Parallel Algorithms in Linear Algebra

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Abstract

This paper provides an introduction to algorithms for fundamental linear algebra problems on various parallel computer architectures, with the emphasis on distributed-memory MIMD machines. To illustrate the basic concepts and key issues, we consider the problem of parallel solution of a nonsingular linear system by Gaussian elimination with partial pivoting. This problem has come to be regarded as a benchmark for the performance of parallel machines. We consider its appropriateness as a benchmark, its communication requirements, and schemes for data distribution to facilitate communication and load balancing. In addition, we describe some parallel algorithms for orthogonal (QR) factorization and the singular value decomposition (SVD).

1. Introduction – Gaussian elimination as a benchmark

Conventional benchmarks are often inappropriate for parallel machines. A good benchmark needs to be a well-defined problem with a verifiable solution, as well as being representative of the problems which are of interest to users of the machine. The problem should be scalable because the power of the machine may be wasted on problems which are too small.

For these reasons, the solution of a system of n nonsingular linear equations in n unknowns, by the method of Gaussian elimination with partial pivoting, has become a popular benchmark [15]. For conventional serial machines we can use n = 100, but for more powerful machines n can be increased to 1000 or more.

In Section 2 we introduce some basic concepts such as *speedup* and *efficiency* of parallel algorithms, and *virtual processors*. Various parallel computer architectures are outlined in Section 3.

On parallel machines with distributed memory, questions of data distribution and data movement are very important. In deciding how to partition data over the processors of a distributed-memory machine, attention must be paid both to the data

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distribution patterns implicit in the algorithm and the need to balance the load on the different processors. In Section 4 we consider the data movement required for Gaussian elimination, and how this maps to the movement of data between processors. This allows us to reach some conclusions about the appropriateness of different machine architectures for linear algebra computations.

To illustrate the key issues, Section 5 considers in more detail the problem of parallel solution of a nonsingular linear system by Gaussian elimination with partial pivoting on a distributed-memory MIMD machine.

Because of the difficulties and communication overheads associated with pivoting, it is tempting to try to avoid pivoting, but omission of pivoting in Gaussian elimination leads to numerical instability. One solution is to implement a parallel version of the orthogonal (QR) decomposition instead of the triangular (LU) decomposition obtained by Gaussian elimination. This permits a stable solution without pivoting, but at the expense of an increase in the number of floating-point operations. The QR decomposition has many other useful applications, e.g. to the solution of linear least squares problems or as a preliminary step in the singular-value decomposition. Some ways of implementing the QR decomposition in parallel are mentioned in Section 5.3.

Many problems in numerical linear algebra are easy to solve if we can find the singular value decomposition (SVD) of a rectangular matrix, or the eigenvalues and eigenvectors of a symmetric (or Hermitian) matrix. In Section 6 we describe some good parallel algorithms for these problems. Often the parallel algorithms are not just a straightforward modification of the best serial algorithms.

There has been an explosive growth of interest in parallel algorithms (including those for linear algebra problems) in recent years, so we can not attempt to be comprehensive. For more detailed discussions and additional references, the reader is referred to surveys such as those by Gallivan *et al* [23], Dongarra *et al* [16], and Heller [36].

2. Basic concepts

We assume that a parallel machine with P processors is available. Thus P measures the degree of parallelism; P = 1 is just the familiar serial case. When considering the solution of a particular problem, we let T_P denote the time required to solve the problem using (at most) P processors. The speedup S_P is defined by

$$S_P = T_1/T_P,$$

and the efficiency $E_P = S_P/P$.

When converting a serial algorithm into a parallel algorithm, our aim is usually to attain constant efficiency, i.e.

$$E_P \ge c$$

for some positive constant c independent of P. This may be written as

$$E_P = \Omega(1).$$

Equivalently, we want to attain linear speedup, i.e.

$$S_P \ge cP$$
,

which may be written as

$$S_P = \Omega(P).$$

2.1 Amdahl's Law

Suppose a positive fraction f of a computation is "essentially serial", i.e. not amenable to any speedup on a parallel machine. Then we would expect

$$T_P = fT_1 + (1 - f)T_1/P$$

so the overall speedup

$$S_P = \frac{1}{f + (1 - f)/P} \le \frac{1}{f},$$
(2.1)

i.e. the speedup is *bounded*, not linear. The inequality (2.1) is called *Amdahl's Law* [1] and has been used as an argument against parallel computation. However, what it shows is that the speedup is bounded as we increase the number of processors for a *fixed* problem. In practice, it is more likely that we want to solve larger problems as the number of processors increases, because the desire to solve larger problems is a primary motivation for building larger parallel machines.

