Lab 2: Programming a Real Processing-in-Memory Architecture

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1. Introduction

In this lab, you will work hands-on with a real processing-in-memory (PIM) architecture. You will program the UPMEM PIM architecture [1, 2, 3, 4] for several workloads and will experiment with them. Your main goals are (1) to become familiar with the UPMEM PIM system organization (as an example of real-world memory-centric computing system), (2) to understand the UPMEM programming model and write your own code, and (3) to understand the microarchitecture and instruction set architecture (ISA) of UPMEM’s PIM core (called DRAM Processing Unit, DPU).

As we introduced in class, the UPMEM PIM architecture is composed of multiple DPUs (up to 2,560), each of which has access to its own DRAM bank (called Main RAM, MRAM) and its own scratchpad memory (called Working RAM, WRAM). You can find a full description of the UPMEM PIM system in [3, 4].

2. Your Task 0/4: Installing the UPMEM SDK

Your goal is to set up the UPMEM SDK on your machine to compile and run the code in the following subtasks. If you have access to a system with a supported Linux version, you can install the UPMEM SDK natively from the UPMEM website [1, 2]. If you encounter issues with the installation or do not have access to a system with a supported Linux version, you can use the Dockerfile we provide, along with the associated shell scripts for either Windows or Unix-based host systems.

2.1. Using the Dockerfile

Using the Dockerfile requires a containerization tool like Docker [5] or Podman [6] to be installed on your system. With Docker (Podman) installed, you can execute the `container/start_docker.sh` (container/start_podman.sh) shell script on Linux or `container\start_docker.bat` (container\start_podman.bat) on Windows.

```
$ container/start_docker.sh
```

The script will automatically build the container image (which will take a few minutes the first time) and then start an interactive shell within it. The working directory of the host machine where the container was started will be mounted inside the container (try running `ls` inside the container). The code for this lab can then be compiled and executed using this interactive shell.

3. Your Task 1/4: Transferring Data between Main Memory and PIM-enabled Memory

Your goal is to get familiar with different types of data transfers between the host main memory and the PIM-enabled memory. You are provided with a template for this task. Find more details in Section 8.

Your tasks are as follows:

1. Write a host program that exercises all types of data transfers between the host main memory and one
or multiple MRAM banks. Concretely, there are three types of data transfers: (1) serial, (2) parallel, and (3) broadcast. Serial and parallel transfers move data from main memory to the MRAM banks or vice versa. Broadcast transfers can only happen from the main memory to the MRAM banks.

4. Your Task 2/4: AXPY

Your goal is to get familiar with the DPU kernel launch and execution, and the performance scaling for different numbers of PIM threads (called tasklets in the UPMEM architecture). You are provided with a template for this task. Find more details in Section 8.

Your tasks are as follows:

1. Write a DPU kernel that executes the AXPY operation \( y = y + \alpha x \) on every element of a vector. You have to (1) transfer two input vectors, \( Y \) and \( X \), to the MRAM bank/s, (2) perform the AXPY operation with a variable number of tasklets, (3) write the results to the output vector, \( Y \), and (4) transfer the output vector back to the host main memory.

2. Allocate as much WRAM as needed, and use \texttt{mram\_read} and \texttt{mram\_write} to move data between MRAM and WRAM.

3. Your code should produce correct results for any data type (e.g., \texttt{char}, \texttt{short}, \texttt{int}, \texttt{long long}, \texttt{int}, \texttt{float}, \texttt{double}) and vector size. Hint: Make sure all data transfers are 8-byte aligned.

4. Run your kernel for all numbers of tasklets between 1 and 24, and make sure that the kernel produces correct results in all cases (i.e., for any number of tasklets and DPUs). Use input vectors of, at least, 8 MB (per DPU).

Based on your analysis, submit answers to the following questions with your lab report.

1. Use the performance counters to count the number of executed instructions per tasklet (see Section 8) for different numbers of tasklets and generate a plot. Analyze the results and provide your observations.

5. Your Task 3/4: Operations and Data Types

Your goal is to analyze how the DPU performs for different types of data and operations.

Your tasks are as follows:

1. Modify your AXPY DPU kernel to make it a vector addition \( y = y + x \) and to support other operations besides addition (i.e., subtraction, multiplication, division).

2. Evaluate the performance of your new kernel for different operations (addition, subtraction, multiplication, division) and data types (\texttt{char}, \texttt{short}, \texttt{int}, \texttt{long long}, \texttt{int}, \texttt{float}, \texttt{double}).

Based on your analysis, submit answers to the following questions with your lab report.

1. Using the performance counters (see Section 8), measure the number of instructions for each data type and operation. For each operation and data type, you can calculate the average number of instructions.

2. Report your observations about your evaluation.

3. We recommend you use the LLVM object file dumper (see Section 8) to read the assembly code of your program, and use this information to explain your observations.

6. Your Task 4/4: Vector Reduction

Your goal is to get familiar with the synchronization primitives for intra-DPU communication (i.e., across tasklets). To do so, you will write a DPU kernel that performs the parallel reduction of an input vector.

Your tasks are as follows:
1. Your vector reduction DPU kernel should have four different versions: (1) final reduction with a single tasklet, (2) final tree-based reduction with barriers, (3) final tree-based reduction with handshakes, (4) final reduction with mutexes.

2. Run the different versions of your kernel for different numbers of tasklets. Make sure all versions of your code produce correct results for any number of tasklets and DPUs. Count the number of executed instructions. Use vector sizes of 16 KB, 1 MB, 16 MB.

Based on your analysis, submit answers to the following questions with your lab report.

