Fluid Co-processing: GPU Bloom-filters for CPU Joins

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ABSTRACT
It has so far been unclear which data-intensive CPU tasks can be accelerated with GPUs, as GPUs are bottlenecked by the slow bus connection to the CPU and the limited size of GPU memories.

In this paper we demonstrate a database workload where co-processing actually helps: accelerating large join pipelines where the join condition is selective, by pushing down a Bloom filter test for early pruning. GPUs are more powerful than CPUs for computing hash functions needed in Bloom filter tests, have a local memory with significantly more random-access bandwidth than the CPU, and since only keys (or extracts thereof) have to be moved to the GPU, data transfers over the bus are relatively small. Our micro-benchmarks show that raw Bloom filter lookups are up to \(6 \times\) faster on the GPU than on the CPU in case the Bloom filter is larger than the CPU cache.

The next quest is for a database architecture that allows efficient CPU-GPU co-processing. We present a new heterogeneous query processing framework based on fluid co-processing. In fluid co-processing, tasks of different sizes – that fit the device – are dynamically co-processed. Early results show that fluid co-processing consistently improves end-to-end CPU performance of early pruning in join queries thanks to the GPU, by factors up to 2-3\(\times\).

ACM Reference Format:

1 INTRODUCTION
As the performance of modern processors is increasingly limited by power consumption and heat dissipation, the so called power wall, modern hardware progressively evolves into a landscape of heterogeneous devices. Besides traditional CPUs, these devices include GPUs, FPGAs and specialized hardware. Particularly, the acceleration of compute-intensive tasks [20] using GPUs has gained tremendous traction.

GPUs significantly outperform CPUs in compute-intensive tasks, as they combine a massively parallel architecture with high bandwidth memory. However, since analytic workloads are data-intensive rather than compute-intensive, GPUs typically fail to accelerate database workloads. The main problems are the limited size of the local GPU memory and the limited PCI bus bandwidth between CPU and GPU. GPUs typically have less than 16 GiB of memory, while main memories attached to a CPU are typically an order of magnitude larger (and flash or other persistent memories are yet another order of magnitude bigger). The current generation of bus architectures in practice is limited to sustained transfer of 12 GiB/s; whereas CPU memory bandwidth is well over 100 GiB/s and internal GPU memory can be over 400 GiB/s.

In scientific literature where advances of GPU-accelerated database systems are demonstrated, the experiments are typically performed on small datasets, cached in the local GPU memory, such that significant CPU-GPU data transfers are avoided; or the CPU baselines that are compared with are not state-of-the-art systems (e.g. VectorWise [3, 23] or Hyper [17, 19] would qualify our definition). While there are a few start-up companies with GPU-accelerated database solutions, these have little traction yet in the industry – a far cry from the situation in Machine Learning (ML) where GPUs are in extreme demand by industry (ML being a compute rather than data-intensive workload). Given that heterogeneous hardware acceleration would be especially useful for Big Data analysis tasks where queries take a lot of time, the search was still on for database use-cases where GPUs fit in.
This paper contributes a use-case where GPUs can truly contribute: early pruning in large selective joins.

