Reducing overheads of dynamic scheduling on heterogeneous chips

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ABSTRACT
In recent processor development, we have witnessed the integration of GPU and CPUs into a single chip. The result of this integration is a reduction of the data communication overheads. This enables an efficient collaboration of both devices in the execution of parallel workloads.

In this work, we focus on the problem of efficiently scheduling chunks of iterations of parallel loops among the computing devices on the chip (the GPU and the CPU cores) in the context of irregular applications. In particular, we analyze the sources of overhead that the host thread experiences when a chunk of iterations is offloaded to the GPU while other threads are executing concurrently other chunks on the CPU cores. We carefully study these overheads on different processor architectures and operating systems using Barnes Hut as a study case representative of irregular applications. We also propose a set of optimizations to mitigate the overheads that arise in presence of oversubscription and take advantage of the different features of the heterogeneous architectures. Thanks to these optimizations we reduce Energy-Delay Product (EDP) by 18% and 84% on Intel Ivy Bridge and Haswell architectures, respectively, and by 57% on the Exynos big.LITTLE.

1. INTRODUCTION
Recently, we have seen a trend towards the integration of GPU and CPUs on the same die. Examples include recent Intel processors (Ivy Bridge, Haswell), the AMD APUs, or more power constrained processors targeted at mobile and embedded devices, like the Samsung Exynos 5 or the Qualcomm Snapdragon 800, among others. The integration allows the sharing of the memory system, what reduces the communication overheads between the devices and enables a more effective cooperation among all the computing devices. In contrast to discrete GPUs (connected through a slower PCI bus), integrated GPUs also enable the implementation of more dynamic strategies to distribute the workload among the cores and the GPU. In fact, achieving maximum performance and/or minimum energy consumption often requires simultaneous use of both, the GPU and the CPU cores.

In this context, one problem that has received attention lately is the efficient execution of the iterations of a parallel loop on both devices, the CPU cores and the integrated GPU. However, the optimal division of work between CPU and GPU is very application and input data dependent so it is required a careful partitioning of the workload across the CPU cores and the GPU accelerator. In the case of regular applications, the main difficulty is to determine how to partition the load between the two devices to avoid load imbalance. This is usually accomplished by running a few chunks of iterations in both devices to determine the speed difference between them. The appropriate chunk for each device is then computed and scheduled to run. In the case of irregular applications the challenge is bigger because, in this case, GPU performance can be suboptimal when the application workload is distributed among all the cores and the GPU without considering the size of the chunk assigned to the GPU, even when the load is balanced among the devices. For instance, using the irregular Barnes Hut benchmark as our case of study, we have found that the size of the chunk of iterations assigned to the GPU has a significant impact on its performance, as we will show in Section 2. Thus, for each application and input data, the scheduling strategy must be aware of the optimal chunk of iterations to offload to the GPU. Since the workload assigned to the GPU and the CPU cores must also be balanced, the best approach is a dynamic scheduling strategy that assigns the optimal chunk of iterations to each device. This means that the cores and the GPU will repeatedly receive a chunk of iterations to be computed until the end of the iteration space. This scheduling strategy is described in Section 3.

The intensive use of the GPU offloading mechanism will reveal several sources of overheads that have to be carefully considered. For instance, not only the traditionally studied overheads like the data transfers ones (host-to-device and device-to-host) have to be taken into account, but also kernel launching and host thread dispatch overheads gain relevance. In this paper, we study the impact of each one of these overheads on different heterogenous architectures with an integrated GPU. More precisely, in Section 4, we conduct our experiments on two Intel processors (Ivy Bridge...
2. MOTIVATION

We have found that, for irregular applications, offloading big chunk sizes to the GPU can hinder performance. This is illustrated in Figure 1 that shows the evolution of different GPU hardware metrics on the Haswell architecture (described in section 4.1) through the iteration space of one time step of the irregular benchmark Barnes Hut. An input set of 100,000 bodies was used to collect these results. Each subfigure represents the evolution of the metric of interest for different chunk sizes assigned to the GPU (see chunk sizes legend in subfigure 1c). We have used Intel VTune Amplifier 2015 [1] to trace the ratio of cycles when EU (Execution Units) are active (∑all EU all cycles) and EU idle (∑all EU all cycles); the ratio of cycles when EU are idle (∑all EU all cycles); and the L3 cache misses due to GPU memory requests. Subfigure 1c also shows the GPU effective throughput, measured as the number of iterations per ms., through the iteration space. Data transfer and kernel offloading overheads have been included in the computation of the throughput.

