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Automatic identification of embedded network rows in large-scale optimization models

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AUTOMATIC IDENTIFICATION OF EMBEDDED NETWORK ROWS
IN LARGE-SCALE OPTIMIZATION MODELS

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November 1980

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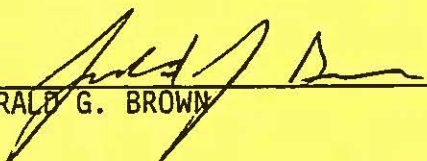
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
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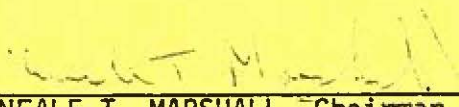
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

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AUTOMATIC IDENTIFICATION OF EMBEDDED NETWORK ROWS
IN LARGE-SCALE OPTIMIZATION MODELS

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ABSTRACT

The solution of a contemporary large-scale linear, integer, or mixed-integer programming problem is often facilitated by the exploitation of intrinsic special structure in the model. This paper deals with the problem of identifying embedded pure network rows within the coefficient matrix of such models and presents two heuristic algorithms for identifying such structure. The problem of identifying the maximum-size embedded pure network is shown to be among the class of NP-hard problems; therefore, the polynomially-bounded, efficient algorithms presented here do not guarantee network sets of maximum size. However, upper bounds on the size of the maximum network set are developed and used to evaluate the algorithms. Computational tests with large-scale, real-world models are presented.

Key words: Networks, Large-Scale Optimization, Basis Factorization,
Computational Complexity, Mixed Integer Optimization
Generalized Upper Bounds

Abbreviated Title: Identifying Embedded Networks

1. INTRODUCTION

The success of mathematical optimization and the increase in size and speed of digital computers have led to the formulation of very large and complex systems as mathematical programming models. The direct solution of the associated linear programming (LP) problems using the classical simplex method is often prohibitively expensive, if not impossible in a practical sense.

Large-scale models are predominantly characterized by sparse coefficient matrices and inherent special structure. If special structure can be identified, it can often be used to reduce the apparent problem monolith to components of more manageable size, or to admit enhancement of solution procedures. We are concerned with structures which can be used in factorization algorithms, for which all simplex bases share a common structure under row partition, and with structures which invite decomposition. The details of actual exploitation of special structure, once identified, will not be discussed here (e.g., see [6], or [7]).

Useful factorizations (even for a subset of columns) include simple bounds, generalized (upper) bounds (GUB), and embedded network rows, among others. Simple bound rows have only one non-zero coefficient. GUB refers to a set of rows for which each column (restricted to those rows) has at most one non-zero coefficient. Embedded network rows refers to a set of rows for which each column (restricted to those rows) has at most two non-zero coefficients of opposite sign. If the non-zero coefficients in the embedded network rows are restricted to at most one +1 and one -1 in each column, then the structure is referred to as an embedded pure network (NET).

Various methods are available to identify and exhibit special structure in the coefficient matrix. These range from simple permutation of rows and

columns to full (linear) transformations of the coefficient matrix. An intermediate method allows simple scaling (multiplication by a non-zero constant) of each row and/or column. Generally, entire transformation methods are used in an attempt to convert the complete coefficient matrix to one having a very special structure, such as a node-arc incidence matrix for a network. Partial transformation methods look for large subsets on the coefficient matrix which exhibit the desired structure, with the implicit presumption that large subsets are more efficiently exploited than small subsets.

Much of the computational improvement of the specialized simplex algorithms is obtained when logic can be substituted for arithmetic in simplex operations. This is most conveniently accomplished when the coefficient values in the special structure set are restricted to 0, ± 1 . This restriction can be satisfied by considering only subsets of the coefficients with intrinsic 0, ± 1 entries. In practice, however, it is often possible, through row and/or column scaling, to induce the desired 0, ± 1 values. For simple upper bounds, row scaling will suffice. GUB sets can be converted with row and column scaling (except that columns corresponding to integer variables are not customarily scaled). To produce pure network rows, however, the scaling problem is non-trivial due to the existence of two non-zero coefficients in many columns as well as the requirement that unit elements in the same column be of opposite sign.

The use of GUB has received much attention since the concept was introduced in 1964 by Dantzig and Van Slyke [4]. Some form of GUB has been implemented in many commercial LP systems, though restrictions on what constitutes an admissible (i.e., implemented) GUB set vary. Work has been done in the automatic identification of GUB sets [2], [9]; computational results on large-scale problems indicate that this is not only feasible, but can be extremely advantageous [3], [14].

Although some elegant work has been done in the theory of entire conversion of a linear program to a network problem ([1], [11]), few practical results have been achieved which reliably identify a subset (of rows) which forms a network structure if entire conversion fails. An efficient algorithm for doing so is of considerable value since a model usually fails to be completely convertible, and since the expense of attempting entire conversion may be prohibitive.

