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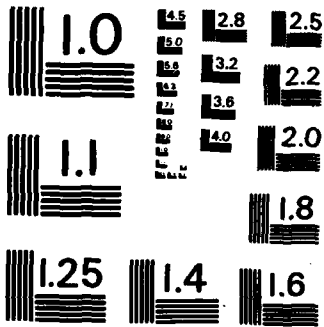
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THESIS

MINIMIZING FILL-IN FOR GAUSSIAN ELIMINATION
OF
SYMMETRIC SYSTEMS

by

Young Ho Seo

September 1985

Thesis Advisor:

R. K. Wood

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ABSTRACT

The new projective linear programming algorithm by Karmarkar requires solution of symmetric, positive definite systems of equations. One key to solving these systems efficiently is minimizing the fill-in, i.e., the number of new nonzero elements in the reduced matrix, in the various forms of Gaussian elimination used to solve these systems. Fill-in is affected only by symmetric reordering of rows and columns of the system of equations. Various heuristic ordering algorithms are tested and compared with the heuristic minimum degree algorithm of George and Liu. Computational results are reported for sixteen large-scale real-world and artificial problems. The minimum degree algorithm of George and Liu is the most effective of six tested methods.

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Y. INTRODUCTION

A. INTRODUCTION

The new projective linear programming algorithm by Karmarkar [Ref. 1] requires solution of symmetric, positive definite systems of equations. One of the keys to solving these systems efficiently is controlling the fill-in, i.e., the number of new nonzero elements in the reduced matrix, in the various forms of Gaussian elimination used to solve these systems. Under the assumption of noncancellation of terms, fill-in created by direct solution techniques is affected only by symmetric reordering of the rows and columns of the system to be solved.

The problem of minimizing fill-in by symmetric reordering is known to be NP-complete [Ref. 9] so, in this thesis, we examine a number of heuristic reordering algorithms for this problem.

Several different algorithms for reducing fill-in in Gaussian elimination are compared for sixteen large-scale matrices derived from linear and integer programming problems.

B. PROJECTIVE LP ALGORITHM

Consider the linear programming problem

$$\begin{aligned} \min \quad & c^T x \\ \text{s.t.} \quad & Ax = b \\ & x \geq 0 \end{aligned}$$

where A is an n -by- n matrix, c is an n -element column vector and b is an n -element column vector. . . Karmarkar's algorithm requires the solution of $BB^T z = Bc$, where

$$B = \begin{bmatrix} AR \\ \mathbf{1}^T \end{bmatrix}, \text{ } E \text{ is an } n\text{-by-}n \text{ diagonal matrix, and } \mathbf{1} \text{ is}$$

a n -element column vector of 1s. Thus,

$$BB^T = \begin{bmatrix} AR^2A^T & AR\mathbf{1} \\ \mathbf{1}^T EA^T & n \end{bmatrix}.$$

We will not be concerned with the bottom row and right-most column; they will typically be 100% dense when using an artificial starting solution and will be placed last in any ordering. Under the assumption of noncancellation of terms, the nonzeros created in AA^T will be the same as in AR^2A^T , both when creating these matrices and when performing the elimination. Thus, we may restrict study to solving systems of the form $AA^T z = Ac = b'$.

C. DIRECT SOLUTION METHODS

Symmetry and positive definiteness are important in computation with the matrices. The matrix D is symmetric when $D^T = D$. Otherwise, D is asymmetric. A symmetric matrix D is said to be positive definite if $z^T Dz > 0$ for all nonzero vectors z .

Three basic methods are available for direct solution of positive definite systems of the form $AA^T z = Dz = b'$. These methods are basic Gaussian elimination [Ref. 6], Cholesky factorization [Ref. 4] and QR factorization [Ref. 7]. All three methods end up creating an upper or lower triangular matrix which is used with b' , or a modified version of b' , to solve for z by backward and/or forward substitution.

One key to success of these methods is keeping the triangular matrix as sparse as possible by symmetrically reordering the rows and columns of D . This keeps computation time down and round-off errors to a minimum.

1. Basic Gaussian Elimination

The Gaussian elimination method we will consider for the solution of a set of equations is just a generalization of the familiar method of eliminating one unknown by substitution between a pair of simultaneous equations and is the basic pattern of a large number of solution methods that can be classed as direct methods.

