Abstract

The communication cost plays a key role in the performance of many parallel algorithms. In the particular case of the one-sided Jacobi method for symmetric eigenvalue and eigenvector computation the communication cost of previously proposed algorithms is mainly determined by the particular ordering being used. In this paper we proposed two novel Jacobi orderings: the permuted-BR ordering and the degree-4 ordering, aimed at efficiently exploiting the multi-port capability of a hypercube. It is shown that the former is nearly optimal for some scenarios and the latter outperforms previously known orderings by a factor of two.

1. Introduction

The one-sided Jacobi method for symmetric eigenvalue computation is very suited for its application on a multiprocessor since it exhibits a high parallelism and potentially low communication requirements [5]. The one-sided method uses a series of similarity transformations to make the original symmetric matrix converge to a diagonal form. Every similarity transformation zeroes one off-diagonal element and its symmetric. The elements of the matrix can be zeroed in any order, giving place to different Jacobi orderings and, as a result, to different one-sided Jacobi algorithms. In addition to the convergence rate, different Jacobi orderings differ in the communication requirements of the resulting parallel algorithm.

In multicomputers, the communication overhead plays an important role on the performance of any particular algorithm [1]. In this paper, we focus on multicomputers with a hypercube interconnection topology and with multiple ports per node [14]. In such scenario one may design algorithms that communicate multiple messages simultaneously through different links of the same node (communication parallelism), which may result in a significant reduction in the communication overhead. One-sided Jacobi algorithms previously proposed for hypercubes make a poor utilization of the multi-port capability because, at any given time, the information to be communicated by every node is sent through one (or at most two) link, constraining in this way the exploitation of communication parallelism.

In [9] a method was developed to design parallel algorithms that efficiently exploit the multi-port capability in hypercubes. The method requires the specification of the original problem in the form of a CC-cube algorithm, whose properties will be described later. The method reorganizes the computation in a systematic way to introduce the appropriate level of communication parallelism in order to efficiently exploit the multi-port capability.

It will be shown in this paper that one-sided Jacobi computation can take the form of a CC-cube algorithm. Thus, the method in [9] can be used to reduce the communication cost in multi-port hypercubes. We will show first that the impact of the method when applied to known Jacobi algorithms is limited by the structure of the Jacobi ordering used in these algorithms. The key contribution of this paper is the proposal of two novel Jacobi orderings, the permuted-BR and the degree-4 orderings, which enable an efficient exploitation of the multi-port capability, through the use of the method described in [9]. The permuted-BR ordering has a performance that tends asymptotically (for large matrices) to 80% of a lower bound. The degree-4 ordering has a worse asymptotic performance but it behaves better for small matrices. In this case, it reduces the communication overhead of the algorithm to the half when compared with previous Jacobi orderings.

The rest of this paper is organized as follows. Section 2 provides a brief review of the one-sided Jacobi method and the orderings proposed for its implementation on a hypercube multicomputer. Section 3 presents the novel orderings. Section 4 evaluates the performance of the proposed orderings. Finally, the main conclusions are summarized in section 5.

2. Preliminaries

In this paper we focus on efficient algorithms for eigenvalues and eigenvectors computation of symmetric matrices through the one-sided Jacobi method on multiprocessors with a hypercube interconnection topology and
multiple ports per node. Below we summarize what it is implied by each one of these terms. Then, we review the most relevant related work and point out the motivation for this work.

2.1. Target architecture

The target architecture on which the proposed algorithm is to be executed is a multi-port hypercube multicomputer. A hypercube multicomputer of dimension $d$, which is also called a $d$-cube multicomputer, consists of $2^d$ processors such that they can be labelled from $0$ to $2^d-1$ in such a way that all processors that differ in exactly one bit in the binary representation of their label are neighbors (they are connected by a link) [10]. The link that connects neighbor nodes whose labels differ in the $i$-th bit will be referred to as link $i$. The label of the link, which ranges from $0$ to $d-1$, will also be called the dimension of the link. As an example, node 2 uses link 1 (or dimension 1) to send messages to node 0.

A multi-port multicomputer is distinguished by the fact that every node can simultaneously be transmitting/receiving messages from more than one link. In particular, in an all-port configuration every node can send and receive a message through each of its $d$ links simultaneously. A configuration in which every node can send and receive a message from just one of its links is called one-port configuration [14].

2.2. The one-sided Jacobi method and parallel Jacobi orderings

The one-sided method for symmetric eigenvalue and eigenvector computation works with two matrices: $\bar{A}$ (initially set to the original matrix $A$) and $U$ (initially set to the identity matrix $I$). In every iteration of the method one similarity transformation is applied to zero one off-diagonal element of $A$ and its symmetric one [15]. The key feature of the one-sided method is that the computation and application of the similarity transformation that zeroes elements $(i,j)$ and $(j,i)$ of $A$ uses only columns $i$ and $j$ of $\bar{A}$ and $U$. For this reason, such a transformation will be also referred to as the pairing of columns $i$ and $j$. According to this, transformations that use disjoint pairs of columns can be applied in parallel, since they do not share any data. This is what makes the one-sided Jacobi method to be very suitable for parallel implementations [5].

In the literature, the process of zeroing every off-diagonal element exactly once is called a sweep. Since a similarity transformation may fill in elements that had been zeroed in previous transformations, several sweeps are required until the resulting matrix converges to a diagonal matrix. Since similarity transformations preserve the eigenvalues, the elements in the diagonal of the resulting matrix coincide with the eigenvalues of the original matrix.