Recent work [13] shows that GPUs are able to efficiently prune using Bloom filters [2]. In database systems, Bloom filters can be exploited in queries with selective joins, to quickly eliminate disqualifying tuples, before they even enter the join. Concretely, after completing the build phase of a hash-join, a bloom filter can be created that holds the set of keys in the hash table. This Bloom filter is smaller than the hash table and is cheaper to probe. Also, it can be probed very early in the query (early pruning) even in the scan operator on the probe side of the query plan. This could mean that when whole stretches of keys fail to qualify, whole blocks of data holding the non-key columns can be skipped (saving I/O and decompression effort in column stores). Also, other operators in between the scan and the selective join (e.g., other joins or aggregations) will receive less tuples and will be accelerated, thanks to the early pruning. This is depicted in the left of Figure 1. However, large hash tables that hold many keys require larger Bloom filters. Once the Bloom filter size exceeds CPU caches, L3 cache size in particular, the Bloom filter lookup throughput drops significantly due to additional main memory access. The right side of Figure 1 illustrates that this is a situation where the higher memory bandwidth of GPUs will significantly improve lookup performance. A Bloom filter is much smaller than a hash table storing the same amount of keys and some payloads – therefore it is reasonable that to assume that if a hash table fits in the CPU memory, the corresponding Bloom filter should fit in GPU memory. GPUs are more powerful than CPUs for computing hash functions needed in Bloom filter tests, have a local memory with significantly more random-access bandwidth than the CPU, and since only keys (or extracts thereof) have to be moved to the GPU, data transfers over the bus are relatively small. All of this makes CPU offloading of large Bloom filtering a very promising task. Our microbenchmarks show that raw Bloom filter lookups are up to 6× faster on the GPU than on the CPU in case the Bloom filter is larger than the CPU cache (skip to Figure 3).

The follow-up question is what a database architecture should look like that can efficiently offload tasks to the GPU. On the one hand, it has been shown that many-core CPUs can perfectly scale joins using shared hash-tables and an adaptive, fine-grained (“morsel driven”) task allocation [17]. On the other hand, for Bloom filtering, keys need to be transferred to the GPU on-the-fly; and these transfers have to be relatively coarse-grained in order to achieve good transfer speeds over the bus. Given the fact that bus transfer speeds are a bottleneck (an order of magnitude slower than memory access), this is a critical detail. We therefore propose the fluid co-processing framework, wherein (i) query pipelines are cut into dependent fragments that may run on a specific device or on both (ii) both CPU and GPU adaptively accept work from a shared queue and (iii) different granularities (amounts of tuples) are used for work-distribution: the GPU getting larger work units at-a-time than the CPU. Our early results show that fluid co-processing consistently improves end-to-end CPU performance of early pruning in join queries thanks to the GPU, by factors up to 2-3×.

2 FAST CPU AND GPU BLOOM FILTERS

The Bloom filter [2] represents a collection of \( n \) keys with an initially-zeroed array of \( m \) bits, setting for each inserted key \( k \) bits to 1, using as many hash functions to identify the positions \( \{0, m\} \) where the bits are set in the array. This structure allows for fast true-negative tests, but it can produce false-positives at some probability that drops with larger \( m \) and \( k \). In their classic implementation, multiple (up to \( k \)) random memory accesses are necessary to check the bits of a key set. A blocked Bloom filter [21] represents the data structure as a set of \( m/B \) blocks, each of which contains a small Bloom filter of size \( B \) – a hash function determines in which block a key must be stored. To optimize in-memory processing, the size of a block is equal to the size of a cache line, because this allows a key to be looked up with at most one cache miss. On GPUs, cache lines are typically 128 bytes and aligned in global memory: we therefore use up to \( B=1024 \) bits in our blocked GP Bloom Filters.

Recently, an experimental study was published on (CPU-based) Bloom filtering [15], which also introduced a number of innovations. One of these is the idea to use minuscule blocks that are as small as a 32-bit register, because this allows to test all \( k \) bits in one load-compare instruction sequence. That is, the lookup code first computes a bit mask with \( k \) bits set, loads one word and tests that against the cache line, because this allows a key to be looked up with at most one cache miss. On GPUs, cache lines are typically 128 bytes and aligned in global memory: we therefore use up to \( B=1024 \) bits in our blocked GPUs Bloom Filters.

Figure 2: Bloom filter variants

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(b) Blocked & Sectorized [15]

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Figure 3: Bloom filter probe throughput varying the bloom filter size. For larger Bloom filters, the throughput on GPU is up to $6 \times$ the CPU lookup performance. This allows efficient support of (a) larger inner relations (hence, bigger Bloom filters) and/or (b) more precise Bloom filters.

with an early-out if a bit is found to be 0. Getting rid of the loop and early-out branch is actually beneficial for the GPU, because control-divergence is avoided.

