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Porting a PCA-based Hyperspectral Image Dimensionality Reduction Algorithm for Brain Cancer Detection on a Manycore Architecture


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Abstract—This paper presents a study of the parallelism of a Principal Component Analysis (PCA) algorithm and its adaptation to a manycore MPPA (Massively Parallel Processor Array) architecture, which gathers 256 cores distributed among 16 clusters. This study focuses on porting hyperspectral image processing into manycore platforms by optimizing their processing to fulfill real-time constraints, fixed by the image capture rate of the hyperspectral sensor. Real-time is a challenging objective for hyperspectral image processing, as hyperspectral images consist of extremely large volumes of data and this problem is often solved by reducing image size before starting the processing itself. To tackle the challenge, this paper proposes an analysis of the intrinsic parallelism of the different stages of the PCA algorithm with the objective of exploiting the parallelization possibilities offered by an MPPA manycore architecture. Furthermore, the impact on internal communication when increasing the level of parallelism is also analyzed.

Experimenting with medical images obtained from two different surgical use cases, an average speedup of 20 is achieved. Internal communications are shown to rapidly become the bottleneck that reduces the achievable speedup offered by the PCA parallelization. As a result of this study, PCA processing time is reduced to less than 6 seconds, a time compatible with the targeted brain surgery application requiring 1 frame-per-minute.

Keywords—Dimensionality Reduction; Hyperspectral Imaging; Massively Parallel Processing; Real-time processing

I. INTRODUCTION

Hyperspectral imaging (HI) collects both spatial and spectral information from across the electromagnetic spectrum, covering a wide range of wavelengths. This new technology aims at identifying elements in an image by distinguishing among their spectral signatures, which represent the reflectance measured by the sensor for each wavelength [1]. Although the original application field for this technology was remote sensing [2]-[3], its use has spread over several research fields, such as astronomy, security, forensics and medicine [4]-[7].

Regarding the medical field, the ability to distinguish among materials has become crucial for cancer detection applications. This technology has already been applied in two different scenarios: ex-vivo and in-vivo studies –i.e., with images captured from a resected sample and directly taken from the patient, respectively.

Related literature shows an increasing research interest concerning the performance of in-vivo HI processing during medical procedures to assist surgeons in discerning between tumor tissues and healthy tissues [8]-[9]. Furthermore, to help surgeons in determining the margins of the tumor during surgery, a real-time analysis of the hyperspectral image becomes compulsory, considering this real-time as the time needed for the hyperspectral sensor to capture a new image. As nowadays hyperspectral sensors usually present a push-broom scanning mechanism, real-time in this context can be set to a maximum of 1 picture per minute. Neurosurgeons have stated that a processing time of one image per minute is sufficient to assist them during an operation [8]. Processing a diagnostic helping image in less than 1 minute is not possible with the existing alternative to HI, which is the Intraoperative Magnetic Resonance Imaging (iMRI) that usually needs more than thirty minutes to acquire one image [10].

Hyperspectral sensors generate large amounts of data, which makes meeting real-time constraints challenging. As a result, it is advisable to reduce the volume of data before beginning with high-level processing. Therefore, a dimensionality reduction stage is often performed as an essential step during image preprocessing. This dimensionality reduction method is usually accomplished through a principal components transformation [11], which selects and retains the most relevant information for classification.
Principal Component Analysis (PCA) is the most widely used technique in remote sensing applications, specifically in those using hyperspectral images. In this kind of images, the adjacent bands are profoundly correlated, thus providing no new information. PCA reduces the volume of information by removing the dependencies among the different bands. To do so, an eigenvector decomposition of the covariance matrix of the original data is computed [11].

Sequential implementations of this algorithm do not have the performance required to achieve the real-time constraint of this use case. Hence, to reach real-time performance, this study analyzes the intrinsic parallelism of the PCA algorithm and exploits the resulting parallelized model to minimize the time needed for an image to be processed. As this processing requires an extensive usage of computational resources, High Performance Computing (HPC) architectures are targeted.

HPC platforms are evaluated based on two criteria: processing time and energy consumption. Although the former has frequently been the metric chosen to assess HPC platforms, the latter is gaining importance as a first-class performance criterion.

Even though current medical applications rarely work under energy requirements, it is not difficult to foresee future clinical applications where portable and real-time processing becomes a crucial tool to support medical decisions. In that sense, manycore processors are today some of the most efficient architectures [12] [13] for the task. For instance, the Kalray Massively Parallel Processor Array (MPPA) in its Bostan version (MPPA-256-N) requires only 5W in average operating mode [14].

The main contribution of this paper is the study and implementation of a PCA algorithm and the evaluation of its performance on an HPC MPPA manycore architecture. Additionally, this research also aims at studying the effect of the internal communications within the manycore architecture when the degree of parallelism is increased. This paper extends the results of [15] with a new discussion section to compare the current work with the state-of-the-art. In addition, the results of a new parallel approach and a better exploitation of the platform parallelism are analyzed. At last, a new data set has been described and employed to obtain the results.

The rest of the paper is structured as follows. First, Section II describes the target MPPA platform together with the studied PCA algorithm. Secondly, Section III focuses on the implementations. Afterwards, Section IV shows the experimental results, and Section V provides a comparison with some state-of-the-art implementations. Finally, Section VI draws the main conclusions of this research work.