If $A$ is an $m \times m$ matrix, a total of $m(m-1)/2$ similarity transformations are required to complete a sweep. Assuming $m$ even, these transformations can be organized into a maximum of $m-1$ groups of $m/2$ independent transformations each (transformations that involve disjoint pairs of columns). Such an organization of the sweep is called a parallel Jacobi ordering and every group of independent transformations is usually referred to as a step.

Parallel Jacobi orderings offer a good framework for the design of parallel implementations, since the computation involved in every step can be evenly distributed among the nodes of the multicomputer, and carried out without any communication. Communication is required however at the end of every step, when the columns of matrices $\bar{A}$ and $U$ must be exchanged to obtain the pairs required for the next step. Such a communication will be called a transition. A parallel Jacobi ordering is suitable for a multicomputer if the transitions between steps use a communication pattern that matches the topology of the interconnection network of the machine.

2.3. Related work

Jacobi orderings for different parallel architectures can be found in the literature [4,6,8,13]. Regarding hypercube multicomputers, which is the target architecture considered in this work, the most relevant proposals are [3,7,12]. All these proposals share the following features: (a) they use a minimum number of steps per sweep, achieving in this way a perfect load balance, and (b) the transitions between steps can be implemented through communication between neighbors in the hypercube. Particularly relevant to our work is the Block Recursive (BR) ordering proposed in [12]. The distinguishing feature of this ordering is that in every transition all nodes exchange information through the same dimension of the hypercube. This is one of the requirements for an algorithm to belong to the CC-cube class, and therefore, to enable the application of the communication pipelining technique. Below we describe the BR ordering in detail since some of the orderings proposed in this paper are obtained through the application of some transformations to it.

2.3.1. BR ordering

The BR ordering was originally proposed in [7] and reviewed later in [12], where the ordering was completely specified and its correctness was shown.

Given a $d$-cube and a $m \times m$ matrix, the $m$ columns of matrices $\bar{A}$ and $U$ are grouped into $2^{d+1}$ blocks of $m/2^{d+1}$ columns each. Then, a pair of blocks are allocated to each node of the $d$-cube. The algorithm to perform the first sweep proceeds as follows (all the nodes are executing the following code in parallel):

1. If $m$ is not a power of 2, then the number of columns per block will differ in one unit at most. This will result in a slight load imbalance.
1) Pair each column of a block with the remaining columns of the same block.

2) Pair each column of a block with all the columns of the other block allocated to the same processor.

3) Exchange one of the blocks with a neighbor along a given dimension of the hypercube (transition).

4) Go to (2) if not all the columns have been paired (i.e., (2) and (3) are repeated $2d+1$ times).

In order to achieve that every pair of blocks is paired exactly once, the transitions (i.e., the exchange operations required by (3)) must be performed in a particular order. To precisely define a exchange operation it suffices to identify the link (i.e., the dimension) that is used by it. Below we describe the sequence of transitions implied by the BR ordering as described in [12].

The $2d+1$ steps of the ordering are grouped into three different types of phases: exchange, division and last transition. A phase is just one or more consecutive steps, each of them followed by a transition. A sweep consists of $d$ exchange phases, each one of them followed by a division phase, and a final last transition phase. The exchange phases are numbered from $d$ to 1. The exchange phase $e$ ($e \in [d,1]$) consists of $2^{d_e}-1$ steps and transitions. The sequence of links (i.e., dimensions) that define the transitions is denoted by $D_e^{BR}$ and can be systematically generated as follows:

$$D_0^{BR} = \langle 0 \rangle$$

$$D_i^{BR} = \langle D_{i-1}^{BR}, i-1, D_{i-1}^{BR} \rangle \quad 1 < i \leq d$$

For instance, the sequence of links for $e=4$ is $D_4^{BR} = \langle 010201030102010 \rangle$.

The exchange phase $e$ is followed by a division phase that consists of just one step and one transition through link $e$. Finally, the last transition phase consists of one step and one transition through link $d-1$.

The second and next sweeps use the same algorithm but after applying a permutation to the links. In particular, the permutation corresponding to sweep $s$ (assuming $s=0$ for the first one) is defined as follows:

$$\sigma_0(i) = i$$

$$\sigma_s(i) = (\sigma_{s-1}(i) - 1) \mod d \quad \text{for } i=0...d-1$$

After $d$ sweeps, the links are used again in the order described for the first sweep.

2.4. Motivation

The previously summarized BR algorithm is efficient for a one-port hypercube since the system is using all the available ports for every transition. However, for a multi-port architecture it achieves the same performance as for a one-port because it uses just one out of the $d$ links of each node at the same time.

In [9] we proposed a systematic transformation of a class of algorithms that is called CC-cube algorithms in order to optimize them for a multi-port environment. Such transformation is called communication pipelining. Every process of the CC-cube algorithm executes a loop that iterates $K$ times. Every iteration consists of a certain computation and an exchange of information through one of the hypercube dimensions (all the processes use the same dimension in a given iteration of the loop). The key idea of communication pipelining is to decompose the computation in every iteration into $Q$ packets and reorganize the computation as follows. Every process of the new algorithm (the pipelined CC-cube algorithm) compute the first packet of the computation in the first iteration of the original CC-cube and exchanges the result with the corresponding neighbor. Then, it computes the second packet of the first iteration and the first packet of the second one. The results of these packets can be sent in parallel to the corresponding neighbors, using simultaneously two hypercube dimensions. Proceeding in this way, the pipelined CC-cube can send an increasing number of messages in parallel, being able to exploit the multi-port feature.