The problem of finding a maximum GUB set (in terms of number of rows) within a general coefficient matrix has been shown by Thomen to be NP-hard [14]. We prove the same result for the maximum embedded pure network problem. The implication is that currently only exponential-time algorithms exist to solve these types of problems and the hope of finding a more efficient algorithm is dim.

Therefore, the efficient identification methods we have developed have been heuristic algorithms--they find large, sometimes even maximum structures, but they cannot guarantee a maximum result. Since the size of the maximum structure is not known for the large-scale problems with which we work, we develop upper bounds on this size to evaluate our heuristics [14].

Computational results are given for a number of large-scale, real-world problems. They show the NET identification algorithms to be very effective and efficient in identifying large sets of pure network rows.

Some of this research has been summarized in [16].

2. PROBLEM DEFINITION AND REPRESENTATIONS

The Linear Programming Problem is defined here as:

$$\begin{aligned} \text{(L)} \quad & \text{minimize } c^t x \\ \text{s.t.} \quad & \underline{r} \leq Ax \leq \bar{r} \quad (\text{ranged constraints}) \\ & \underline{b} \leq x \leq \bar{b} \quad (\text{simple bounds}) \end{aligned}$$

where \underline{r} and \bar{r} are m -vectors, x , c , \underline{b} and \bar{b} are n -vectors and A is an $m \times n$ matrix. Consider for the moment the case where all x variables are real-valued; the integer and mixed integer cases are admitted later.

The (maximum) GUB problem (a well-known paradigm of embedded structure) for (L) can be stated as:

(GUB) Find a (maximum) subset of rows in A which can be scaled to contain only 0, +1 entries and which satisfy the property that each column of A (restricted to those rows) has at most one non-zero entry.

The real values of the non-zero coefficients in A do not make a difference in the GUB problem, since any non-zero entry in a GUB row can be scaled to +1 by column scaling alone. Therefore, it is convenient to replace A by a binary (0,1) matrix, K , of the same dimension where each non-zero entry of A is replaced by +1 with all other entries zero.

Using the matrix K , with entries k_{ij} , the (maximum) GUB problem can be formulated as the binary integer program

$$\begin{aligned} \text{(GUBI)} \quad & \text{(maximize)} \quad z_1 + z_2 + \dots + z_m \\ \text{s.t.} \quad & \sum_i k_{ij} z_i \leq 1; \quad j = 1, \dots, n \\ & \text{where } z_i \in \{0, 1\} . \end{aligned}$$

(z_i is an indicator variable for GUB inclusion.)

Alternative representations of the GUB problem have been developed as the basis for various heuristic algorithms and for theoretical considerations such as determining the complexity of the problem and developing bounds on the maximum achievable size of a GUB set. These include graphical conflict, conflict matrix, and vector space representations [14].

Two rows in A are said to *conflict* if there is at least one column of A with non-zero entries in both rows. If each row of A is considered as a vertex in an undirected graph with two vertices connected by an edge whenever the corresponding rows conflict, then the (maximum) GUB problem becomes one of finding a (maximum) independent set of vertices in the graph. An independent set of vertices in a graph is a subset of the total vertex set with no two vertices adjacent (connected by an edge) in the graph.

The conflict matrix representation of the GUB problem uses an $m \times m$ symmetric binary matrix M with each row and column representing a row of A . M has +1 values in those i, j entries where row i and row j conflict in A . By definition, every row conflicts with itself so the main diagonal of M has all +1 entries. The (maximum) GUB problem then becomes one of finding (through permutation of the rows of A) an embedded identity matrix (of maximum size) in the conflict matrix M .

The vector space representation [13] considers each row of K as a vector in n -space having unit length in those directions corresponding to its non-zero entries. The vector R is formed as the sum of each of the row vectors. A unit hypercube in n -space situated at the origin with length 1 in all positive directions represents the feasible GUB region. If R extends beyond this region, the set of rows is not a GUB set and at least one row must be removed to bring R into the feasible region. The (maximum) GUB problem becomes one of determining (the minimum number of) rows which must be removed

in order to bring R into the feasible region. The heuristics based on this representation compute gradient vectors which indicate the direction of shortest distance to the feasible region and remove first those rows which produce the greatest movement in that direction; these methods produce GUB sets of comparable quality to other heuristics, but have proven to be more computationally efficient.

The (maximum) Embedded Pure Network problem for (L) can be stated as:

(NET) Find a (maximum) subset of rows in A which can be scaled to contain only $0, \pm 1$ entries and which exhibit the property that each column of A (restricted to those rows) has at most two non-zero entries, and if the column has two non-zero entries, the (scaled) entries must be of opposite sign.

The real values of the non-zero coefficients in A cannot be ignored as they were in the GUB problem since simple column scaling is no longer sufficient to produce the required ± 1 entries in columns containing two non-zero entries. The addition of row scaling may help, but even this is not sufficient to guarantee that a network set of rows obtained by considering only the signs of the non-zero elements can be scaled to the required $0, \pm 1$ values.

