

Divide-and-Conquer Bidirectional Search

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Abstract

We present a new algorithm to reduce the space complexity of heuristic search. It is most effective for problem spaces that grow polynomially with problem size, but contain large numbers of cycles. For example, the problem of finding a lowest-cost corner-to-corner path in a D -dimensional grid has application to gene sequence alignment in computational biology. The main idea is to perform a bidirectional search, but saving only the OPEN lists and not the CLOSED lists. Once the search completes, we have one node in the middle of an optimal path, but don't have the solution path itself. The path is then reconstructed by recursively applying the same algorithm between the initial node and the middle node, and also between the middle node and the goal node. If n is the length of the grid in each dimension, and d is the number of dimensions, this algorithm reduces the memory requirement from $O(n^d)$ to $O(n^{d-1})$. For example, in a two-dimensional problem, the space complexity is reduced from quadratic to linear. The time complexity only increases by a constant factor of π in 2 dimensions, and $\pi\sqrt{3}/3 \approx 1.8$ in 3 dimensions.

1 Introduction

Consider an $n \times m$ grid, where each edge has a different cost associated with it. We want a lowest-cost path from say the upper-lefthand corner to the opposite lower-righthand corner, where the cost of a path is the sum of the costs of the edges in it. One important application of this problem is finding the best alignment of two genes of lengths n and m , represented as sequences of amino acids [Carrillo and Lipman, 1988]. We omit the details of the mapping between these two problems, and present it only as evidence that the problem is of practical importance. If we want to align d different sequences simultaneously, the problem generalizes to finding a lowest-cost path in a d -dimensional grid.

Another example of this problem is that of finding a shortest path in a maze. In this case, an edge between

two nodes either has a fixed cost such as zero, indicating that there is a direct path between the two nodes, or an infinite cost, indicating there is a wall between them.

In this paper, we consider only the orthogonal moves up, down, left and right, and want a lowest-cost path from one corner to another. The extension to include diagonal moves is straightforward. While our experiments use a two-dimensional grid, the algorithms we discuss trivially generalize to multiple dimensions. In general, a lowest-cost path will include some moves that do not go in the direction of the goal. For example, in going from upper-left to lower-right, a lowest-cost path may include some up and left moves, as our experiments will show.

2 Previous Work

2.1 Problems that Fit in Memory

If the grid is small enough to fit into memory, the problem is easily solved by existing methods, such as Dijkstra's single-source shortest path algorithm [Dijkstra, 1959]. It requires $O(nm)$ time, since each node has a constant number of neighbors, and $O(nm)$ space, in the worst case. Since we are only interested in the cost to the goal, we can terminate the algorithm when this value is computed, but in practice almost the entire grid will usually be searched, as our experiments will show.

If we restrict the problem to paths that only include down and right moves, it can be solved by a much simpler dynamic programming algorithm. We simply scan the grid from left to right and from top to bottom, storing at each point the cost of a lowest-cost path from the start node to that grid point. This is done by adding the cost of the edge from the left to the cost of the node immediately to the left, adding the cost of the edge from above to the cost of the node immediately above, and storing the smaller of these two sums in the current node. This also requires $O(nm)$ time and $O(nm)$ space.

The difference between these two algorithms is that in the general case, any of the neighbors of a node can be its predecessor along an optimal path, and hence Dijkstra's algorithm must maintain an Open list of nodes generated but not yet expanded, and process nodes in increasing order of their cost from the root.

Since both these algorithms store the entire graph

in memory in the worst case, the amount of available memory becomes the main bottleneck, and the time required is unlikely to be significant, unless the generation of individual nodes is very expensive. We implemented Dijkstra's algorithm for a two-dimensional grid using randomly generate edge costs, on a SUN ULTRA-SPARC Model 1 workstation with 128 megabytes of memory. The largest problem that we can solve without exhausting memory is a 3500×3500 grid, which contains 12,250,000 nodes. Our implementation requires about five and a half minutes to solve problems of this size.

2.2 Problems that Don't Fit in Memory

The problem is much more difficult when it is large enough that the entire grid cannot fit in memory. This can happen if all the edge costs are not explicitly listed, but are implicitly generated according to some rule. For example, in the gene sequence alignment problem, the edge cost is computed based on the particular pair of amino acids being aligned at a particular point in the two genes, and hence the number of different edge costs is only the square of the number of amino acids. In an even simpler example, finding the longest common subsequence in a pair of character strings [Hirschberg, 1975], the edge costs are just one or zero, depending on whether a pair of characters at a given position are the same or different, respectively.