The value of $Q$ is called pipelining degree and in [9] it is shown how to determine the pipelining degree that minimizes the execution time of any particular CC-cube algorithm and any particular hypercube architecture. In this section we only summarize the result of the application of the communication pipelining technique and omit any detail that is not strictly necessary to understand this paper. The interested reader is referred to the original paper for such details.

If $Q$ is not higher than $K$, the pipelined CC-cube algorithm is said to work in shallow pipelining mode. Every process can send up to $Q$ messages simultaneously through different links if the architecture supports it. These $Q$ links correspond to all the subsequences of $Q$ elements of the sequence that describes the link usage in the CC-cube algorithm.

For instance, if in the original CC-cube $K=7$ and communications are carried out through links 0, 1, 0, 2, 0, 1 and 0, the pipelined CC-cube with $Q=3$ will perform a series of computations each one followed by communication through links 0-1-0, 1-0-2, 0-2-0 and 2-0-1 respectively. This part of the pipelined CC-cube is called the kernel and has $K-Q$ stages (computation followed by communication).

Like in the software pipelining technique [11], the kernel is preceded by a prologue and followed by an epilogue. The prologue consists of $Q-1$ stages and each of these stages consists of a computation followed by a communication though an increasing number of the first links of the original sequence. In the previous example, the prologue

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1. The pairing of two blocks is defined as all the pairings that consist of two columns from different blocks.

2. $a-b-c$ means that three messages are sent in parallel through links $a$, $b$, and $c$. If several messages use the same link, they are packed into a single message and sent together.
consists of two stages that use the following links: 0 and 0-1 respectively. Similarly, the epilogue consists of \( Q - 1 \) stages and each of these stages consists of a computation followed by a communication through a decreasing number of the last links of the original sequence. In the previous example, the epilogue consists of two stages that use the following links: 1-0 and 0.

If \( Q \) is higher than \( K \), the pipelined CC-cube is said to work in deep pipelining mode. In this case, the prologue and epilogue have \( K-1 \) stages each and the kernel has \( Q\cdot K+1 \) stages. In each kernel stage, \( K \) messages can be sent in parallel (one packet from every iteration of the original CC-cube). For instance, if \( K=3 \) and the links are used in the order 0, 1, and 0, the pipelined CC-cube with \( Q=100 \) has a prologue with two stages that use links 0 and 0-1. Every one of the 98 stages of the kernel use links 0-1-0. Finally, the two stages of the epilogue use the links 1-0 and 0.

Communication pipelining can be applied to any CC-cube algorithm that meets some requirements as specified in [9]. In particular, it cannot be applied to the whole BR algorithm but it can be applied to every exchange phase, which are the most time-consuming part of the BR algorithm. Communication pipelining can reduce the communication overhead of the algorithm by a factor of \( \alpha \), which produces very important improvements in some problems where the communication operations have an important weight. Unfortunately, the application of communication pipelining to the BR algorithm can reduce the communication overhead by a factor not higher than 2 regardless of the value of \( d \). This is basically due to the properties of the BR ordering: in the sequence \( D_e^{BR} \), which defines the order in which the links of each node are used in exchange phase \( e \), any subsequence of \( Q \) consecutive elements has at least \( \left\lfloor \frac{Q}{2} \right\rfloor \) elements equal to 0. In consequence, the capability to send simultaneously messages through multiple links can provide a reduction in the communication cost by a factor not higher than 2, since about half of the messages must be sent through the same link (link 0).

In this paper we propose novel Jacobi orderings that exhibit a more balanced use of the links of each node and thus, communication pipelining can provide a much higher improvement than for the BR ordering.

### 3. New Jacobi orderings

All the new orderings proposed in this section are obtained from the BR ordering by replacing the sequence \( D_e^{BR} \) (\( e \in \{d,1\} \)) by alternative sequences that, beside generating all the required pairings of columns, exhibit a more balanced use of the links of every node, enabling a more efficient exploitation of the multi-port capability.

The first proposal is optimal but it is only defined for small hypercubes. The second one has a good asymptotic performance for large problem sizes. Finally, third one behaves better for small problem sizes. This section also includes a discussion on the convergence of the new orderings.

#### 3.1. Minimum–\( \alpha \) ordering

Let us denote by \( D_e \) the sequence of links used in exchange phase \( e \) for the first sweep (e.g. \( D_e^{BR} \) for the BR ordering). Communication pipelining allows each node to send messages through all or several consecutive links in \( D_e \), depending on the degree of pipelining (value of \( Q \)). In the case of deep pipelining, all the links of the sequence \( D_e \) are used in each communication operation (except for the prologue and epilogue). A message of a fixed size \( S \) must be sent per each element of \( D_e \) through the corresponding link. If there are repeated elements in \( D_e \), then the corresponding messages are combined into a single message. Since all the elements in \( [0,e-1] \) must appear at least once in \( D_e \), otherwise, it would be impossible to generate all the pairings required by exchange phase \( e \) the time to perform the communication operation in every kernel stage, in an all-port hypercube is \( e T_w + \alpha ST_w \), where \( T_w \) is the start-up time to initiate the communication through one link, \( S \) is the fixed message size, \( T_w \) is the transmission time per data element, and \( \alpha \) is the maximum number of messages that must be combined into a single message and sent through the same link, that is, \( \alpha \) is the maximum number of repetitions of the same link in the sequence \( D_e \).