Let N be a measure of the problem size. For many problems it is reasonable to assume that

$$f \le K/N \tag{2.2}$$

for some constant K. For example, in problems involving N by N matrices, we may have $\Omega(N^3)$ arithmetic operations and $O(N^2)$ serial input/output.

Suppose also that N increases at least linearly with P, with the same constant as in (2.2), i.e.

$$N \ge KP. \tag{2.3}$$

(2.2) and (2.3) imply that $fP \leq 1$, so from (2.1) we have

$$S_P = \frac{P}{fP + (1-f)} \ge \frac{P}{2-f} \ge \frac{P}{2}.$$

Thus we get linear speedup, with efficiency $E_P \ge 1/2$.

For further discussion of Amdahl's law and the scaling of problem size, see [33].

2.2 Virtual processors

In practice any parallel machine has a fixed maximum number (say P) of processors imposed by hardware constraints. When analysing parallel algorithms it is often convenient to ignore this fact and assume that we have as many (say p) processors as desired. We may think of these p processors as *virtual processors* or *processes*. Each

real processor can simulate $\lceil p/P \rceil$ virtual processors (provided the real processor has enough memory). Thus, ignoring overheads associated with the simulation, we have

$$S_P \ge S_p / \lceil p/P \rceil. \tag{2.4}$$

If our analysis for p processors gives us a lower bound on S_p , then (2.4) can be used to obtain a lower bound on S_P . In practice this may be an oversimplification, because the hardware/software system may or may not provide good support for virtual processors.

3. Parallel architectures

Many varieties of parallel computer architecture have been proposed in recent years. They include –

- Pipelined vector processors such as the Cray 1, Fujitsu VP 100, or NEC SX/2, in which there is a single instruction stream and the parallelism is more or less hidden from the programmer. (More recent descendants such as the Cray 2S, Cray Y-MP/8, and Fujitsu VP 2000 series incorporate parallel vector processors.)
- Single-instruction multiple-data (SIMD [19]) machines such as the Illiac IV, ICL DAP, or MasPar MP-1, in which a number of simple processing elements (PEs or cells) execute the same instruction on local data and communicate with their nearest neighbours on a square grid or torus. There is usually a general-purpose controller which can broadcast instructions and data to the cells.
- Multiple-instruction multiple-data (MIMD) machines such as those constructed from transputers, the Carnegie-Mellon CM*, the Fujitsu AP 1000, and hypercube machines such as the Caltech "Cosmic Cube", Intel iPSC, and nCUBE2. In the hypercube machines 2^k processors are connected like the vertices of a k-dimensional cube, i.e. the processors are identified by k-bit binary numbers, and are connected to the processors whose numbers differ by exactly one bit from their own.
- Massively parallel SIMD machines such as the CM-200 Connection Machine (which may also be regarded as a hypercube machine).
- Shared-memory multiprocessors such as the Alliant FX/80, Cedar, Encore Multimax, and Sequent Symmetry.
- Systolic arrays [41], which are 1 or 2-dimensional arrays of simple processors (cells) connected to their nearest neighbours. The cells on the edge of the array are usually connected to a general-purpose machine which acts as a controller. Examples are the Warp and iWarp machines [2, 6] and several machines described in [53]. Variations on the idea of systolic arrays are wavefront arrays [45] and instruction systolic arrays [58].

The categories listed above are not mutually exclusive. For example, there is an overlap between vector processors and shared-memory multiprocessors. Also, the distinction between SIMD and MIMD machines is orthogonal to the distinction between hypercube, grid and torus topologies.