Let us consider the system $Dz = b'$, represented by the augmented matrix $[D, b']$, where D is an n -by- n positive definite symmetric matrix and b' is an n -vector. Gaussian elimination by columns consists of n steps. The purpose of the k th step is to eliminate all the nonzero elements of the augmented matrix which lie on column k below the diagonal of D . At the k th step, $d_{kk}^{(k)}$ which is the diagonal of the active submatrix (formed by rows k to n and column k to n of $D^{(k)}$) is selected to be the pivot. Row k of the augmented matrix is normalized and the elements in column k below the diagonal of $D^{(k)}$ are eliminated by subtraction of multiples of the normalized row k from all those rows which have a nonzero in column k below the diagonal. The augmented system $[D^{(k)}, b'^{(k)}]$ is obtained where $D^{(k)}$ has zeros in its first k columns below the diagonal and is on the first k positions of the diagonal.

The process is continued until the upper triangular system $[D^{(n)}, b'^{(n)}]$ is obtained and z can be solved for by backward substitution.

2. Cholesky Factorization

There exists a unique lower triangular matrix L such that $LL^T = D$. Cholesky factorization creates L from D directly without consideration of b' . Then, $LL^T z = Dz = b'$ is solved for by:

(1) Solve $Ly = b'$ for y .

(2) Solve $L^T z = y$ for z .

Suppose a positive definite symmetric m -by- m matrix D is given. The way in which fill-in occurs when computing L is best illustrated by showing a proof, following George and Liu [Ref. 4], of one of the Cholesky factorization techniques.

$$\text{Let } D = \begin{bmatrix} d_1 & a_1^T \\ a_1 & \kappa_1 \end{bmatrix} \text{ where } d_1 \text{ is a scalar, } a_1 \text{ is an } m-1$$

vector and κ_1 is an $(m-1)$ -by- $(m-1)$ submatrix. Then we can write

$$\begin{aligned} D &= \begin{bmatrix} d_1^{1/2} & 0 \\ a_1/d_1^{1/2} & \tau_{m-1} \end{bmatrix} \begin{bmatrix} 1 & 0 \\ 0 & H_1 \end{bmatrix} \begin{bmatrix} d_1^{1/2} & a_1^T/d_1^{1/2} \\ 0 & \tau_{m-1} \end{bmatrix} \\ &= L_1 \begin{bmatrix} 1 & 0 \\ 0 & H_1 \end{bmatrix} L_1^T \\ &= L_1 D_1 L_1^T \end{aligned}$$

where $H_1 = \kappa_1 - a_1 a_1^T / d_1$, and I_k denotes an identity

of the submatrix K_i by the outer product $a_i a_i^T / d_i$ to give H_i , which is the submatrix remaining to be factored. Thus, the structure of H_i is different from K_i due to $a_i a_i^T / d_i$. $a_i a_i^T / d_i$ creates new fill-ins if $(a_i)_k (a_i)_j \neq 0$ but $(H_i)_{kj} = 0$ where $(a_i)_k$ is a k th element of vector a_i and $(H_i)_{kj}$ is the scalar element in row k and column j if H_i .

It can be seen from the above derivation that the matrices $D^{(k)}$ and H_k correspond in their patterns of nonzeros, and if the normalization step of standard Gaussian elimination is replaced by "divide row k by $(d^{(k)})_{kk}^{1/2}$," then the upper triangular matrix produced is actually L^T . Under the assumption of noncancellation of terms, the pattern of nonzeros in L^T and the upper triangular portion of $D^{(n)}$ will be the same.

3. QR Factorization

For completeness let us also consider QR orthogonal factorization. Orthogonal factorization methods compute the triangular Cholesky factor L directly from A rather than AA^T and thus avoid explicit formation of the cross product used in basic Gaussian elimination and Cholesky factorization. In QR factorization, an orthogonal matrix Q of order n is computed which reduces A and c to the form

$$Q^T A^T = \begin{bmatrix} L^T \\ 0 \end{bmatrix} \quad \text{and} \quad Q^T c = \begin{bmatrix} q \\ r \end{bmatrix},$$

where q is a vector of dimension m , and r is a vector of dimension $m-n$.

Then, $Dz = b'$ is equivalent to

$$AA^T z = Ac,$$

$$\Rightarrow AQQ^T A^T z = AQQ^T c,$$

$$\Rightarrow [L \ 0] \begin{vmatrix} L^T \\ 0 \end{vmatrix} z = [L \ 0] \begin{vmatrix} q \\ 0 \end{vmatrix},$$

$$\Rightarrow LL^T z = Lq,$$

$$\Rightarrow \text{and } L^T z = q.$$

Thus, after computing L^T and q using the orthogonal matrix Q , we need only solve for z by backward substitution.