Since \( T_w \) and \( S \) are fixed for a given architecture and \( S \) and \( e \) are fixed for a particular problem, the cost of each communication operation just depends on \( \alpha \), which is a function of the sequence \( D_e \).

In the BR ordering, the value of \( \alpha \) corresponding to \( D_e^{BR} \) is \( 2^{e-1} \). The communication overhead can be minimized by finding an alternative sequence \( D_e \) whose \( \alpha \) is minimum.

The sequence \( D_e \) can be also regarded as the definition of a hamiltonian path in an \( e \)-cube. This is so because the half of the columns of \( U \) initially allocated to each node must visit a different node after each exchange operation and thus, they must have visited all the nodes after the \( 2^{e-1} \)-1 exchange operations that make up the exchange phase \( e \). For instance, the exchange phase \( e=3 \) of the BR ordering is defined by \( D_3 = <0102010> \). Starting at any node of the hypercube and moving through the links that form this sequence in the order they appear, all the nodes of a 3-cube are visited exactly once.

We can then conclude that the problem of defining alternative sequences \( D_e \) can also be stated as the problem of finding alternative hamiltonian paths in an \( e \)-cube. The resulting sequence consists of the link identifiers that have been used for traversing the hypercube. Since there are many different hamiltonian paths in a hypercube, there are many alternative Jacobi orderings that can be generated in this way.
Minimizing the communication cost of exchange phase $e$ when communication pipelining is used can be achieved by computing all the hamiltonian paths of an $e$-cube and choosing that with the minimum $\alpha$. Since any sequence $D_e$ has $2^{e-1}$ elements and all values in $[0, e-1]$ must appear in $D_e$ at least once, the minimum value for $\alpha$ is given by:

$$\alpha \geq \left\lceil \frac{2^e - 1}{e} \right\rceil$$

Unfortunately, finding a hamiltonian path with minimum $\alpha$ is an NP-hard problem and thus, it can be solved only for small values of $e$. In particular, following this approach we could compute the sequences with minimum $\alpha$ for values of $e$ lower than 7. These sequences are called $D_e^{\min-\alpha}$ and are the following:

- $D_2^{\min-\alpha} = \langle 010 \rangle$, $\alpha = 2$
- $D_3^{\min-\alpha} = \langle 010210 \rangle$, $\alpha = 3$
- $D_4^{\min-\alpha} = \langle 010203120312 \rangle$, $\alpha = 4$
- $D_5^{\min-\alpha} = \langle 01020103010214123213234123 \rangle$, $\alpha = 7$
- $D_6^{\min-\alpha} = \langle 01020103010201040102131252131232312343 \rangle$, $\alpha = 11$

The Jacobi ordering that uses the sequences $D_e^{\min-\alpha}$ is called the minimum-$\alpha$ ordering. However, it is only defined for hypercubes with $d < 7$.

3.2. Permuted-BR ordering

The Jacobi ordering proposed in this section, which is called the permuted-BR ordering, uses the sequence denoted by $D_e^{p-BR}$. This sequence is defined for any value of $e$ and it will be shown that the value of $\alpha$ for the sequence is close to the minimum. As reflected in the name of the ordering, the sequence $D_e^{p-BR}$ is obtained by applying some permutations to the link identifiers in $D_e^{BR}$. This transformation is based on the following property.

Property 1. Let $D_e$ be a sequence of links that define a hamiltonian path in an $e$-cube. Let $\sigma$ be any permutation of the link identifiers. If we apply $\sigma$ to any subsequence of $D_e$ that corresponds to a hamiltonian path of a subcube, then the resulting sequence defines also a hamiltonian path of the $e$-cube.

Proof. Given a $n$-cube, if we apply any permutation to all its links and re-label the nodes accordingly, the resulting graph is isomorphically equivalent to the initial one. Thus, if a link permutation is applied to a sequence of link identifiers that define a hamiltonian path in a $n$-cube, the resulting sequence is also a hamiltonian path in the same $n$-cube.

Examples

- $\langle 010 \rangle$ is a hamiltonian path in a 2-cube. If links 0 and 1 are exchanged in the whole sequence, the resulting sequence $\langle 101 \rangle$ is also a hamiltonian path.

- $\langle 0102010 \rangle$ is a hamiltonian path in a 3-cube. If we apply the previous permutation to the subsequence made up of the last 3 elements, which is a hamiltonian path in a 2-cube, we obtain the sequence $\langle 010210 \rangle$, which is also a hamiltonian path in a 3-cube.

Definition 1. A sequence that corresponds to a hamiltonian path of a $d$-cube will be called a $d$-sequence, or a $d$-subsequence if it appears within a larger sequence.

Obviously, $D_e^{p-BR}$ is an $e$-sequence. Otherwise, it could not be used to implement exchange phase $e$ (see [12] for a proof of the correctness of the BR ordering). Due to the way $D_e^{p-BR}$ is built, (see section 2.3.1), it is an $e$-sequence that consists of two $(e-1)$-subsequences separated by the link $e-1$. Each one of these subsequences is in turn composed of two $(e-2)$-subsequences separated by the link $e-2$. In general, each $(e-i)$-subsequence is composed of two $(e-i-1)$-subsequences separated by the link $e-i-2$. This is a useful property of the $D_e^{p-BR}$ sequence that will be used in the following.