In view of the diversity of parallel computer architectures, it is difficult to describe practical parallel algorithms in a machine-independent manner. In some cases an algorithm intended for one class of parallel machine can easily be converted for another (more general) class. For example, an algorithm designed for a systolic array can easily be mapped onto a hypercube, but not conversely (in general). An algorithm designed for a distributed-memory machine can easily be implemented on a shared memory machine, but the converse may not be true. As a rough approximation, it is easier to implement algorithms on the more general machines, but the more specialised machines may be more efficient (or cost-effective) for certain classes of problems. For example, systolic arrays are sufficient and cost-effective for many problems arising in digital signal processing [7, 31, 41, 44, 45, 53].

In the following sections we describe algorithms for distributed-memory messagepassing machines or systolic arrays – the reader should be able to translate to other appropriate architectures.

4. Data movement and data distribution

Before considering how to map data (e.g. vectors and matrices) onto distributedmemory machines, it is worth considering what forms of data movement are common in linear algebra algorithms. For the sake of example we focus our attention on Gaussian elimination with partial pivoting, but most other linear algebra algorithms have similar data movement requirements. To perform Gaussian elimination we need –

- Row/column broadcast. For example, the pivot row needs to be sent to processors responsible for other rows, so that they can be modified by the addition of a multiple of the pivot row. The column which defines the multipliers also needs to be broadcast.
- Row/column send/receive. For example, if pivoting is implemented by explicitly interchanging rows, then at each pivoting step two rows have to be interchanged. (This could be done by broadcasting both rows, but it might be less efficient than explicitly sending the rows to the appropriate destinations.)
- Row/column scan. Here we want to apply an associative operator θ to data in one (or more) rows or columns. For example, when selecting the pivot row it is necessary to find the index of the element of maximum absolute value in (part of) a column. This may be computed via an associative operator θ defined on pairs:

$$(a,i) \ \theta \ (b,j) = \begin{cases} (a,i), & \text{if } |a| \ge |b|;\\ (b,j), & \text{otherwise.} \end{cases}$$
(4.1)

Other useful associative operators are addition (of scalars or vectors) and concatenation (of vectors).

4.1 Data distribution

On a distributed-memory machine where each processor has a local memory which is accessible to other processors only by explicit message passing, it is customary to partition data such as matrices and vectors across the local memories of several processors. This is essential for problems which are too large to fit in the memory of one processor, and in any case it is usually desirable for load-balancing reasons. (The exception is for very small problems which may as well be solved in a single processor.) Since vectors are a special case of matrices, we consider the partitioning of an m by n matrix A. It is desirable for data to be distributed in a "natural" manner, so that the operations of row/column broadcast/send/receive/scan described above can be implemented efficiently. This is possible if a square grid is a subgraph of the connection graph of the parallel machine. For example, it is true for machines whose connection topology is an s by s torus or a hypercube of even dimension. (On machines for which a rectangular grid of moderate aspect ratio can be embedded, say an s by ks grid for some small positive integer k, we can use a square *virtual* ks by ks grid by having each processor simulate k virtual processors.)

The simplest mapping of data to processors is the *column-wrapped* (or *row-wrapped*) representation. Here column (or row) i of a matrix is stored in the memory associated with processor $i \mod P$, assuming that the P processors are numbered $0, 1, \ldots, P-1$. (A Fortran programmer might prefer $i-1 \mod P$, but we find that C array conventions are more convenient.)

Although simple, and widely used in parallel implementations of Gaussian elimination (e.g. [24, 25, 46, 52]), the column-wrapped (or row-wrapped) representation has some disadvantages –

- \cdot Lack of symmetry rows are treated differently from columns. It is instructive to consider the data communication involved in transposing a matrix.
- Poor load-balancing for moderate-sized problems if n < P some processors store no columns, so presumably perform no useful work. On the other hand, if *n* increases from *P* to *P* + 1 then the load on processor 0 doubles. Thus, the performance curve as a function of *n* (the number of columns in the matrix) will be "jagged" – there will be jumps at each multiple of *P*.
- Poor communication bandwidth for column broadcast since each column is stored in the memory associated with only one processor, the speed of column broadcast is constrained by the communication bandwidth of a single processor (compare the blocked/scattered storage representations below).