D. GRAPH THEORETIC MODEL

Consider the two matrices below where x indicates a nonzero. If we perform Gaussian elimination on the two matrices, we get the upper triangular forms shown.

$$\begin{array}{c|cccc} 1 & x & x & x & x \\ 2 & x & x & & \\ 3 & x & & x & \\ 4 & x & & & x \end{array} \Rightarrow \begin{array}{c|cccc} 1 & x & x & x & x \\ 2 & & x & x & x \\ 3 & & & x & x \\ 4 & & & & x \end{array} \quad (1)$$

$$\begin{array}{c|cccc} 1 & x & & x & \\ 2 & & x & & x \\ 3 & & & x & x \\ 4 & x & x & x & x \end{array} \Rightarrow \begin{array}{c|cccc} 1 & x & & & x \\ 2 & & x & & x \\ 3 & & & x & x \\ 4 & & & & x \end{array} \quad (2)$$

Under assumption of noncancellation of terms, the Cholesky factor of (1) is 100% dense. If the first and last columns of (1) are interchanged, and the first and last rows of (1) are interchanged, then Gaussian elimination will produce no fill-in at all. Below, a graph theoretic model is described

[Ref. 3] which helps clarify the process of creating fill-ins and how this can be modified by reordering.

An undirected graph $G=(V, E)$ consists of a finite set of vertices V together with a set E of edges, where an edge is an unordered pair of vertices (u, v) such that $u \neq v$. An undirected graph may be associated with any symmetric matrix D of order n . The "adjacency graph" of D is a graph $G=(V, E)$ such that $V=\{v_1, \dots, v_n\}$ and $(v_i, v_j) \in E$ if and only if $d_{ij} \neq 0$ for all $i \neq j$. The adjacency graph $G^{(k)}$ associated with the active submatrix of $D^{(k)}$ is called an "elimination graph." Note that an adjacency graph G associated with D is unaffected by any symmetric reordering of the rows and columns D .

If (u, v) is an edge, u and v in G are "adjacent." For $S \subset V$, the adjacent set of S , denoted by $Adj(S)$, is the set of all vertices not in S which are adjacent to vertices in S . That is,

$$Adj(S) = \{u \in V-S \mid (u, v) \in E \text{ for some } v \in S\}$$

where $V-S$ is the set of all vertices of V which are not in S .

A subgraph $G' = (V', E')$ of $G = (V, E)$ is a graph which consists of some or all vertices of G and some of the edges of G : $V' \subseteq V, E' \subseteq E$. A subgraph is a clique if every pair of vertices in a subgraph is adjacent.

Let us consider a symmetric sparse matrix and its associated undirected graph. The matrix is of order 8.

$$D = D^{(0)} = \begin{array}{c|cccccccc} & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 \\ \hline 1 & x & & & & & & & \\ 2 & & x & & & & & & \\ 3 & & & xx & & & & & \\ 4 & & & & xx & & & & \\ 5 & & & & & x & & & \\ 6 & & & & & & x & & \\ 7 & & & & & & & xx & \\ 8 & & & & & & & & x & xx \end{array} \quad G^{(0)} = \begin{array}{c} 1 \text{---} 5 \text{---} 2 \\ | \hspace{1.5cm} | \\ 6 \text{---} \hspace{1.5cm} 7 \\ | \hspace{1.5cm} | \\ 3 \text{---} 4 \text{---} 8 \end{array}$$

If we perform Gaussian elimination at column 1, we obtain $D^{(1)}$.

$$D^{(1)} = \begin{array}{c|cccccccc} & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 \\ \hline 1 & - & - & - & - & - & - & - & - \\ 2 & x & x & x & - & - & - & - & - \\ 3 & & xx & x & - & - & - & - & - \\ 4 & & & xx & x & - & - & - & - \\ 5 & & & & x & xo & - & - & - \\ 6 & & & & & x & oxx & - & - \\ 7 & & & & & & x & xxx & - \\ 8 & & & & & & & x & xx \end{array} \quad G^{(1)} = \begin{array}{c} & & & & 5 & \text{---} & 2 \\ & & & & / & & | \\ 6 & & & & & \text{---} & 7 \\ | & & & & & & | \\ 3 & \text{---} & 4 & \text{---} & & & 8 \end{array}$$

where new fill-ins are noted by circles and - corresponds to a nonzero in the nonactive part of $D^{(k)}$. Fill-in in $D^{(1)}$ corresponds to edges of $G^{(1)}$ which were not in $G^{(0)}$. $G^{(1)}$ is created from $G^{(0)}$ by deleting v_1 and all incident edges and by then connecting all vertices which were adjacent to v_1 and not already connected. This is formalized below.