II. HARDWARE AND ALGORITHM

A. MPPA-256-N Kalray Platform

The HPC platform selected for this research is the Kalray MPPA-256-N, whose structure is shown in Fig. 1. This platform is a single-chip manycore processor that gathers 256 cores organized in 16 clusters running at up to 600MHz. It also contains two quad-core Input/Output (I/O) subsystems responsible for handling the communications between a host processor and the clusters, which are interconnected by a Network-On-Chip (NoC).

Each cluster gathers 2 MB of memory shared among the 16 cores within the cluster. In addition, there is also a Direct Memory Access (DMA) engine managing the communications between this memory and the NoC, and a Resource Management (RM) core responsible for starting the NodeOS operating system and handling events and interrupts for the whole cluster.

B. Principal Component Analysis

As mentioned in Section I, Principal Component Analysis is the most well-known and widely used technique for data shrinking in HI applications, proving to be a powerful tool for hyperspectral image processing [16].

As hyperspectral images are composed of spectral information gathered from an extensive number of narrow bands, this information is frequently deeply correlated, thus containing a large amount of redundancies. Hence, these redundancies should be eliminated, reducing the image size and, therefore, its processing cost.

Specifically, PCA reduces the data volume by converting the original data into a subspace of smaller dimension where the image is rearranged as a decreasing function of its spectral information – i.e., accumulating the useful spectral information in the first bands—. To do so, PCA computes the covariance matrix of the original data, extracts its associated eigenvectors and projects the image onto these eigenvectors. Finally, the algorithm finishes by selecting the number of bands—or principal components—to retain. Algo. 1 provides the pseudocode of the algorithm, which is divided in four stages:

i. Image preprocessing: It is the first step of the algorithm, and it centers the image by computing and removing the average of each spectral band of the original image, composed by N pixels per M spectral bands. It shall be noted that the monochromatic image associated to a frequency band is treated as a vector, ignoring the spatial relationship among pixels.

ii. Covariance computation: This stage computes the covariance matrix associated to the original image multiplying the preprocessed image by its transpose.

iii. Eigenvector decomposition: This step extracts the eigenvectors associated to the covariance matrix computed in the previous stage.
iv. **Projection and reduction:** This stage combines steps 4 and 5, and it projects the original image onto the set of eigenvectors to store then the first $P$ principal components, where $P$ represents the number of principal components—or bands—to retain.

<table>
<thead>
<tr>
<th>Algorithm 1: Principal Component Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Input:</strong> Hyperspectral image $Y$ ($N \times M$ matrix)</td>
</tr>
<tr>
<td><strong>Step 1:</strong> $X$ = Remove the average of each band of $Y$</td>
</tr>
<tr>
<td><strong>Step 2:</strong> Covariance matrix $C = X^T \cdot X$</td>
</tr>
<tr>
<td><strong>Step 3:</strong> $E$ = Eigenvector decomposition of $C$</td>
</tr>
<tr>
<td><strong>Step 4:</strong> Projection $Q = Y \cdot E$</td>
</tr>
<tr>
<td><strong>Step 5:</strong> Reduce $Q$ to $P$ principal components</td>
</tr>
<tr>
<td><strong>Output:</strong> Reduced hyperspectral image $Q'$ ($N \times P$ matrix)</td>
</tr>
</tbody>
</table>

Related with the eigenvector decomposition, traditionally the conventional method for extracting them has involved computing the inverse matrix and finding the roots of its characteristic polynomial [17]. However, for extensively large matrices—e.g. hyperspectral images—this procedure is not feasible, therefore other methods shall be considered.

In [17], Panju summarizes some of the iterative methodologies for addressing this issue. These approaches work by refining approximations of the eigenvectors in each iteration and, consequently, their convergence depends on the criterion set for the approximation accuracy. However, iterative methods usually work in detriment of real-time, as they are very demanding in terms of processing time. In order to minimize this effect, Jacobi method has been selected in this research due to its high degree of parallelism. The convergence of this algorithm has been demonstrated for two different strategies [18], regarding the order in which the elements are chosen to be zeroed.

1) **Classical method:** As described before, this method zeroes the largest off-diagonal element in each rotation.
2) **Cyclic method:** This method zeroes the off-diagonal elements in a given order, e.g. row by row.

The first method has been proven to guarantee the least number of rotations, but the second one is typically faster, as it avoids the location of the largest element in each iteration, which is a quadratic order operation.

Specifically, each Jacobi iteration performs the following steps:

- First, the next off-diagonal element to be zeroed is selected, following one of the two described methods (classical or cyclic Jacobi).
- Secondly, the Jacobi rotation matrix ($P$) is calculated, which is similar to the model shown in equation (1). The dimensions of this matrix equal those of the covariance matrix $C$, which is the input for the Jacobi method.
- The $\alpha$ value depends on the element selected to be zeroed $-C_{ij}$, where $i$ represent the rows of the covariance matrix and $j$ represents the columns, so it must be recalculated at the beginning of each iteration. In each iteration, equations (2) to (5) are applied to calculate this value and, hence, the Jacobi rotation matrix.