As we can see in subfigure 1e for the time step studied, the optimal chunk size is 640 (see red line in the figure). Increasing the chunk size beyond this value degrades the throughput. The hardware metrics indicate that small chunk sizes (e.g. 320) do not effectively fill the GPU computing units, as the ratio EU Idle indicates in subfigure 1b (see blue line). However, when the chunk size is large enough to fill the EUs (EU Idle <0.1), the EUs might stall when the chunk size increases, due to the increment in L3 cache misses (see green and pink lines in subfigure 1d). This is what happens in our irregular benchmark in which the majority of memory accesses are uncoalesced. In our case, chunk sizes larger than 1280 dramatically increase L3 misses, which in turn increases the ratio of EU Stalled (>0.9) and reduces the ratio of EU Active (<0.08), causing a reduction of the effective throughput.

Therefore, in the quest of finding the optimal distribution of work between the GPU and the CPU, if we assign a big chunk of iterations to each device we can end up by not exploiting the GPU EUs optimally. Thus, the partitioning strategy must be aware of the optimal chunk of iterations that must be offloaded to the GPU for the corresponding application. On the other hand, we have observed that the effective throughput of the CPU cores is not so sensible to the chunk size. As long as the chunk size is bigger than a threshold value, the CPU throughput tends to be constant.
For instance, when using the Threading Building Blocks library (TBB) [15], it is recommended to have CPU chunk sizes that take 100,000 clock cycles at least. By allowing a dynamic scheduling of the iteration space in such a way that each device gets an optimal chunk of iterations while balancing the workload among them, we can minimize the execution time. However, some overheads are involved in this type of scheduling, as we will see in the next sections. Anyway, we explore here the applicability of a dynamic scheduling strategy on a heterogeneous architecture using Barnes Hut as example.

In Figure 2 we compare the time, energy and EDP (Energy-Delay-Product) [6] for Barnes Hut on three different heterogeneous architectures using 4 CPU cores and 1 integrated GPU: Ivy Bridge, Haswell and Exynos. The input set also has 100,000 bodies, but now 75 time-steps were computed. For each platform, two scenarios are shown: 3+1, which corresponds to the case in which 4 threads are scheduled in the processor (3 CPU threads + 1 host thread dedicated to offload the work to the GPU) and 4+1 which corresponds to 5 threads (4 CPU threads + 1 host thread). The later represents the case in which 1 thread of oversubscription is allowed. We explore this case because we want to make the most of our computing resources: since the host thread is most of the time blocked while the GPU takes care of its chunk of iterations, we add an additional thread to avoid one core becoming idle.

In the figure, we call Dynamic to our dynamic scheduling strategy (it works similarly to the OpenMP dynamic scheduling policy of the pragma omp parallel for). In this strategy, first we have to find the optimal chunk size for the GPU (the GPU chunk). This is done through an offline training phase, where we explore different chunk sizes, and choose the value that maximizes the effective throughput of the application in the corresponding GPU. Table 1 shows the optimal size for each platform. This size is passed to our scheduler which dynamically assigns a new chunk to the GPU each time it finishes the computation. Similarly, each CPU core gets a new chunk of iterations (the CPU chunk) every time it finishes the previous one. The size of the CPU chunk is selected to balance the load with the GPU computation (details are provided in the next section).

We compare this dynamic strategy with a static scheduling that assigns a single GPU chunk to the GPU and the rest of the iterations to the CPU cores. Previously, for the static scheduling we carry out an exhaustive offline profiling that looks for the static partitioning of the iteration space between the CPU and GPU that minimizes the execution time. We vary the percentage of the iteration space offloaded to the GPU (between 0% -only CPU execution- and 100% -only GPU execution- using 10% steps). We call this strategy Bulk-Oracle. Again, Table 1 shows the optimal percentage of iterations offloaded to the GPU for each platform under the static partition strategy. Notice that with the static approach the whole chunk is offloaded to the GPU at the beginning of each time step.

