## Multidimensional Spatial Data Structures

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### 16.1 Introduction

The representation of multidimensional data is an important issue in applications in diverse fields that include database management systems (Chapter 60), computer graphics (Chapter 54), computer vision, computational geometry (Chapters 62, 63 and 64), image processing (Chapter 57), geographic information systems (GIS) (Chapter 55), pattern recognition, VLSI design (Chapter 52), and others. The most common definition of multidimensional data is a collection of points in a higher dimensional space. These points can represent locations and objects in space as well as more general records where only some, or even none, of the attributes are locational. As an example of nonlocational point data, consider an employee record which has attributes corresponding to the employee's name, address, sex, age, height, weight, and social security number. Such records arise in database management systems and can be treated as points in, for this example, a seven-dimensional space (i.e., there is one dimension for each attribute) albeit the different dimensions have different type units (i.e., name and address are strings of characters, sex is binary; while age, height, weight, and social security number are numbers).

When multidimensional data corresponds to locational data, we have the additional property that all of the attributes have the same unit which is distance in space. In this case, we can combine the distance-denominated attributes and pose queries that involve proximity. For example, we may wish to find the closest city to Chicago within the two-dimensional space from which the locations of the cities are drawn. Another query seeks to find all cities within 50 miles of Chicago. In contrast, such queries are not very meaningful when the attributes do not have the same type.

[^0]When multidimensional data spans a continuous physical space (i.e., an infinite collection of locations), the issues become more interesting. In particular, we are no longer just interested in the locations of objects, but, in addition, we are also interested in the space that they occupy (i.e., their extent). Some example objects include lines (e.g., roads, rivers), regions (e.g., lakes, counties, buildings, crop maps, polygons, polyhedra), rectangles, and surfaces. The objects may be disjoint or could even overlap. One way to deal with such data is to store it explicitly by parameterizing it and thereby reduce it to a point in a higher dimensional space. For example, a line in two-dimensional space can be represented by the coordinate values of its endpoints (i.e., a pair of $x$ and a pair of $y$ coordinate values) and then stored as a point in a four-dimensional space (e.g., [33]). Thus, in effect, we have constructed a transformation (i.e., mapping) from a two-dimensional space (i.e., the space from which the lines are drawn) to a four-dimensional space (i.e., the space containing the representative point corresponding to the line).

The transformation approach is fine if we are just interested in retrieving the data. It is appropriate for queries about the objects (e.g., determining all lines that pass through a given point or that share an endpoint, etc.) and the immediate space that they occupy. However, the drawback of the transformation approach is that it ignores the geometry inherent in the data (e.g., the fact that a line passes through a particular region) and its relationship to the space in which it is embedded.

For example, suppose that we want to detect if two lines are near each other, or, alternatively, to find the nearest line to a given line. This is difficult to do in the four-dimensional space, regardless of how the data in it is organized, since proximity in the two-dimensional space from which the lines are drawn is not necessarily preserved in the four-dimensional space. In other words, although the two lines may be very close to each other, the Euclidean distance between their representative points may be quite large, unless the lines are approximately the same size, in which case proximity is preserved (e.g., [69]).

Of course, we could overcome these problems by projecting the lines back to the original space from which they were drawn, but in such a case, we may ask what was the point of using the transformation in the first place? In other words, at the least, the representation that we choose for the data should allow us to perform operations on the data. Thus when the multidimensional spatial data is nondiscrete, we need representations besides those that are designed for point data. The most common solution, and the one that we focus on in the rest of this chapter, is to use data structures that are based on spatial occupancy. Such methods decompose the space from which the spatial data is drawn (e.g., the twodimensional space containing the lines) into regions that are often called buckets because they often contain more than just one element. They are also commonly known as bucketing methods.

In this chapter, we explore a number of different representations of multidimensional data bearing the above issues in mind. While we cannot give exhaustive details of all of the data structures, we try to explain the intuition behind their development as well as to give literature pointers to where more information can be found. Many of these representations are described in greater detail in $[60,62,63]$ including an extensive bibliography. Our approach is primarily a descriptive one. Most of our examples are of two-dimensional spatial data although the representations are applicable to higher dimensional spaces as well.