Register-blocked Bloom filters are quite radical as they maximally reduce computational work, but this goes at the expense of accuracy; because mini Bloom filters of just 32 bits can get unlucky and receive many keys, in which case their selective power deteriorates. In order to use larger blocks that occupy the full 1024 bits of a GPU cache-line (or 512-bits in case of the CPU), one can divide the block in multiple sectors, e.g. 32 sectors of 32-bits. The sectorized Bloom filter (see Figure 2) still tests multiple bits in each sector at once to reap the aforementioned computational benefits; but, it evenly spreads the $k$-bits over multiple sectors in the block. In fact, not all sectors need to receive bits; rather, a few of them are selected with yet another few random bits from the hash number. For instance with $k=8$, one could set/test 4 bits concentrated in two 32-bits words (=sectors) out of 32. This approach thus combines computational and memory efficiency, and can achieve high-precision and very low lookup latency [15].

Figure 3 shows micro-benchmarks of our C++ and CUDA implementation of Bloom filters on GPUs and CPUs ($k=2$, $n/m=8$) for varying Bloom filter sizes ($m$). The hardware is described in Section 4. For fairness, the CPU uses all its cores; and the GPU performance includes copying the keys (input) and the boolean results (output) resp. from and to CPU memory. In all, while the CPU is competitive with the GPU for smaller Bloom filters; when they start to exceed the CPU L3 cache (beyond 16 MiB), a large performance gap widens, up to a factor $6 \times$.

3 FLUID CO-PROCESSING

Commonly, co-processing approaches statically assign specific operations to devices. Notable examples are CoGaDB [5] and HAPE [7]. CoGaDB allows offloading specific operators to devices. HAPE relies on HetExchange [6], an extension of Volcano-model exchange operators [9] that allow offloading of complete pipelines. However, static resource allocation has two well-known caveats:

(a) It requires accurate cost-models to decide what to offload. In practice, building "good" cost-models is very hard and cost-models very often not precise [18].

(b) Optimal parallel performance requires load-balance, or available hardware will become under-utilized. Imbalance can have many causes, e.g. by interference (another process running, processors clocking up/down), sub-optimal data distributions or different hardware characteristics. However, once a static parallel plan is created, load-balancing is hard to achieve [10, 17].

3.1 Morsel-driven Parallelism is not enough

For CPU-only systems, morsel-driven parallelism [17] is known to eliminate, or at least mitigate, these issues. In a morsel-driven system, workers repeatedly consume tasks from a global queue and execute them. The task granularity, the morsel size, leads to the trade-off between scheduling overhead (removing tasks from a queue) and load-balancing. Small morsel sizes will lead to high scheduling overhead whereas large morsel sizes will severely worsen opportunities for load-balancing.

However, extending the traditional morsel-driven parallelism into a co-processing system is more complex:

(a) The optimal unit of work - the morsel size - differs from device to device. For parallel CPU-based systems Leis
et al. [17] suggest morsel sizes of roughly 10K – 100K. CPU-GPU communication latencies force 10x – 100x larger morsel sizes, to minimize scheduling overheads and achieve efficient data transfers. The absence of an efficient common morsel size prevents the naive approach of dividing the work into equally sized chunks and pushing them into a queue.

(b) The classical morsel-driven parallelism executes a query pipeline-by-pipeline. However, for co-processing, a pipeline can contain multiple pipeline fragments. When executing our join query on the GPU, we have two such fragments: The first fragment, scans the table and evaluates the Bloom filter on GPU. As we want to parallelize the following join operation, we need a second fragment which consumes morsels from the Bloom filter results and executes the join followed by the aggregation.

Because of these reasons, we concluded that plain morsel-driven parallelism is not "good" enough for Co-Processing and therefore we needed to extend it.