$$
P = \begin{pmatrix} 1 & \cdots & \cdots & 0 \\ \vdots & \cos \alpha & \sin \alpha & \vdots \\ \vdots & -\sin \alpha & \cos \alpha & \vdots \\ 0 & \cdots & \cdots & 1 \end{pmatrix} \tag{1}
$$

$$
m = \frac{2 \cdot C_{ij}}{C_{ij} - C_{ii}} \tag{2}
$$

$$
t = \frac{-1 + \sqrt{1 + m^2}}{m} \tag{3}
$$

$$
\cos \alpha = \frac{1}{\sqrt{1 + t^2}} \tag{4}
$$

$$
\sin \alpha = t \cdot \cos \alpha \tag{5}
$$

- Once the matrix $P$ has been computed, the operation provided in (6) is performed. As a result, the off-diagonal element is now zeroed in $C_j$, and also its symmetric counterpart, as $C$ has to be symmetric. For the next iteration, $C_j$ will be considered as the input of Jacobi algorithm.

$$
C_j = P^T \cdot C \cdot P \tag{6}
$$

Each iteration repeats the described steps, zeroing one element at a time. However, the algorithm does not stop when all the elements have been chosen to be zeroed once, due to the fact that, in each iteration, several previous zeros can be undone. That is the reason why a stop factor is needed.
Once the last iteration finishes –K– and the convergence is reached, the eigenvalues are stored in the diagonal of $C_K$, as shown in (7), where $P_i$ are the successive Jacobi rotation matrices.

$$C_K = P_1^T \cdot \cdots \cdot P_i^T \cdot C \cdot P_{i+1} \cdot \cdots \cdot P_K \quad 1 < i < K \quad (7)$$

Likewise, the eigenvectors associated to these eigenvalues can be calculated as depicted in (8), where the eigenvectors are placed in the columns of $E$.

$$E = P_1 \cdot P_2 \cdot \cdots \cdot P_K \quad (8)$$

As each rotation affects only a couple of rows and columns, several rotations can be calculated simultaneously, thus providing a parallel method.

Specifically, in each iteration the elements that are changed in the input matrix are the ones belonging to the concerned rows and columns, so several elements can be processed in parallel if they do not share any of those positions. For instance, elements $C_{23}$ and $C_{35}$ could be processed in parallel; on the contrary, $C_{23}$ and $C_{14}$ could not be simultaneously zeroed.

Next section will provide a detailed description of the adaptation of this method –along with the adaptation of the rest of the operations involved in PCA algorithm– to the architecture under study.

### III. Implementation

#### A. Database

The conducted experiments have been tested upon hyperspectral images extracted from the HELICoD project database [22]. The in-vivo human brain surface images were captured during neurosurgical operations performed at the University Hospital Doctor Negrin of Las Palmas de Gran Canaria (Spain) and at the University Hospital of Southampton (UK).

To capture these images, the HELICoD setup described in [23] has been used. Two hyperspectral sensors compose this setup: one in the visible and near infrared spectral range (VNIR), covering from 400 nm to 1000 nm, and the other in the near infrared range (NIR), covering from 900 nm to 1700 nm of the electromagnetic spectrum. Both cameras are attached to a push-broom scanning unit, together with an illumination system that provides a cold light to protect the exposed brain surface from the heat generated by the lamp. As the cameras take the images with a push-broom mechanism, each image needs from 1 to 2 minutes to be captured –depending on the spatial size of the image–, which can be considered as the definition of the surgery real-time constraint.

Specifically, two different hyperspectral images have been used to assess the algorithm. A preprocessing stage has spatially limited the image to the area of interest and it has reduced the spectral resolution to 128 bands [24]. As a result, the first image –hereafter case 1– presents a spatial resolution of 377 lines and 329 samples –i.e., 124033 pixels–, while the second –hereafter case 2– contains 479 lines and 552 samples –i.e., 264408 pixels–. Fig. 2 gathers both hyperspectral images, case 1 on the left and case 2 on the right.

Considering that these images are stored as float numbers (4 Bytes), the required memory for storing each image is 60.6 MB for case 1 –i.e. 124033 pixels x 128 bands– and 129.1 MB for case 2 –i.e. 264408 pixels x 128 bands–, respectively.

![RGB representation of the hyperspectral images extracted from the HELICoD database: case 1 (left) and 2 (right)](image)

#### B. Target environment

As mentioned in Section I, the aim of this work is to achieve real-time while processing hyperspectral images during a surgical procedure. Due to the large amount of information contained in these images, the computational complexity is such that sequential implementations of the algorithms do not provide enough performance to fulfill these requirements; thus, HPC platforms become a necessity.

The Kalray MPPA-256-N appears to be an optimal solution, as it is particularly competitive in terms of energy efficiency, which is a parameter of growing interest. Specifically, the target environment is a workstation that includes an MPPA-256-N chip, whose simplified block diagram when connected to a host PC is provided in Fig. 3. The MPPA architecture presents three different levels: the host module, the I/O interface and the 256 processing units, which are organized in 16 clusters. The host module is responsible for managing the global functioning, and it communicates with the I/O interface through a PCI express (PCIe) connection; it also presents the largest memory space, with more than 10 GB of available memory. Similarly, the I/O interface handles both the communications with the host –through the PCIe connection– and with the clusters –through a NoC interface–; as for the available memory, it contains a 4 GB external DDR. Finally, the processing cores are responsible for the processing itself. As this chip contains 256 processing cores, the potential level of parallelism is very high. Nevertheless, this platform also presents an important restriction for hyperspectral image processing, which is the reduced amount of memory within each cluster –a 2 MB block of shared memory among the 16 internal cores of each cluster–. Furthermore, some of this memory is reserved for both program code and operating system, so the available memory for storing data cannot represent more than 1.5 MB, approximately.