3.2.1. Methodology to generate the $D_e^{p-BR}$ sequence

The $D_e^{p-BR}$ sequence is obtained by applying some transformations to the $D_e^{BR}$ sequence. Such transformations meet property 1 stated above and in consequence, the resulting sequences are guaranteed to be valid in the sense that they also correspond to a hamiltonian path of an $e$-cube. The objective of this transformations is to obtain a sequence in which the number of repetitions of each link identifier is about the same in order to minimize the value of $\alpha$.

The transformation is applied to $D_e^{p-BR}$ in order to obtain $D_e^{BR}$. Each transformation consists of a link permutation that is applied to some subsequences of the whole sequence. In particular, transformation $k$, $k$ being an integer from 0 to $\lfloor \log_2 (e-1) \rfloor - 1$, consists of a link permutation applied to every other $(e-k-1)$-subsequence, starting at the second one. That is, the first transformation is applied to the second $(e-1)$-subsequence. The second transformation is applied to the second and forth $(e-2)$-subsequences and so on. Since property 1 holds, the resulting sequence is also a valid sequence to implement the exchange phase $e$.

For each transformation $k$, the link permutation applied to the second $(e-k-1)$-subsequence is defined as the transposition of the following pairs of link identifiers:

$$i \leftrightarrow \left\lfloor \frac{(e-1)}{2^k} \right\rfloor - 1 - i$$

$$i \in [0, \lfloor (e-1)/2^k \rfloor - 1], \quad k \in [0, \lfloor \log_2 (e-1) \rfloor - 1]$$

The link permutation applied to any of the remaining subsequences (just to every other) is derived by compounding the permutation applied to the second $(e-k-1)$-subsequence with all the permutations applied in previous transformations to any subsequence that includes the current one.
3.3. Degree-4 ordering

The permuted-BR ordering proposed in the previous section is expected to work well when the pipelined CC-cube works in deep pipelining mode and the cost is dominated by the kernel phase. In other words, when \( Q \) is very large. In the case that the matrix size is not large enough to enable large values of \( Q \) the pipelined CC-cube will work in shallow pipelining mode. In this mode of operation the quality of sequence \( D_e \) is no longer related to the value of \( \alpha \) since only a subsequence of the links in \( D_e \) are used in every communication operation. In this context, the sequence \( D_5^{p-BR} \) proposed before is not adequate since when considering small subsequences of links, nearly half of the elements are equal (see for instance the \( D_5^{p-BR} \) sequence at the end of section 3.2.1).

What would be desirable for shallow pipelining is a sequence such that any subsequence of length \( Q \) consist of different elements if \( Q \) is not higher than \( e \), or it consists of all the elements from 0 to \( e-1 \) repeated the same number of times, with a maximum difference of one if \( Q \) is not multiple of \( e \). Finding such optimal sequences is still an open problem with unknown solution. The Jacobi ordering proposed in this section, called degree-4 ordering, uses the sequence \( D_e^{D4} \) whose main feature is that most subsequences of length 4 consist of different elements and thus, it outperforms previous orderings for shallow pipelining.

Definition 2. We say that a sequence \( D_e \) has degree \( n \) if the majority of subsequences of size \( n \) consists of different elements but the majority of subsequences of size \( n+1 \) have less than \( n+1 \) different elements. For instance, \( D_5^{BR} \) has degree 2 for any \( e \).

Shallow pipelining can reduce the communication cost of the Jacobi method by a factor near to \( n \) if the ordering being used is defined by sequences \( D_i \) of degree \( n \). The new ordering proposed in this section is called degree-4 ordering since it uses sequences of degree 4, enabling a reduction of the communication cost by a factor of about 4.

Definition 3. The sequence \( D_e^{D4} \) can be systematically generated for any value of \( e \) as follows:

\[
E_3 = \langle 0123012 \rangle \\
E_i = \langle E_{i-1}, i, E_{i-1} \rangle \\
D_e^{D4} = \langle E_{e-1}, 1, E_{e-1} \rangle \\
e \geq 4
\]

For instance, \( D_5^{D4} = \langle 01230124012301240123012401 \rangle \). Note that the sequence \( D_e^{D4} \) has degree four because only four central subsequences of length 4 have different elements (\( \langle 0121 \rangle, \langle 1210 \rangle, \langle 2101 \rangle \) and \( \langle 1012 \rangle \) in the previous example). This is true for any \( e \geq 3 \). When \( e \) is large, these subsequences have a negligible effect on performance.

It remains to be shown that \( D_e^{D4} \) is an \( e \)-sequence and, therefore, it can be used to implement the exchange phase \( e \). This is shown in the following theorem, that requires a previous lemma.

\[
\begin{array}{|c|c|c|c|}
\hline
e & \alpha & \text{lower bound} & \alpha / \text{lower-bound} \\
\hline
7 & 23 & 19 & 1.21 \\
8 & 43 & 32 & 1.34 \\
9 & 67 & 58 & 1.16 \\
10 & 131 & 103 & 1.28 \\
11 & 289 & 187 & 1.55 \\
12 & 577 & 342 & 1.69 \\
13 & 776 & 631 & 1.23 \\
14 & 1543 & 1171 & 1.32 \\
\hline
\end{array}
\]

Table 1. Value of \( \alpha \) corresponding to the permuted-BR ordering and compared to a lower bound.
Lemma 1. Let \(i\) be an arbitrary node of an \(e\)-cube. Let \(f\) be the node that is reached from \(i\) by following the path identified by the links in \(D^4_e\). Then, nodes \(i\) and \(f\) are neighbors in dimension 1.