Another conceptually simple mapping is the *blocked* representation. Assume that the processors form an s by s grid $(P = s^2)$. The matrix A is padded with zero rows and columns if necessary, so that m and n can be assumed to be multiples of s. A is partitioned into an s by s matrix of blocks. Each block, of dimension m/sby n/s, is assigned to one processor in the natural way. This avoids the lack of symmetry inherent in the row/column-wrapped representation. It also improves the communication bandwidth for column broadcast, because each column is shared by $s = P^{1/2}$ processors. However, it suffers from a load-balancing problem –

• Poor load balancing for triangular and band matrices – if A is upper triangular then about half the processors (those storing the strict lower triangle of blocks) are only storing zeros and can probably not do any useful computation. Similarly (but even worse) if A is a band matrix with bandwidth small relative to m and n.

Harder to visualize, but often better than the row/column-wrapped or blocked representations, is the *scattered* representation [21] (also called *dot mode* in image-processing applications [39]). Assume as above that the processors form an s by s grid, and let the processors be numbered $(0,0), (0,1), \ldots, (s-1,s-1)$. Then the matrix element $a_{i,j}$ is stored in processor ($i \mod s, j \mod s$). Now the matrices stored

locally on each processor have the same shape (e.g. triangular, band, \ldots) as the global matrix A, so the computational load on each processor is approximately equal.

It is sometimes useful to regard a blocked representation of a matrix as a scattered representation of the same matrix with its rows and columns permuted. Formally, if s|k, define a k by k permutation matrix π_k whose (i, j)-th element is 1 iff

$$j = \begin{cases} i.s \mod k - 1, & \text{if } i < k - 1; \\ k - 1, & \text{if } i = k - 1 \end{cases}$$
(4.2)

(assuming C conventions – indices run from 0 to k-1). If $P = s^2$, s|m, s|n, then the scattered representation of the m by n matrix A is the same as the block representation of $\pi_m A \pi_n^{-1}$. Similarly, if B is n by p, s|p, then the scattered representation of B is the same as the block representation of $\pi_n B \pi_p^{-1}$, and the scattered representation of AB is the same as the block representation of $\pi_m A B \pi_p^{-1} = (\pi_m A \pi_n^{-1})(\pi_n B \pi_p^{-1})$. This shows formally that a matrix multiplication algorithm which works for matrices stored using the blocked representation should also work for matrices stored using the scattered representation, and vice versa.

The blocked and scattered representations do not actually require a square processor array – rectangular would suffice. The reason why we ask for the processor array to be square is that this makes matrix multiplication and matrix transposition much simpler than in the general case.

4.2 Implications for architectures

First consider distributed memory machines. In order to have a natural mapping of global matrices to a scattered storage representation, the inter-processor connection graph should have a square grid as a subgraph. A square torus is safisfactory, though slightly more general than necessary. A hypercube of even dimension is also satisfactory, but considerably more general than necessary.

A hypercube does have some advantages over a simple square s by s grid. Consider communication along one row (or column) of the grid. With a simple grid the maximum distance between processors in one row is s - 1, so time required to send or broadcast a short message is of order s in the worst (and average) case. With a hypercube this is reduced to order $\log s$.

In practice, for small and moderate values of s, the reduction from order s to order log s may not be so important as the overhead incurred at each intermediate node in the path from source to destination. For example, early hypercube machines used *store and forward*, which imposed a considerable processing load on the intermediate nodes. Consider sending a message of length n. At each intermediate node the delay is of order n, so the overall delay is of order $n \log s$.

Some more recent hypercube and torus machines (e.g. the Fujitsu AP 1000) have used wormhole routing [14] rather than store and forward. With wormhole routing the overall delay is of order $n + \log s$ (for a hypercube) or n + s (for a grid). Also, the wormhole routing protocol can be implemented in hardware so as not to impose a load on the processors at intermediate nodes.

For shared memory machines the problems are different. Clearly the memory must have sufficiently high overall bandwidth, but this is not sufficient. The problem of *hotspots*, i.e. memory locations accessed intensively by several processors, needs to be overcome. For example, in Gaussian elimination the memory locations where the pivot row is stored will be hotspots. On a distributed memory machine the programmer (or compiler) has to solve this problem by explicitly broadcasting the pivot row, but on a shared memory machine the hardware needs to be able to provide for simultaneous read-only access to the pivot row by several processors.