In our experiments, each edge of a grid is assigned a random number, from a given position in the sequence of values returned by a pseudo-random number generator. In particular, we use the pseudo-random number generator on page 46 of [Kernighan and Ritchie, 1988]. This requires the ability to efficiently jump to an arbitrary point in the random sequence, an algorithm for which is given in [Korf and Chickering, 1996].

One approach to the memory problem is to use a heuristic search, such as A* [Hart, Nilsson, and Raphael, 1968] to reduce the amount of the problem space that must be searched. This assumes that a lower-bound heuristic, which estimates the lowest cost from a given node to the goal, can be efficiently computed, and has been applied to the gene sequencing problem [Ikeda and Imai, 1998]. If we establish a non-zero minimum edge cost in our random grids, we can use the manhattan distance to the goal times this minimum edge cost as a lower-bound heuristic. Unfortunately, A* must still store every node that it generates, and ultimately is limited by the amount of available memory.

The memory limitation of algorithms like Dijkstra's and A* has been addressed extensively by AI researchers over the last 15 years. See [Korf, 1995] for a brief survey. Many of these algorithms, such as iterative-deepening-A* (IDA*) [Korf, 1985], rely on a depth-first search to avoid the memory problems of best-first search. The key idea is that a depth-first search only has to keep in memory the path of nodes from the start to the current node, and as a result only requires memory that is linear in the maximum search depth.

While this technique is highly effective on problem

spaces that are trees, or only contain a small number of cycles, it becomes hopeless on a problem space with a large number of short cycles, such as a grid. The reason is that a depth-first search must generate every distinct path to a given node. In an $n \times m$ grid, the number of different paths of minimum length from one corner to the opposite corner is $(n+m)!/(n! \cdot m!)$. For example, a 10×10 grid, which contains only 100 nodes, has 184,756 different minimum-length paths from one corner to another, and a 30×30 grid, with only 900 nodes, has over 10^{17} such paths. A minimum-length path only includes moves that go toward the goal, meaning down and right moves in a path from upper-left to lower-right.

Another technique, based on finite-state-machines [Taylor and Korf, 1993], has been used to avoid this problem in regular problem spaces such as grids. Unfortunately, this method assumes that all shortest paths to a given node are equivalent, and does not apply when different edges have different costs.

Other techniques, such as caching some nodes that are generated, have been applied to these problems, with very limited success [Miura and Ishida, 1998]. The problem with these techniques is that they can only cache a small fraction of the total nodes that must be generated on a large problem.

We implemented IDA* on our random grid problems, with pitiful results. The largest problems that we ran were of size 10×10 . In addition to the problem of duplicate node generations, since most paths had different costs, each iteration on average only expanded about four new nodes that weren't expanded in the previous iteration. As a result, five problems of size 10×10 expanded an average of 1.128 billion nodes each, and took an average of about an hour each to run.

3 Divide & Conquer Bidirectional Search (DCBDS)

We now present our new algorithm. While we discovered it independently, a subsequent search of the literature revealed a special case of the main idea [Hirschberg, 1975]. For pedagogical purposes, we first describe our general algorithm, and then Hirschberg's special case.

A best-first search, such as Dijkstra's algorithm or A*, stores both a Closed list of nodes that have been expanded, and an Open list of nodes that have been generated, but not yet expanded. The Open list corresponds to the frontier of the search, while the Closed list corresponds to the interior region. Only nodes on the Open list are expanded, assuming the cost function is consistent, and thus in theory we could execute a best-first search without storing the Closed list at all.

In an exponential problem space, with a branching factor of two or more, the Open list will be larger than the Closed list, and not storing the Closed list doesn't save much. In a polynomial problem space, however, the dimensionality of the frontier or Open list is one less than that of the interior or Closed list, resulting in significant memory savings. For example, in a two-

dimensional problem space, the size of the Closed list is quadratic, while the size of the Open list is only linear.

There are two problems with this approach that must be addressed. The first is that normally duplicate node expansions are eliminated by checking new nodes against the Open and Closed lists. Without the Closed list, we have to prevent the search from “leaking” back into the closed region. This can be done by storing with each Open node a list of forbidden operators that would take us back into the closed region. Initially, for each node this is just the operator that generates its parent. When each new node is generated, it is compared against the nodes on the Open list, and if it already appears on Open, only the copy arrived at via the lowest-cost path is saved. When this happens, the new list of forbidden operators for the given state becomes the union of the forbidden operators of each copy.