Proof. The proof is based on induction. It is easy to check that the lemma holds for \(D^4_e\). We show now that if it holds for \(D^4_e\), then it holds also for \(D^{e+1}_e\). First, rewrite \(D^{e+1}_e\) as follows:

\[
D^{e+1}_e = \langle E_{e-1}, E_e, E_{e-1}, 1, E_{e-1}, e, E_{e-1}, e, E_{e-1} \rangle = \langle E_{e-1}, e, D^4_e, e, E_{e-1} \rangle
\]

Any \((e+1)\)-cube can be decomposed into two \(e\)-cubes that are joined by links in dimension \(e\). As shown in figure 1, the path described by \(D^{e+1}_e\) consists of path \(E_{e-1}\) through subcube 0, since link \(e\) does not belong to \(E_{e-1}\), a jump to subcube 1 (through dimension \(e\)), a path \(D^4_e\) through subcube 1, a return to subcube 0 and a path \(E_{e-1}\) through subcube 0. By induction hypothesis, nodes \(b\) and \(c\) in subcube 1 are neighbors in dimension \(e\). Since nodes \(a\) and \(b\) are neighbors in dimension \(e\), and nodes \(d\) and \(c\) too, \(a\) and \(d\) must be neighbors in dimension \(1\). Notice that the path from \(i\) to \(a\), then to \(d\) through dimension \(1\), and from \(d\) to \(f\) uses the links as in \(D^4_e\) (see definition 3). Again, by induction hypothesis, \(i\) and \(f\) must be neighbors in dimension 1.

Theorem 1. \(D^4_e\) is an \(e\)-sequence.

Proof. We use again induction. The theorem holds for \(D^4_e\). Assume that \(D^4_e\) is an \(e\)-sequence. We refer again to figure 1, which shows the structure of path \(D^{e+1}_e\). As observed in the previous lemma, the path from \(i\) to \(a\), then to \(d\) through dimension \(1\), and from \(d\) to \(f\) uses the links as in \(D^4_e\). Therefore, \(D^{e+1}_e\) is a hamiltonian path in subcube 1 that is interrupted in the middle to jump to subcube 1, which is traversed by a hamiltonian path before returning to subcube 0. As a result \(D^{e+1}_e\) is a hamiltonian path of an \((e+1)\)-cube, that is, it is an \((e+1)\)-sequence.

3.4. Convergence of the new orderings

The detailed analysis of the convergence of the proposed Jacobi orderings is out of the scope of this work. In this section, we give the results of some convergence tests. Table 2 shows the number of sweeps required by BR, permuted-BR and degree-4 orderings, for different matrix sizes \((m)\) and different number of nodes \((P)\). The test matrices have been generated with random numbers on the interval \([-1,1]\) having a uniform distribution. Since 30 different matrices have been tested for every value of \(m\) and \(P\), the average number of sweeps is shown. The results suggest that the convergence rates of the proposed orderings appear to be practically the same as that of the BR ordering.

<table>
<thead>
<tr>
<th>(m)</th>
<th>(P)</th>
<th>BR</th>
<th>permuted-BR</th>
<th>degree-4</th>
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<td>3.76</td>
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<td>4.03</td>
<td>4.03</td>
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<td>5.03</td>
<td>5.16</td>
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<td></td>
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<td>5.00</td>
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</tr>
</tbody>
</table>

Table 2. Convergence rate of the different orderings

4. Performance evaluation

In the previous section we have argued that the performance of the permuted-BR ordering is nearly optimal for deep pipelining and that the degree-4 ordering can reduce the communication cost by a factor near to 4 for shallow pipelining. We confirm these conclusions in this section with some performance figures obtained through analytical models of performance.

Figure 2 shows the communication cost of the different orderings for a varying size of a hypercube multicomputer; a varying matrix size; and assuming that the transmission time per element and the start-up time, \(T_w\) and \(T_v\), are 100 and 1000 time units respectively. The communication cost has been computed through the models developed in [9]. The communication cost is given in relation to the cost of the CC-cube algorithm using the BR ordering. As a reference, the figure 2 also includes the lower bound on the communication cost.

When communication pipelining is used, the optimum degree of pipelining for each particular hypercube dimension and matrix size was chosen, using the procedure presented in [9] to compute it. In the case of the permuted-BR ordering, the filled symbols indicate that deep pipelining has been used in all the exchange phases, while unfilled symbols indicate that shallow pipelining is used in the first (the most time consuming) exchange phases.

It can be seen in figure 2 that the communication cost of the pipelined CC-cube algorithm when the BR ordering is used is about one half of that of the original CC-cube.
The performance of the permuted-BR ordering approaches the lower bound when deep pipelining is used. However, when the hypercube size forces the use of shallow pipelining, it tends to be similar to the BR ordering. On the other hand, the degree-4 ordering exhibits a more stable performance behavior, since its communication cost is about one forth of the cost of the CC-cube BR algorithm in all the considered scenarios.