As hyperspectral images gather extremely large volumes of data, this is an important limitation. As described before, the images used during this work require 60.6 MB –case 1– and 129.1 MB –case 2–. Therefore, it seems obvious that, to process an entire image, the algorithm must be split into several iterations. Subsequently, iterating the processing of the image highlights another important limitation, which is the...
communication between the I/O subsystems and the clusters. This communication is very demanding in terms of performance, so it seems obvious that the processing time will be proportional to the number of iterations needed to complete the algorithm execution.

![Block diagram](image)

**C. MPPA-256-N implementation**

Taking into consideration the previous features, this section provides a detailed description of the implementation of each part of the PCA algorithm, highlighting its intrinsic parallelism. It shall be noted that this implementation is an extended work of that presented in [15].

1) **Preprocessing:** To compute the covariance matrix of the original data, first the image ought to be mean-centered. To do so, the average of each spectral band is calculated and removed. This operation is band-wise parallelizable: the original matrix is divided among the clusters, and the internal cores compute and remove the average of the bands received in each cluster. Due to the high dimensionality of the image and the cluster memory limitations, each cluster can only compute the average of one band at a time; therefore, each iteration processes 16 bands simultaneously. As a result, each core computes one portion of the average of the band; then, the master thread –i.e., the core executing the main function– computes the global average and subtracts it.

2) **Covariance matrix computation:** The resulting matrix is then multiplied by its transpose, thus generating the covariance matrix. As neither of these matrices fit into a cluster, this computation needs to be iterated; further, due to the dimensions of the matrices involved in the multiplication, this operation is the main bottleneck of the algorithm. Considering that the memory needed to store a row -i.e., a band– of the matrix is 0.48 MB in case 1 and 1 MB in case 2, there are two different cases:

- For the smallest image –case 1–, two bands fit into a cluster, so each cluster can compute one element of the covariance matrix at a time. This means that, in each iteration, 16 elements of the covariance matrix can be computed simultaneously. As the dimensions of the resulting matrix are 128×128, 1024 iterations are required to complete this computation.

- On the other hand, for the largest image, even two bands exceed the memory restriction, so at least two clusters must be used to compute just one element of the covariance matrix. To do so, instead of sending a whole row to each cluster, all the bands are divided in half, and each cluster only receives the halves that should be multiplied (as a matrix multiplication is just a concatenation of dot products). With this method, the number of iterations is thus doubled.

In both cases, the cores within each cluster compute their corresponding share of the vector multiplication –as they are a concatenation of dot products– and send it to the I/O subsystem, which adds them and stores the resulting value.

Nevertheless, it should be noted that, as multiplying a matrix by its transpose generates a symmetric matrix, only the upper triangle of the covariance matrix needs to be computed, thus reducing the iterations in half for both cases.

For that reason, another method has also been proposed for avoiding these data transfers. This method consists of taking advantage of the host of the platform to remove the memory restriction and, thus, all the I/O-cluster communications. Although the processing time will certainly increase –as the multiplication will not be parallelized–, the time saved in the communications will hypothetically compensate this increase. These two solutions will be evaluated in Section IV.

3) **Eigenvector decomposition:** This stage deals with the implementation of the Jacobi method introduced in Section II. Among all the existing variations, the cyclic Jacobi [21] has finally been implemented. As described in Section II, the main variation of this method is that, instead of choosing the elements to be zeroed by finding the largest off-diagonal element, it chooses the next element in a given order, e.g., row by row. As the covariance matrix (128×128) fits into a cluster, the use of only one of them for implementing this step removes all the internal communications, as all the cores within a cluster share the same memory. In addition, as Jacobi method is highly parallelizable, all cores of the chosen cluster have been used for implementing this step. As a result, this method has been implemented as follows:
The master thread—i.e., the core executing the main function—is responsible for handling the search of the next element to be zeroed. In each iteration, this thread verifies that the stop condition is not fulfilled, chooses a maximum of 15 different new elements to be processed in parallel, calculates the rotation matrix and sends it to the processing cores. This process is repeated until the stop condition is reached by all of the off-diagonal elements. Once this happens, this thread sorts the eigenvalues in a descending order, as well as their associated eigenvectors.

