

A Dynamic Programming Algorithm to Find All Solutions in a Neighborhood of the Optimum

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ABSTRACT

Just after he introduced dynamic programming, Richard Bellman with R. Kalaba in 1960 gave a method for finding K th best policies. Their method has been modified since then, but it is still not practical for many problems. This paper describes a new technique which modifies the usual backtracking procedure and lists all near-optimal policies. This practical algorithm is very much in the spirit of the original formulation of dynamic programming. An application to matching biological sequences is given.

INTRODUCTION

Finding near-optimal policies for dynamic programming models is usually stated in terms of searching for the first, second, ..., K th shortest paths between a specified origin and destination through a directed, acyclic network. A near-optimal solution is such a path whose length is within a specified distance or neighborhood of the optimum. Thus, the set of near-optimal paths includes the optimal path(s), whereas a set of paths generated from heuristic or approximate methods may not. Recently we presented a new algorithm for finding all paths from an origin to a destination whose length is within a specified distance of the shortest path(s) [2]. Details of implementation and comparison with K th shortest path methods will also be given here.

This algorithm was motivated by a study of the evolutionary distance problem in molecular biology. In this context, dynamic programming meth-

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ods are used to investigate evolutionary relationships between two DNA sequences [20]. Analysis by K th shortest path methods was not practical. This application is discussed below.

Pollack [16, 17] and Dreyfus [4] provide surveys of currently available K th shortest path methods for directed, acyclic networks. Bellman and Kalaba [1], later modified by Fan and Wang [7], use forward dynamic programming to find K th best paths. Dreyfus [4] modifies the minimum tree technique of Hoffman and Pavley [12] to yield a new algorithm. Dreyfus also demonstrates its superiority to the algorithm of Bellman and Kalaba. The algorithm is essentially a forward dynamic approach and uses both node labels and arc labels to find K best paths. Fox [9–11] and Lawler [13] present slight improvements of the Hoffman-Pavley-Dreyfus algorithm for certain networks by suggesting the use of special data structures called heaps in order to increase computational efficiency. Elmaghraby [6] presents another dynamic programming formulation, which is reviewed by Yen [23]. Minieka and Shier [14] develop, and Shier [18,19] refines, a path algebra approach. Neither algorithm is as good as that of Hoffman and Pavley (or its modification) in terms of computational and storage requirements.

The new algorithm described in this paper is at least as fast as any previously described method and requires much less storage. It is, in fact, the first practical algorithm for these problems. The importance of this new algorithm is to allow sensitivity analysis, robustness studies, or simple approximations to complex solutions to actually be obtained. This has not previously been possible.

ALGORITHM

The object of the "shortest path problem" is to locate the shortest path from node 1 to node N in an acyclic network of N nodes and A arcs. Each arc (i, j) has an associated weight $t(i, j)$. Dynamic programming, as described in Dreyfus and Law [5] and Denardo [3], solves this problem by recursively solving many optimization problems. Nodes i are labeled with $f(i)$, the length of the shortest path from node i to node N . Bellman's insight was his famous principle of optimality: "subpaths of optimal paths are themselves optimal." This is embodied in the recursion

$$f(i) = \min\{t(i, j) + f(j) : (i, j) \text{ an arc}\}.$$

The idea is that to reach i from N , the last step is from some node j . The node j must be reached in an optimal manner if j is on an optimal path from N to i . It only remains to note that $f(N) = 0$ is required to start the recursion. The subject of this paper is to give a new algorithm for near-optimal paths.

Whereas previous algorithms find K shortest paths, the new algorithm requires some percentage corresponding to an interval e above the optimal length $f(1)$ from the user. If a $p\%$ class of near-optimal paths is desired, then e is equal to $(p/100)f(1)$. All paths less than or equal to quantity $f(1) + e$ are then found by the algorithm. This is convenient in problems where the corresponding K is unknown and/or large.

While recursively calculating the node labels $f(i)$, no "pointer" or decision information needs to be kept. These node labels are found by working backwards from node N until node 1 is labelled. The new algorithm then performs a depth-first search with stacking, starting at node 1 and continuing until all near-optimal paths are output.

