ADAPTIVE GRIDS FOR GEOMETRIC OPERATIONS
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INTRODUCTION

Spatial and topological relationships are integral to cartography, and an efficient use of computer data structures is essential in automated cartography. A new data structure, the adaptive grid, is presented here. It allows the efficient determination of coincidence relationships, such as ‘Which pairs of edges intersect?’ and enclosure relationships like ‘Which region contains this point?’ A good reference for computer data structures is (Aho 1974). For cartographic data structures, see (Peucker 1975). Some other useful algorithms from the area of computational geometry are (Bentley 1979), (Bentley 1980), and (Dobkin 1979).

DATA STRUCTURE

The adaptive grid data structure (in 2-D) is based on a single level uniform grid superimposed on the data. For example, suppose that we are given N small straight line segments or edges scattered within a square of side one. (See Figure 1, where N = 4). The grid has G cells (G = 3 in Figure 1), each of side B = 1/G. Let L be a measure describing the edges’ length, such as the average. The data structure consists of a set of ordered pairs of a cell number and an edge number, i.e.,

\{(cell, edge)\}

Each pair represents an edge passing through a cell. For Figure 1, we obtain the following data structure:

\{(2,1), (1,1), (4,1), (2,2), (5,2), (4,3), (5,3), (6,3), (8,4)\}

This data structure bears a superficial resemblance to several others: unlike Warnock’s hidden line algorithm (see Sutherland 1974), the adaptive grid does not clip an edge into several pieces if it passes through several cells. In the example, edge #1 passes through cells #2, 1, and 4, but we merely obtain 3 ordered pairs in the set. Unlike a tree (see Nievergelt 1974), or k-d tree (see Bentley 1975a, and Willard), it divides the data coordinate space evenly independently of the order in which the data occur. Unlike a quad tree (see Bentley 1975b, and Finkel 1974) or octree (see Meagher 1982), the adaptive grid is one level, and does not subdivide in the regions where the data are denser. Although such a hierarchy is the most obvious ‘improvement’ to an adaptive grid, it will be shown later that if the data are reasonably distributed, then a hierarchy would not increase the speed. Further, such a hierarchy would make it harder to determine which cells a given edge passes through.

When the adaptive grid is described as a set, the term is used precisely. The only operations to be performed on it are:

1. Insert a new element, and
2. Retrieve all the elements in some order so that they may be sorted, adjacent elements combined under certain circumstances, and a new set created.

This gives a great freedom in implementing this abstract data structure. For example, on a microcomputer, the adaptive grid can be a sequential disk file, assuming that a file sorting routine exists.

Finding Intersections

We will now see how to determine all the pairs of edges that intersect. The operations are as follows:

1. Determine the optimal G, or resolution of the grid, from the statistics of the input edges. This will be described in more detail later, but letting G = 1/L is reasonable. Initialize an empty grid data structure for this G.
2. Make a single sequential pass through the input edges. For each edge, determine which grid cells it passes through, and for each such cell, insert a (cell, edge) pair into the grid data structure.

Since determining exactly which cells an edge passes through requires an extension of the Bresenham algorithm (see Foley 1982), which is a little complicated, in practice, an enclosing box is placed around the edge. The edge is considered to pass through all cells in this box. Considering an edge to be in
some extra cells speeds up this section and slows down the pair by pair comparison in section 5 below.

Retrieve the (cell, edge) pairs and sort them by cell number.

Make a sequential pass through the sorted list. For each cell mentioned in the list, determine all the edges that pass through that cell, and combine them into a set. Now we have a new set:

\[
\{(\text{cell, \{edge, edge, \ldots\}})\}
\]

with one element for each cell that has at least one edge passing through it.

Each element has the cell number and a smaller set of the edges that pass through that cell. Continuing the example of Figure 1, we have at this stage:

\[
\begin{align*}
&\{(1, \{1\}\}, \\
&(2, \{1,2\}\}, \\
&(4, \{1,3\}\}, \\
&(5, \{2,3\}\}, \\
&(6, \{3\}\}, \\
&(8, \{4\}\} \\
\end{align*}
\]

Note that since cells 3, 7, and 9 have no edges passing through them, they do not appear at all here. An empty cell does not use even one word of storage.

For each element of this set, i.e., for each cell with at least one edge passing through it, test all the edges in the cell pair by pair to determine which intersect. Since two edges that intersect must do so in some cell, and so must appear together in that cell, this will find all intersections. In the example, edges 1 and 3 both pass through cell 4, but an exact test shows that they do not intersect. On the other hand, cell 5 contains edges 2 and 3, and they do intersect.

If a pair of edges that do intersect appear together in more than one cell, then that intersection will be reported for each such cell. To avoid this duplication, when an intersection is found, it can be ignored unless it falls in the current cell. For this strategy to work, the cells must partition the space exactly, i.e., each point must fall in exactly one cell. This can be satisfied by considering each vertical grid line between two cells to be inside its right neighbor, and considering each horizontal grid line between two grid cells to be inside its upper neighbor.

**Timing**

His method is useful only because it executes efficiently. We will now analyze the time and determine the optimal \(G\).

Let \(U\) = the average number of cells that each edge falls in. Then, approximately

\[
U = 1 + 2 \, LG
\]

However, assuming that we place a box around the edge and count all cells in hat, as described above, we will get a higher figure:

\[
U \approx (1 + L/B)^2 = (1 + LG)^2,
\]

since \(BG = 1\)

We will use this higher number since it is more conservative and does not affect the rate of growth of the final time. Next, \(V = \text{total number of (cell, edge) pairs}\)

\[
= NU = N(1 + LG)^2
\]

Then, \(W = \text{the average number of edges in each cell}\)

\[
= V/G^2 = N(B + L)^4
\]

For the execution times, we will use the notation \(\theta(x)\), which means proportional to \(x\), as \(x\) increases.