Table 1: Optimal GPU chunk size for Dynamic and optimal percentage of the iteration space offloaded to the GPU for Bulk-Oracle

| Platform   | Dynamic | Bulk-Oracle |
|------------|---------|-------------|
| Ivy        | 3+1     | 4+1         |
| Haswell    | 1536    | 2048        |
| Exynos     | 50%     | 20%         |

In Figure 2 each parameter (time, energy and EDP) has been normalized with respect to the value obtained for the Bulk-Oracle 3+1 execution on each architecture. As we see, the dynamic strategy outperforms the static one (Bulk-Oracle), except in the case of Haswell for 4+1, where overheads associated to the host thread degrade performance (both in time and energy). This issue is discussed in the next section.

Another interesting result is that oversubscription (4+1) improves the execution times on Ivy, for both the static and dynamic approaches, but it also increases the energy consumption. This is due to the fact that 4 threads compute a larger number of chunks on the CPU cores and, since the CPU is less energetically efficient computing the chunks than the GPU, this results in higher energy consumption. For this architecture and this benchmark, the increment of the energy on the CPU is not amortized by the reduction of time. On the other hand, on Exynos, a static partitioning does not scale when going from 3+1 to 4+1, while the dynamic strategy reduces time and energy consumption.

3. SCHEDULING STRATEGY

In this section, we present in more detail our scheduler (Section 3.1), how the partitioner works (Section 3.2), and the
#include <BScheduler.h>

class Body {
public:
  void operatorCPU(int begin, int end) {
    for (i = begin; i != end; i++) {...}
  }
  void operatorGPU(int begin, int end) {
    hostToDevice(begin, end)...
  }
  void opDeviceToHost(int begin, int end)...
  clFinish();
  deviceToHost(begin, end) {...}
}

Figure 3: Using the parallel_for template

1. potential sources of overhead (Section 3.3).

3.1 Scheduler description

Our scheduler, that we call Dynamic, considers loops with independent iterations (parallel_for) and features a work scheduling policy with a dynamic GPU and CPU chunk partitioning. Our approach dynamically partitions the whole iteration space into chunks or blocks of iterations. The goal of the partitioning strategy is to evenly balance the workload of the loop among the compute resources (GPU and CPU cores) as well as to assign to each device the chunk size that maximizes its throughput. This is key because, as shown in Figure 1, the chunk size can have a significant impact on the performance of heterogeneous architectures, especially when dealing with irregular codes.

Our scheduler builds on top of an extension of the TBB parallel_for template for heterogeneous GPU-CPU systems by Navarro et al. 13. Figure 3 shows the pseudo-code to use the extended parallel_for construct in such heterogeneous systems. As in any TBB program, the scheduler is initialized (line 20). In this step, the developer sets the number of OS threads, nThreads, that the TBB runtime will create, which can vary from 1 to the number of CPU cores plus one additional thread to host the GPU (the host thread). Then, the developer can invoke the parallel_for (line 22), which has the following parameters: the iteration space (the range begin, end), the body object of the loop (body), and the partitioner object (Partitioner_H). The latter argument, effectively overloads the native TBB parallel_for function so that the heterogeneous version is invoked. Besides, the Partitioner_H method takes care of the dynamic partitioning strategy that gets the optimal chunk size for the GPU (parameter G provided by the user) and for the CPU cores as described in Section 3.2.

The user also writes the class Body that processes the chunk on the CPU cores or on the GPU (lines 2 to 13 in Figure 3). Two methods (operators) must be coded. One for the CPU (lines 4 to 6) and one for the GPU (lines 7 to 9). For the GPU, the user has to define two functions to perform the asynchronous host-to-device (line 8) and device-to-host (line 10) memory transfers, as well as the kernel launching (line 11). Since these functions are all asynchronous, we finish the GPU part of the body with the synchronous clFinish() call that does not return until all the previous steps have been completed.