Considering, for the moment only, matrices with $0, \pm 1$ entries (or a subset of m rows with $0, \pm 1$ entries in a general matrix) with no scaling allowed, the (maximum) NET problem can be formulated as the binary integer program:

$$\begin{aligned}
 \text{(NETI)} \quad & \text{maximize} \quad z_1 + z_2 + \dots + z_m \\
 & \text{s.t.} \quad \sum_{i: a_{ij} = -1} z_i \leq 1; \quad j = 1, \dots, n \\
 & \quad \quad \sum_{i: a_{ij} = +1} z_i \leq 1; \quad j = 1, \dots, n \\
 & \quad \quad \text{where} \quad z_i \in \{0, 1\} .
 \end{aligned}$$

(z_i is an indicator variable for inclusion in the network set.)

Unfortunately, NET does not lend itself to the many representations which GUB admits. The primary reason for this is that the scaling problems associated with NET make it impossible to disregard the real values of the non-zero coefficients in A . Also, the concept of pairwise row conflicts so useful in the GUB algorithms does not apply directly to network rows when row scaling is allowed.

To efficiently confront the scaling dilemma, we are currently forced to restrict the eligibility of rows for membership in the network set. The most obvious restriction is to allow no scaling and consider only those rows with intrinsic $0, \pm 1$ entries. Two less restrictive options are employed in the algorithms described later. These are:

1. Admit only rows with intrinsic $0, \pm 1$ entries but allow row *reflection* (multiplication of a row by -1).
2. Admit only rows whose non-zero entries can be row-scaled to $0, \pm 1$. This includes rows with all non-zero entries of the same absolute value.

Two representations of the NET problem are developed for the algorithms presented.

As suggested by Thomen [14], GUB heuristics can be used to produce a bipartite-network row factorization which can be partitioned into two subsets, G_1 and G_2 , such that each column of the matrix has at most one non-zero entry in G_1 and at most one non-zero entry in G_2 . Additionally, the entries must be of opposite sign. To produce such a (D-GUB) factorization, a GUB heuristic can be applied to the eligible rows of A producing G_1 , and then applied again to remaining eligible rows (not selected in the first pass and compatible for NET inclusion, allowing row reflection if necessary) giving G_2 .

If we consider only the rows of A with $0, \pm 1$ entries, or those which have been scaled to $0, \pm 1$, a vector space representation for NET can be developed

similar to that developed for GUB. The representation can also allow reflection of rows, if desired.

With each row in the eligible set, we associate two vectors in n -space, V_i^+ and V_i^- , each consisting of ± 1 in those dimensions corresponding to ± 1 entries in row i and zero in all other dimensions. For example, if row i is $(1,0,-1,1,0)$, then $V_i^+ = (1,0,0,1,0)$ and $V_i^- = (0,0,-1,0,0)$.

We define R^+ (R^-) as the resultant vector from the sum of all V_i^+ (V_i^-). These vectors extend from the origin into the orthants of n -space corresponding to all positive dimensions and all negative dimensions, respectively. A unit hypercube in each of these orthants constitutes the feasible NET region. Should either R^+ or R^- extend beyond its feasible region then the rows in the eligible set do not currently form an admissible set of network rows.

The reflection (multiplication by -1) of a row merely results in the switching of the V^+ and V^- vectors for the row. That is, when row i is reflected, the negative of V_i^+ becomes V_i^- and the negative of V_i^- becomes V_i^+ . This in turn will change the vectors R^+ and R^- . In fact, it is possible that just the reflection of these rows in an infeasible set may bring R^+ and R^- into their feasible regions without deletion of any rows.

If either R^+ or R^- extends beyond the feasible region, a row penalty for each is row is computed as the dot product of V_i^+ and R^+ plus the dot product of V_i^- and R^- . The row with the greatest row penalty is identified and the revised penalty for that row, if reflected, is computed. If this reflected penalty is less than the original row penalty, the row is reflected, otherwise it is deleted. When both R^+ and R^- fall within the feasible region, the set of rows which remain constitutes an admissible network set [15].

3. IMPLEMENTATION OF AUTOMATIC NETWORK IDENTIFICATION HEURISTICS

The D-GUB Algorithm:

Step 0. *Determine Eligible Rows.* Using the scaling scheme desired, determine which rows of the matrix are eligible for selection as network rows.

Step 1. *Find First GUB Set.* Apply a GUB heuristic to the eligible set.

Step 2. *Determine Eligibility for Second GUB Set.* For each eligible row not included in the first GUB set, check the columns in which the row has non-zero entries. In each of these columns, if the first GUB set has no non-zero entries or one non-zero entry of opposite sign then the row is eligible for inclusion in the second GUB set in its present form. If the first GUB set has no non-zero entries or a non-zero entry of like sign in each column, then the row is eligible for inclusion in reflected form. Otherwise, the row is not eligible and is discarded.