At times, we discuss bounds on execution time and space requirements. Nevertheless, this information is presented in an inconsistent manner. The problem is that such analyses are very difficult to perform for many of the data structures that we present. This is especially true for the data structures that are based on spatial occupancy (e.g., quadtree (see Chapter 19 for more details) and R-tree (see Chapter 21 for more details) variants). In particular, such methods have good observable average-case behavior but may have very bad
worst cases which may only arise rarely in practice. Their analysis is beyond the scope of this chapter and usually we do not say anything about it. Nevertheless, these representations find frequent use in applications where their behavior is deemed acceptable, and is often found to be better than that of solutions whose theoretical behavior would appear to be superior. The problem is primarily attributed to the presence of large constant factors which are usually ignored in the big $O$ and $\Omega$ analyses [46].

The rest of this chapter is organized as follows. Section 16.2 reviews a number of representations of point data of arbitrary dimensionality. Section 16.3 describes bucketing methods that organize collections of spatial objects (as well as multidimensional point data) by aggregating the space that they occupy. The remaining sections focus on representations of non-point objects of different types. Section 16.4 covers representations of region data, while Section 16.5 discusses a subcase of region data which consists of collections of rectangles. Section 16.6 deals with curvilinear data which also includes polygonal subdivisions and collections of line segments. Section 16.7 contains a summary and a brief indication of some research issues.

### 16.2 Point Data

The simplest way to store point data of arbitrary dimension is in a sequential list. Accesses to the list can be sped up by forming sorted lists for the various attributes which are known as inverted lists (e.g., [45]). There is one list for each attribute. This enables pruning the search with respect to the value of one of the attributes. It should be clear that the inverted list is not particularly useful for multidimensional range searches. The problem is that it can only speed up the search for one of the attributes (termed the primary attribute). A widely used solution is exemplified by the fixed-grid method [10, 45]. It partitions the space from which the data is drawn into rectangular cells by overlaying it with a grid. Each grid cell $c$ contains a pointer to another structure (e.g., a list) which contains the set of points that lie in $c$. Associated with the grid is an access structure to enable the determination of the grid cell associated with a particular point $p$. This access structure acts like a directory and is usually in the form of a $d$-dimensional array with one entry per grid cell or a tree with one leaf node per grid cell.

There are two ways to build a fixed grid. We can either subdivide the space into equalsized intervals along each of the attributes (resulting in congruent grid cells) or place the subdivision lines at arbitrary positions that are dependent on the underlying data. In essence, the distinction is between organizing the data to be stored and organizing the embedding space from which the data is drawn [55]. In particular, when the grid cells are congruent (i.e., equal-sized when all of the attributes are locational with the same range and termed a uniform grid), use of an array access structure is quite simple and has the desirable property that the grid cell associated with point $p$ can be determined in constant time. Moreover, in this case, if the width of each grid cell is twice the search radius for a rectangular range query, then the average search time is $O\left(F \cdot 2^{d}\right)$ where $F$ is the number of points that have been found [12]. Figure 16.1 is an example of a uniform-grid representation for a search radius equal to 10 (i.e., a square of size $20 \times 20$ ).

Use of an array access structure when the grid cells are not congruent requires us to have a way of keeping track of their size so that we can determine the entry of the array access structure corresponding to the grid cell associated with point $p$. One way to do this is to make use of what are termed linear scales which indicate the positions of the grid lines (or partitioning hyperplanes in $d>2$ dimensions). Given a point $p$, we determine the grid cell in which $p$ lies by finding the "coordinate values" of the appropriate grid cell. The linear


FIGURE 16.1: Uniform-grid representation corresponding to a set of points with a search radius of 20 .
scales are usually implemented as one-dimensional trees containing ranges of values.
The array access structure is fine as long as the data is static. When the data is dynamic, it is likely that some of the grid cells become too full while other grid cells are empty. This means that we need to rebuild the grid (i.e., further partition the grid or reposition the grid partition lines or hyperplanes) so that the various grid cells are not too full. However, this creates many more empty grid cells as a result of repartitioning the grid (i.e., empty grid cells are split into more empty grid cells). The number of empty grid cells can be reduced by merging spatially-adjacent empty grid cells into larger empty grid cells, while splitting grid cells that are too full, thereby making the grid adaptive. The result is that we can no longer make use of an array access structure to retrieve the grid cell that contains query point $p$. Instead, we make use of a tree access structure in the form of a $k$-ary tree where $k$ is usually $2^{d}$. Thus what we have done is marry a $k$-ary tree with the fixed-grid method. This is the basis of the point quadtree [22] and the PR quadtree $[56,63]$ which are multidimensional generalizations of binary trees.