In fact, this technique can be applied to the standard Dijkstra’s or A* algorithm with a Closed list as well. This speeds up the algorithms, since it is much faster to not generate a node at all, than to generate it and then search for it in the Open and Closed lists. In our grid experiments, this technique alone sped up our implementation of Dijkstra’s algorithm by over 25%.

The main value of this technique, however, is that it executes a best-first search without a Closed list, and without expanding the same state more than once. When the algorithm completes, we will have the cost of an optimal path to a goal node, but unfortunately not the path itself. If we store the path to each node with the node itself, each node will require space linear in its path length, eliminating all of our space savings. In fact, this approach requires more space than the standard solution of storing the paths via pointers through the Closed list, since it doesn’t allow us to share common subpaths.

The way to reconstruct the path is the following. We perform a bidirectional search from both the initial state and the goal state simultaneously, until the two search frontiers meet, at which point a node on a solution path has been found. Its cost is the sum of the path costs from each direction. We continue the search, keeping the best solution node found so far, until its cost is less than or equal to the sum of the lowest-cost nodes on each search frontier. At this point we are guaranteed to have a node roughly in the middle of a lowest-cost solution path. We add this middle node to the solution vector. Then, we recursively apply the same algorithm to find a path from the initial state to the middle node, and from the middle node to the goal state. Each of these searches will add another node to the final solution path, and generate two more recursive subproblems, etc, until we have built up the entire solution. We call this algorithm divide-and-conquer bidirectional search, or DCBDS.

Figure 1 shows an idealized view of DCBDS. The first node of the solution found is node *a*, and the two arcs that intersect at node *a* represent the final search horizons of the first bidirectional search. Next, we find node *b*, at the intersection of the arc centered at the initial state, and a circle centered at node *a*. The reason this

frontier looks like a circle is that it represents an uniformed Dijkstra’s algorithm, which doesn’t know the direction of the goal. The remaining nodes are shown in the order in which they would be found by DCBDS.

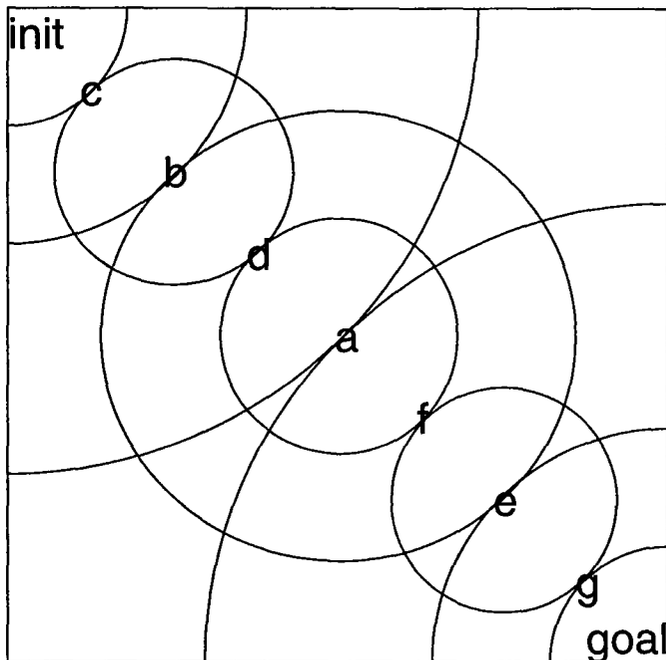


Figure 1: Divide and Conquer Bidirectional Search

3.1 Hirschberg’s Algorithm

In 1975, Hirschberg published an algorithm for computing a maximal common subsequence of two character strings in linear space [Hirschberg, 1975]. The algorithm computes a two-dimensional matrix, with each of the original strings placed along one axis. An element of the matrix corresponds to a pair of initial substrings of the two original strings, and contains the length of the maximal common subsequence of the two initial substrings.

If n and m are the lengths of the strings, the standard dynamic programming algorithm for this problem computes the matrix by scanning from left to right and top to bottom. This requires $O(nm)$ time and $O(nm)$ space. However, to compute the value of any point in this matrix, we only need the value immediately to its left and immediately above it. Thus, we can compute the length of the maximal common subsequence by only storing two rows of the matrix at a time, deleting each row as soon as the next row is completed. In fact, only one row needs to be stored, since we can replace elements of the row as soon as they are used. Unfortunately, this only yields the length of the maximal common subsequence, and not the subsequence itself.