Step 3. *Find Second GUB Set.* If there are any rows eligible for the second pass, reapply the GUB heuristic to those rows.

The D-GUB Algorithm uses a two-phase, one-pass, non-backtracking GUB algorithm which is feasibility seeking (i.e., [3]). Phase 1 attempts to delete as few rows as possible in order to produce a feasible GUB set. Phase 2 examines the rows deleted in Phase 1 and reincludes rows which do not violate the GUB restriction.

Computational experience with many real-world models indicates that Phase 2 of the GUB heuristic rarely adds additional rows to the GUB sets obtained in either pass. For the second GUB set, Phase 2 is especially ineffectual. This suggests that the algorithm, which is already extremely fast, can be made even faster by the elimination of Phase 2 with minimal loss of solution quality.

The NET Algorithm is also two-phased, one-pass, non-backtracking and a deletion heuristic which is feasibility seeking. As such, it begins with an eligible set of rows which normally do not form an admissible network set and attempts to delete as few rows as possible to obtain a feasible set. Deleted rows are then considered for reinclusion if they do not violate the feasibility requirements.

The measure of infeasibility at any point is a matrix penalty computed as the sum of individual row penalties. Rows in the eligible set are examined in order of decreasing row penalty and either reflected, if the row penalty would be reduced, or removed and placed in a candidate set for later use. This guarantees that the matrix penalty will be reduced at each iteration. Thus, the number of iterations in Phase 1 is limited by the initial matrix penalty, which is *polynomially* bounded. In Phase 2, the rows in the candidate set are examined for reinclusion in the eligible set if they do not increase the matrix penalty. Those not reincluded are discarded.

Statement of the Problem:

Let $A = \{a_{ij}\}$ be an $m \times n$ matrix with $a_{ij} = 0, \pm 1 \forall i, j$.

Problem: Find a matrix $N = \{n_{ij}\}$ with $(m-k)$ rows and n columns which is derived from A by

1. Deleting k rows of A where $k \geq 0$,
2. Multiplying zero or more rows of A by -1 , where N has the property that each column of N has at most one $+1$ element and at most one -1 element. We wish to find a large N in the sense of containing as many rows as possible, i.e., minimize k .

Terminology and Notation:

1. E is the set of row indices for rows eligible for inclusion in N and is called the eligible set.

2. C is the set of row indices for rows removed from E in Phase I (Deletion). Some rows in C may be readmitted to E in Phase II. C is called the candidate set.
3. The phrase "reflect row i ' of A " means to multiply each element in row i ' by -1 , i.e., $a_{i',j} \leftarrow -a_{i',j} \forall j$.
4. Other notation will be defined in the algorithm itself.

The NET Algorithm:

Phase I - *Deletion of Infeasible Rows*

Step 0: *Initialization.* Set $E = \{1, 2, \dots, m\}$, $C = \phi$. For each column j of A compute the $+$ penalty (K_j^+) and the $-$ penalty (K_j^-) as follows:

$$K_j^+ = \left(\sum_{i \in E: a_{ij} > 0} 1 \right) - 1, \quad K_j^- = \left(\sum_{i \in E: a_{ij} < 0} 1 \right) - 1.$$

These penalties represent the number of excess $+1$ and -1 elements, respectively, in column j which prevent the rows in E from forming a valid N matrix. A penalty value of -1 for $K_j^+(K_j^-)$ indicates that the column does not contain a $+1(-1)$ element.

Step 1: *Define Row Penalties.* For every $i \in E$, compute a row penalty (p_i) as follows:

$$p_i = \sum_{j: a_{ij} > 0} K_j^+ + \sum_{j: a_{ij} < 0} K_j^-.$$

This is simply the sum of $+$ penalties for all columns in which row i has a $+1$ plus the sum of $-$ penalties for all columns in which row i has a -1 .

Step 2: *Define Matrix Penalty.* Compute the penalty (h) for the matrix by summing the row penalties as follows:

$$h = \sum_{i \in E} p_i.$$

If $h = 0$, then go to Step 7. Otherwise, go to Step 3.

Step 3: *Row Selection.* Find the row $i' \in E$ with the greatest penalty, i.e.,

$$\text{Find } i' \in E \text{ such that } p_{i'} = \max_{i \in E} p_i .$$

(If there is a tie, choose i' from among the tied values.) Compute the reflected row penalty $\bar{p}_{i'}$, for i' as follows:

$$\bar{p}_{i'} = \sum_{j: a_{i',j} > 0} (K_j^- + 1) + \sum_{j: a_{i',j} < 0} (K_j^+ + 1) .$$

This would be the row penalty for row i' if it were to be reflected.

Step 4: *Delete, or Reflect Row.*

Case i) $\bar{p}_{i'} \geq p_{i'}$. Let $E \leftarrow E - \{i'\}$, $C \leftarrow C \cup \{i'\}$. Go to Step 5.