The difference between the point quadtree and the PR quadtree is the same as the difference between trees and tries [25], respectively. The binary search tree [45] is an example of the former since the boundaries of different regions in the search space are determined by the data being stored. Address computation methods such as radix searching [45] (also known as digital searching) are examples of the latter, since region boundaries are chosen from among locations that are fixed regardless of the content of the data set. The process is usually a recursive halving process in one dimension, recursive quartering in two dimensions, etc., and is known as regular decomposition.

In two dimensions, a point quadtree is just a two-dimensional binary search tree. The first point that is inserted serves as the root, while the second point is inserted into the relevant quadrant of the tree rooted at the first point. Clearly, the shape of the tree depends on the order in which the points were inserted. For example, Figure 16.2 is the point quadtree corresponding to the data of Figure 16.1 inserted in the order Chicago, Mobile, Toronto, Buffalo, Memphis, Omaha, Atlanta, and Miami.


FIGURE 16.2: A point quadtree and the records it represents corresponding to Figure 16.1: (a) the resulting partition of space, and (b) the tree representation.

In two dimensions, the PR quadtree is based on a recursive decomposition of the underlying space into four congruent (usually square in the case of locational attributes) cells until each cell contains no more than one point. For example, Figure 16.3a is the partition of the underlying space induced by the PR quadtree corresponding to the data of Figure 16.1, while Figure 16.3b is its tree representation. The shape of the PR quadtree is independent of the order in which data points are inserted into it. The disadvantage of the PR quadtree is that the maximum level of decomposition depends on the minimum separation between two points. In particular, if two points are very close, then the decomposition can be very deep. This can be overcome by viewing the cells or nodes as buckets with capacity $c$ and only decomposing a cell when it contains more than $c$ points.

As the dimensionality of the space increases, each level of decomposition of the quadtree results in many new cells as the fanout value of the tree is high (i.e., $2^{d}$ ). This is alleviated by making use of a $k-d$ tree [8]. The $\mathrm{k}-\mathrm{d}$ tree is a binary tree where at each level of the tree, we subdivide along a different attribute so that, assuming $d$ locational attributes, if the first split is along the $x$ axis, then after $d$ levels, we cycle back and again split along the $x$ axis. It is applicable to both the point quadtree and the PR quadtree (in which case we have a $P R k$ - $d$ tree, or a bintree in the case of region data).

At times, in the dynamic situation, the data volume becomes so large that a tree access structure such as the one used in the point and PR quadtrees is inefficient. In particular, the grid cells can become so numerous that they cannot all fit into memory thereby causing them to be grouped into sets (termed buckets) corresponding to physical storage units (i.e., pages) in secondary storage. The problem is that, depending on the implementation of the tree access structure, each time we must follow a pointer, we may need to make a disk access. Below, we discuss two possible solutions: one making use of an array access structure and one making use of an alternative tree access structure with a much larger fanout. We assume that the original decomposition process is such that the data is only associated with the leaf nodes of the original tree structure.

The difference from the array access structure used with the static fixed-grid method


FIGURE 16.3: A PR quadtree and the points it represents corresponding to Figure 16.1: (a) the resulting partition of space, (b) the tree representation, and (c) one possible $\mathrm{B}^{+}$-tree for the nonempty leaf grid cells where each node has a minimum of 2 and a maximum of 3 entries. The nonempty grid cells in (a) have been labeled with the name of the $\mathrm{B}^{+}$-tree leaf node in which they are a member.
described earlier is that the array access structure (termed grid directory) may be so large (e.g., when $d$ gets large) that it resides on disk as well, and the fact that the structure of the grid directory can change as the data volume grows or contracts. Each grid cell (i.e., an element of the grid directory) stores the address of a bucket (i.e., page) that contains the points associated with the grid cell. Notice that a bucket can correspond to more than one grid cell. Thus any page can be accessed by two disk operations: one to access the grid cell and one more to access the actual bucket.