5. The solution of linear systems

Suppose we want to solve a nonsingular n by n linear system

$$Ax = b \tag{5.1}$$

on a parallel machine for which a 2-dimensional mesh is a natural interconnection pattern. It is easy to implement Gaussian elimination without pivoting, because multipliers can be propagated along rows of the augmented matrix [A|b], and it is not necessary for one row operation to be completed before the next row operation starts. Unfortunately, as is well-known [32, 60, 63], Gaussian elimination without pivoting is numerically unstable unless A has some special property such as diagonal dominance or positive definiteness. Thus we consider the implementation of Gaussian elimination with partial pivoting on a parallel machine.

5.1 Gaussian elimination with partial pivoting

For the sake of definiteness, we assume that Gaussian elimination is to be performed on a distributed-memory machine with an s by s grid, and that the augmented matrix [A|b] is stored in the scattered representation. The reader should be careful to distinguish between a row (or column) of *processors* and a row (or column) of the *matrix*. Generally, each row of processors stores several rows of the matrix.

It is known [32, 60, 63] that Gaussian elimination is equivalent to triangular factorization. More precisely, Gaussian elimination with partial pivoting produces an upper triangular matrix U and a lower triangular matrix L (with unit diagonal) such that

$$PA = LU \tag{5.2}$$

where P is a permutation matrix (not the number of processors here !). In the usual implementation A is overwritten by L and U (the diagonal of L need not be stored). If the same procedure is applied to the augmented matrix $\bar{A} = [A|b]$, we obtain

$$P\bar{A} = L\bar{U} \tag{5.3}$$

where $\overline{U} = [U|\overline{b}]$ and (5.1) has been transformed into the upper triangular system

$$Ux = \bar{b} \tag{5.4}$$

In the following we shall only consider the transformation of A to U, as the transformation of b to \bar{b} is similar.

If A has n rows, the following steps have to be repeated n-1 times, where the k-th iteration completes computation of the k-th column of U –

- 1. Find the index of the next pivot row by finding an element of maximal absolute value in the current (k-th) column, considering only elements on and below the diagonal. With the scattered representation this involves s processors, which each have to find a local maximum and then apply the associative operator (4.1).
- 2. Broadcast the pivot row vertically.
- 3. Exchange the pivot row with the current k-th row, and keep a record of the row permutation. Generally the exchange requires communication between two rows of s processors. Since the pivot row has been broadcast at step 3, only the current k-th row needs to be sent at this step. (Alternatively, the exchanges could be kept implicit, but this would lead to load-balancing problems and difficulties in implementing block updates, so explicit exchanges are usually preferable).
- 4. Compute the "multipliers" (elements of L) from the k-th column and broadcast horizontally.
- 5. Perform Gaussian elimination (using the portion of the pivot row and the other rows held in each processor). If done in the obvious way, this involves saxpy operations (a *saxpy* is the addition of a scalar multiple of one vector to another vector), but the computation can also be formulated as a rank-1 update.

We can make an estimate of the parallel time T_P required to perform the transformation of A to upper triangular form. There are two main contributions –

- A. Floating-point arithmetic. The overall computation involves $2n^3/3 + O(n^2)$ floating-point operations (counting additions and multiplications separately). Because of the scattered representation each of the $P = s^2$ processors performs approximately the same amount of arithmetic. Thus floating-point arithmetic contributes a term $O(n^3/s^2)$ to the computation time.
- B. Communication. At each iteration of steps 1-5 above, a given processor sends or receives O(n/s) words. We shall assume that the time required to send or receive a message of w words is $c_0 + c_1 w$, where c_0 is a "startup" time and $1/c_1$ is the transfer rate. (For real machines the time may depend on other factors, such as the distance between the sender and the receiver and the overall load on the communication network.) With our assumption, the overall communication time is $O(n^2/s) + O(n)$, where the O(n) term is due to startup costs.