Hirschberg’s linear-space algorithm computes the first half of the matrix from the top down, storing only one row at a time, and the second-half from the bottom up, again only storing one row. Then, given the two differ-

ent versions of the middle row, one from each direction, he finds the column for which the sum of the two corresponding elements from each direction is a maximum. This point splits both original strings in two parts, and the algorithm is then called recursively on the initial substrings, and on the final substrings as well.

The most important difference between DCBDS and Hirschberg's dynamic programming algorithm is that the latter scans the matrix in a predetermined systematic order, while DCBDS expands nodes in order of cost. The dynamic programming algorithm can only be used when we can distinguish the ancestors of a node from its descendents a priori. For example, it can find a lowest-cost path in a grid only if we restrict ourselves to shortest paths that only include moves toward the goal. DCBDS generalizes Hirschberg's dynamic programming algorithm to best-first search of arbitrary graphs.

4 Complexity of DCBDS

If the original problem is of size $O(n^d)$, DCBDS reduces the space complexity from $O(n^d)$ in the worst case to $O(n^{d-1})$, a very significant improvement. For example, if we can store ten million nodes in memory, this increases the size of two-dimensional problems we can solve from about $3,000 \times 3,000$ to about $2,500,000 \times 2,500,000$ before memory is exhausted, since the maximum size of each search frontier is roughly the sum of the lengths of the axes. In practice, time becomes the limiting factor for these larger grids, eliminating the space constraint.

The asymptotic time complexity of Hirschberg's algorithm, which only considers moves directly toward the goal, is the same as for the standard dynamic programming algorithm, or $O(n^2)$ on a two-dimensional grid.

To analyze the time complexity of DCBDS, we model the search frontiers as circles and arcs of circles. A search frontier represents an Open list, and consists of a set of nodes whose costs from the root node are approximately equal, since the lowest-cost node is always expanded next. In our experiments, we only consider the moves up, down, left, and right. Thus, a set of nodes whose distance from the root are equal, in terms of number of edges, would have a diamond shape, with points at the four compass points. In this diamond, however, the nodes at the points only have a single path of minimal distance to them, but the nodes closest to the diagonals through the center have a great many different paths to them, all of minimal distance. Thus, the lowest-cost path to a node near the diagonal is likely to be much smaller than the lowest-cost path to a node near a point of the diamond. Since the frontier represents a set of nodes of nearly equal lowest-path cost, the frontier near the diagonals bows out relative to the points of the diamond, approximating the circular shape. In fact, our graphic simulation of best-first search on a grid shows that the search frontiers are roughly circular in shape.

The time complexity is proportional to the number of nodes expanded, which is proportional to the area contained within the search frontier. Assume that we have

a square grid of size $r \times r$, whose lowest-cost path is along the diagonal, which is of length $r\sqrt{2}$. The first bidirectional search, to determine point a in Figure 1, will cover two quarter circles, each of which is of radius $r\sqrt{2}/2$, for a total area of $2 \cdot 1/4\pi(r\sqrt{2}/2)^2$, which equals $\pi r^2/4$. At the next level, we need two bidirectional searches, one to determine point b , and one for point e . This will generate two quarter circles from the initial and goal corners, plus the full circle centered at node a and reaching nodes b and e . This full circle will be generated twice, once to find node b , and once for node e . Thus, we generate $2 + 1/4$ circles, each of which are of radius $r\sqrt{2}/4$, for a total area of $5/2\pi(r\sqrt{2}/4)^2$ or $5\pi r^2/16$. At the third level, which generates nodes c , d , f , and g , we generate three full circles twice each, plus two quarter circles, all of radius $r\sqrt{2}/8$. In general, the set of searches at the n^{th} level of recursion sweep out a total area of $2(2^{n-1} - 1) + 2(1/4)$ circles, each of radius $r\sqrt{2}/2^n$, for a total area of

$$(2(2^{n-1} - 1) + 2(1/4))\pi \left(\frac{r\sqrt{2}}{2^n}\right)^2 = \pi r^2 \frac{2^{n+1} - 3}{2^{2n}}$$

The total area of all the searches is the sum of these terms from $n = 1$ to the number of levels. Since the sum converges, we can write it as the infinite sum

$$\pi r^2 \sum_{i=1}^{\infty} \frac{2^{n+1} - 3}{2^{2n}}$$

It is easy to show that this sum converges to one, so the total area, and hence time complexity of DCBDS, is πr^2 .