Internally, our scheduler is implemented as a pipeline that consists of two stages or filters: Filter1, which selects the computing device and the chunk size (number of iterations) assigned, and Filter2, which processes the chunk on the corresponding device. Filter2 firstly checks if the GPU device is available. In that case, a G_token is created and initialized with the range of the GPU chunk. If there is no idle GPU device, then a CPU core is idle; thus, a C_token is created and initialized with the range of the CPU chunk. In both cases, the partitioner extracts a chunk of iterations from the range of the remaining iteration space. Next, Filter2 processes the chunk in the corresponding device and records the time it takes to compute the corresponding chunk. This is necessary to compute the device’s throughput, which is used by the partitioner described next.

3.2 Partitioning strategy

We assume that the execution time can be seen as a sequence of scheduling intervals \{tG0, tG1, ..., tGi-1, tGi, tGi+1, ...\} for the GPU and \{tC0, tC1, ..., tCi-1, tCi, tCi+1, ...\} for each CPU core. Each computing device at the current interval, tGi or tCi, can get a chunk of iterations. The running time T(tGi), for each GPU’s chunk size G(tGi) = G, or the time T(tCi) for a CPU’s chunk size C(tCi), is recorded. This time is used to compute the throughput, \(\lambda_G(tGi)\) for the GPU or \(\lambda_C(tCi)\) for a CPU core, in the current scheduling interval as:

\[
\lambda_G(tGi) = \frac{G}{T(tGi)}
\]

\[
\lambda_C(tCi) = \frac{C(tCi)}{T(tCi)}
\]

In order to compute the chunk size for the GPU, an offline training phase explores different chunk sizes, and chooses the value that maximizes the effective throughput of the application for a given input data. To reduce the number of runs of this offline training phase, we set the GPU chunk size to the smallest number of iterations that fully occupy the GPU resources. For example, on the integrated GPU of the Intel Haswell we have 20 EU (execution units) each one running a SIMD-thread (aka waveform) at a time, and 8, 16 or 32 work-items per SIMD-thread (decided at compile time). In OpenCL these values can be queried reading two variables.\(^1\) Thus, the product of these two arguments is used as the initial GPU chunk size. For Barnes Hut the compiler chooses 16 SIMD and therefore we select an initial GPU chunk size of 20 \times 16 = 320 iterations. Then, chunk sizes that are multiple of this initial chunk size are tried.

\(^1\)Note that for the GPU, the time includes the kernel execution time as well as the hostToDevice and deviceToHost times.

\(^2\)CL_DEVICE_MAX_COMPUTE_UNITS, and CL_KERNEL_PREFERRED_WORK_GROUP_SIZE_MULTIPLE
We keep trying different chunk sizes while the throughput increases. Once the throughput decreases or remains stable for 2 or more chunk sizes, the training phase ends and the GPU chunk size that obtained the highest throughput is chosen. This approach requires only a few runs.

Regarding the policy to set the CPU core chunk size, our partitioner follows the heuristic by Navarro et al. [13]. Basically, this heuristic is based on a strategy to minimize the load imbalance among the CPU cores and the GPU, while maximizing the throughput of the system. To that end, the optimization model described there recommends that: each time that a chunk is partitioned and assigned to a device, its size should be selected such that it is proportional to the device’s effective throughput. Therefore, we implemented a greedy partitioning algorithm based on the following key observation: while there are sufficient remaining iterations, the chunk size assigned to a GPU at the scheduling interval \( t_G_i \) should be the optimal for the GPU \( (G) \), as explained in the previous paragraph), whereas at the scheduling interval \( t_C_i \) the chunk size assigned to a CPU core, \( C(tC_i) \), should verify:

\[
\frac{C(tC_i)}{\lambda_C(tC_{i-1})} = \frac{G}{\lambda_G(tG_{i-1})}
\]

where \( \lambda_C(tC_{i-1}) \) and \( \lambda_G(tG_{i-1}) \) are the CPU and GPU throughputs in the previous scheduling intervals, respectively. So we have:

\[
C(t_i) = G \cdot \frac{\lambda_C(tC_{i-1})}{\lambda_G(tG_{i-1})}
\]

### 3.3 Sources of overhead

Figure 4 shows the different phases that our Dynamic framework has to perform to offload a chunk of iterations to the GPU.