If arithmetic and communication can not be overlapped, the overall time T_P is simply the sum of A and B above, i.e.

$$T_P \simeq \alpha n^3 / s^2 + \beta n^2 / s + \gamma n, \qquad (5.5)$$

where α depends on the floating-point and memory speed of each processor, β depends mainly on the communication transfer rate between processors, and γ depends mainly on the communication startup time. We would expect the time on a single processor to be

$$T_1 \simeq \alpha n^3, \tag{5.6}$$

although this may be inaccurate for various reasons - e.g. the problem may fit in memory caches on a parallel machine, but not on a single processor.

From (5.5) and (5.6), the efficiency E_P is

$$E_P \simeq \frac{1}{1 + (1 + \bar{\gamma}/\bar{n})\bar{\beta}/\bar{n}},$$
 (5.7)

where $\bar{\beta} = \beta/\alpha$ is proportional to the ratio of communication to computation speed, $\bar{\gamma} = \gamma/\beta$ measures the importance of the communication startup time, and $\bar{n} = n/s$ is the number of rows or columns of A stored in a single processor. From (5.7), the efficiency is close to 1 only if $\bar{n} \gg \bar{\beta}$.

We have ignored the "back-substitution" phase, i.e. the solution of the upper triangular system (5.4), because this can be performed in time much less than (5.5) (see [21, 46]).

5.2 Blocking

On many machines it is impossible to achieve peak performance if the Gaussian elimination is performed via saxpys or rank-1 updates. This is because performance is limited by memory accesses rather than by floating-point arithmetic, and saxpys or rank-1 updates have a high ratio of memory references to floating-point operations. Closer to peak performance can be obtained for matrix-vector or (better) matrixmatrix multiplication.

It is possible to reformulate Gaussian elimination so that most of the floatingpoint arithmetic is performed in matrix-matrix multiplications, without compromising the error analysis. Partial pivoting introduces some difficulties, but they are surmountable. The idea is to introduce a "blocksize" or "bandwidth" parameter ω . Gaussian elimination is performed via saxpys or rank-1 updates in vertical strips of width ω . Once ω pivots have been chosen, a horizontal strip of height ω can be updated. At this point, a matrix-matrix multiplication can be used to update the lower right corner of A. The optimal choice of ω depends on details of the machine architecture, but

$$\omega \sim n^{1/2} \tag{5.8}$$

is a reasonable choice.

The effect of blocking is to reduce the constant α in (5.5) at the expense of increasing the lower-order terms. Thus, a blocked implementation should be faster for sufficiently large n, but may be slower than an unblocked implementation for small n.

5.3 Orthogonal factorization

On machines whose architecture makes pivoting difficult, we can avoid it at the expense of increasing the amount of arithmetic. For example, on systolic arrays it is possible to compute the orthogonal (QR) factorization of A efficiently and in a numerically stable manner using Givens transformations [5, 27, 47]. The cost is an increase by a factor of four in the number of arithmetic operations (though this factor may be reduced if "fast" Givens transformations [26] are used). In any case, the QR factorization is of independent interest because it can be used to solve the linear least squares problem

$$\min \|Ax - b\|_2 \tag{5.9}$$

where A is an m by n matrix of rank n.

On a single processor Householder transformations are cheaper than Givens transformations. The steps involved in implementing a Householder QR factorization on a parallel machine are similar to those involved in implementing Gaussian elimination. Although pivoting is not usually required, the vectors u which define the Householder transformations $I - 2uu^T$ need to be broadcast in the same way as the pivot row and multiplier column in Gaussian elimination.

As for Gaussian elimination, optimal performance for large matrices may require blocking. Several Householder transformations can be combined [3, 57] and then applied together so that most of the arithmetic is done in matrix-matrix multiplication.

6. The SVD and symmetric eigenvalue problems

A singular value decomposition (SVD) of a real m by n matrix A is its factorization into the product of three matrices:

$$A = U\Sigma V^T, \tag{6.1}$$

where U is an m by n matrix with orthonormal columns, Σ is an n by n nonnegative diagonal matrix, and V is an n by n orthogonal matrix (we assume here that $m \ge n$). The diagonal elements σ_i of Σ are the *singular values* of A. The singular value decomposition has many applications [29, 44].