Consider a node x not equal to the destination. Some path P with cumulative distance d has led us to node x from node 1. The test for entry of the arc (x, y) and distance d onto the stack now takes the general form, for all $(x, y) \in A$,

$$d + t(x, y) + f(y) \leq f(1) + e, \quad (*)$$

where d is the cumulative distance to node x from node 1 by path P (not necessarily by shortest path), $t(x, y)$ is the distance from node x to node y , and $f(y)$ is the optimal remaining distance to node N from node y .

The algorithm eventually constructs a path P of length d from node 1 to node N . Then P and d are output and the stack is examined to see if other near-optimal paths exist. Hence the algorithm performs a last in, first out or depth-first search.

To justify the algorithm, suppose the algorithm has generated a path P of length d from node 1 to node x . The arc (x, y) added to path P is on at least one near-optimal path if and only if $d + t(x, y) + f(y) \leq f(1) + e$, since the best path from y to N has length $f(y)$.

Note that ties in path lengths present no special problems. It is important to see that an arc is not stacked unless it lies on some near-optimal path. Also, the test $(*)$ is never performed more than once for each node on any particular near-optimal path.

EXAMPLE

Consider the acyclic network given in Figure 1 with arc lengths $t(i, j)$ and nodes A through I , where nodes A and I represent origin and destination. Numbers above each node $[f(i)]$ represent the shortest length from that node to node I . Suppose the user requests all paths within 20% of the optimal path length 13, which is the node label at node A . This percentage implies an upper bound of 15.6. The ordered path array P contains the nodes of the

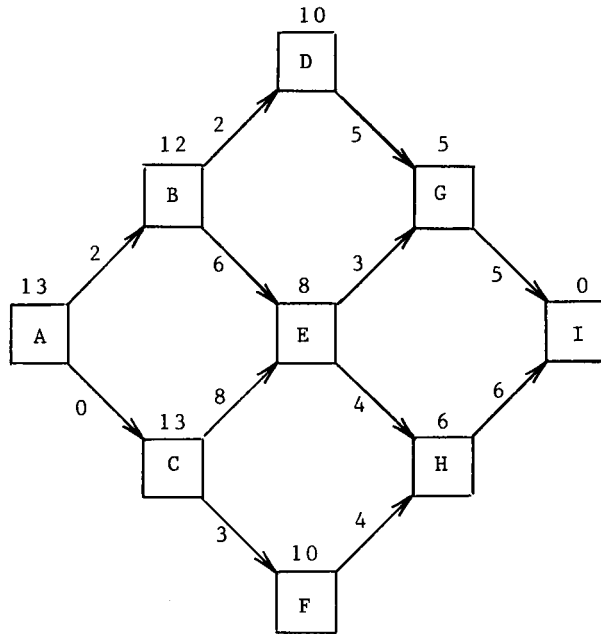


FIG. 1.

near-optimal path currently being traced out, and is initialized with node *A*.
An element of the stack contains three items of information:

- #1: The current node.
- #2: The next node.
- #3: The distance d from the origin (node *A*) to the node in #2.

The algorithm proceeds as follows:

Step 1. At node *A*, $d = 0$.

$t(A, B) + f(B) = 14$; PUSH element to stack.

$t(A, C) + f(C) = 13$; PUSH element to stack.

Stack contains:

#1	#2	#3
<i>A</i>	<i>B</i>	2
<i>A</i>	<i>C</i>	0

Step 2. POP last element of stack and put "next node" stored in #2 into P array = (A, C) . At node C , d equals the distance stored in #3 or $d = 0$.

$$d + t(C, E) + f(E) = 16; \text{ no action taken.}$$

$$d + t(C, F) + f(F) = 13; \text{ PUSH element.}$$

Stack contains:

#1	#2	#3
A	B	2
C	F	3

Step 3. POP last element. $P = (A, C, F)$. At node F , $d = 3$.

$$d + f(F, H) + f(H) = 13; \text{ PUSH element.}$$

Stack contains:

#1	#2	#3
A	B	2
F	H	7

Step 4. POP last element. $P = (A, C, F, H)$. At node H , $d = 7$.

$$d + f(H, I) + f(I) = 13; \text{ PUSH element.}$$

Stack contains:

#1	#2	#3
A	B	2
H	I	13

Step 5. POP last element. $P = (A, C, F, H, I)$. Node I has been reached, so path P with $d = 13$ is output. Stack now contains:

#1	#2	#3
A	B	2

Step 6. POP element. $P = (A, B)$, $d = 2$.

$$d + t(B, D) + f(D) = 14; \text{ PUSH element.}$$

$$d + t(B, E) + f(E) = 16; \text{ no action taken.}$$

Stack contains:

#1	#2	#3
B	D	4

Step 7. POP element. $P = (A, B, D)$, $d = 4$.

$$d + t(D, G) + f(G) = 14; \text{ PUSH element.}$$

Stack contains:

#1	#2	#3
D	G	9

Step 8. POP element. $P = (A, B, D, G)$, $d = 9$.

$$d + t(G, I) + f(I) = 14; \text{ PUSH element.}$$

Stack contains:

#1	#2	#3
G	I	14

Step 9. POP element. $P = (A, B, D, G, I)$. Node I has been reached, so path P with $d = 14$ is output. Stack is empty, so terminate.

To summarize, the near-optimal paths from node A to node I are:

$$\begin{aligned} A \rightarrow C \rightarrow F \rightarrow H \rightarrow I, & \quad \text{length 13,} \\ A \rightarrow B \rightarrow D \rightarrow G \rightarrow I, & \quad \text{length 14.} \end{aligned}$$

DISTANCE BETWEEN BIOLOGICAL SEQUENCES

Dynamic programming methods to compare macromolecular sequences started with Needleman and Wunsch [15], who introduced a similarity measure. Subsequently many modifications and advances have been made. A recent review by Waterman [22] discusses these methods as well as several which are not based on dynamic programming. The basic problem is to find the minimum weighted sum of substitutions and insertion/deletions to change the sequence $x = x_1 x_2 \dots x_n$ into $y = y_1 y_2 \dots y_m$. Let $d(x, y)$ denote the distance between the letter x and the letter y ; $d(x, y)$ can also be considered to be the weight associated with a substitution of y for x . Also let

$w(k)$ be the weight associated with a deletion (or insertion) of k consecutive letters.

Let $\rho(x,y)$ denote the distance between x and y . Define

$$\rho_{ij} = \rho(x_1 x_2 \dots x_i, y_1 y_2 \dots y_j), \quad \rho_{0,j} = w(j), \quad \rho_{i,0} = w(i).$$

The utility of dynamic programming for sequence comparisons is that it allows recursive calculation of $\rho_{i,j}$:

$$\begin{aligned} \rho_{i,j} = \min & \left\{ \min_{1 \leq k \leq i-1} \{ \rho_{k,j} + w(i-k) \}, \right. \\ & \rho_{i-1,j-1} + d(x_i, y_j), \\ & \left. \min_{1 \leq k \leq j-1} \{ \rho_{i,k} + w(j-k) \} \right\}. \end{aligned}$$

Since $\rho_{n,m} = \rho(x,y)$, it can be seen that $\rho(x,y)$ can be found in $O(n^3)$ steps when $n = m$. If $w(k) = ak + b$, the computation can be reduced to $O(n^2)$ steps.

The biologically correct values for $d(\cdot)$ and $w(\cdot)$ are not precisely known, although there are attempts to infer them from data. In addition, unknown biological constraints frequently act so that a "nonoptimal" sequence alignment is the biologically correct alignment. The algorithm described in this paper was developed to help with these difficulties.

As above, the algorithm is to find all alignments within e of $\rho_{n,m}$, the optimal alignment. It is possible to view $\rho_{n,m}$ as the length of the shortest path from $(0,0)$ to (n,m) . The algorithm described above solves this problem and was first presented with a specific application [21].

Fitch and Smith [8] studied sensitivity of alignments to the weighting functions. The example they used was taken from chicken hemoglobin mRNA sequences, 57 bases from the β chain and 39 bases from the α chain. A correct alignment is known here from analysis of many amino acid sequences for which these mRNA sequences code.

With a mismatch weight of 1 [that is, $d(xy) = 1$ if $x \neq y$] and $w(k) = 2.5 + k$, the correct alignment is found in the list of 14 optimal alignments. Here 14 alignments are in 0% of the optimum, 14 alignments in 1%, 35 alignments in 2%, 157 alignments in 3%, 579 alignments in 4%, and 1317 alignments in 5%.