Let \(T_1 = \text{the time to calculate the (cell, edge) pairs}\)

\[
= \theta(V)
\]

and \(T_2 = \text{the time to determine the intersections}\)

\[
= \theta(G^2W^2)
\]

since in each cell, each edge must be tested against each other edge. So \(T = \text{total time}\)

\[
T = T_1 + T_2 = \theta(N(1 + LG) + N^2G^2(B + L)^4)
\]

Now, if we let \(B = \theta(L)\), i.e., the grid size is proportional to the edge length, we get

\[
T = \theta(N + S)
\]

where

\[
S = N^2L^2/4
\]

since with \(\theta\) notation, which considers only the rate of growth, constant multipliers can be added freely. But \(S\) is approximately the expected number of edge intersections for a given \(N\) and \(L\). Since a routine that finds all edge intersections must examine each edge and each intersection at least once, setting \(B = cL\), for any \(c\), gives an optimal time, up to a constant factor. The actual \(c\) which minimizes \(T\) should be determined heuristically, since it depends on the relative speeds of various parts of the program, and this may depend on the model of the computer.

We sometimes have even more freedom to select \(B\), that is, there may be a range of functions for \(B\) that give the same minimum time, depending on the dependency of \(L\) on \(N\) as the problems get bigger. We can use this extra freedom to also minimize space if we wish.

It might be objected that the execution time for any cell depends on the average of the square of the number of edges in that cell, whereas we have used the square of the average. However, if the edges are independently and identically distributed, then each edge has the same independent probability of passing through any given cell. Thus the number of edges in a given cell is Poisson distributed, so the square of the mean equals the mean of the square.
The adaptive grid has been implemented and tested in various applications. First, the edge intersection algorithm described above was implemented as a Fortran preprocessor (see Beyer 1975) program on a Prime mid-computer. The largest example had 50,000 edges and 47,222 intersections. See Table 1 for a list of execution times.

<table>
<thead>
<tr>
<th>N</th>
<th>L</th>
<th>B</th>
<th>S</th>
<th>T1</th>
<th>T2</th>
<th>T</th>
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<td>364.72</td>
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</table>

**Column Labels**

- **N**: Number of edges
- **L**: Average length of edges, assuming screen is 1 by 1
- **B**: Length of side of each grid cell
- **S**: Number of intersections found
- **T1**: CPU time (sec.) to put edges in cells
- **T2**: CPU time to find the intersections among the edges
- **T**: Total CPU time

From the table, we see that this program takes about \((N + S)/300\) CPU seconds to execute, even for the largest case. Some other facts about the largest case tested are: 98,753 (cell, edge) pairs, and 11,534 duplicate intersections.

Figure 2 shows the case with \(N = 1000, G = 10, L = 0.5\) approximately. In these examples, the coordinates of the edges are generated with a pseudo-random number generator. Now, manufacturer-supplied random number generators are all linear congruential which has the effect that if you pair up successive random numbers, then the resulting 2-D points will fall on a comparatively small number of parallel lines. This does not mean that these edges will be parallel if a linear congruential generator is used since the random numbers are used for \(x_1, y_1\), and the angle of inclination. Still, it is better to use a non-linear generator.

The adaptive grid was next used in a hallowed line program designed and implemented by Varol Akman (see Akman 1981, and Franklin 1980). A hallowed line drawing is a means of displaying a 3-D wire frame model (i.e., edges but not faces) of an object with front lines cutting gaps in real lines where they cross. This was tested on scenes with over 10,000 edges. It can process such a scene in about 10 minutes on a Prime, depending on the edges' lengths.

The Spheres program (see Franklin 1981), is a third test. Here spheres are projected on top of one another. A case of ten thousand spheres of radius 0.02 overlaid on the average 10 deep through the scene, could be processed in 6.4 seconds. Here not only the intersections were determined, but also the visible segments of each sphere's perimeter were found.

Finally, the adaptive grid is an essential part of the simplified map overlay algorithm (see Franklin 1985), currently under implementation.

**Testing Whether a Point is in a Polygon**

The adaptive grid can also be used to test whether a polygon contains a point. If we preprocess the polygon first, this method is very efficient even if the polygon has many edges. The execution time per point depends on the depth complexity of the polygon, i.e., the average number of edges that a random scan line would cut. The total number of edges has no effect on the time.

This method is an extension of the method where a semi-infinite ray is
As before, we put the planar graph's edges into a 1-D adaptive grid and test the ray against only those edges in the same cell as P's projection.

**SUMMARY**

Techniques from computational geometry have been shown, both by theoretical analysis and by implementation, to lead to more efficient means of solving certain common operations in automated cartography.

**ACKNOWLEDGEMENT**

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**REFERENCES**


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**LOCATING A POINT IN A PLANAR GRAPH**

The obvious extension of the above problem is to take a planar graph and a point, P, and to determine which polygon of the graph contains P. This can be done by testing P in turn against each polygon in turn, unless one polygon completely contains another, but that is slow. A more efficient method is this:

1. Extend a ray up from P.
2. Record all the edges that it crosses, along with those edges' neighboring polygons.
3. Sort those edges by their ray crossings from P.
4. Finally, P is contained in the lower polygonal neighbor of the closest crossing edge to P.

**FIGURE 3.** Point in polygon testing.