Case ii) $\bar{p}_{i'} < p_{i'}$. Reflect row i' . Go to Step 6.

Step 5: *Reduce column penalties* as follows:

For all j such that $a_{i',j} > 0$, $K_j^+ \leftarrow K_j^+ - 1$

For all j such that $a_{i',j} < 0$, $K_j^- \leftarrow K_j^- - 1$

Go to Step 1.

Step 6: *Change column penalties* as follows:

Using the $a_{i',j}$ values after reflection of row i' ,

For all j such that $a_{i',j} > 0$, $K_j^+ \leftarrow K_j^+ + 1$ and $K_j^- \leftarrow K_j^- - 1$

For all j such that $a_{i',j} < 0$, $K_j^+ \leftarrow K_j^+ - 1$ and $K_j^- \leftarrow K_j^- + 1$

Go to Step 1.

Phase II - *Reinclusion of Rows from C*

Step 7: *Eliminate Conflicting Rows.* The rows in E , some possibly reflected

from the original A matrix, form a valid N matrix. However, some of

the rows removed from E and placed in C may now be reincluded in E

if they do not make $h > 0$. Remove from C (and discard) all rows which,

if reincluded in E in present or reflected form, would make $h > 0$.

I.e., remove i from C if

a) $\exists j_1$ such that $a_{ij_1} > 0$ and $K_{j_1}^+ = 0$

or $a_{ij_1} < 0$ and $K_{j_1}^- = 0$

and

- b) $\exists j_2$ such that $a_{ij_2} > 0$ and $K_{j_2}^- = 0$
or $a_{ij_2} < 0$ and $K_{j_2}^+ = 0$

If $C = \phi$, STOP, otherwise go to Step 8.

Step 8: *Select Row for Reinclusion.* At this point a row from C may be reincluded in E . There are several possible schemes for selecting the row. After the row is reincluded, the column penalties are adjusted. Then go to Step 7.

No dominating rule has been discovered for breaking ties in maximum row penalty encountered in Step 3. The rule used herein is to select the row with the minimum number of non-zero entries in an attempt to place a larger number of non-zero entries in the network set. Other possible rules are "first-come, first-served," maximum number of non-zero entries, type of constraint, or modeler preference.

Although the algorithm described above is presented for a matrix with strictly $0, \pm 1$ entries, it can be generalized to any matrix by simply letting E be the set of rows with strictly $0, \pm 1$ entries or which can be scaled to contain only $0, \pm 1$ entries.

Prespecified network rows can also be accommodated with the following modifications:

Let $P = \{i \mid \text{row } i \text{ is prespecified}\}$.

Then $E \leftarrow E - P$.

After computation of K_j^+ and K_j^- in Step 0, for each column j ,

if $\exists i \in P$ such that $a_{ij} = 1$ then $K_j^+ \leftarrow K_j^+ + 1$,

if $\exists i \in P$ such that $a_{ij} = -1$ then $K_j^- \leftarrow K_j^- + 1$.

Rows in P are not eligible for deletion or reflection. At the termination of the algorithm, the rows in N are given by EUP.

Computational experience on real-world models indicates that Phase 2 of the NET algorithm is even less productive than that of the GUB algorithm. In only two of sixteen cases were any rows eligible for reinclusion and the maximum number eligible was three. This indicates that the expense of examining the rows in the candidate set for eligibility is probably not justified for the occasional small improvement in quality.

It is easy to modify the NET algorithm to detect other embedded structures.

4. PROBLEM COMPLEXITY

Analysis of the inherent complexity of a problem can reveal whether there is a possibility of developing an efficient algorithm to completely solve all cases of the problem (e.g., [5], [10]). Unfortunately, analysis of the NET problem indicates that it cannot be solved optimally by an efficient algorithm at this time [15].

The problem of finding a GUB set of specified size (i.e., number of rows) is NP-complete, while that of finding a maximum GUB set is NP-hard [3], [14]. The corresponding maximum D-GUB problem with no scaling, since it represents a composition of two disjoint GUB problems, is also NP-complete (for a D-GUB set of specified size) and NP-hard (for a maximum D-GUB set).

The problem of *entire* conversion by general linear transformation of any matrix to the node-arc incidence matrix of a pure network is possible, has been shown to be polynomial in complexity [1], [11]. This, however, does not apply to the problem of finding the maximum embedded pure network should entire conversion fail. (The *entire* GUB problem is polynomial, too.) A slightly less complicated problem than finding the maximum size embedded network set is the following:

(NETD) Given an $m \times n$ matrix A and an integer $p < m$, determine whether A contains a set of p or more rows such that each column of A (restricted to those rows) has at most two non-zero entries, where entries in the same column must be of opposite sign.