Let $G^{(k)}$ be an elimination graph. If vertex v_i corresponding to row i and column i is next selected for elimination, then the new fill-in created will be equal to the number of edges in the complement of the graph induced from $G^{(k)}$ by $\text{Adj}(v_i)$. A graph $G'(V', E')$ is "induced" from $G(V, E)$ by $V' \subseteq V$ if $E' = E - \{(u, v) \in E \text{ s.t. } u \notin V' \text{ or } v \notin V'\}$. $G'' = (V, E'')$ is the "complement" of G if $E'' = \{(u, v) \text{ s.t. } u, v \in V, u \neq v\} - E$.

The next elimination step is shown below for completeness.

$$D^{(2)} = \begin{array}{c|cccccccc} & 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 \\ \hline 1 & - & - & - & - & - & - & - & - \\ 2 & & - & - & - & - & - & - & - \\ 3 & & & xx & x & - & - & - & - \\ 4 & & & & xx & x & - & - & - \\ 5 & & & & & xoo & - & - & - \\ 6 & & & & & & x & oxx & - \\ 7 & & & & & & & oxxx & - \\ 8 & & & & & & & & x & xx \end{array} \quad G^{(2)} = \begin{array}{c} & & & & 5 & & & & \\ & & & & / & & \backslash & & \\ 6 & & & & & \text{---} & & & 7 \\ | & & & & & & | & & | \\ 3 & \text{---} & 4 & \text{---} & & & & & 8 \end{array}$$

The notion of "reachable sets" is useful for the study of Gaussian elimination (George and Liu, 1978a). Given a graph and a subset S of vertices, if u and v are two distinct vertices which do not belong to S , we say that v is

reachable from u through S when u and v are connected by a path which is either of length 1 (u and v are adjacent) or is composed entirely of vertices which belong to S . Given S and $u \notin S$, the reachable set $\text{Reach}(u, S)$ of u through S is defined to be the set of all vertices which are reachable from u through S . When S is empty, or when u does not belong to $\text{Adj}(S)$, then $\text{Reach}(u, S) = \text{Adj}(u)$.

A tree is an undirected acyclic graph which is connected and has no cycles. A matrix may have a graph which is a tree. If the tree is labelled in increasing order of vertex degree and the matrix is permuted accordingly, then Gaussian elimination with diagonal pivoting can be performed without any fill-in being produced. When the graph of a matrix is not a tree, then it is possible to find a tree partitioning of the graph, i.e., a partitioning such that the "quotient graph" is a tree. Let the set of vertices V of a graph G be partitioned into the disjoint subsets $\Pi_1, \Pi_2, \dots, \Pi_k$. The quotient graph has the k sets Π_i as composite vertices, and its edges are defined by the condition that (Π_i, Π_j) is an edge if and only if two vertices $u \in \Pi_i$ and $v \in \Pi_j$ can be found such that u and v are adjacent in G . When the quotient graph is a tree, it is called a "quotient," and the corresponding partitioning a "tree partitioning."

E. CURRENT METHODS

A number of reordering methods are known which dramatically reduce the fill-in resulting from factorization. These include the minimum degree algorithm, various dissection schemes, and various bandwidth or profile ordering schemes.

Band and profile methods regard the fill-in in a global manner, and confine it to certain areas, i.e., the band or profile of the matrix.

The dissection ordering method is a method for systematically partitioning the graph associated with a matrix using separators. In the dissection method, the fill-in caused by Gaussian elimination with diagonal pivoting is confined to nested partitioned areas of the matrix.

It is difficult to compare methods because the efficiency of a method depends on the problem or class of problems for which it is being used. Thus, we will only be concerned with algorithms which do not exploit special structure except implicitly. The method which is currently accepted as best for "unstructured" problems is the "minimum degree algorithm" of George and Liu [Ref. 4]. This thesis tests for efficiency the George and Liu algorithm and several other algorithms including variants of the George and Liu algorithm.

F. TEST PROBLEMS

The algorithms described in this thesis are tested on sixteen linear programming and mixed integer programming problems. These problems consist of fourteen real-world problems (AIR, COAL, COPS, BUS, ELPC, FOAM, JCAP, NETTING, PASTOR1, PAD, PIES, PILOT, STEEL, TIGER) and two artificial problems (STEINER1, STEINER2). The dimensions of these problems are shown in Table V.

All programs are written in FORTRAN, compiled under FORTRAN H-Extended (Opt 2) and run on an IBM 3033A² under VM/CMS. Computation times presented in the following chapters are accurate to the number of decimal places shown.