The SVD is usually computed by a two-sided orthogonalization process, e.g. by two-sided reduction to bidiagonal form (possibly preceded by a one-sided reduction [11]), followed by the QR algorithm [28, 30, 63]. It is difficult to implement this Golub-Kahan-Reinsch algorithm efficiently on a parallel machine. It is much simpler (though perhaps less efficient) to use a one-sided orthogonalization method due to Hestenes [37]. The idea is to generate an orthogonal matrix V such that AV has orthogonal columns. Normalizing the Euclidean length of each nonnull column to unity, we get

$$AV = U\Sigma \tag{6.2}$$

As a null column of U is always associated with a zero diagonal element of Σ , there is no essential difference between (6.1) and (6.2).

The cost of simplicity is an increase in the operation count, compared to the Golub-Kahan Reinsch algorithm.

6.1 Implementation of the Hestenes method

Let $A_1 = A$ and $V_1 = I$. The Hestenes method uses a sequence of plane rotations Q_k chosen to orthogonalize two columns in $A_{k+1} = A_k Q_k$. If the matrix V is required, the plane rotations are accumulated using $V_{k+1} = V_k Q_k$. Under certain conditions $\lim Q_k = I$, $\lim V_k = V$ and $\lim A_k = AV$. The matrix A_{k+1} differs from A_k only in two columns, say columns i and j. In fact

$$\left(a_{i}^{(k+1)}, a_{j}^{(k+1)}\right) = \left(a_{i}^{k}, a_{j}^{k}\right) \left(\begin{array}{cc}\cos\theta & \sin\theta\\-\sin\theta & \cos\theta\end{array}\right)$$

where the rotation angle θ is chosen so that the two new columns $a_i^{(k+1)}$ and $a_j^{(k+1)}$ are orthogonal. This can always be done with an angle θ satisfying

$$|\theta| \le \pi/4,\tag{6.3}$$

see for example [9].

It is desirable for a "sweep" of n(n-1)/2 rotations to include all pairs (i, j) with i < j. On a serial machine a simple strategy is to choose the "cyclic by rows" ordering

$$(1, 2), (1, 3), \dots, (1, n), (2, 3), \dots, (n - 1, n).$$

Forsythe and Henrici [20] have shown that the cyclic by rows ordering and condition (6.3) ensure convergence of the Jacobi method applied to $A^T A$, and convergence of the cyclic by rows Hestenes method follows. In practice only a small number of sweeps are required. The speed of convergence is discussed in [9].

6.2 The Go tournament analogy

On a parallel machine we would like to orthogonalize several pairs of columns simultaneously. This should be possible so long as no column occurs in more than one pair. The problem is similar to that of organizing a round-robin tournament between n players. A game between players i and j corresponds to orthogonalizing columns i and j, a round of several games played at the same time corresponds to orthogonalizing several pairs of (disjoint) columns, and a tournament where each player plays each other player once corresponds to a sweep in which each pair of columns is orthogonalized. Thus, schemes which are well-known to Go players (or players of other two-person games such as Chess, Sumo, ...) can be used to give orderings amenable to parallel computation. It is usually desirable to minimize the number of parallel steps in a sweep, which corresponds to the number of rounds in the tournament.

On a parallel machine with restricted communication paths there are constraints on the orderings which we can implement efficiently. A useful analogy is a tournament of lazy Go players. After each round the players want to walk only a short distance to the board where they are to play the next round.

Using this analogy, suppose that each Go board corresponds to a virtual processor and each player corresponds to a column of the matrix (initially A but modified as the computation proceeds). A game between two players corresponds to orthogonalization of the corresponding columns. Thus we suppose that each virtual processor has sufficient memory to store and update two columns of the matrix. If the Go boards (processors) are arranged in a linear array with nearest-neighbour communication paths, then the players should have to walk (at most) to an adjacent board between the end of one round and the beginning of the next round, i.e. columns of the matrix should have to be exchanged only between adjacent processors. Several orderings satisfying these conditions have been proposed [8, 9, 50, 56].