Given a set of p rows from A , it is easy to verify, in polynomial time, whether the set satisfies the above criterion. Given an integer $p < m$, it is not easy to determine whether there exists a set of p or more rows in A which satisfies the criterion--in general, there does not currently exist an algorithm which can do so in polynomial time.

Two rows conflict if they both contain a non-zero element of like sign in a common column. The absence of such pairwise conflicts in a subset of rows from A is not a necessary condition for the rows to form a valid network set if row reflection is allowed. However, that condition is necessary and sufficient for that purpose when no scaling is allowed. With no scaling, it is evident that the absence of pairwise conflicts is necessary in a valid network set, for the existence of a conflict violates the opposite-sign requirement for columns containing two non-zero elements. It is also sufficient, because the violation of the criterion for a valid network set would require at least one column of A to contain at least two non-zero entries of like sign in rows of the set. This, in turn, would imply that the two rows in which this occurs are in conflict.

Consider a graph in which the nodes represent the rows of A and two nodes are connected by an edge if and only if the rows conflict in A . The problem of finding a set of p or more rows in A which do not conflict is then equivalent to finding an independent set of size p or more in the graph so defined. This problem, known as the independent set decision problem, is known to be NP-complete [5]. Furthermore, the problem of finding a maximum independent set, and therefore, a maximum GUB set or network set, is NP-hard.

The addition of row reflection to the problem simply means that each row can exist in one of two states, namely, unreflected or reflected. Clearly then, in a set of m rows, there are 2^m distinct states for the set, each corresponding to a different subset of reflected rows. The problem of finding a maximum network set in A , allowing row reflection, is equivalent to finding a maximum network set with no scaling allowed (shown above to be NP-hard) for 2^m distinct matrices. As a result, this problem is also NP-hard. For a general matrix in which non-zero entries may be of any magnitude, and allowing

simple row and column scaling, the problem of finding a maximum subset of rows which can be scaled to produce a pure network set is shown in [15] to be NP-hard, as well.

This analysis of network identification algorithms has only addressed the worst-case bound. No conclusions can be made about the average performance of an optimal algorithm--it may be possible to develop an optimal algorithm with good average performance, but having an exponential worst-case bound.

5. UPPER BOUNDS ON MAXIMUM NETWORK SET SIZE

The problem of finding a maximum-size, pure-network set of rows in a matrix, regardless of scaling restrictions, has been shown to be NP-hard. This also applies to the problem of determining just the size of a maximum set. Upper bounds on the maximum set size, computed in polynomial time, can be useful in evaluating the quality of network sets produced by heuristic algorithms.

The bounds developed here apply to the maximum set size obtainable from the set of eligible rows, and thus depend on the scaling restrictions employed. Clearly, the maximum set size can be no greater than the number of rows in the eligible set, but this bound is of little practical use.

Each column of the matrix (restricted to the eligible set) is allowed at most two non-zero entries. If k represents the maximum number of non-zero entries in any column of A (considering only entries in eligible rows), then it is clear that at least $k-2$ rows must be deleted from the eligible set in order to make this "worst column" feasible. Since the column counts are readily available in the form of the column penalties (K_j^+ and K_j^-), the upper bound on the network set size for a matrix with m eligible rows is:

$$u_1 = m - \max_j (K_j^+ + K_j^-) .$$

This bound is evidently *sharp* in that matrices can be constructed for which it is achieved.

A tighter bound is based on a matrix penalty computed from column penalties, rather than row penalties as in the NET algorithm. This penalty is:

$$H = \sum_{j:K_j^+>0} K_j^+ + \sum_{j:K_j^->0} K_j^- .$$

Clearly, as long as $H > 0$, the rows remaining in the eligible set do not form a valid network set. The reflection of a row in the eligible set may decrease H , increase H , or leave it unchanged. The deletion of a row from the eligible set may decrease H , or leave it unchanged. The actual effect of a reflection or deletion depends on the rows remaining in the eligible set and their state (unreflected or reflected) at the time. However, it is possible to compute for each row the maximum possible reduction in H obtainable by reflection or deletion of the row, regardless of the other rows remaining in the eligible set. These maximum possible reductions are called the *reflection potential* and *deletion potential* for the row, respectively.

The bound is determined by finding the minimum number of row deletions necessary to reduce H to zero. This cannot, of course, be specified exactly; however, the result will be conservative in that it will guarantee that at least that number of rows must be deleted.

Case	K_j^L	K_j^U
1	0	-1
2	0	0
3	0	>0
4	>0	-1
5	>0	0
6	>0	>0

K_j^L = column penalty of like sign to a_{ij}

(K_j^+ if $a_{ij} > 0$; K_j^- if $a_{ij} < 0$)

K_j^U = column penalty of unlike sign to a_{ij}

Table 1

Consider the possible states of a column j of A in which row i has a non-zero entry (i.e., $a_{ij} \neq 0$). The six possible cases are summarized in Table 1.

The non-zero entries in each column are counted only when they occur in the initial eligible set. The penalties used are those computed before any row reflections or deletions have occurred.