### 3.2 Fluid Parallelism for Co-processing

To allow multiple morsel sizes, we replaced the task queue (morsel queue) with a lock- and wait-free linear allocator. It works similar to the following pseudo-code:

```c
bool get_range(Range& out, const Table& t, int morsel_size) {
    offset = __sync_add_and_fetch(&t.current_offset, morsel_size);
    if (offset >= t.size()) return false; // failed getting range
    // use up to 'morsel_size' starting from offset
    out.offset = offset;
    out.num = MIN(t.size() - offset, morsel_size);
    return true;
}
```

The allocator returns a range which points into the table t. The linear range allocation relies on the atomic increment (__sync_add_and_fetch) to move the starting offset forward. If a valid offset was allocated, the actual size of the morsel is determined. In practice, all morsels, except the last one, will have the given morsel_size.

In contrast to morsel-driven parallelism’s one active pipeline, co-processing can work with multiple alternative pipelines. In our case study, we have (1) the whole CPU-only Bloom-filter + join-probe + aggregation pipeline that starts with CPU Bloom filtering, (2a) GPU Bloom filtering and (2b) CPU join-probe + aggregation. Tuples are either processed with (1) or by (2a) followed by (2b). Figure 4 illustrates our framework. Generally, we differentiate between GPU+CPU workers and CPU workers. Their main difference being that GPU+CPU workers can also schedule GPU work whereas CPU workers only operate on CPU-only pipeline fragments.

A CPU worker runs in a loop. In each iteration it will try to consume a morsel that either (a) evaluates the full pipeline for a whole CPU morsel or, if a GPU bloom filter probe is ready, (b) using the GPU bloom filter result disqualify non-matching tuples and execute the join. GPU+CPU workers extend CPU workers with the ability to also handle GPU work. They run a loop similar to the pseudo-code in Listing 1. To allow full utilization of the GPU, we prioritize GPU work over CPU work. Additionally, we allow the use of multiple streams. Each stream consists of copying to the GPU, executing the Bloom filter lookup kernel and copying the results back. Using multiple streams allows the GPU to efficiently overlap computation kernels and data movement.

If a free/idle GPU stream is detected, it will first try to consume a large GPU morsel from the table and schedule another GPU Bloom filter probe. This probing process will run asynchronously and is handled by the CUDA implementation. While the GPU+CPU worker waits for a probe to finish, it will execute regular CPU work. When a GPU Bloom filter lookup is finished, it will be schedule for work-sharing i.e. we add it to a global queue of finished probe from which CPU and GPU+CPU workers can consume morsels and execute the remaining pipeline fragment.

### 4 EXPERIMENTAL EVALUATION

We implemented the proposed techniques in a prototype which evaluates the query: SELECT SUM(a.k), SUM(b.p1), SUM(b.p2), ..., SUM(b.p32) FROM a, b WHERE a.k = b.k.

In detail, our prototype operates on cache-resident vectors in a columnar fashion, so called "vectorized execution" [3]. Its join operator executes a flat, non-partitioned, primary-key-foreign-key hash join (i.e. either zero or one match in

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1. Source code can be found under: https://github.com/t1mm3/1fluid-coprocessing
the hash table) on a bucket-chained hash table with records in row-wise layout [22]. Additionally, we simulated an expensive operator (P in Figure 1) between scan and join. In our experiments, we use a Bloom filter with one sector and set \( k = 2 \) bits. We used a probe relation with a cardinality of 1 billion tuples. However, when scaling the inner relation we increased the outer relation to 4 billion tuples. Our co-processing framework uses 4 GPU streams and morsel sizes of 16 Ki tuples for CPU and 1 Mi tuples for GPU. Our experiments were conducted on a 10-core i9-7900X with 13.75 MiB L3 cache and 32 GiB main-memory. We always use all 10 cores. As GPU, we used one GeForce GTX 1080 with 8 GiB memory. We used Fedora 28 with CUDA 10.1 and driver version 418.56.

In this section, we first evaluate the performance of our fluid co-processing framework. Afterwards, we investigate the influence of a varying pipeline cost \( c_P \) and cardinality of the inner relation on the query time, followed by a discussion of the influence of the number of streams on the performance. Last but not least, we discuss the scheduling behaviour of our fluid co-processing framework.