Likewise, the processing cores perform the operation shown in (9), where \( i \) represents the current iteration, \( P_i \) is the rotation matrix of the iteration \( i \), \( C_{i-1} \) is the covariance matrix modified in the previous iteration and \( C_i \) is the resulting matrix—i.e., the covariance matrix with several elements already zeroed. It shall be noted that (9) is a generalization of equation (6) described in Section II.

\[
C_i = P_i^T \cdot C_{i-1} \cdot P_i \tag{9}
\]

4) Projection and principal components selection: In this step, the original matrix is projected onto the set of eigenvectors and the first P bands—i.e., principal components—are stored. To reduce the complexity of the projection, instead of using the whole set of eigenvectors only the subset of the first P ones is utilized. Specifically, as for this application only the first principal component is required [25], the set of eigenvectors is thus reduced from 128 to 1. This also reduces the projection complexity, since instead of multiplying two matrices, just one matrix—the original data—by a vector—the first eigenvector—multiplication is required. Related with the parallelization, the method is similar to the one applied in step 2, but much simpler. As the eigenvector fits into a cluster, it is broadcasted to all of them and then the original matrix is split and sent to the clusters in a pixel-wise order iteratively—i.e., in groups of 128 elements—, until all the matrix is multiplied by the eigenvector. From the point of view of the parallelization within the clusters, each one receives blocks of 1024 pixels, so each core projects 64 pixels onto the eigenvector and returns the results to the I/O subsystem.

Consequently, concerning the MPPA-256-N resource usage, the PCA algorithm is divided into two blocks: steps 1, 2 and 4 use all the available resources (16 clusters and 16 cores per cluster), while step 3 only uses the 16 cores of one cluster. It should be noted that, for the alternative solution to step 2, all platform resources are idle, as the processing is performed in the host.

IV. RESULTS

In this section, the results obtained while exploring the proposed solutions are evaluated in terms of processing time. First, the sequential version of the PCA algorithm on the MPPA-256-N is assessed; then, the parallelism is exploited, locating the main bottlenecks. After that, the communication cost is evaluated in order to analyze its effects on the overall computation time. Finally, the second proposed approach for computing the covariance matrix is evaluated. It should be noted that the results presented in this section have been measured when setting the stop factor, \( \epsilon \), to the maximum value that provides a final result with a relative error lower than 1% when compared to a Matlab version.

A. First approach

In order to address the analysis of the obtained results, the different stages of the PCA algorithm described before are characterized. To do so, as the MPPA architecture presents two different levels of parallelism, first the sequential implementation will be studied; after that, the first level of parallelism will be evaluated by distributing the computational load among the 16 clusters, but using just 1 core in each one. Finally, the second level of parallelism is assessed by using all the cores of each cluster, instead of just one.

Table I provides the execution time for each of the stages when they are processed in a sequential way—i.e., using only one core of one cluster. As expected, the main bottleneck of the algorithm is the covariance computation stage, as it consumes more than a 92% of the global execution time.

TABLE I. AVERAGE TOTAL EXECUTION TIME (MS) - SEQUENTIAL

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Step 1</th>
<th>Step 2</th>
<th>Step 3</th>
<th>Steps 4-5</th>
<th>Global</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1</td>
<td>900.7</td>
<td>40,870.5</td>
<td>1,759.8</td>
<td>840.2</td>
<td>44,305.2</td>
</tr>
<tr>
<td>Case 2</td>
<td>1,908.9</td>
<td>81,503.4</td>
<td>1,849.7</td>
<td>1,759.8</td>
<td>87,321.8</td>
</tr>
</tbody>
</table>

Likewise, Table II presents the execution time of each stage of the algorithm when their computational load is divided among the 16 available clusters—i.e., using the 16 clusters, but just 1 core of each of them. As a result, it can be noticed that, although the execution time has decreased, the global speedup achieved is rather low, as ideally it should grow up to 16 but in reality it is, approximately, 1.5. This limitation is basically due to data broadcasting: as hyperspectral images contain extremely high volumes of information, the transmission of the images to the clusters could be such that it could overcome any speedup the processing itself may achieve. To check whether this hypothesis is correct, first the second level of parallelization is studied and, after that, the effect of communications is evaluated.

TABLE II. AVERAGE TOTAL EXECUTION TIME (MS) – FIRST LEVEL OF PARALLELISM

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Step 1</th>
<th>Step 2</th>
<th>Step 3</th>
<th>Steps 4-5</th>
<th>Global</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1</td>
<td>402.7</td>
<td>26,699.4</td>
<td>1,759.8</td>
<td>380.3</td>
<td>29,244.2</td>
</tr>
<tr>
<td>Case 2</td>
<td>860.4</td>
<td>52,990.1</td>
<td>1,849.7</td>
<td>839.6</td>
<td>56,540.2</td>
</tr>
</tbody>
</table>

Table III gathers the equivalent results to those presented in Tables I and II, but exploiting the whole architecture, i.e., using the 16 cores of the 16 clusters. As can be observed, the speedup achieved is almost negligible when compared to the previous one, since it only accelerates the computation by a factor of 1.1, approximately. Again, the reason to explain this behavior is the communications: supposing that the previous
hypothesis was correct, it explains this new behavior, as it means that the processing itself cannot be further parallelized, and hence all the measured time is being wasted for communications. To prove whether this hypothesis is true, next subsection evaluates the impact of communications.