Consider first the effect on column j , and thus H , of reflecting row i . In cases 1, 5, and 6, reflection of row i would not change H . In case 4, reflection of row i would decrease H by 1, unless another row with a non-zero in column j was previously reflected. In cases 2 and 3, reflection of row i would actually increase H by 1, unless enough other rows with non-zero entries in column j were reflected or deleted to produce a -1 value for K_j^u . Since we cannot be sure that reflection in cases 2 and 3 would actually increase H , we must consider H unchanged by reflection in these cases. In summary, we allow H to be decreased only by reflection of rows with non-zero entries in columns exhibiting case 4. The reflection potential for row i is computed by summing the effects for each column in which row i has a non-zero element, with the condition that only one row reflection is allowed to decrease H for each column exhibiting case 4.

Row deletions provide greater opportunity for reducing H . In cases 1 and 2, deletion of row i has no effect on H , while in cases 4, 5, and 6, deletion of row i directly decreases H by 1. In case 3, deletion of row i does not directly decrease H , but it allows reflection of another row with a non-zero in column j , producing a net decrease of 1 in the value of H . In summary, we allow H to be decreased by deletion of rows with non-zero entries in columns exhibiting case 3, 4, 5, or 6. The deletion potential for row i is computed by summing the effects for each column in which row i has a non-zero entry.

To obtain this bound, the reflection and deletion potentials for each row in the eligible set are computed. Then the maximum possible reduction of H by row reflections alone is computed by summing the individual row reflection potentials. If $H > 0$ at this point, then rows must be deleted. Rows are deleted in order of decreasing deletion potential until $H \leq 0$. The upper bound is then computed as:

$$u_2 = m - \text{number of rows deleted,}$$

where m is the number of rows in the initial eligible set.

This bound is evidently sharp, since examples can be constructed which satisfy the bound exactly.

Similar arguments can be used to construct even better bounds, but the additional computation cost may not be justified for routine use with every model.

6. COMPUTATIONAL RESULTS

The D-GUB and NET algorithms were coded in FORTRAN IV and were tested on the set of real-world models with characteristics shown in Table 2.

The results obtained for the D-GUB algorithm are given in Table 3. The row eligibility criterion used was that each contain only $0, \pm 1$ entries, or be able to be scaled to $0, \pm 1$ entries by row scaling only. The number of eligible rows as a fraction of the total row count ranged from 9% to 100% (the objective row(s) not being eligible in any case). The number of GUB rows obtained in each pass is indicated. In two cases, the entire eligible set was determined to be a GUB set, so no second pass was required. The times given are in CPU seconds for the IBM 360/67 with the program compiled using FORTRAN H (Extended) with code optimization (OPT = 2).

The results for the NET algorithm are given in Table 4. Also included are the upper bounds on the maximum pure network set size computed from the problem data. The times given for determining the eligible set should be nearly the same as those for the D-GUB algorithm since the same eligibility criterion and code were used in both cases. The eligibility of rows in the candidate set for reinclusion in Phase 2 was determined, but Phase 2 was not included due to the absence of eligible rows in nearly every case. The solution time does not include the time required to determine eligibility for Phase 2. The NET quality value is the number of rows in the network set, expressed as a percentage of the better upper bound on the pure network set size. As explained earlier, the actual maximum network set size is, in general, unknown and thus the actual NET quality may be better than this conservative estimate. In particular, the bounds are almost certainly too high for problems with a large number of eligible rows (e.g., PAPER) and for problems with dense, or unstructured coefficient matrices (e.g., TRUCK, which is included in this study as a deliberate torture-test).

TABLE 2

SAMPLE LP (MIP) MODEL CHARACTERISTICS

<u>MODEL</u>	<u>DESCRIPTION</u>	<u>ROWS</u>	<u>COLUMNS</u>		<u>NON-ZERO COEFFICIENTS</u>
			<u>TOTAL</u>	<u>BINARY</u>	
NETTING	Currency Exchange	90	177	114	375
AIRLP	Distribution	171	3,040	0	6,023
COAL	Energy Development	171	3,753	0	7,506
TRUCK	Fleet Dispatch (Set Covering)	220	4,752	4,752	30,074
CUPS	Production Scheduling	361	582	145	1,341
FERT	Production & Distribution	606	9,024	0	40,484
PIES	Energy Production & Consumption	663	2,923	0	13,288
PAD	Energy Production & Consumption	695	3,934	0	13,459
ELEC	Energy Production & Consumption	785	2,800	0	8,462
GAS	Production Scheduling	799	5,536	0	27,474
PILOT	Energy Production & Consumption	976	2,172	0	13,057
FOAM	Production Scheduling	1,000	4,020	42	13,083
LANG	Equipment & Manpower Scheduling	1,236	1,425	0	22,028
JCAP	Production Scheduling	2,487	3,849	560	9,510
PAPER	Econometric Production	3,529	6,543	0	32,644
ODSAS	Manpower Planning	4,648	4,683	0	30,520