To find a lowest-cost corner-to-corner path, the search frontier of Dijkstra's algorithm will spread in a circular arc from the initial corner to the goal corner, at which point the entire grid will generally be covered. Since the area of the grid is r^2 , the overhead of DCBDS compared to Dijkstra's algorithm is a constant factor of π .

We can perform the same analysis in three dimensions, the differences being that the searches sweep out volumes of spheres instead of areas of circles, the main diagonal of a cube is $r\sqrt{3}$ instead of $r\sqrt{2}$, and the searches from the initial and goal states only generate eighths of a sphere, instead of quarters of a circle. In three dimensions, DCBDS generates a constant factor of $\pi\sqrt{3}/3 \approx 1.8138$ more nodes than Dijkstra's algorithm.

5 Experiments

We tested DCBDS on the problem of finding a lowest-cost corner-to-corner path on a two-dimensional grid. Each edge of the grid is assigned a random cost, and the cost of a path is the sum of the edge costs along it. We considered the general lowest-cost problem, which allows moves away from the goal as well as toward the goal. Using a technique that allows us to efficiently jump around in a pseudo-random number sequence without generating all the intermediate values [Korf and Chickering, 1996], we can search much larger random grids than we can store in memory.

Size	Shortest Path	Solution Length	Total Nodes	Dijkstra Nodes	DCBDS Nodes	Ratio
1,000	1,998	2,077	1,000,000	999,995	3,051,861	3.052
2,000	3,998	4,175	4,000,000	3,999,998	12,335,057	3.084
3,000	5,998	6,251	9,000,000	8,999,999	28,048,471	3.116
4,000	7,998	8,362	16,000,000		50,034,676	3.127
5,000	9,998	10,493	25,000,000		78,430,448	3.137
10,000	19,998	20,941	100,000,000		316,354,315	3.164
20,000	39,998	41,852	400,000,000		1,274,142,047	3.185
30,000	59,998	62,787	900,000,000		2,877,505,308	3.197
40,000	79,998	83,595	1,600,000,000		5,118,240,659	3.199
50,000	99,998	104,573	2,500,000,000		8,001,200,854	3.200

Table 1: Experimental Results for Corner-to-Corner Paths on Square Grids

Table 1 shows our experimental results. For each grid size, we tested DCBDS on three different random problem instances, generated from different random seeds, and averaged the results for the problem instances. The results from one problem instance to the next are very similar, so a sample size of three seems reasonable. The first column gives the length of the grid in each dimension, and the second gives the number of edges in a shortest corner-to-corner path, which is twice the grid size minus two. The third column gives the average number of edges in the lowest-cost corner-to-corner path, given the random edge costs. The reason some of these values are odd, even though all corner-to-corner paths are of even length, is because these values are averages of three trials. The data shows that in general a lowest-cost path is generally not a shortest path.

The fourth column gives the total number of nodes in the grid, which is the square of the grid size. The fifth column shows the average number of nodes expanded by Dijkstra’s algorithm, for problems small enough to fit in memory. This data shows that Dijkstra’s algorithm generates almost all the nodes in the grid. Since grids of size 4000 and greater are too large to fit in 128 megabytes of memory, we were unable to run Dijkstra’s algorithm on these problems, and hence those entries are empty.

The sixth column shows the average number of nodes expanded by DCBDS, and the sixth column shows the ratio of the number of nodes expanded by DCBDS divided by the number of nodes that would be expanded by Dijkstra’s algorithm, given sufficient memory. Even though we can’t run Dijkstra’s algorithm on problems greater than 3500 nodes on a side, we still compute a ratio on the assumption that Dijkstra’s algorithm would generate the entire grid, if there were sufficient memory. As predicted by our analysis, the number of nodes expanded by DCBDS is approximately $\pi = 3.14$ times the total number of grid points. This factor seems to increase slightly with increasing problem size, however.

The asymptotic running time of both algorithms is $O(n^2 \log n)$, where n is the size of the grid in one dimension. The n^2 term comes from the total number of nodes in the grid that must be examined. The $\log n$ term comes from the fact that both algorithms store the Open list as a heap, and the size of the Open list is $O(n)$, resulting

in $O(\log n)$ time per node to access the heap.