TABLE I
PROBLEM DIMENSIONS

Problem	Rows	Columns	¹ NZEL	Density	NCE	² m*(m-1)/2	Model
AIR	170	3040	6023	1.27	2.0	14365	Physical Distribution
BUS	56	330	3365	11.3	6.3	1540	Bus routing (Set Covering)
COAL	170	3753	7506	11.2	2.0	14365	National Energy Planning
CUPS	360	618	1341	0.6	2.2	64620	Production Scheduling
ELEC	784	2800	8462	0.4	3.0	306936	Energy Production & Consumption
FCAN	999	4020	13083	0.3	3.3	499501	Production Scheduling
JCAP	2486	3049	19510	0.1	2.5	3088655	Prod. & Shift Scheduling
NETTING	89	190	388	2.3	2.0	3916	International Currency Exchange
PASTOR1	657	2074	15854	0.8	5.2	215496	Combinatorics Problem
PAD	694	3297	15541	0.1	4.7	240471	Energy Alloc. & Cspn.
PIES	662	3011	13376	0.7	4.4	218791	Energy Prdn. & Dist.
PILOT	974	2112	12927	0.6	6.0	473851	Energy Development Planning
STEEL	831	1276	19808	0.9	7.7	344865	Economic Data
STEEL#1	117	27	352	11.1	13.0	6786	Artificial Data
STEEL#2	330	45	991	6.7	22.0	54285	Artificial Data
TIGER	160	636	4134	4.1	6.7	12720	Set Covering

¹ NZEL is the total number of nonzeros in the m by n constraint matrix.

² NCE is the average number of nonzeros in each column.

II. THE MINIMUM DEGREE ALGORITHM

A. . INTRODUCTION

In this chapter we discuss the minimum degree algorithm, with details of the implementation by George and Liu [Ref. 4]. This algorithm is the most commonly used ordering algorithm which does not explicitly exploit special matrix structures. It is used here as a standard for comparison against the algorithms given in chapter III.

The main idea of the minimum degree algorithm is that when we perform Gaussian elimination we choose the vertex of minimum degree from the elimination graph at every step. This typically minimizes the total fill-in versus the new fill-in. If there are several minimum degree vertices at a particular step, ties are broken arbitrarily.

B. BASIC ALGORITHM

Algorithm : The Basic Minimum Degree Algorithm.

Input : Elimination graph $G^{(0)}$ associated with a positive definite symmetric matrix.

Output : A permutation vector $PERM(\)$ for the vertices of $G^{(0)}$

Step 1. (Initialization)

Set $i = 1$.

Step 2. (Select minimum degree vertex)

In the elimination graph $G^{(i-1)} = (V_{i-1}, E_{i-1})$,

select a vertex $v_k \in V_{i-1}$ which is of minimum

degree.

Step 3. (Transform the graph)

Form the new elimination graph $G^{(i)} = (V_i, E_i)$

by eliminating the vertex v_k from $G^{(i-1)}$,

where all vertices adjacent to v_k in $G^{(i-1)}$

are connected to form a clique.

Set $PERM(i) = k$.

Step 4. (Loop or stop)

Set $i = i + 1$.

If $i > |V|$, go to step 5.

Otherwise, go to step 2.

Step 5. (Output)

For $j = 1$ to $|V|$, print $PERM(j)$.

End of The Basic Minimum Degree Algorithm.

C. ENHANCED ALGORITHM

We can enhance the minimum degree algorithm by using the concept of "indistinguishable" vertices by George and Liu(1981). Two unlabelled vertices $u, v \in V-S$ are said to be indistinguishable if $Reach(u, S) \cup \{u\} = Reach(v, S) \cup \{v\}$. At any step k , if we find indistinguishable vertices they can be labelled next without any minimum degree search.

Algorithm : The Enhanced Minimum Degree Algorithm.

Input : Elimination graph $G^{(0)}$ associated with a positive definite symmetric matrix.

Output : A permutation vector $PERM()$ for the vertices of $G^{(0)}$

Step 1. (Initialization)

Set $i = 1$.

Step 2. (Select minimum degree vertex)

In the elimination graph $G^{(i-1)} = (V_{i-1}, E_{i-1})$,

select a vertex $v \in V_{i-1}$ which is of minimum

degree.

Step 3. (Find the indistinguishable vertices)

Find the vertices which are indistinguishable from v . Let Y be the set of v and all vertices indistinguishable from v .

Step 4. (Elimination)

For each vertex $u \in Y$: Eliminate u in $G^{(i-1)}$

to obtain $G^{(i)}$, set $PERM(i) = u$, and set $i = i + 1$.

Step 5. (Loop or stop)

If $i > |V|$, go to step 6.

Otherwise go to step 2.

Step 6. (Output)

For $j = 1$ to $|V|$, print $PERM(j)$.

End of The Enhanced Minimum Degree Algorithm.

D. IMPLEMENTATION

The implementation of the minimum degree algorithm uses the basic ideas of elimination graphs, reachable sets, indistinguishable vertices and is further enhanced through a

quotient graph partitioning of the initial elimination graph.

