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<tr>
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</thead>
<tbody>
<tr>
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<td>R6</td>
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</tr>
</tbody>
</table>

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Data Structures for Generalized Network Algorithms

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Abstract
Generalized network problems involve the optimization of a flow through a network. In contrast to normal networks, generalized networks include multipliers which alter the flow as it passes through the arcs. This enables the modelling of changes in flow caused by factors such as interest rates, as well as by processes such as energy conversion.

Generalized network problems can be solved by using the simplex algorithm. However, if the simplex bases are represented as forests of quasi-trees, connected graphs with a single cycle, a far more efficient algorithm can be obtained. Matrix multiplications required by the simplex algorithm can be interpreted as traversals of the quasi-trees, and consequently many of the arithmetic operations can be replaced by logical operations. This increases the numerical stability of the algorithm.

In this paper we outline this approach and discuss some of the data structures that can be used in its implementation.

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1. Introduction
Generalized network models include the assignment, transportation and transshipment problems. The major advantage over these models is that the introduction of multipliers permits the modelling of problems where material appreciates, depreciates or changes from one form to another. The utility of generalized networks is illustrated by the many applications [1,2,3].

A generalized network model consists of a set of nodes N and a set of arcs A. The problem is to find a flow \( x_k \) (\( k \in A \)) along each of the arcs such that:

- (a) it is of minimum cost (i.e. it minimizes \( \sum c_i x_i \));
- (b) it satisfies the capacity constraints placed on the arcs (i.e. \( f_k \leq c_i \leq w_k \)); and
- (c) it is sufficient to meet certain demands \( d_i \) (\( i \in N \)) at each of the nodes. When \( d_i \) is negative, this is interpreted as a supply of \( (-d) \) units available at node \( i \).

For generalized networks this last requirement may be formulated as:

\[
\sum_{\text{Arcs leading to node } i} m_k x_k = \sum_{\text{Arcs leading from node } i} x_k = d_i \quad (3)
\]

The coefficients \( m_k \) are the multipliers, and they indicate by which factor the flow will increase (or decrease) as it passes along the arc. The above equation states that the total flow into node \( i \) minus the total flow from the node should equal the demand, or the negative of the supply, at that node.

The algorithm used to solve these models is derived from the simplex algorithm for linear programs, but, because of the special structure of generalized networks, the solution can be obtained thirty to fifty times faster than by means of a comparably sized linear program [4].

Past experience with this and a similar problem, namely the transshipment problem, has shown that the speed of the algorithm depends on the data structures used in its implementation.

2. Linear Programs and the Simplex Algorithm
The generalized network problem, as formulated above, corresponds to the capacitated linear program

\[
\text{Min } \quad c^T x \\
\text{subject to } \quad N x = d \\
\quad f \leq x \leq w \quad (4)
\]

The coefficient matrix \( N \) has one column corresponding to each arc in the network. Each column has at most two non-zero elements; if \( k = (i,j) \) is an arc leading from node \( i \) to node \( j \), then \( n_{ik} = -1 \) and \( n_{jk} = m_{ij} \); and if \( k = (i,i) \) is a loop on node \( i \), then \( n_{ik} = m_{ii} - 1 \).

The simplex algorithm, which is used to solve the linear program, maintains a set of \( n \) basic variables. The non-basic variables are taken to be at either their lower limit \( f_k \), or at their upper limit \( w_k \). Consequently, the values of the \( n \) basic variables are determined by the \( n \) linear equations in [4].

At each iteration the simplex algorithm determines whether movement of one of the non-basic variables \( x_k \) from its lower or upper limit will reduce the total cost. If there is no such variable, the current solution is optimal. Otherwise, the algorithm determines how far that non-basic variable can move before it, or one of the basic variables \( x_b \), reaches its upper or lower limit.

In the former case the non-basic variable \( x_k \) goes to its opposite limit, and in the latter case the basic variable \( x_b \) becomes non-basic while the non-basic variable \( x_k \) becomes basic. This is referred to as a pivot. The algorithm pivots until an optimal solution is obtained.

In brief the simplex algorithm employs three major steps:

(a) Firstly it establishes an initial basis;
(b) in subsequent iterations it determines whether a non-basic variable is eligible for entry to the basis; and if so,
(c) it computes the effect of changing the value of that variable on the variables that are currently basic.