Even though DCBDS expands over three times as many nodes as Dijkstra’s algorithm, it takes less than twice as long to run. The main reason is that by saving the operators that have already been applied to a node, and not reapplying them, expanding a node takes less time than if all operators were applied and duplicates rejected by searching for them in the Closed list. The grids of size 10,000 take DCBDS about two hours to run, and those of size 50,000 take about 3 days. The grids of size 50,000 only require the storage of about 200,000 nodes, and even with only 128 megabytes of memory, we can store over twelve million nodes. Thus, memory is no longer a constraint.

6 Further Work

An obvious source of waste in DCBDS is that most of the individual searches are performed twice. For example, in Figure 1, the full circle centered at node a and including nodes b and e is searched once to locate node b , and then again to locate node e . By performing the search for nodes b and e simultaneously, we would only have to search the circle centered at a once. The same optimization can be applied to all the full-circle searches. Since most of the searches are full circles, this would reduce the time complexity by up to a factor of two, making DCBDS run almost as fast as Dijkstra’s algorithm.

The drawback of this optimization is that it complicates the implementation of the algorithm. In particular, all the searches at the same level of recursion must be performed simultaneously. For example, in Figure 1, the searches corresponding to the equal-sized circles centered at nodes a , b , and e must all be interleaved. This destroys the simple recursive structure of the algorithm, replacing it with an iterative outer loop with increasing numbers of interleaved searches in each iteration. In addition, some of these searches will terminate before others, and some searches may not be necessary at all.

Our current implementation continues executing recursive bidirectional searches until the initial and goal nodes are the same. Another obvious optimization would be to terminate the recursion when the problem size is small enough that there is sufficient memory to hold the entire subgrid in memory, and then execute Dijkstra’s al-

gorithm at that point. Since our analysis suggests that the lower-level searches in the recursion hierarchy contribute diminishingly less to the overall time, this optimization may not result in significant savings.

The idea of storing only the Open list in memory suggests yet another algorithm for this problem. What we could do is to execute a single pass of Dijkstra's algorithm, saving only the Open list, but writing out each node as it is expanded or closed to a secondary storage device, such as disk or magnetic tape, along with its parent node on an optimal path from the root. Then, once the search is completed, we reconstruct the solution path by scanning the file of closed nodes backwards, looking for the parent of the goal node, then the parent of that node, etc. Since nodes are expanded in nondecreasing order of their cost, we are guaranteed that the complete solution path can be reconstructed in a single backward pass through the file of closed nodes.

The advantage of this approach is that the capacity of most secondary storage devices is considerably greater than that of main memory, and we only have to access this device sequentially rather than randomly. Unfortunately, we aren't aware of any devices that can be read backwards very efficiently. The best we could probably do would be to simulate this by reading a disk file or tape file in blocks large enough to fit in main memory, and then access these blocks in reverse order. Given the efficiency of DCBDS and the slow speed of secondary storage devices, however, it seems unlikely that this will lead to a faster algorithm.

7 Conclusions

We generalized Hirschberg's dynamic programming technique to reduce the memory requirement of best-first search in arbitrary graphs with cycles. The most important difference between DCBDS and the dynamic programming algorithm is that the latter only works when we know a priori which neighbors of a node can be its ancestors and descendents, respectively, while DCBDS requires no such knowledge. For example, Hirschberg's algorithm can find a lowest-cost path in a grid if we only allow edges in the direction of the goal, whereas DCBDS allows arbitrary solution paths. Our experiments show that in general the lowest-cost path in a two-dimensional grid contains moves away from the goal as well.

DCBDS is most effective on polynomial-sized problems that are too big to fit entirely in memory. In such problems, it reduces the memory requirement from $O(n^d)$ to $O(n^{d-1})$. In practice, this removes the memory constraint from the problem. Our analysis shows that the time cost of this memory reduction is only a constant factor of π in two dimensions, which is supported by our experimental results. In three dimensions, our analysis predicts a constant overhead of $\pi\sqrt{3}/3 \approx 1.8138$. Further optimizations suggest that these constants could be reduced by up to a factor of two, and that additional constant savings may reduce the actual running time to no more than traditional best-first search. While our ex-

periments were done with Dijkstra's algorithm using no heuristics, everything trivially generalizes to A* with an admissible heuristic evaluation function as well.

The traditional drawback of bidirectional search has been its memory requirements. Ironically, DCBDS shows that bidirectional search can be used to save memory, and has the potential to revive study of this area.

8 Acknowledgements

Thanks to Toru Ishida and Teruhisa Miura for introducing me to the gene sequence alignment problem. This research was supported by NSF grant IRI-9619447.

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