As we explained in the previous section, our framework uses nThreads, from which one of them is called the host thread and just takes care of serving the GPU. This host thread runs in one of the available cores and first executes the code associated with the scheduler and the partitioner (the code of Filter1 explained earlier). Then, the same host thread executes the Filter2 stage that includes the function calls listed in Figure 5 (hostToDevice(), launchKernel(), deviceToHost, and clFinish()). The first three calls just asynchronously enqueue the corresponding operation on the GPU’s command queue, whereas the latter is a synchronous wait. In Figure 6 we can see that the enqueued operations are sequentially executed on the GPU where we consider the times taken by the “Host-to-Device”, “Kernel launching”, “Kernel execution” and “Device-to-Host”. When this last operation is done, the host thread is notified but some time may be taken by the OS to re-schedule the host thread. This time is illustrated in the figure with the label “Thread dispatch”.

In order to measure the relevant overheads involved in the execution of the code some time stamps are taken on the CPU (Tc1, Tc2 and Tc3) and on the GPU (Tg1 to Tg5) as depicted in Figure 5. To get the CPU time stamps we rely on TBB’s tick_count class, whereas for the GPU we configure the OpenCL command queue in the profile mode so that we can read the “start” and “complete” time stamps of each of the enqueued commands.

With this information we compute the overhead of Scheduling and Partitioning, \( O_{sp} \), Host-to-Device operation, \( O_{hd} \), Kernel Launching, \( O_{kl} \), Device-to-Host, \( O_{dh} \), and Thread Dispatch, \( O_{td} \), as follows:

\[
O_{sp} = \frac{\sum_{\#GPUChunks} (Tc2 - Tc1)}{\text{TotalExecutionTime}}
\]

\[
O_{hd} = \frac{\sum_{\#GPUChunks} (Tg2 - Tg1)}{\text{TotalExecutionTime}}
\]

\[
O_{kl} = \frac{\sum_{\#GPUChunks} (Tg3 - Tg2)}{\text{TotalExecutionTime}}
\]

\[
O_{dh} = \frac{\sum_{\#GPUChunks} (Tg5 - Tg4)}{\text{TotalExecutionTime}}
\]

\[
O_{td} = \frac{\sum_{\#GPUChunks} (Tc3 - Tc2) - (Tg5 - Tg1)}{\text{TotalExecutionTime}}
\]

### 4. ANALYSIS OF OVERHEADS

In this section we analyze the sub-optimal performance of the Dynamic scheduler for the 4+1 configuration, as shown in Figure 2. To better understand the underlying reasons for that performance degration we first describe our environment setup (Section 4.1) and then discuss the overheads that we measure (Section 4.2).

#### 4.1 Experimental Settings

We run our experiments on three different platforms: two Intel-based desktops and an Odroid XU3 bare-board. More
It is possible to monitor energy (in Joules) consumed by the Cortex A15 cores, the Cortex A7 cores, DRAM and GPU, separately. The energy figures on the Exynos 5 we present in this paper are the sum of the four power monitors aforementioned.

The sample rate used is 10 Hz. Thus, one sampled power value is obtained every 100 milliseconds. The power value read is multiplied by the sampling period and the product is integrated in a cumulated energy value (in Joules). We have chosen this sample rate because the power sensors actualize their values every 260 milliseconds approximately, so a sample rate two times faster is good enough for getting accurate measurements (sampling rate is below Nyquist rate).

### 4.2 Discussion

Figure 5 shows the overheads that we measured for the corresponding terms defined in the previous section (eqs. 6-9) for the three platforms that we consider, Ivy Bridge, Haswell and Exynos when running the Barnes Hut benchmark with 100,000 bodies and 15 time steps for the 3+1 and 4+1 scenarios. This section covers only the Dynamic results (shown on the left side) of the three subfigures in Figure 5.

The smallest overhead, on average, is the one due to Scheduling and Partitioning, $O_{sp}$, that represents 0.02% on Ivy Bridge, and less than 0.004% on Haswell and Exynos.