**TABLE III. AVERAGE TOTAL EXECUTION TIME (MS) – SECOND LEVEL OF PARALLELISM**

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Step 1</th>
<th>Step 2</th>
<th>Step 3</th>
<th>Steps 4-5</th>
<th>Global</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1</td>
<td>393.3</td>
<td>25,726.1</td>
<td>350.9</td>
<td>369.9</td>
<td>26,840.2</td>
</tr>
<tr>
<td>Case 2</td>
<td>831.9</td>
<td>51,039.5</td>
<td>370.3</td>
<td>779.8</td>
<td>53,021.5</td>
</tr>
</tbody>
</table>

Before getting into the communication assessment, the behavior of the Jacobi algorithm should be highlighted. As described in Section III, Jacobi has only been implemented using one cluster due to the communications cost, so there is no difference in the time measured for Jacobi in Tables I and II. As for the results presented in Table III, it can be observed that, in this case, the Jacobi algorithm is accelerated by a factor of 5, approximately. This result supports the communications hypothesis: as described before, the input of the Jacobi algorithm – i.e., the covariance matrix – fits into a cluster; as a result, this step is the only one in which there are not any communications that could cause a delay in the processing of the algorithm. Additionally, there are two main reasons as to why the speedup for Jacobi step is still not near the ideal –16. Firstly, the Jacobi algorithm is an iterative process that aims at zeroing one off-diagonal element at a time, and the order in which these elements are chosen affects the number of iterations, as each iteration can undo the zeroes achieved in the previous one and, hence, the number of iterations needed to reach the convergence criterion may vary. Secondly, the Jacobi algorithm has data dependencies, so its convergence may vary depending on the nature of the data – e.g., its dynamic range.

**B. Communications assessment**

This subsection deals with the evaluation of communications, so as to check whether they become a bottleneck or not. To do so, the measurements previously shown have been divided in two transmission (hereafter, Tx) and processing (hereafter, Px). These measurements are gathered in Tables IV and V – case 1 and case 2, respectively. As can be observed, each table contains the three configurations described before: the sequential execution (1 cluster – 1 core), the first level of parallelism (16 clusters – 1 core) and the second one (16 clusters – 16 cores).

**TABLE IV. DETAILED SYSTEM EXECUTION TIME (MS) – CASE 1**

<table>
<thead>
<tr>
<th>Cluster-Core</th>
<th>1-1</th>
<th>16-1</th>
<th>16-16</th>
<th>Speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 1</td>
<td>Tx</td>
<td>329.8</td>
<td>329.8</td>
<td>329.8</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>541.4</td>
<td>50.9</td>
<td>5.1</td>
</tr>
<tr>
<td>Step 2</td>
<td>Tx</td>
<td>25,609.9</td>
<td>25,609.9</td>
<td>25,609.9</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>15,257.3</td>
<td>1,089.4</td>
<td>89.9</td>
</tr>
<tr>
<td>Step 3</td>
<td>Tx</td>
<td>0.34</td>
<td>0.34</td>
<td>0.34</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>1,759.8</td>
<td>1,759.8</td>
<td>350.9</td>
</tr>
</tbody>
</table>

The most important conclusion that can be extracted from these tables is that, indeed, the time needed for the data to be transmitted to the clusters is extremely high. The most representative example of this behavior is the covariance matrix computation (step 2): as shown in both tables, even in the sequential execution the transmission time already consumes more than 60% of the covariance execution time. As a result, when the processing is fully parallelized, more than 99% of the covariance execution time is dedicated to data transmission.

The main reason why the communications delay is so large is that the I/O interface is executed sequentially, that is, the communications are not parallelized. This is shown in both tables, where there are no speedups for any communications. For instance, in step 1, the whole image is needed. As a result, when only one cluster is used, the entire image is sent to it. Conversely, when several clusters are involved, the image is distributed among them – i.e., the same amount of data is transmitted –, but the transmission time remains the same because the I/O core sends the information sequentially. Furthermore, each transmission is blocking, so until the transmission to the first cluster has finished, the second transmission cannot begin.

**TABLE V. DETAILED SYSTEM EXECUTION TIME (MS) – CASE 2**

<table>
<thead>
<tr>
<th>Cluster-Core</th>
<th>1-1</th>
<th>16-1</th>
<th>16-16</th>
<th>Speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 1</td>
<td>Tx</td>
<td>721.9</td>
<td>721.9</td>
<td>721.9</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>1,177.4</td>
<td>86.1</td>
<td>11.2</td>
</tr>
<tr>
<td>Step 2</td>
<td>Tx</td>
<td>50,796.8</td>
<td>50,796.8</td>
<td>50,796.8</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>30,706.2</td>
<td>2,193.3</td>
<td>178.2</td>
</tr>
<tr>
<td>Step 3</td>
<td>Tx</td>
<td>0.34</td>
<td>0.34</td>
<td>0.34</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>1,849.7</td>
<td>1,849.7</td>
<td>370.3</td>
</tr>
<tr>
<td>Steps 4-5</td>
<td>Tx</td>
<td>721.9</td>
<td>721.9</td>
<td>721.9</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>999.6</td>
<td>68.0</td>
<td>9.2</td>
</tr>
<tr>
<td>Global</td>
<td>Tx</td>
<td>52,240.9</td>
<td>52,240.9</td>
<td>52,240.9</td>
</tr>
<tr>
<td></td>
<td>Px</td>
<td>34,732.9</td>
<td>4,247.1</td>
<td>642.3</td>
</tr>
</tbody>
</table>

Finally, related with the speedups achieved on the processing itself, it can be observed that, in general, they are rather large –more than 100, without taking into consideration the Jacobi step, as it is not distributed among the 16 clusters–, which proves that the parallelization potential of this architecture is very high.