In this paper we will concentrate on step (b) and show how this can be efficiently implemented for generalized network models. The simplifications obtained for this step are comparable to simplifications that can be obtained in the other two steps.

3. The Structure of the Basis and the Effects of a Pivot
It can be shown that the arcs, associated with the basic variables in a generalized network, form a forest of graphs, each having a single cycle and a number of attached sub-trees. These graphs are called quasi-trees [5].

As an example consider the network in Figure 1. This shows the basic arcs as solid lines and the non-basic arcs as dotted lines. The basic arcs form two separate quasi-trees.

When the algorithm pivots, either of the following may happen:

(a) the non-basic arc selected may move from its current level...
to its opposite level without any of the basic arcs becoming non-basic; or
(b) the non-basic arc replaces one of the basic arcs in the basis.
In the former case the structure of the basis is unaffected, but in the latter case a number of interesting occurrences may take place. When the basic arc leaves the basis, it either splits the cycle of a quasi-tree, transforming it into a tree as in Figure 2, or it disconnects a sub-tree from a quasi-tree as in Figure 3. In either case we are left with one tree and a number of quasi-trees.

The arc entering the basis changes this tree into a quasi-tree, either by forming a new cycle as in Figure 4, or by linking it onto an existing quasi-tree as in Figure 5.

4. The Predecessor Array
The basis can be represented by means of a predecessor array p(.). For a node on the cycle of quasi-tree this indicates the previous node encountered when circling the cycle in a clockwise direction. For a node in a tree attached to a cycle, the predecessor indicates the father of that node.

In Figures 1-5, the predecessors of each node is indicated by means of the arrows drawn on the basic arcs.

These figures also show that only those arcs on the path joining the incoming arc to the outgoing arc alter during a pivot and these changes merely involve a reversal of the direction of the arrows. Updating of the predecessor array during each pivot is thus very simple.

5. Finding a Variable Eligible for Entry to the Basis
In this section we consider the problem of finding a variable that will reduce the total cost when introduced into the basis.

Most books on linear programming contain an equation, which, apart from changes in notation, reads

$$c_k = c_k - c_B B^{-1} N_k$$  \(5\)

The quantity \(c_k\) is the reduced cost and indicates how the total cost will alter when the value of \(x_k\) is changed. If \(c_k\) is negative, then the total cost will decrease when \(x_k\) is increased. Similarly, if \(c_k\) is positive, we should decrease \(x_k\) to reduce the total cost. Consequently, the non-basic variables eligible for entry to the basis are those which are at their lower limit and have \(c_k\) negative, as well as those which are at their upper limit and have \(c_k\) positive.

Expression (5) involves
(a) the cost \(c_k\) associated with the variable \(x_k\);
(b) the costs \(c_B\) associated with all basic variables;
(c) the matrix \(B\) which comprises the columns of \(N\) corresponding to the basic variables; and
(d) \(N_k\) which is the \(k\)th column of \(N\).

Efficient implementations of the simplex algorithm normally store \(B^{-1}\) as a product of elementary matrices

\(B^{-1} = E_1 E_2 \ldots E_r\)

and then compute the quantity \(c_B B^{-1} N_k\) by using matrix multiplications.

For generalized networks an approach without matrix multiplications is used. Firstly, we write \(c_k B^{-1} N_k\) as \(u N_k\) by letting \(u = c_k B^{-1}\). The values \(u\) are the node potentials. We then observe that for an arc \(k=(i,j)\) leading from node \(i\) to \(j\), \(N\) has at most two non-zero values, \(n_{ik} = -1\) and \(n_{jk} = m_{kj}\). Consequently, the formula for the reduced cost simplifies to \(c_k = c_k + u_i - m_{kj} u_j\). For a loop on node \(i\), \(k=(i,i)\), the formula reduces to \(c_k = c_k + (1-m_{ii}) u_i\).

It is thus simple to compute the reduced costs — provided that we know the values of the node potentials.

6. Computing the Node Potentials
Since it is known that the reduced costs of all basic arcs are zero, we can, in principle, determine the node potentials by setting the reduced costs of the \(n\) basic arcs to zero and solving the resulting \(n\) linear equations.

The problem can be simplified by observing that we can com-

FIGURE 1. A Typical Network. The basic arcs, shown as solid lines, form two quasi-trees.