### 4.1 Fluid Co-Processed Join

The effectiveness of early pruning depends on the selectivity of the join itself. The more tuples eliminated by the join, the more effectively can a Bloom filter prune them already in the scan. Therefore, we evaluated the overall performance of the join (probe) pipeline over varying selectivity. Figure 5 visualizes our results.

We noticed that, during CPU-only execution (CPU, BF) the benefit of using the Bloom filter to prune is limited, compared to CPU-only execution without a Bloom filter (CPU, no BF). This is due to the size of the Bloom filter. As the Bloom filter exceeds the CPU caches, it introduces additional cache misses and put further stress on the memory subsystem. With rising selectivity (less tuples pruned) the benefit of early filtering goes to zero and only the filtering overhead, additional cache misses in particular, remains.

The GPU+CPU execution (GPU+CPU, BF), however, benefits from the GPU’s additional computational power and memory bandwidth. For low selectivities this leads to a performance improvement up to \( 3 \times \) compared to the non-Bloom CPU implementation and roughly \( 2 \times \) compared to the CPU-only version. With rising selectivity, similar to the CPU-only scenario, the benefit of early pruning gradually disappears. However, the risk of slowing down the join is much lower because the Bloom filter is much faster i.e. it uses CPU and GPU in parallel.

We now focus on the performance of the join. Figure 6 displays the join time with the probe key column already residing in the GPU memory (GPU+CPU, BF (cached)). Caching the keys speeds up the GPU pipeline (Bloom filter and sending results back) which, in turn allows more tuples to be processed on GPU. This leads to a significant speedup of up to 27% which additionally boost the already existing performance gain. However, caching the keys only allows the GPU to consume \( \approx 5\% \) more tuples (Figure 8) and is therefore not explaining the 27% boost (the lion’s share, 95%, did not change). This indicates that the memory transfer from CPU to GPU (HtoD) and back (DtoH) is not free and affects the memory bandwidth of the host (CPU).

### 4.2 Scaling Pipeline Cost and Inner Relation

In joins, the efficiency of early pruning using Bloom depends on a variety of factors: (a) the size of the inner (build) relation, (b) the size of the Bloom filter, (c) the number of hash functions \( k \), (d) the Bloom filter type (naive, blocked, counting etc.) and (e) cost of the pipeline \( c_P \).
Table 2: Performance-optimal Bloom filter false-positive ratio (accuracy) for combinations of inner relation cardinality and additional pipeline cost $c_A$ with Bloom filter size $m$ (bits) and $k$ hash functions. GPU Bloom filters efficiently support larger filters with more hash functions ($k$ up to 14-15) than CPU Bloom filters.

(a) CPU  

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<th>Additional Pipeline Cost $c_A$</th>
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<th>100</th>
<th>200</th>
<th>400</th>
<th>800</th>
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(b) GPU  

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<th>Inner Relation Size</th>
<th>Additional Pipeline Cost $c_A$</th>
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Figure 7: Influence of inner relation’s cardinality and pipeline cost $c_p$ on query time at 1% selectivity. Colors represent the total runtime (s). Variations in $c_p$ are simulated through an additional pipeline cost $c_A$. Bloom filters on GPU allow to accelerate the upper half (large inner relations) and are able to speed up expensive pipelines (high $c_p$).

In this experiment, we demonstrate the effects of the pipeline cost $c_p$ and inner relation size on the run time. For each combination of CPU/GPU, inner relation and $c_p$, we determined the performance-optimal Bloom filter, as defined by Lang et al. [15], using a hardware-calibrated cost-model. Note that a Bloom filter might not always provide a benefit. In particular, a combination of a small $c_p$ and an expensive Bloom filter might lead to inferior performance. We marked such cases as invalid values.