5. Conclusions

We have proposed two novel Jacobi orderings: the permuted-BR ordering and the degree-4 ordering. These new orderings significantly outperform previous orderings in a hypercube with multiple ports when using the communication pipelining technique. The permuted-BR ordering is obtained by applying a series of permutations to the BR ordering. The result is an ordering that makes a nearly balanced use of all the hypercube links, which results in a performance close to the optimum when all the links of the hypercube are used simultaneously. Depending on the start-up cost and the transmission cost there are cases in which the most efficient solution is to use just a few number of links simultaneously. In this scenario, the permuted-BR ordering is not nearly optimal anymore. For such cases, we have proposed the degree-4 ordering, which outperforms by a factor of 2 the BR ordering.

References


1. In the case of the permuted-BR ordering we have used the sequences \( D_{e}^{BR} \) for all values of \( e \). For \( e < 7 \) we could have used the optimal sequences \( D_{e}^{opt} \). However this would have had a negligible impact on the performance since it would affect only the less time-consuming exchange phases.
In the following it is shown that the value of $\alpha$ corresponding to the $D_{e}^{p-BR}$ sequence tends to be 1.25 times the lower bound given by $\left\lceil \frac{2^{e-1}}{2} \right\rceil$ for large values of $e$.

To simplify the expressions we assume that $e-1$ is a power of two. For any other value of $e$, the corresponding $\alpha$ will take an intermediate value between the $\alpha$ of the two surrounding powers of two and thus, the asymptotic behavior observed for powers of two will also apply to any other value.

Let $e = 2^{k}+1$. In this case, the sequence $D_{e}^{p-BR}$ will be obtained after applying $S$ transformations to the sequence $D_{e}^{BR}$. To illustrate the following development we will use the particular case of $e=17$. This case requires four transformations, which are shown in figure 3.

We denote by $D_{e}^{p-BR}(k)$ the sequence that is obtained after transformation $k$, with $k \in [0, S-1]$. In general, some of the elements in $D_{e}^{p-BR}(k)$ will be affected by further transformations while some others will not. This second group of elements will be referred to as the elements that have been fixed after transformation $k$. We denote by $r_{k}(i)$ the number of repetitions of link $i$ in $D_{e}^{p-BR}(k)$ (i.e. the amount of elements in $D_{e}^{p-BR}(k)$ that are equal to $i$) that have been fixed after transformation $k$. We denote by $p_{k}(i)$ the number of repetitions of link $i$ in $D_{e}^{p-BR}(k)$ (k) that will be affected by further transformations. Obviously, the number of repetitions of link $i$ in the final sequence $D_{e}^{p-BR}$ is $r_{k}(i) + \ldots + r_{S-1}(i) + p_{S-1}(i)$. In the following, we derive analytical expressions for $r_{k}(i)$ and $p_{k}(i)$ that will enable the analysis of the asymptotic behavior of $\alpha$.

Notice that the first transformation is a permutation that is applied to the second $(e-1)$-subsequence and it consists of a series of transpositions, as defined in section 3.2.1. Such transpositions are defined by pairing all elements from 0 to $e-2$ in such a way that the most frequent element is exchanged with the least frequent one. The same with the second-most and second-least frequent elements, and so on. Since the first and second $(e-1)$-subsequences are exactly equal but the permutation is only applied to the second one, after the permutation, the number of repetitions of the most frequent element will decrease to the half it was before plus the half of the number of repetitions of the least frequent element before the permutation. The number of repetitions of the least frequent element will increase in the same amount as the number of repetitions of the most frequent element is decreased.

Notice that any element $i \in [0,(e-1)/2-1]$ appearing in the second $(e-1)$-subsequence is fixed after the first transformation. The same is true for any element $i \in [(e-1)/2,e-2]$ in the first $(e-1)$-subsequence. In addition the number of repetitions of elements $i \in [0,(e-1)/2-1]$ in the second $(e-1)$-subsequence is $2^{e-2}, 2^{e-1}, \ldots, 2^{e-1}/2-1$ respectively. For elements $i \in [(e-1)/2,e-2]$ in the first $(e-1)$-subsequence we have the same number of repetitions but in reverse order.

Something similar happens after the second permutation. In this case, any element $i \in [0,(e-1)/2^{2}-1]$ in the second $(e-2)$-subsequence is fixed. The same is true for all the following cases:

- Any element $i \in [(e-1)/2^{2},(e-1)/2-1]$ in the first $(e-2)$-subsequence.
- Any element $i \in [(e-1)/2,3(e-1)/2-1]$ in the third $(e-2)$-subsequence.
- Any element $i \in [3(e-1)/2,e-2]$ in the forth $(e-2)$-subsequence.

The number of repetitions of elements $i \in [0,(e-1)/2^{2}-1]$ of the second $(e-2)$-subsequence that have been fixed is:

$$2^{(e-1)/2-1}, 2^{e-1}/2-2$$

For the other intervals of elements and their corresponding $(e-2)$-subsequences, the number of repetitions is the same as those of the first interval but in reverse order for some of them.

In general, due to the recursive nature of $D_{e}^{p-BR}$, we have that after each transformation $k \in [0, S-1]$, the number of repetitions of any element $i \not\in [0,(e-1)/2^{k+1}-1]$ that have been fixed is the same as that of some other element $j \in [0,(e-1)/2^{k+1}-1]$.