With a mismatch weight of 1 and $w(k) = 2.5 + 0.5k$, the correct alignment is not one of the two optimal alignments. Not until the list of 31 alignments within 4% of the optimum is examined is the correct alignment found.

TABLE 1

Computational and Storage Requirements for
 $N^{0.5}$ Stages or Columns of Nodes,
 $N^{0.5}$ States or Rows of Nodes,
and R Decisions or Arcs/Node

	Computations	Storage
HP	$O(RN + KRN^{0.5})$	$O(N + KN^{0.5} + K)$
HPD without heaps	$O(RN + KRN^{0.5})$	$O(N + KN^{0.5} + RN)$
HPD with heaps	$O(RK + KN^{0.5} \log R)$	$O(RN)$
New algorithm	$O(RN + KRN^{0.5})$	$O(N + K)$

Not only does this example illustrate the sensitivity of alignments to weighting functions; it also shows the need for the new method. The problem described here has a network of 2200 nodes and 110,000 arcs. Analysis by K th shortest path techniques is not practical.

COMPARISON OF METHODS

Table 1 summarizes the computational and storage requirements of the Hoffman-Pavley (HP) algorithm, the Hoffman-Pavley-Dreyfus (HPD) without heaps, HPD with heaps, and the new algorithm. For comparative purposes only, the requirements are given for a square, acyclic network of N nodes and RN arcs, where the average number of arcs emanating from each node is denoted by R and each path contains $N^{0.5}$ nodes. Columns and rows of nodes can be viewed as dynamic programming stages and states, respectively. The set of paths within the specified percentage of the optimal length has exactly K members.

Computational requirements are estimates of the number of additions and comparisons necessary to find the K shortest paths. Storage requirements are estimates of the number of computer words needed, not including storage of the network itself.

Each estimate of computation appearing in Table 1 contains a term RN for computing the original node labels. As suggested by Dreyfus (personal communication), the published estimates for HPD in Fox [11] have been revised for the special case of a specified origin and destination. Only the $N^{0.5}$ nodes along each of the K near-optimal paths are updated during each iteration. As K becomes large, the heaps allow the algorithm to become more efficient than HP and HPD without heaps.

In estimating the new algorithm's requirements, a worst case situation has been assumed—none of the K paths use the same beginning arc. The K paths leave node 1 in K distinct directions, requiring the search to continue for all $N^{0.5}$ nodes of each path. This situation occurs in the simple path

problem of the previous section, but is unlikely to occur in most networks. If some of the K paths begin on the same arc or set of arcs, the requirements for the new algorithm are considerably less. Even in the worst case, the new algorithm remains computationally competitive to HPD with heaps and at least as good as HP and HPD without heaps.

The new algorithm requires significantly less storage than HP or either implementation of HPD. HP must store the original node labels, an ordered list of unused deviations, and the actual K paths (each of length $N^{0.5}$). HPD without heaps must store a set of arc labels, the original node labels, and any additional node labels along each path of $N^{0.5}$ nodes. HPD with heaps must store the arc labels and a heap of size R at each node N .

The new algorithm stores the original node labels and may require a stack of height K . The stack could become as large as the product of the average number of arcs in a path and the average number of arcs emanating from a node. This maximum height would only occur if K is greater than this product, a situation which is highly unlikely. For reasonable K , the stack length is negligible in comparison with storing the original node labels. In addition, since some near-optimal paths usually share beginning arcs, the requirements are considerably less. The key feature of the new algorithm remains its minimal storage requirements.

CONCLUSION

The new algorithm is easy to understand and install, which increases the likelihood of successful implementation and subsequent user acceptance. Only the backtracking or traceback procedure of a previously installed shortest path problem need be modified, leaving the major component of the model unaffected. As described in the introduction, K th best path methods are inadequate to solve most practical problems, while the new technique is practical for many of these problems.

Although the set of near-optimal solutions can be very large, it may contain information that is of interest. Frequently, the algorithm gives a sequence of paths where the (structural) differences between adjacent paths is small. In this situation, only every M th path, say, need output to the user. In some situations, it will be possible to only produce every M th path and reduce the running time accordingly. These problems can only be resolved in specific cases by careful consideration of the dynamic programming problem and the corresponding space of near-optimal solutions.

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