Bloom Filter Configurations & Accuracy. Using the cost-model, we obtained performance-optimal Bloom filter configurations for a variety of inner relational cardinalities (i.e., join build size $n$) and pipeline cost. For the latter, we increase the basic scan-filter-join cost $c_p$ by adding an additional cost $c_A$ that corresponds to operators (such as aggregations or other joins) that would be in between the scan-filter and the join. The Bloom filter size $m$ and the number of hash functions $k$ for each filter is shown in Table 1. We noticed that, compared to the CPU, the GPU efficiently allows larger Bloom filters and roughly 2x more hash functions. On the CPU, such configurations would suffer from low memory throughput and high computational cost for the hashing.

The false-positive ratio $f$ of each filter is depicted in Table 2. With increasing pipeline cost, higher $c_A$, we noticed a trend to higher accuracy. The reason is that, with a high penalty for a false-positive, it makes more sense to pay the
price of a more expensive Bloom filter with higher precision to prevent many false-positives.

Compared to CPU Bloom filters, GPU Bloom filters tend to have a very low false-positive ratio because (a) the GPU has a higher memory bandwidth (less penalty for larger filters) and (b) more computational power - hence a GPU can compute more hash functions than a CPU, before becoming compute-bound rather than memory-bound. A configuration with size \( m=4GB \) and \( k=14 \) hash functions is not practical on a CPU, but can be performance-optimal on a GPU.

**Bloom Filter Lookup Performance.** After determining the optimal Bloom filters, we ran the experiment. Figure 7 visualizes our results. The lookup performance of CPU-only Bloom filters, as visible in Figure 7a, is more sensitive to the size of the build relation i.e. fast lookup is only achieved for Bloom filters that fit into cache. Consequently, the benefits of early pruning decreases with the Bloom filter size, as the lookup becomes more expensive with increasing build sizes.

In contrast to CPU-only filtering, the GPU+CPU Bloom filtering (see Figure 7b) is much less sensitive to the build size, and consequently the Bloom filter. This allows efficient pruning in selective joins even for large dimension tables (i.e. larger inner relations).

Higher pipeline costs \( (c_p+c_A) \) tend to justify more accurate Bloom filters. The early pruning becomes more costly in more accurate filters, but if offset by the much more expensive pipeline cost, that can be saved for every pruned tuple. We notice again that the overhead of more accurate Bloom filters on GPU+CPU is much lower than on CPU and, hence, allows much more effective pruning.

### 4.3 Optimal Number of Streams

In our fluid co-processing, the term *stream* refers to a work queue, where the CPU enqueues tasks for the GPU, such as kernel executions and memory transfers from the host to the device (HtoD) or from the device back to the hosts memory (DtoH). Modern GPUs are capable of performing two simultaneous (DMA) memory transfers, one in each direction. At the same time the GPU can execute kernels. Due to the fact that the streams are processed sequentially, a co-processing system needs multiple concurrent streams, associated with the same GPU, to overlap memory transfers and kernel execution and thus to maximize performance. In our query, each stream delivers tuples to the GPU, probes the Bloom filter, and sends the resulting bitmap back to the host memory. We aim to find the lowest number of streams which (a) keeps the GPU busy and (b) leads to the lowest run time. Therefore, we executed our co-processed join query with a selectivity of 1% on 1 billion tuples and measured the fraction of tuples sent to the GPU (*GPU Utilization*), as well as the runtime of the probe pipeline.

(a) ... when transferring keys \hspace{1cm} (b) ... when keys are cached

Figure 8: More than 4 streams do not provide a better GPU utilization.

(a) ... when transferring keys \hspace{1cm} (b) ... when keys are cached

Figure 9: Influence of more than 2 streams onto total query runtime is negligible.

**GPU Utilization.** The results are visualized in Figure 8a. We noticed that GPU utilization heavily depends on the number of streams. For only one stream, we can only filter up to \( \approx 80\% \) of the tuples on the faster device (the GPU) and, hence, leave resources idle. With lower selectivities this effect is more extreme. However, with more than 2 streams we noticed that almost all (\( \approx 98\% \)) are being filtered by the GPU. When using 4 or 8 streams, we observed a better behaviour for lower selectivities (\( \leq 5\% \)). However, for higher selectivities we measured a degraded utilization. This indicates that, besides memory footprint, there is also CUDA overhead involved. We also noticed that using more streams are not necessarily the better choice for GPU utilization, as compared to using 4 streams, 8 streams lead to inferior GPU utilization. We argue that, in our case, 4 streams appears to be the optimal choice, as it provides high GPU utilization for low selectivities, as well as, negligible degradation for high selectivities.