Regarding the data transfer overheads, $O_{hd}$ and $O_{dh}$, on Ivy Bridge and Haswell, they are always below 0.3%. However, on Exynos, these overheads are significantly larger, around an order of magnitude higher: on average $O_{hd} = 2.8\%$ and $O_{dh} = 1.6\%$. After testing our platforms with memory bound micro-benchmarks, we found that Exynos exhibits an order of magnitude higher data transfers times than the Intel architectures. This explains the impact of Host-To-Device and Device-To-Host overheads on the Exynos. More precisely, the `hostToDevice()` operation implicitly copies the host buffer onto a different region of memory that can be accessed by the GPU and that is non-pageable (pinned memory). Similarly, `deviceToHost()` does another memory-to-memory copy operation in the other direction. Therefore, lower memory bandwidth on the Exynos results in more appreciable Host-To-Device and Device-To-Host overheads with respect to the Intel architectures. As future work we will study how Barnes Hut memory accesses could be reorganized so that the cores and the GPU could share the same buffer, avoiding the copy operations using the zero-
copy-buffer capability of OpenCL.

With respect to the Kernel Launch overhead, it goes from up to 0.4% on Haswell to up to 3% and 2% on Ivy Bridge and Exynos, respectively. Average times consumed on this operation are 1.8 ms, 1 ms, and 3.6 ms on Ivy Bridge, Haswell and Exynos, respectively.

However, on the desktop platforms one of the most noticeable overheads is \(O_{td}\) (Thread Dispatch) especially for the 4+1 scenario. For that case, it represents 22% and 33% of the total execution time on Ivy Bridge and Haswell, respectively. Notice that this happens only for the oversubscription cases and under Windows OS. Actually, on Exynos under the Linux OS, the Thread Dispatch overhead represents less than 0.09% in all cases. The overhead in Windows is explained because the OpenCL driver ends up blocking the host thread once it has offloaded the kernel. It does that to avoid wasting a core with a busy-waiting host thread. In the 4+1 scenario, 4 threads are already intensively using the 4 cores and when the OpenCL driver notifies the OS about the GPU completion, the OS wakes up the host thread. However, if this host thread does not have enough priority it is unlikely it will be dispatched to the running state straightforward. The core of the Windows scheduling policy is Round Robin, so the just awaken host thread has to wait on the ready queue to the next available time slice. This does not happen in the Linux scheduler, where a just awaken thread gains more priority than the other CPU-bound threads that are intensively computing CPU chunks. This scheduling decision in Linux rewards interactive threads (IO-bound). As a consequence, in Linux, the host thread will be immediately dispatched after being woken up. This result is corroborated thanks to the experiment described in the next section.

5. OPTIMIZATIONS

After identifying the main sources of overhead of our Dynamic approach, in this section we discuss the optimizations that can be implemented to address them. The overarching goal is not only reduce the impact of the overhead, but also to reduce the energy consumption. To that end, we propose one strategy for the Windows based platforms (Section 5.1) and a different one for the Exynos architecture (Section 5.2).

5.1 Increasing the priority of the host thread

As it has been shown in the previous section, on the Windows based desktops, the highest overhead appears for the 4+1 configuration, where we have measured that up to 22% and 33% of the execution time is wasted on the \texttt{c1Finish()} operation on the Ivy Bridge and Haswell, respectively. This can be solved by assigning a higher priority to the host thread so that another thread can be immediately preempted and the just awaken host thread can take up a core and start feeding the GPU again. This is key when the GPU processes the chunks more efficiently than the CPU, as it happens in our benchmark. To boost the host thread priority we rely on the \texttt{SetThreadPriority()} Windows API. The framework obtained with this optimization is called Dynamic Pri.

The right part of the three subfigures of Figure 6 shows the overheads incurred by Barnes Hut when using Dynamic Pri and the arguments described in the previous section. The figure shows that on Ivy Bridge and Haswell, the \(O_{td}\) overhead for the 4+1 configuration has been reduced as is now similar to that of the 3+1 one. This confirms that increasing the priority of the host thread for the 4+1 configuration reduces this particular overhead. Note that on the Exynos platform increasing the host thread priority barely affects the measured overheads.