As a result, the initial hypothesis has been proven correct: although the potential parallelism offered by an architecture
such as the MPPA-256-N is rather large, the communications within the chip quickly become the main bottleneck of data consuming algorithms, as PCA.

C. Second approach

The previous subsection has proven right the initial hypothesis about the communications. Now, as a proof of concept, the second approach proposed in Section III is analyzed.

This approach aims at proving that the main bottleneck of the system is the communications by removing them from the most consuming stage of the algorithm, which is the covariance matrix computation. To do so, instead of performing computation on the clusters, the host is used for the task, where the memory limitation is lower and, thus, there is no need for communicating data.

Specifically, Table VI gathers the results obtained when applying this solution, comparing them with those obtained in the previous one. As only the covariance matrix computation is involved, just the results of step 2 are provided. Additionally, the global execution times for both cases and in all the configurations are also presented, as well as the speedup obtained in each configuration when the mentioned computation is moved to the host.

As can be noticed, performing this operation on the host clearly provides considerably better results in terms of processing time, as, for instance, a speedup of more than 35 is achieved when comparing a sequential execution –1 cluster, 1 core– with the equivalent version in the host. Specifically, the speedup achieved with this change is such that even using all the resources of the chip provides far worse results than those generated by the sequential version of the system –but with the covariance computation in the host–, as the speedup achieved with the latter is still more than 22.

TABLE VI. COVARIANCE MATRIX COMPUTATION COMPARISON (MS)

<table>
<thead>
<tr>
<th>Configuration</th>
<th>Case 1</th>
<th>Case 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Step 2</td>
<td>Global</td>
</tr>
<tr>
<td>Clusters</td>
<td>40,870.5</td>
<td>41,405.2</td>
</tr>
<tr>
<td>Host</td>
<td>1,061.1</td>
<td>1,818.9</td>
</tr>
<tr>
<td>Speedup</td>
<td>98.5</td>
<td>98.5</td>
</tr>
<tr>
<td>Clusters</td>
<td>26,699.4</td>
<td>29,244.2</td>
</tr>
<tr>
<td>Host</td>
<td>1,061.1</td>
<td>3,602.2</td>
</tr>
<tr>
<td>Speedup</td>
<td>25.2</td>
<td>8.1</td>
</tr>
<tr>
<td>Clusters</td>
<td>25,726.1</td>
<td>26,840.2</td>
</tr>
<tr>
<td>Host</td>
<td>1,061.1</td>
<td>2,196.6</td>
</tr>
<tr>
<td>Speedup</td>
<td>24.2</td>
<td>12.2</td>
</tr>
</tbody>
</table>

To conclude this section, Fig. 4 provides a graphical comparison, for case 1, of the processing times for the four different implementations described before: (i) the sequential implementation –1 cluster, 1 core–; (ii) the first parallelization approach –16 clusters, 1 core–; (iii) the second parallelization –16 clusters, 16 cores–; and (iv) the one that moves the covariance computation to the host, while the rest of the algorithm maintains the configuration of the second approach.

Fig. 4 also provides the speedups of each implementation when compared to the sequential one; as a result, it can be observed that the speedup achieved when moving the covariance computation to the host is, approximately, 20; on the other hand, the speedup achieved when using only the MPPA resources is 12.5.

Finally, to graphically observe the generated results, Fig. 5 displays the one-band representation of the image obtained when applying the PCA algorithm to both cases 1 and 2 described before. As can be seen, these images contain a spatial representation of the most relevant spectral information, which drastically reduces data dimensionality.

Fig. 4. PCA performance comparison for case 1: execution time and speedup for different implementations

Fig. 5. PCA result (one band representation of the original image) for case 1 (left) and 2 (right)

V. DISCUSSION

After analyzing the results obtained during this research work, the present section compares the performance achieved using the MPPA architecture with studies presented in [26] and [27]. The former deals with the implementation of PCA on a Xilinx Virtex-7 XC7VX690T FPGA, while the latter uses the dataflow programming language RVC-CAL [28] to highlight...
the intrinsic parallelism of the algorithm and automatically distribute the workload among the available cores of an Intel Core i7-4790 — which gathers four processors running at 3.6 GHz — with 32 GB of RAM.

Before starting the comparison itself, it should be noted that, as HI technology initially aimed remote sensing, when working with hyperspectral images the most common datasets used in the related literature are the well-known AVIRIS Cuprite scene — available online1 and hereafter AV_Cuprite — and the AVIRIS Jasper Ridge scene — hereafter AV_JRidge. Both images have been widely used for validating the accuracy of dimensionality reduction algorithms, such as PCA. They are composed of several reference ground signatures of well-known minerals, which can be also extracted from the United States Geological Survey (USGS) database. These two images have been the ones used for evaluation purposes in [26].

However, the pervasive development of HI has led to its application to several other fields, such as the medical one. In this line, this work and the one in [27] use the same input dataset as introduced in Section III, which has been obtained from the HELICoiD project database [22].

As a result, these implementations cannot be compared directly, so the comparison will be performed in terms of the size of the images. In [26], the images have an approximate size of 50 MB (AV_Cuprite, 350×350×224) and 140 MB (AV_JRidge, 614×512×224)– lines×samples×bands–. Likewise, the images used in this work need 60.6 MB (case 1, 377×329×128) and 129.1 MB (case 2, 479×552×128)– again lines×samples×bands– to be stored.