When the keys are cached on the GPU, see Figure 8b, we noticed similar behaviour. However, as the keys do not need to be transferred to the GPU (i.e. GPU kernels are faster), we measured slightly higher GPU utilization.

**Runtime.** In Figure 9 we plotted the runtime of the probe pipeline with a given number of streams. For 1 stream, we observed a penalty because the GPU is not fully utilized. For 2 or more streams, we did not notice any significantly
4.4 Optimal Morsel Sizes

The performance of our fluid co-processing strongly depends on the morsel sizes used for CPU and GPU work. The choice of morsel sizes is typically a trade-off between load-balancing behaviour and scheduling overhead as small morsels lead to good load balance between workers but higher scheduling overhead, and vice versa. Hence, the optimal morsel sizes is the smallest one which still allows efficient execution. We, therefore, experiment with various combinations of morsel sizes for CPU and GPU. Our results can be seen in Figure 10.

We noticed that too small CPU morsels increase total runtime as they incur higher scheduling overhead. Too large CPU morsels also increase the runtime as they leave resources idle (most workers are done whereas the last ones are still processing their large morsels). Similarly we noticed a tendency that small GPU morsels incur additional scheduling and GPU-specific overhead, i.e. GPU-internal scheduling and less efficient data transfers lead to lower throughput. In the other extreme large GPU morsels fully utilize GPU resources but leave CPU resources idle.

We found that 16 Ki tuples for CPU and 1 Mi tuples for GPU are sufficient to allow efficient processing and are, therefore, the optimal choice in our case.

Figure 11: Timeline with 1 GPU+CPU worker. GPU+CPU worker is mostly busy issuing GPU work. Depicted are 6 out of 10 workers. Thickness of the bars Post GPU Join and CPU Bloom + Join represents their processing speed (higher means faster).
Our fluid co-processing schedules tasks dynamically. To give an insight into the scheduling decisions being made, we ran the same experiment as in Section 4.1, profiled scheduling decisions and plotted onto a timeline. First, we discuss the behaviour of our prototype with only 1 GPU+CPU worker and, afterwards, the behaviour with 2 GPU+CPU workers.

1 GPU+CPU Worker. We ran the experiment for three different selectivities (1%, 5% and 10%). Figure 11 visualizes the results we obtained. We noticed that the 1 GPU+CPU was mostly busy issuing GPU work (in the following experiment we parallelize this part using 2 GPU+CPU workers).

At 1% selectivity (Figure 11a), we observed that most time is spent on the CPU Bloom + Join. The reason is that the GPU-accelerated pipeline (Post GPU Join) runs in negligible time as it eliminates 99% of tuples in a very lightweight fashion. The remaining time each worker is trying to prevent going idle by "stealing" CPU morsels and executing the CPU-only pipeline.

When moving to 5% selectivity (Figure 11b), we found that there is roughly an equal amount of time spent on the GPU-accelerated and the CPU-only pipeline.

Moving to 10% selectivity (Figure 11c), we observe the opposite of the 1% case: Most time is spent on the GPU-accelerated Join (Post GPU Join) because the join, including fetching payloads, is very expensive. However, we also noticed a startup lag for the GPU-accelerated Join. This is caused by the initial latency of issuing GPU work until the results from the GPU are received.

2 GPU+CPU Workers. In the previous experiment, it appeared that the one single GPU+CPU worker might be overloaded. Therefore, we modified the experiment and parallelize issuing GPU work over 2 GPU+CPU workers which can concurrently schedule GPU work. The results can be seen in Figure 12. We noticed that allowing multiple GPU+CPU workers tends to worsen initial load-balance (at the start of the query). The cause of this behaviour is the preference of GPU+CPU workers to issue GPU work. When creating GPU+CPU workers, each will first schedule GPU work which involves locking inside the CUDA library. However, we want to highlight, that our fluid co-processing framework is still able to achieve load-balance and, hence, the negative influence of such delayed starts, or other interference, is significantly reduced.