TABLE 3

D-GUB ALGORITHM RESULTS

<u>MODEL</u>	<u>ELIGIBILITY</u>		<u>NETWORK ROWS FOUND</u>						<u>ROWS REFLECTED</u>	<u>NONZEROS IN SET</u>
	<u>ROWS</u>	<u>TIME</u>	<u>PASS 1</u>	<u>TIME</u>	<u>PASS 2</u>	<u>TIME</u>	<u>TOTAL</u>	<u>TIME</u>		
NETTING	59	0.01	36	0.03	18	0.04	54	0.07	18	89
AIRLP	150	0.09	150	0.41	ALL GUB		150	0.41	0	3,000
COAL	111	0.11	111	0.50	ALL GUB		111	0.50	0	3,753
TRUCK	219	0.76	29	3.96	18	4.44	47	8.40	18	1,755
CUPS	300	0.05	150	0.14	101	0.15	251	0.29	1	710
FERT	585	0.62	559	3.00	13	3.03	572	6.03	13	16,291
PIES	142	0.09	128	0.51	0	0.05	128	0.56	0	1,392
PAD	174	0.10	160	0.52	0	0.06	160	0.58	0	1,552
ELEC	322	0.12	266	0.47	6	0.52	272	0.99	6	2,691
GAS	752	0.58	607	2.61	75	2.39	682	5.00	75	9,008
PILOT	109	0.10	96	0.44	13	0.48	109	0.92	1	479
FOAM	966	0.33	917	0.95	34	0.94	951	1.89	1	8,001
LANG	850	0.26	342	2.91	243	0.83	585	3.74	1	1,804
JCAP	1,811	0.26	517	1.49	357	1.01	874	2.50	201	2,622
PAPER	2,324	0.79	1,016	3.53	468	3.71	1,484	7.24	433	8,176
ODSAS	410	0.61	195	1.84	122	1.55	317	3.39	92	5,344

TABLE 4

NET ALGORITHM RESULTS

MODEL	ELIGIBILITY		UPPER BOUNDS		NETWORK ROWS FOUND			ROWS REFLECTED	NONZEROS IN SET
	ROWS	TIME	U1	U2	NUMBER	TIME	QUALITY		
NETTING	59	0.01	58	57	54	0.08	94.74%	18	89
AIRLP	150	0.09	150	150	150	0.35	100%	0	3,000
COAL	111	0.11	111	111	111	0.43	100%	0	3,753
TRUCK	219	0.76	214	137	46	19.82	33.58%	18	1,781
CUPS	300	0.05	299	297	295	0.14	99.33%	1	862
FERT	585	0.62	584	572	572	6.15	100%	13	16,291
PIES	140	0.09	139	132	128	0.59	96.97%	0	1,392
PAD	174	0.10	171	164	160	0.59	97.56%	0	1,552
ELEC	322	0.14	310	306	286	2.07	93.46%	34	2,915
GAS	752	0.60	750	710	668	9.71	94.08%	33	11,002
PILOT	109	0.10	109	109	109	0.36	100%	1	479
FOAM	966	0.34	965	955	951	1.16	99.58%	1	8,001
LANG	850	0.29	836	758	661	14.82	87.20%	2	2,239
JCAP	1,811	0.26	1,801	1,092	917	44.07	83.97%	200	2,540
PAPER	2,324	0.66	2,316	2,072	1,627	94.16	78.52%	603	8,995
ODSAS	410	0.61	406	369	286	14.55	77.51%	45	6,207

7. CONCLUSION

The identification algorithms are very fast (especially when compared with computer time expended in any attempt to solve these large problems) and they consistently produce maximum or near maximum pure network sets (from the eligible rows) as evidenced by the upper bounds.

Better yet, they provide independent insights which can be used to explain and improve the model at hand, or make it easier to solve. For instance, several models in Table 2 have been revealed as multi-commodity production/transportation problems, a totally unexpected perspective for the model proponents. Further, these results have yielded prescriptive benefits for model solution, especially via decomposition.

Many problems exhibiting intrinsic network structure are disguised by their formulation and resist the simplistic attempts used here to rescale them. In particular, the COAL model is known to be an entire network if appropriately restated, but it is not yet evident how this is to be discovered using efficient, general, problem-independent automatic identification. Methods used to scale an entire matrix to $0, \pm 1$ values (see [1], [11]) can be attempted, but failing entire conversion the next step is not evident.

Using the conflict matrix method of Greenberg and Rarick [8], Schrage [12] reports finding several other embedded structures. Our experience with this method at large scale has not been encouraging. Its principal disadvantage is the requirement for some representation of the conflict matrix. We feel that the superior speed and modest region demands of the gradient method exhibited in both GUB and NET identification will carry over to the identification of other special structures. This approach is currently being pursued.

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