1. Report your observations about your evaluation. What is the best performing version for different vector sizes?

7. **Bonus Task: Implement RGB to Grayscale Conversion**

In this task, your goal is to implement the RGB to grayscale conversion of an image on a variable number of DPUs. The image should have 8-bit red, green, and blue channels. For each image pixel, you perform a conversion to obtain a grayscale value from the R, G, and B values. Compare the following three methods:

1. **Lightness method:**
   \[\text{grayscale} = \frac{\min(R,G,B) + \max(R,G,B)}{2}\]

2. **Average method:**
   \[\text{grayscale} = \frac{R + G + B}{3}\]

3. **Luminosity method:**
   \[\text{grayscale} = 0.3 \times R + 0.59 \times G + 0.11 \times B\]

Based on your analysis, submit answers to the following questions with your lab report.

1. How do the different methods scale for different numbers of tasklets?
2. What is the fastest of the methods? Explain.
3. What is the slowest of the methods? Explain.

Think about potential optimizations of your code, implement them, and reevaluate. A general recommendation is to try to reduce the number of executed instructions.

**You can receive up to 20% extra points for this lab, if you implement correct versions and explore additional optimizations with successful results.**

8. **Lab Resources**

8.1. **Source Code**

The source code that we provide contains templates for tasks 1 (Section 3) and 2 (Section 4). For the rest of tasks, you can use the same template as for task 2. You can find the templates in the folder `template`. Look for `//@@` to find the places where you need to insert code. Do **NOT** modify any files or folders unless explicitly specified in the list below.

- **task1**
  - Makefile
  - host
    - app.c: Host CPU code (**modifiable**).
  - dpu
    - task.c: DPU kernel code. It is empty in this template because it is not needed for task 1.
  - support
    - common.h: Common definitions. Note that `T` is `int64_t` for this task.
    - params.h: Functions to read input parameters from command line.
8.2. Compilation

**Task 1.** The Makefile for task 1 contains the following input parameters and default values:

- `NR_DPUS`\(\equiv 1\): Number of DPUs that the program will use.
- `NR_TASKLETS`\(\equiv 16\): Number of tasklets per DPU that the program will use.
- `TRANSFER`\(\equiv PARALLEL\): Type of data transfer (SERIAL, PARALLEL, BROADCAST).
- `PRINT`\(\equiv 0\): Print log from the DPU kernel.

For task 1, you will only have to use `TRANSFER` and `NR_DPUS`. To compile with the default parameters:

```bash
$ make
```

To compile, for example, with serial transfers and 64 DPUs:

```bash
$ NR_DPUS=64 TRANSFER=SERIAL make
```

The compiled binaries will be in the `bin` folder. You can run the program with the default input:

```bash
$ ./bin/host_code
```

You can check the possible input arguments with `-h`:

```bash
$ ./bin/host_code -h
```

For example, you can run the program to transfer 2MB of data (i.e., 262144 64-bit elements) between the host main memory and the MRAM banks, and vice versa, and repeat the experiment 10 times after 2 times of warm-up:

```bash
$ ./bin/host_code -w 2 -e 10 -i 262144
```

**Task 2 and Rest of Tasks.** The Makefile for task 2 contains the following input parameters and default values:

- `NR_DPUS`\(\equiv 1\): Number of DPUs that the program will use.
- `NR_TASKLETS`\(\equiv 16\): Number of tasklets per DPU that the program will use.
**BLOCK** ?= 10: Size of the blocks (chunks) of data moved between MRAM and WRAM. The actual size in bytes is $2^{\text{BLOCK}}$.

**TYPE** ?= INT32: Datatype of the input arrays (CHAR, SHORT, INT32, INT64, FLOAT, DOUBLE).

**TRANSFER** ?= PARALLEL: Type of data transfer (SERIAL, PARALLEL).

**PRINT** ?= 0: Print log from the DPU kernel.

**PERF** ?= NO: Use of performance counters for cycle or instruction count (CYCLES, INSTRUCTIONS)

To compile with the default parameters:

```bash
$ make
```

To compile, for example, with parallel transfers, 64 DPU, 12 tasklets per DPU, 32-bit floating point datatype, 512-byte MRAM-WRAM and WRAM-MRAM data transfers, and counting instructions executed by the tasklets:

```bash
$ NR_DPU=64 NR_TASKLETS=12 BLOCK=9 TYPE=FLOAT TRANSFER=PARALLEL PERF=INSTRUCTIONS make
```

As in task 1, the compiled binaries will be in the bin folder. For example, you can run the AXPY program to operate on 4MB arrays (e.g., 1048576 32-bit floating-point elements), with a value of $\alpha$ equal to 20, and repeat the experiment 10 times after 2 times of warm-up:

```bash
$ ./bin/host_code -w 2 -e 10 -i 1048576 -a 20
```

### 8.3. LLVM Object File Dumper

You can use this tool to print the contents of object files [9]. The -S option displays the source code interleaved with the disassemble code:

```bash
$ llvm-objdump -S ./bin/dpu_code
```

Another useful tool with a similar purpose is Compiler Explorer [10].

### 9. Tips

- Please do not distribute the provided program files. These are for exclusive individual use of each student of the Computer Architecture course. Distribution and sharing violates the copyright of the software provided to you.
- Read this handout in detail.
- Read the UPMEM programming guide [2].
- If needed, please ask questions to the TAs using the online Q&A forum in Moodle.
- When you encounter a technical problem (e.g., a compilation error), please first read the error messages.

### 10. Submission

Use the corresponding assignment in Moodle (https://moodle-app2.let.ethz.ch/). You should submit:

- All the files needed to compile your code (including files that you did not change).
- One folder per task with the same organization as the provided templates.
- A report as a single PDF file that contains your answers to the tasks indicated in this handout.
- Also, please do not submit compiled files (e.g., binary files).

Please submit the above files in a single tarball (with the name ‘lab2_⟨YourSurname⟩_<YourName>.tar.gz’).
References