Since A has n columns and at most $\lfloor n/2 \rfloor$ pairs can be orthogonalized in parallel, a sweep requires as least n-1 parallel steps (n even) or n parallel steps (n odd). The ordering of [9] attains this minimum, and convergence can be guaranteed if n is odd

[50, 59]. It is an open question whether convergence can be guaranteed, for any ordering which requires only the minimum number of parallel steps, when n is even – the problem in proving convergence is illustrated in [34]. However, in practice lack of convergence is not a problem, and it is easy to ensure convergence by the use of a "threshold" strategy [63], or by taking one additional parallel step per sweep when n is even [51].

As described above, each virtual processor deals with two columns, so the columnwrapped representation is convenient. However, the block or scattered representations can also be used. The block representation involves less communication between real processors than does the scattered representation if the standard orderings are used. However, the two representations are equivalent if different orderings are used. (As in the discussion of matrix multiplication at the end of Section 4.1, the algorithm can assume that the block representation is used, since the SVD of $\pi_m A \pi_n^{-1}$ is just a reordering of the SVD of A.) The scattered representation does not have a loadbalancing advantage here, since the matrix does not change shape.

6.3 The symmetric eigenvalue problem

There is a close connection between the Hestenes method for finding the SVD of a matrix A and the Jacobi method for finding the eigenvalues of a symmetric matrix $B = A^T A$. Important differences are that the formulas defining the rotation angle θ involve elements $b_{i,j}$ of B rather than inner products of columns of A, and transformations must be performed on the left and right instead of just on the right (since $(AV)^T(AV) = V^T BV$). Instead of permuting columns of A as described in Section 6.2, we have to apply the same permutation to both rows and columns of B. An implementation on a square systolic array of n/2 by n/2 processors is described in [9], and could easily be adapted to other parallel architectures. If less than $n^2/4$ processors are available, we can use the virtual processor concept described in Section 2.2.

6.4 Other SVD and eigenvalue algorithms

In Section 6.2 we showed how the Hestenes method could be used to compute the SVD of an m by n matrix in time $O(mn^2S/P)$ using P = O(n) processors in parallel. Here S is the number of sweeps required (conjectured to be $O(\log n)$). In Section 6.3 we sketched how Jacobi's method could be used to compute the eigen-decomposition of a symmetric n by n matrix in time $O(n^3S/P)$ using $P = O(n^2)$ processors. It is natural to ask if we can use more than $\Omega(n)$ processors efficiently when computing the SVD. The answer is yes – Kogbetliantz [40] and Forsythe & Henrici [20] suggested an analogue of Jacobi's method, and this can be used to compute the SVD of a square matrix using a parallel algorithm very similar to the parallel implementation of Jacobi's method. The result is an algorithm which requires time $O(n^3S/P)$ using $P = O(n^2)$ processors. Details and a discussion of several variations on this theme may be found in [10].

In order to find the SVD of a rectangular m by n matrix A using $O(n^2)$ processors, we first compute the QR factorization QA = R (see Section 5.3), and then compute the SVD of the principal n by n submatrix of R (i.e. discard the m - n zero rows of R). It is possible to gain a factor of two in efficiency by preserving the upper triangular structure of R [48].

The Hestenes/Jacobi/Kogbetliantz methods are not often used on a serial computer, because they are slower than methods based on reduction to bidiagonal or tridiagonal form followed by the QR algorithm [63]. Whether the fast serial algorithms can be implemented efficiently on a parallel machine depends to some extent on the parallel architecture. For example, on a square array of n by n processors it is possible to reduce a symmetric n by n matrix to tridiagonal form in time $O(n \log n)$ [4]. On a serial machine this reduction takes time $O(n^3)$. Thus, a factor $O(\log n)$ is lost in efficiency, which roughly equates to the factor O(S) by which Jacobi's method is slower than the QR algorithm on a serial machine. It is an open question whether the loss in efficiency by a factor $O(\log n)$ can be avoided on a parallel machine with $P = \Omega(n^2)$ processors. When P = O(n), "block" versions of the usual serial algorithms are attractive on certain architectures [17], and may be combined with the "divide and conquer" strategy [18]. Generally, these more complex algorithms are attractive on shared memory MIMD machines with a small number of processors, while the simpler algorithms described above are attractive on systolic arrays and SIMD machines.

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