5 RELATED WORK
Co-processing can be seen as an intra-query parallelization problem where different devices work together. Many database systems use Volcano parallelism [9] to adapt to modern multi-core hardware. Notable examples are SQLServer [16], Oracle [14] and Vectorwise [23]. Recently, it found application in Co-processing, as HetExchange [6] which allows to offload whole pipelines. Volcano originally became popular because it allowed an easy integration of parallelism into existing single-threaded systems by abstracting the parallelization into Exchange operators. This also means that parallelization and partitioning decisions have to be made at query optimization time rather than dynamically at query runtime. This, consequently, creates imbalance between threads and leaves resources idle. These issues have been fixed through dynamic morsel-driven parallelism [17]. It splits table ranges into tasks. These tasks are inserted into a global queue from which multiple workers can consume and process this task. Recently, a dynamic virtual machine architecture was proposed to allow cross-device load-balancing [10]. This idea inspired our fluid co-processing framework to perform adaptive work placements for different devices, but instead of placing the work on the unit, here we adapt the granularity of work per device.

A number of GPU co-processing database systems have been developed, most notably Ocelot [12], CoGaDB [5] and HAPE [7]. Ocelot was proposed as a hardware-oblivious database engine to abstract away operators from CPU and GPU devices [12]. CoGaDB [5] places operators on devices at runtime, and allows pipeline parallelism. However, compared to our Fluid Co-processing, CoGaDB's operator placement is neither fully dynamic, nor elastic. HAPE [7] is based on HetExchange [6]. Entire pipelines are compiled for different architectures (CPUs and GPUs). Thereby, CPUs and GPUs perform the same tasks and Exchange operators "route" the
tuples between the executing devices. In contrast, our prototype offloads different tasks (i.e., Bloom filter probe) to a different device, that eventually assists query execution on the CPU which leads to fully dynamic partitioning.

Our case study uses Bloom filters to accelerate selective joins. Bloom filters were introduced by Bloom [2]. Similar data-structures for approximate membership queries have been discovered. Notable examples are Quotient filters [1], Cuckoo filter [8] and Morton filters [4]. However their performance on GPU has not been studied deeply.

6 CONCLUSIONS & FUTURE WORK
This work (finally) identified in Early Pruning of Selective Joins using Bloom filters a relevant database workload where CPU database systems can be significantly accelerated by GPU co-processing. We realized this in a prototype of the dynamic, adaptive, heterogeneous Fluid co-processing framework. This framework splits pipelines into pipeline fragments, such that a pipeline can be co-processed on different devices. It also allows multiple equivalent pipeline fragments to co-exist, so certain tuple ranges use a CPU-only pipeline, whereas others follow a CPU-GPU pipeline consisting of multiple fragments. Its shared work queue is inspired by the elastic and dynamic co-processing model based of morsel-driven parallelism [17], but allows for different morsel granularities, tailored to each device. Note that the queue-based architecture also helps solve the problem of handling concurrent queries on a GPU, as the queue can coordinate scheduling of morsels belonging to different queries; and if the GPU is too busy, queries can just execute their morsels mostly (or completely) on the CPU-pipeline.

To measure the effectiveness of our techniques, we implemented a simple multi-threaded join query. Our experiments show performance improvements of up to 3× by dynamically offloading Bloom filtering.

In the future, we plan to deeply investigate the design space of GPU-based Bloom filtering; as our work so far just increases the Bloom filter block size to the correct GPU cache line size. We are experimenting with multiple GPU-specific Bloom filtering optimizations as well.

As for the Fluid Co-Processing framework, we intend to mature our prototype into a system that can execute generic and concurrent queries and perform adaptive offloading to heterogeneous hardware [11].

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