the performance of the DataScale node-local and remote inference. For disaggregated accelerators, the remote inference is the measurement important to CogSim workflows.

Figure 17 shows a comparison of the mini-batch latency measured for different configurations of the A100 and DataScale, from Sections V-B and V-C. The included configurations represent the fastest and slowest performance for each and the fastest remote inference for DataScale. We observe that at mini-batch sizes below 1K, the node-local RDU provides a lower latency than the A100. At mini-batch sizes in the range [4, 256] the measured latency of the remote inference on the DataScale is lower than the latency of the most optimized node-local A100. As the mini-batch size increases above 256, the node-local performance of the A100 exceeds first remote and then node-local performance of the DataScale System.

From the measured latency of inference on the A100 and DataScale presented in Figure 17, we find that neither GPU or dataflow architectures dominate the performance landscape. However, for the CogSim application of the Hermit model, small mini-batch size performance is most important. Therefore, we observe that our disaggregated system out-performs traditional GPUs for our specific in-the-loop inference.

More generally, we observe the Hermit model at smaller mini-batch sizes is more performant on the DataScale System while larger mini-batch sizes are faster on the A100. Given the different behaviors of optimized configurations for Hermit and MIR models in Section V-B, we find that the viability of disaggregated systems for CogSim workloads is heavily driven by choice of surrogate model and latency tolerances.

E. Throughput tests

Throughput tolerances are also important for CogSim workloads. Figure 18 shows the Hermit inference throughput for the same configurations of the A100 and DataScale as in Figure 17. In the left panel of the figure, we see that in all configurations at mini-batch sizes below 1K, the DataScale is measured to have the largest throughput. As the mini-batch size increases above 1K, the A100 throughput exceeds the DataScale throughput. We also observe remote inference has less bandwidth than the node-local inference on a single RDU. This drop in throughput bandwidth is a result of additional latency and overhead associated with the high-speed network connecting the remote client to the DataScale System.

The relative performance measured on a single RDU of the DataScale compared to the A100 for different configurations with the Hermit model is shown in Figure 19. This plot
shows the speedup factor of the DataScale System on the Y-axis across all tested mini-batch sizes. We compare 3 sets of configurations between the A100 and RDU: (1) naïve PyTorch implementations (i.e., the slowest), (2) optimized node-local implementations (i.e., the fastest), and (3) optimized A100 node-local and optimized DataScale remote (i.e., CogSim workload). We include another set of datapoints for configuration (3) where we normalize the DataScale throughput by transistor count. The A100 has 1.3x the transistor count of the DataScale RDU. Above the horizontal dotted line of Figure 19 indicates higher throughput bandwidth on the DataScale compared to the A100. These comparisons show that the DataScale dominates performance at the lower mini-batch sizes with the Hermit model. The largest difference in performance is between the most highly optimized node-local measurements, with a more than 7X speedup. In the context of CogSim and in-the-loop inference, we find that the remote inference DataScale measured performance is more than 3X that of the most highly optimized node-local A100 for the smallest mini-batch sizes. As the mini-batch sizes increase above 1K, the DataScale System lags behind the A100. At the largest mini-batch sizes, the A100 offers better performance.

Switching to the MIR model, which has a target throughput of at least 100K samples/s, we compare DataScale and Nvidia GPU performance in Figure 20. This comparison is done on a version of the MIR model without layernorm to ensure the model would compile optimally on both architectures. In this Figure, we show the target throughput with a horizontal dashed line. We observe that at low mini-batch sizes, throughput is similar between the A100 and DataScale. The DataScale system reaches the target throughput bandwidth at a mini-batch size of 128 while the A100 reaches it at size 256. As the mini-batch size increases toward 8K, the DataScale system reaches a maximum throughput of over 140K while the A100 struggles to achieve a throughput much larger than 100K. This result is contrasting to the results with Hermit in Figures 18 and 19, where the DataScale provided the largest advantage over the A100 at small mini-batch sizes.

Our comparisons of measured latency in Section V-D and throughput in this section for the DataScale and A100 reveal that new dataflow architectures are viable for disaggregated CogSim workloads. Specifically, we demonstrated that with the Hermit model at mini-batch sizes below 1K and with the MIR model at large mini-batch sizes the DataScale dominates performance. This indicates that disaggregated systems are not only competitive, but potentially faster than traditional GPU accelerators in these contexts. From differences and similarities in the performance landscapes between the two tested models, we conclude that the DataScale and A100 are more performant based on several contributing factors, including model size and throughput/latency requirements. For example, in the context of tight latency requirements for large mini-batches with Hermit, the A100 would be the best option.
Our current results are model specific and cannot capture all the complexities of the contributing factors. We discuss how our results can be generalized to better demonstrate the viability of disaggregated systems in the following section.

VI. CONCLUSION

In this paper, we explored the viability of disaggregated systems for CogSim workloads, including in-the-loop inference. We tested our hypothesis using two surrogate models, Hermit and MIR, across three Nvidia GPUs, two AMD GPUs, and the SambaNova DataScale system. Measuring inference latency and throughput across various configurations of the hardware, we described the performance landscape and determining factors of DL performance. Our results indicate that disaggregated systems are viable and in some cases offer better performance than node-local GPU accelerators for CogSim workloads. We found that the optimal hardware for CogSim workloads is largely determined by the model and latency/throughput requirements.

Throughout this work we engaged very closely with vendor engineering teams to ensure that our applications were mapped as efficiently as possible for each accelerator architecture. We found that that there were opportunities to use these example applications to identify areas for improving the vendor’s tool chains for CogSim SciML workflows. One area that is the subject of ongoing work is a generalized application for remote inference on the DataScale, which supports remote inference to multiple, independent models that is necessary for the Hermit and MIR integration.

While our findings are relevant to CogSim workloads and more specifically applications that use Hermit and MIR models for in-the-loop inference, our results are not extensive enough to generalize to all workloads. The results we obtained indicate that factors like model size, layer types, computational requirements of the simulation, optimizability of the model, as well as inference latency and throughput requirements are contributing factors to determining the viability of a disaggregated system for a particular workload. Our future work aims to explore this space by extending our results to more automatically generated DL models that represent a wide array of CogSim applications. This work would serve as a reference for other researchers to indicate if a disaggregated system is viable for a given CogSim application.

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