Figure 6 shows the normalized metrics (time, energy, and EDP) w.r.t. Bulk-Oracle 3+1, as shown in Figure 2 but now adding a new bar representing the results for Dynamic Pri. As expected, Figure 6 confirms that for Ivy and Haswell, that run Windows OS, boosting the priority of the host thread has almost no impact in reducing time, energy or EDP for the 3+1 configuration. However, with respect to Dynamic, Dynamic Pri has a significant impact on these metrics in the oversubscribed 4+1 scenario. For instance, on the Ivy Bridge, the Dynamic Pri reduces time, energy, and EDP by 10%, 7% and 18%, respectively, when comparing with Dynamic. On Haswell, these reductions are even more significant: 37%, 33% and 84% for time, energy and EDP, respectively. One interesting result appears on Ivy Bridge when comparing the Dynamic 3+1 configuration versus the Dynamic 4+1 as we mentioned in Section 2. Let’s remind that even though the 4+1 configuration is faster than the 3+1, the energy consumed by the 4+1 is higher, which is not true any more for Dynamic Pri. Now, Dynamic Pri uses the GPU more efficiently (it is served quicker and this results in the GPU processing more chunks). Therefore, the more energy consuming CPU cores end up doing less work (and consuming less energy) resulting in smaller total energy consumption figures.

Finally, notice that incrementing the priority of the host thread does not result in further reductions on time or energy on the Exynos platform. However, in the next section we explore a possible strategy to further reduce time and energy consumption on this architecture.

5.2 Exploiting big.LITTLE architecture

One interesting feature of the Exynos 5422 is that it supports global task scheduling (GTS). This enables using the four A15 cores and the four A7 ones at the same time. The Linux scheduler automatically uses the more powerful A15 cores for compute intensive threads, whereas power saving A7 are reserved for system and background tasks.

Figure 7 shows execution time (ms.), energy (J.) and EDP for the Exynos platform with configurations 3+1, 4+1, 7+1, and 8+1. The 7+1 and the 8+1 configurations use both the A15 and A7 cores for the computing threads, while the 3+1 and the 4+1 only use the A15 cores for the computing threads. In all the configurations, the mapping of the host thread to an A15 or an A7 core depends on the pinning strategy described below. Notice that the GPU also computes chunks of iterations, so there is now a larger degree of heterogeneity. Four alternatives are considered for each configuration:

- Dynamic is the baseline approach where the chunk size assigned to the A15 and A7 cores depends on the relative throughput on each one. The host thread is pinned
to one of the A15 cores.

- Dynamic Pri is based on the baseline, but it increases the priority of the host thread.
- Dynamic A7 is based on the baseline, but pins the host thread to one of the A7 cores.
- Dynamic Pri A7 is based on the baseline, but it pins the host thread to one of the A7 cores and increases its priority.

The energy sampling thread (see Section 4.1.1) is always pinned to one of the A7 cores, so it only produces an overhead of around 0.5% of the total execution time when running Barnes Hut under 8+1 scenarios (less than 0.02% otherwise). Regarding the breakdown of energy consumption, A15 cores use between 72% and 76% of the total energy consumed, the GPU between 15% and 20%, and the memory subsystem around 4.5%. Interestingly, A7 cores only take around 2.5% of the total energy when they are idle (3+1 and 4+1 scenarios) and around 7.3% when they are fully utilized (7+1 and 8+1 scenarios). For our benchmark, the four A7 cores consume one order of magnitude less energy than the four A15 cores.