From the dimensions of the images of both datasets can be inferred that, in the first one, the image is stored with a precision of 2 bytes, while in the second one the precision required is 4 bytes. In other words, the precision in the second dataset is twice the one applied in the first one, which is an important feature when comparing the implementations.

For each implementation, Table VII provides the global processing load of each image, as well as its size. Furthermore, it also provides a generic metric to simplify the comparison among them, which is the time needed for each implementation to process 1 MB of information.

As can be observed, the processing time grows linearly with the size of the image, as the time needed for processing 1 MB of information remains constant regardless of the implementation. Additionally, as shown in Table VII, the FPGA implementation achieves a processing cost of almost 30 ms per MB of information, while the MPPA implementation reaches a processing rate of 36 ms per MB. On the other hand, the Intel-i7 implementation (RVC-CAL) obtains a processing rate of 10 ms per MB.

This comparison shows, again, that the memory limitation plays a crucial role in the processing of data consuming algorithms, as the x86 architecture, which is the one with the least memory limitations thanks to its complex cache architecture, achieves the largest processing rate. The reason of this result is also that its cores run at 3.6 GHz, while those of the MPPA run at 600 MHz.

However, the x86 architecture cannot compete with both FPGAs and manycore platforms in terms of power consumption. For this metric, the MPPA-256-N used during this research clearly outperforms both the Xilinx Virtex-7 XC7VX690T FPGA and the Intel Core i7-4790, as they consume 5 W [14], 30-40W [29] and 84W3, respectively in average conditions. As power consumption is an important feature for clinical applications, trying to minimize this value is very important for designing a portable prototype.

Consequently, the implementation developed during this research work can be considered competitive when compared to those described before, as it outperforms its competitors in terms of power consumption. Furthermore, when compared to the FPGA implementations, it can be observed that the results are equivalent, considering that the precision used in this implementation is twice the precision of the FPGA one.

<table>
<thead>
<tr>
<th>Implementation</th>
<th>Database</th>
<th>Global processing time (MB)</th>
<th>Size (MB)</th>
<th>Processing time per MB</th>
</tr>
</thead>
<tbody>
<tr>
<td>FPGA</td>
<td>AV_Cuprite</td>
<td>1,490.0</td>
<td>50.0</td>
<td>29.8</td>
</tr>
<tr>
<td>FPGA</td>
<td>AV_JRidge</td>
<td>4,170.0</td>
<td>140.0</td>
<td>29.8</td>
</tr>
<tr>
<td>INTEL-i7</td>
<td>HELICoiD_C1</td>
<td>614.8</td>
<td>60.6</td>
<td>10.1</td>
</tr>
<tr>
<td>INTEL-i7</td>
<td>HELICoiD_C2</td>
<td>1,265.1</td>
<td>129.1</td>
<td>9.8</td>
</tr>
<tr>
<td>MPPA</td>
<td>HELICoiD_C1</td>
<td>2,196.6</td>
<td>60.6</td>
<td>36.2</td>
</tr>
<tr>
<td>MPPA</td>
<td>HELICoiD_C2</td>
<td>4,644.7</td>
<td>129.1</td>
<td>36.0</td>
</tr>
</tbody>
</table>

VI. Conclusion

This paper has presented an analysis of the parallelism of a PCA algorithm, including its adaptation to the manycore architecture MPPA-256-N from Kalray, and its comparison with other state-of-the-art studies. The proposed implementation aims at adapting the algorithm to process hyperspectral image analysis in real-time, so as to help surgeons in locating brain tumors during surgical procedures.

On the one hand, the obtained results show that, as expected in such a data consuming algorithm, communications among processing units quickly become the main bottleneck of the system. Consequently and, as a proof of concept, two different implementations have been carried out: first, the most consuming stage of the algorithm has been located and has been parallelized over the clusters and then it has been processed sequentially in the host of the platform. The analysis of these implementations has proven that, indeed, communications are the main bottleneck, as they can consume up to a 99% of the processing time. Therefore, a trade-off between the level of parallelism and the increase in internal communications must be met.

3. http://ark.intel.com/products/80806/Intel-Core-i7-4790-Processor-8M-Cache-up-to-4-00-GHz
On the other hand, the exploitation of the available resources on the MPPA-256-N demonstrates that, in favorable cases, the intrinsic parallelism of the algorithm can be fully exploited. Speedups up to 170 have been achieved on the processing of certain stages of the algorithm on a 256-core implementation. Although these speedups decrease to 20 due to communication delays, the real-time objective of one hyperspectral image per minute is still reached for the application at hand. In these conditions, the PCA algorithm consumes less than a 10% of the available processing time, keeping processing resources for other hyperspectral image analysis tasks.

This implementation has been compared to state-of-the-art implementations of the same algorithm on an FPGA and an x86 architecture, the proposed solution proving to be competitive and providing better power efficiency rates.

As a final conclusion, experimental results have demonstrated that manycores are promising architectures for medical hyperspectral image processing. Future studies will investigate the possibilities to increase the effective communication bandwidth, so as to achieve speedups closer to 256x, which is the ideal speedup achievable with this architecture. Furthermore, OpenMP and OpenCL implementations of the algorithm will be compared to the current POSIX results.

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REFERENCES


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