The graph $G = (V, E)$ is stored using the adjacency array pair (XADJ, ADJNCY) in which the amount of storage used is $O(|V|)$ and $O(|E|)$, respectively. The amount of storage of the remaining arrays is $O(|V|)$. ADJNCY gives the adjacent sets $Adj(v)$ in contiguous storage locations for all v . XADJ(v) gives the location of the first element $Adj(v)$ in ADJNCY. The resulting ordering array pair is PRRQ. The algorithm requires the determination of reachable sets for degree update.

Vertices identified as indistinguishable are merged together to form a supervertex. They are treated as one vertex as the algorithm continues. The current degrees of the vertices in the elimination graph are kept in the array DEG.

An initial partitioning of the graph is made to create a quotient graph which is a tree. Then, vertices of the elimination graph are selected only from within subsets of vertices making up a composite degree-1 vertex of the the quotient graph. In this way, fill-in can be guaranteed not to extend outside of the quotient graph. This technique helps keep fill-in low globally.

III. THE MINIMUM FILL-IN ALGORITHM AND VARIANTS

A. INTRODUCTION

In this chapter we introduce the minimum fill-in algorithm. In the minimum degree algorithm, the main concept is to select that row and column which yields the least total fill-in in the next active submatrix, while in the minimum fill-in algorithm the central idea is to select that row and column which introduces the least new fill-in.

Suppose $\{v_1, v_2, \dots, v_{i-1}\}$ have been labelled. It is a simple matter, conceptually, to compute the new fill-in which would be introduced by selecting any vertex as the next vertex to be labelled as v_i . $\text{Fill}(v_i)$ is the number of vertices $v_j, v_k \in \text{Adj}(v_i)$ such that v_j is not adjacent to v_k . In practice, this number can be computed easily if $\text{Adj}(v)$ is kept as a linked list for all unlabelled vertices v . Below, the minimum fill-in algorithm is described followed by more detail on its implementation.

B. THE MINIMUM FILL-IN ALGORITHM

Algorithm : The Minimum Fill-in Algorithm.

Input : Elimination graph $G(0)$ associated with a positive definite symmetric matrix.

Output: A permutation vector $\text{PERM}()$ for the vertices of $G(0)$

Step 1. (Initialization)

Set $i = 1$.

Compute $\text{FILLIN}(v)$ for all $v \in V$, the amount of new fill-in which would be created if v is eliminated.

Step 2. (Select minimum fill-in vertex)

In the elimination graph $G^{(i-1)} = (V_{i-1}, E_{i-1})$,

select a vertex $v_k \in V_{i-1}$ such that $FILLIN(v)$

is minimum.

Step 3. (Transform the graph)

Form the new elimination graph $G^{(i)} = (V_i, E_i)$

by eliminating the vertex v_k from $G^{(i-1)}$,

where all vertices adjacent to v_k in $G^{(i-1)}$

are connected to form a clique.

Set $PERM(i) = k$.

Step 4. (Update FILLIN array)

Update $FILLIN(v)$ for all $v \in V_i$.

Step 5. (Loop or stop)

Set $i = i+1$.

If $i > |V|$, go to step 6.

Otherwise go to step 2.

Step 6. (Output)

For $j = 1$ to $|V|$, print $PERM(j)$.

End of The Minimum Fill-in Algorithm.

C. IMPLEMENTATION OF THE MINIMUM FILL-IN ALGORITHM

The graph $G = (V, E)$ is stored using the linked adjacency list arrays (XADJ, ADJNCY, LINK1) [Ref. 4] where the pointer XADJ(i) starts the adjacency list for vertex i, with ADJNCY containing a neighbor of vertex i and LINK containing the pointer to the location of the next neighbor of vertex i. The number of vertices in V is given by NEQNS. This data structure requires more space than the George and Liu algorithm; but, it requires space proportional to that necessary for carrying out one of the Gaussian elimination techniques. Thus, this is not really a limitation if storage is reused.

The purpose of the above algorithm is to find the minimum fill-in vertex for a general graph. It operates on the input graph as given by NEQNS and the linked adjacency list structure (XADJ, ADJNCY, LINK1), and returns the ordering in vector PERM.

The array FILLIN(v) stores the number of new fill-ins if vertex u is eliminated at the next step. FILLIN(v) is updated after each elimination step. This need be done only for those vertices which are a distance of two edges or less from u before the elimination step. The FILLIN value for vertices that have been eliminated is set negative.

Initial testing of the minimum fill-in algorithm has shown that it is much too time-consuming. Consequently, a restricted version has been developed which only searches among the minimum degree vertices for the minimum fill-in vertex. This is really a hybrid of the minimum degree algorithm and the minimum fill-in algorithm and is called the "min degree / min fill-in algorithm".