On the other hand, after the first transformation, any element $i \in [0,(e-1)/2-1]$ in the first $(e-1)$-subsequence will be changed by any further permutation. The same is true for any element $i \in [(e-1)/2,e-2]$ in the second $(e-1)$-subsequence. In addition the number of repetitions of
the elements of the first interval that appear in the first 
\((e-1)\)-subsequence is \(2e^2, 2e^3, ..., (e-1)e^2\) respectively. For the second interval and the second \((e-1)\)-subsequence
we have the same number of repetitions but in reverse order. In
addition, any further permutation that reduces/increases the
number of repetitions of an element of the first interval will
result in exactly the same reduction/increase in that dimension
of the second interval that has the same number of repetitions.

In general, due again to the recursive nature of \(D_e^{p-BR}\),
we have that after each transformation \(k \in [0, S-2]\), the
number of repetitions of any element \(i \in [0, (e-1)/2^{k+1}-1]\)
that will be affected by further permutations is the same as
that of some other dimension \(j \in [0, (e-1)/2^{k+1}-1]\). Besides,
the number of repetitions of these two elements \(i\) and \(j\)
will be increased/decreased by the same amount by any further
permutation, thus they will be the same.

As a conclusion of the previous analysis, in order to
obtain the range of values of functions \(p(k,i)\) and \(p(BR)\), it is
easy to compute them for elements \(i \in [0, (e-1)/2^{k+1}-1]\)
since its value for any other element will be the same as that
for an element of this interval.

**Lemma 2.** \(p(k,i) = 2^e - 2k - i\) \(\forall k \in [-1, S-1] \\forall i \in \left[0, \frac{e-1}{2^{k+1}} - 1\right]\)

**Proof.** For \(k=-1\), that is, before applying any
permutation, the equation holds since it corresponds to the
number of repetitions of each dimension in \(D_e^{BR}\). Then,
any transformation \(k\) changes every dimension \(i \in \left[0, (e-1)/2^{k+1}-1\right]\) with dimension \((e-1)/2^{k+1}-i\), but just in
the second \((e-k-1)\)-subsequence. Therefore, the number of
remaining repetitions of dimension \(i\) in the first
\((e-k-1)\)-subsequence is halved by each transformation \(k\)
since \(i\) appears the same number of times in the
first and second \((e-k-1)\)-subsequences.

**Lemma 3.** \(r(k,i) = 2^e - 1 - k - i\) \(\forall k \in [0, S-1] \\forall i \in \left[0, \frac{e-1}{2^{k+1}} - 1\right]\)

**Proof.** In transformation \(k\), a given element \(i \in \left[0, (e-1)/2^{k+1}-1\right]\) is exchanged with element \((e-1)/2^{k}-1-i\)
in the second \((e-k-1)\)-subsequence. Therefore, the number of
repetitions of \(i\) in the second \((e-k-1)\)-subsequence that after
this permutation will appear in the second
\((e-k-1)\)-subsequence corresponds to the number of
repetitions of element \((e-1)/2^{k}-1-i\) in that subsequence before this permutation. That is, \(r(k,i) = p(k,i)\) \((e-1)/2^{k}-1-i\)/2,
which results in the expression shown in the lemma.

**Lemma 4.** After transformation \(k\), the maximum
number of repetitions of any element that will not be
affected by any other permutation because they are in the
second \((e-k-1)\)-subsequence is equal to

\[
N_k = \frac{e-1}{2^{k+1}} - k - 2
\]

\(\forall k \in [0, S-1]\)

**Proof.** It is obvious that \(N_k = \max r(k,i)\). Using the
expression of \(r(k,i)\) given in lemma 3, the above result is
obtained.

**Lemma 5.** \(N_k \leq 2^e - 2S + k\) \(\forall k \in [0, S-2]\)

**Proof.** Given the definition of \(N(k)\) provided by
lemma 4, lemma 5 can be proved by showing that:

\[
\frac{e-1}{2^k} \geq 2S - 2k - 2
\]

Since \(e-1 = 2^k\), the previous inequality holds if \(2^{k-1} \geq 2(S-k-1)\). Since the left-hand side expression grows faster than the right-hand side expression when \(k\) decreases, it is enough to prove that the inequality holds for
the highest value of \(k\). By substituting \(k\) by \(S-2\), we obtain
the same value in both sides.

**Lemma 6.**

\[
\sum_{k=0}^{S-2} N_k \leq 2^e - S - 1 - 2^e - 2S
\]

**Proof.** This is a straightforward result that comes from
computing the sum from 0 to \(S-2\) of the bound
computed for \(N(k)\) in the lemma 5.

**Theorem 2.** The value of \(\alpha\) corresponding to the
sequences \(D_e^{p-BR}\) is bounded by the following expression:

\[
\alpha \leq \frac{2^e}{e-1} + \frac{2^e-2}{e-1} - \frac{2^e}{(e-1)^2}
\]

**Proof.** The value of \(\alpha\) is bounded by the sum of
the following terms: the maximum value of \(p_S(i); N_0; N_1; \ldots
\) and \(N_{S-1}\). That is

\[
\alpha \leq p_S(0) + N_0 + \sum_{k=0}^{S-2} N_k
\]

By adding the value of \(p_S(0)\) given by lemma 2; the
value of \(N_{S-1}\) given by lemma 4; and the bound given by
lemma 6, the bound established by this theorem follows.

**Theorem 3.** The upper-bound of \(\alpha\) established by
theorem 2 tends to 1.25 times the lower bound given by

\[
\left(\frac{2^e-1}{e}\right)^\frac{1}{S-1}
\]

**Proof.** This is proved by observing that

\[
\lim_{e \to \infty} \frac{2^e}{e-1} = \frac{2^e}{(e-1)^2} = 1.25
\]