FIGURE 2. Situation after a cycle in Figure 1 has been split.

FIGURE 3. Situation after a tree in Figure 1 has been disconnected from a quasi-tree.

FIGURE 4. A new quasi-tree is formed by completing a cycle in Figure 2.

FIGURE 5. The tree has been connected to an existing quasi-tree in Figure 2.
pute the potential of a node in one of the sub-trees, if we know the potential of its predecessor (since the reduced cost of the intervening arc is zero). The potentials of the nodes in the sub-trees can thus be established by traversing the sub-trees in pre-order — we visit and compute the potential for a node after we have visited its predecessor.

The problem of determining the potentials of nodes lying on a cycle is slightly more complicated. If the potential of some node \( i \) is \( u_i \), we can circle the loop and compute the potentials of the other nodes \( j \) on that cycle in terms of \( u_i \). These potentials will all be a linear function \( a_i + b_i \), of the initial potential \( u_i \). During a first pass around the cycle we establish the values for \( a_i \) and \( b_i \).

The last node \( f \) encountered before returning to node \( i \) will relate \( u_f = a_i + b_i u_i \). This enables us to solve for \( u_i \).

On a second pass around the cycle we compute the potentials for the other nodes \( j \).

Not all the node potentials change at each pivot. When a tree is linked onto an existing quasi-tree, only the nodes on that tree need to be recomputed, and when a new cycle is formed all the potentials on the new quasi-tree need to be evaluated.

7. The Double-Linked Tree and the Thread Successor Array

In order to compute the node potentials we must be able to (a) circle the loop of a quasi-tree; and (b) traverse the sub-trees in pre-order.

The former can be done by using the predecessor array. While it is possible to implement the latter using only predecessor information, it is rather inefficient and for this reason a new data structure is normally introduced.

One method [6] that can be employed is storing for each node \( i \) the set \( S_i \) of its immediate neighbours in the quasi-tree. For each basic arc \((i,j)\) we have \( i \in S_j \) and \( j \in S_i \). Consequently, this is referred to as a double-linked tree.

For a node in a sub-tree its sons are the nodes \( S \cdot \{p(i)\} \). By using this information and a stack-based procedure, we can traverse the sub-tree in pre-order.

During each pivot one arc becomes basic while another arc becomes non-basic. Consequently, two elements are removed from the sets while two are introduced. Minimal updating is thus required.

One disadvantage of this method is that \( 4n \) words of storage are required for its implementation. An alternative [7] is to store the thread successor array, which uses only \( n \) words of storage. The thread successor of a node is the next node to be visited when traversing the tree in pre-order.

The thread successor array is more difficult to implement, since the elements can change drastically from one iteration to the next. Details for updating this structure can be found in Currin [6]. Computational experience shows that, although the thread successor method is slower than the double-linked tree method, the difference is not large and the thread successor method can be employed profitably when storage is at a premium.

8. Conclusions

Solving a generalized network model by using a specialized computer code is far more efficient than using a standard linear programming code.

Because of the special structure, the basis can be stored as a graph (a forest of quasi-trees). In contrast, a linear programming code has to store the inverse basis \( B^{-1} \). Similarly, calculations that are normally done using matrix multiplications can be replaced by simple scalar operations performed while traversing the quasi-trees.

The net result is that many arithmetic computations are omitted. Although there is an increase in the number of logical operations performed, these can be executed faster and more precisely than arithmetic operations. This results in the computer code being more efficient and numerically stable.

References

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Contents/Inhoud

Evaluating the Performance of Computer-Based Information Systems using a Restricted Linear Regression Model..................1
 P J S Bruwer

An improved Implementation of Grimbleby’s Algorithm*...............7
 S R Schach

Nebulas as Structural Data Models*........................................11
 H O van Rooyen and D J Weiermans

Data Structures for Generalized Network Algorithms*..................15
 D C Currin

Modelling Blocking on Admission of Tasks to Computer Systems*..................19
 S Wulf

Analytical Model of a Mixed-Workload MVS Computer System*..................21
 R J Mann

Add: The Automated Database Design Tool*............................25
 S Berman

A Disk Space Management System*...........................................29
 A J Cuthbertson, I J van Nierkerk, T Turton

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