D. MIN DEGREE / MIN FILL-IN ALGORITHM

Algorithm : The Min Degree / Min Fill-in Algorithm.

Input : Elimination graph $G^{(0)}$ associated with a positive definite symmetric matrix.

Output: A permutation vector $PERM(i)$ for the vertices of $G^{(0)}$

Step 1. (Initialization)

Set $i = 1$ and $nd = 0$.

Step 2. (Lazy update of fill-in array FILLIN)

Find minimum degree $mindeg$.

If $nd \neq mindeg$, set $nd = mindeg$ and compute $FILLIN(v)$ for all vertices v with $deg(v) = nd$.

Otherwise go to step 3.

Step 3. (Selection)

In the elimination graph $G^{(i-1)} = (V_{i-1}, E_{i-1})$,

select a vertex $v_k \in V_{i-1}$ which is of minimum

fill-in among the minimum degree vertices.

Step 4. (Transform the graph)

Form the new elimination graph $G^{(i)} = (V_i, E_i)$

by eliminating the vertex v_k from $G^{(i-1)}$,

where all vertices adjacent to v_k in $G^{(i-1)}$

are connected to form a clique.

Set $PERM(i) = k$.

Step 5. (Loop or stop)

Set $i = i+1$.

If $i > |V|$, go to step 6.

Otherwise go to step 2.

Step 6. (Output)

For $j = 1$ to $|V|$, print $PERS(j)$.

End of The Min Degree / Min Fill-in Algorithm.

E. OTHER VARIANTS

The linked adjacency list structure used in the min degree / min fill-in algorithm makes it easy to try other ordering methods. Three additional methods will be tested: a simple minimum-degree algorithm, an initial minimum degree algorithm, and an initial ordering algorithm.

The "simple minimum-degree algorithm" is essentially the George and Liu algorithm but with a different data structure and without the reachable set and quotient graph enhancements. The "initial minimum degree algorithm" computes the degrees of the vertices in $G^{(0)}$ and selects vertices in increasing order of these degrees. No update of the initial degrees is ever made. The "initial ordering algorithm" just takes the vertices of $G^{(0)}$ in the order in which they are given, i.e., in the order in which the rows of the A matrix are given. This is a rather trivial algorithm but the time to create the compact data structure for sparse Gaussian elimination, included for the other algorithms is shown.

IV. COMPUTATIONAL EXPERIENCE AND DIFFICULTIES

To solve the sixteen test problems, we have used six methods to see which method is most effective. The methods are:

- Method 1. The minimum degree algorithm of George and Liu.
- Method 2. The minimum fill-in algorithm.
- Method 3. The min degree / min fill-in algorithm.
- Method 4. The simple minimum-degree algorithm.
- Method 5. The initial minimum degree algorithm.
- Method 6. Initial ordering algorithm.

Results for method 2 are not included because initial run times were excessive even for the smallest problems. As shown in Table III, two problems (JCAP, PAD) were not solved due to lack of computer memory. Time is CPU seconds necessary to obtain the ordering and to then create the compressed data structure used by George and Liu.

The minimum degree ordering algorithm of George and Liu is best. The simple minimum degree algorithm with linked data structures is almost as good - it is sometimes faster, and it is sometimes slower. But, new fill-ins created are always more than that of the minimum degree algorithm of George and Liu. The quotient graph enhancement of the George and Liu algorithm is probably responsible for its superior performance. Although storage requirements using the linked list data structure are greater, this only seemed to be a problem when the fill-in became very large. This structure gives a very flexible framework for trying different heuristics.

TABLE II
RESULTS OF ORDERING ALGORITHMS

Problem	Setup Time	Method 1		Method 3		Method 4		Method 5		Method 6	
		Fill	Time	Fill	Time	Fill	Time	Fill	Time	Fill	Time
AIR	2:3	1:3	13:1	2:9	3:6	2:6	1:8	1:3	1:2	77:8	0:20
BICAL	3:2	20:4	0:7	23:7	0:4	23:1	0:2	21:6	0:2	23:8	0:1
CCPS	1:7	1:6	0:8	11:9	11:5	11:9	2:5	8:4	2:4	11:9	0:0
PIPC	21:4	1:6	1:1	4:3	1:4	3:9	0:2	29:7	0:2	17:5	0:0
PCAP	41:3	0:2	21:6	30:9	16:8	30:5	1:9	29:8	1:7	13:0	0:1
JCAP	-	-	-	2:2	27:0	-	-	0:4	6:3	0:4	0:1
NETTING	0:1	0:5	0:0	0:7	0:1	0:7	0:0	0:8	0:0	37:9	0:0
PASTOR	27:7	4:1	0:6	35:6	8:1	35:4	0:9	30:7	1:3	25:8	0:1
PAAD	-	-	-	-	-	-	-	-	-	-	-
PIES	28:0	9:4	2:2	-	-	-	-	40:1	3:0	68:0	0:2
PILOT	29:1	9:4	12:3	-	-	21:8	6:4	22:1	16:4	41:8	0:4
STIEBEL	21:4	5:6	0:4	59:1	3:5	12:5	8:2	64:0	1:1	22:1	0:0
STEINER1	1:2	56:4	2:3	-	-	61:8	0:9	-	-	56:2	0:2
STEINER2	1:3	69:3	2:3	12:9	1:5	12:9	0:6	15:3	0:6	70:7	0:0
TIGER	1:5	9:9	0:6	-	-	-	-	-	-	20:6	0:0