Results in Figure 7 show that by increasing the number of threads to use the A7 cores (compare 3+1 or 4+1 with 7+1 or 8+1), execution time and energy reduce significantly. In particular, going from Dynamic 4+1 to Dynamic 8+1, we reduce time, energy and EDP by 22%, 19% and 46% respectively. Increasing the priority of the host thread or pinning it to a A7 have small impact. For instance, for the 3+1 and the 4+1 configurations, increasing the priority of the host thread or pinning it to the A7 core have almost no impact. On the 4+1 scenario, where Dynamic A7 is using 5 cores (4 A15 and 1 A7 for the host thread) we appreciate a marginal energy saving of 3% and similar running times. Increasing the priority of the host thread or pinning it to the A7 core has a higher impact on the 7+1 and 8+1 configurations, although their impact is relatively small. We notice that for 7+1 and 8+1 configurations, Dynamic Pri, Dynamic A7, and Dynamic Pri A7, obtain very similar results in terms of time and energy. Again, for 7+1 and 8+1 scenarios and with respect to Dynamic, Dynamic Pri and Dynamic A7 run, on the average, reduce the time, energy and EDP by 4.3%, 3.6% and 7.8%, respectively. All in all, w.r.t. the Dynamic 4+1, Dynamic Pri 8+1 reduces EDP by 57%.

Overall, while we expected that pinning the host thread to the A7 would have a higher impact on time or energy, our experimental results show little impact on either of them.

6. RELATED WORK

The closest work to ours is that of Zhu et al. [16], which address the problem of performance degradation when several independent OpenCL programs run at the same time (co-run) on the CPU and on the GPU of an Ivy Bridge using the Windows OS. The programs running on the CPU use all cores, so they are in a situation similar to our 4+1 configurations (oversubscription). To avoid degradation of the GPU kernel they also propose increasing the priority of the thread that launches the GPU kernel. Our study differs from theirs because we do not run two different programs,
instead we partition the iteration space of a single program to exploit both, the CPU and the GPU. Our study also shows that increasing the priority of the host thread is not necessary when there is no oversubscription (i.e. 3+1) or when the underlying OS is Linux. We also assess the use of a big.LITTLE architecture.

Other works as [12, 7] also address the overhead problems while offloading computation to GPUs. The work of Lustig and Martonosi [12] presents a GPU hardware extension coupled with a software API that aims at reducing two sources of overhead: data transfers and kernel launching. They use a Full/Empty Bits technique to improve data staging and synchronization in CPU-GPU communication. This technique allows subsets of data results being transferred to the CPU proactively, rather than waiting for the entire kernel to finish. Grasso et al. [7] propose several host code optimizations (Use of Zero-copy Buffer, Global Work Size equal to multiples of #EUs) in order to reduce GPU’s computation overheads on embedded GPUs. They present a comparison in terms of performance and energy consumption between an OpenCL legacy version and an OpenCL optimized one. Our work focus on reducing the sources of overhead as well, but in contrast, we focus on CPU-GPU collaborative computation instead of only targeting the integrated GPU.

Several previous works study the problem of automatically scheduling on heterogeneous platforms with a multicore and an integrated or discrete GPU [11, 16, 10, 3, 8]. Among those works, the only one that also uses chips with integrated GPUs is Concord [8]. However, Concord does not analyze the overheads incurred by offloading a chunk of iterations to the GPU.

7. CONCLUSIONS
In this work, we elaborate on the possibility of successfully implementing a dynamic scheduling strategy that automatically distributes the iteration space of a parallel loop among the cores and the GPU of a heterogeneous chip. To that goal it is key to guarantee optimal resource occupation and load balance while reducing the impact of the overhead. Our proposal is evaluated for the Barnes Hut benchmark on mid/low power heterogenous architectures like Ivy Bridge, Haswell, and Exynos 5, where the first two run under Windows OS and the latter under Linux. We have studied the sources of overhead on these systems, finding that, under Windows, the overhead due to re-scheduling the host thread is prohibitive in oversubscribed scenarios (more threads than cores). We solve this issue by increasing the priority of the host thread. Our experimental results show that this reduces Energy-Delay Product (EDP) by 18% and 84% on Intel Ivy Bridge and Haswell architecture, respectively. On the Exynos platform, the Linux scheduler successfully deals with oversubscription when only using the A15 cores. Therefore, for this platform we explore the benefits, in terms of time and energy, that can be achieved when pinning the host thread to a low power core or by the combined usage of the four A15 cores and the four A7 cores included in the Exynos 5. Our experimental results show that using the A7 cores reduces EDP by 46%. Increasing the priority of the host thread or pinning the thread to an A7 reduces EDP by and additional 7.8% in the Exynos big.LITTLE architecture.

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