- means not solvable with 2 megabytes of memory.
"fill" means new fill-in.

TABLE III
FILL-IN RESULTS

Problem	Initial	Final Fill-in		New Fill-in	
	Fill-in	No Order	Best Order	No Order	Best Order
AIR	20.8	98.6	22.1	77.8	1.3
BUS	55.7	79.5	76.4	23.8	20.6
COAL	26.1	38.0	34.5	11.9	2.4
CUPS	1.3	8.8	2.9	7.5	1.6
ELEC	2.0	15.0	3.6	13.0	1.6
FOAM	1.8	2.2	2.0	0.4	0.2
JCAP	-	-	-	-	-
NETTING	3.8	41.8	4.3	37.9	0.5
PASTOR1	1.7	27.4	5.7	25.8	4.1
PAD	-	-	-	-	-
PIES	1.9	69.8	11.0	68.0	9.1
PILOT	2.5	44.3	11.0	41.8	9.4
STEEL	3.0	25.1	8.7	22.1	5.6
STEINER1	31.0	87.2	87.4	56.2	56.4
STEINER2	19.1	89.8	89.4	70.7	69.3
TIGER	14.7	35.3	24.6	20.6	0.0

The min degree / min fill-in algorithm performed poorly. It is always slower, and the new fill-ins created are always more than that of the George and Liu algorithm. It cannot be recommended in any case and it is unclear why it performed so poorly.

Final fill-in for STEINER1 and STEINER2 in Table III is high, but these are artificial combinatoric problems. BUS is a small real-world problem with a best final fill-in of 76.4% which is much higher than the others. The probable reason for this is that BUS is a set partitioning problem.

TIGER with 24.6% final fill-in is also a set partitioning problem. This indicates that the projective algorithm might not work very well on many set partitioning and set covering problems since these seem to have a lot of intersections along rows. The lowest final fill-in found was 2.0% on FOAM. Six of the problems had less than 10.0% final fill-in. An interesting comparison can be made between the initial fill-in and the original densities of these problems. The initial fill-in can be compared to the density of matrix A which is about the same as the density for a basis in the simplex algorithm. This gives an idea of how hard creating L is compared to creating B^{-1} for the simplex algorithm. The initial fill-in of BUS, STEINER1 and STEINER2 was greater than 19.1% and they had densities greater than 6.7%. But, AIR, COAL and TIGER had more than 14.7% initial fill-in, while these had densities of only 1.2%, 1.2% and 4.1% respectively. The simplex algorithm might have a relatively greater advantage in these last cases.

V. CONCLUSIONS AND RECOMMENDATIONS

This thesis examined various heuristic algorithms for ordering symmetric linear systems of equations to minimize fill-in. The algorithms were tested on problems derived from linear programming problems in the manner of the projective algorithm for linear programming.

The minimum degree algorithm of George and Liu is the most effective of six tested methods. Solution times for this algorithm are sometimes better than and sometimes worse than the times using a simple minimum-degree algorithm, but new fill-ins created are always less. As a result of storage limitations, the simple minimum degree algorithm failed to solve two problems which the George and Liu algorithm solved. This should not be a problem in practice if storage is reused.

The minimum fill-in algorithm and the min degree / min fill-in algorithm used here do not perform as well as might be hoped. The minimum fill-in algorithm is too slow to use in most cases. Both algorithms involve more work than the minimum degree algorithm of George and Liu and the ordering produced is rarely such better than the one produced by the minimum degree algorithm of George and Liu.

The quotient graph enhancement of the George and Liu algorithm is probably responsible for its superior performance. Thus, future research should examine the use of the quotient graph and the min degree / min fill-in ideas since it may still be possible to significantly improve upon the new fill-ins obtained without sacrificing such computational speed. Different heuristics for creating quotient graphs should be examined, too.

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