This results in EXCELL [71] when the grid cells are congruent (i.e., equal-sized for locational data), and grid file [55] when the grid cells need not be congruent. The difference between these methods is most evident when a grid partition is necessary (i.e., when a bucket becomes too full and the bucket is not shared among several grid cells). In particular, a grid partition in the grid file only splits one interval in two thereby resulting in the insertion of a $(d-1)$-dimensional cross-section. On the other hand, a grid partition in EXCELL means that all intervals must be split in two thereby doubling the size of the grid directory.

An alternative to the array access structure is to assign an ordering to the grid cells resulting from the adaptive grid, and then to impose a tree access structure on the elements of the ordering that correspond to the nonempty grid cells. The ordering is analogous to using a mapping from $d$ dimensions to one dimension. There are many possible orderings (e.g., Chapter 2 in [60]) with the most popular shown in Figure 16.4.

The domain of these mappings is the set of locations of the smallest possible grid cells (termed pixels) in the underlying space and thus we need to use some easily identifiable pixel in each grid cell such as the one in the grid cell's lower-left corner. Of course, we


FIGURE 16.4: The result of applying four common different space-ordering methods to an $8 \times 8$ collection of pixels whose first element is in the upper-left corner: (a) row order, (b) row-prime order, (c) Morton order, (d) Peano-Hilbert.
also need to know the size of each grid cell. One mapping simply concatenates the result of interleaving the binary representations of the coordinate values of the lower-left corner (e.g., ( $a, b$ ) in two dimensions) and $i$ of each grid cell of size $2^{i}$ so that $i$ is at the right. The resulting number is termed a locational code and is a variant of the Morton ordering (Figure 16.4c). Assuming such a mapping and sorting the locational codes in increasing order yields an ordering equivalent to that which would be obtained by traversing the leaf nodes (i.e., grid cells) of the tree representation (e.g., Figure 16.8b) in the order SW, SE, NW, NE. The Morton ordering (as well as the Peano-Hilbert ordering shown in Figure 16.4d) is particularly attractive for quadtree-like decompositions because all pixels within a grid cell appear in consecutive positions in the ordering. Alternatively, these two orders exhaust a grid cell before exiting it.

For example, Figure 16.3 c shows the result of imposing a $\mathrm{B}^{+}$-tree [18] access structure on the leaf grid cells of the PR quadtree given in Figure 16.3b. Each node of the $\mathrm{B}^{+}$-tree in our example has a minimum of 2 and a maximum of 3 entries. Figure 16.3c does not contain the values resulting from applying the mapping to the individual grid cells nor does it show the discriminator values that are stored in the nonleaf nodes of the $\mathrm{B}^{+}$-tree. The leaf grid cells of the PR quadtree in Figure 16.3a are marked with the label of the leaf node of the $\mathrm{B}^{+}$-tree of which they are a member (e.g., the grid cell containing Chicago is in leaf node Q of the $\mathrm{B}^{+}$-tree).

It is important to observe that the above combination of the PR quadtree and the $\mathrm{B}^{+}$-tree has the property that the tree structure of the partition process of the underlying space has been decoupled [61] from that of the node hierarchy (i.e., the grouping process of the nodes resulting from the partition process) that makes up the original tree directory. More precisely, the grouping process is based on proximity in the ordering of the locational codes
and on the minimum and maximum capacity of the nodes of the $\mathrm{B}^{+}$-tree. Unfortunately, the resulting structure has the property that the space that is spanned by a leaf node of the $\mathrm{B}^{+}$-tree (i.e., the grid cells spanned by it) has an arbitrary shape and, in fact, does not usually correspond to a $k$-dimensional hyper-rectangle. In particular, the space spanned by the leaf node may have the shape of a staircase (e.g., the leaf grid cells in Figure 16.3a that comprise leaf nodes S and T of the $\mathrm{B}^{+}$-tree in Figure 16.3c) or may not even be connected in the sense that it corresponds to regions that are not contiguous (e.g., the leaf grid cells in Figure 16.3a that comprise leaf node R of the $\mathrm{B}^{+}$-tree in Figure 16.3c). The PK-tree [73] is an alternative decoupling method which overcomes these drawbacks by basing the grouping process on $k$-instantiation which stipulates that each node of the grouping process contains a minimum of $k$ objects or grid cells. The result is that all of the grid cells of the grouping process are congruent at the cost that the result is not balanced although use of relatively large values of $k$ ensures that the resulting trees are relatively shallow. It can be shown that when the partition process has a fanout of $f$, then $k$-instantiation means that the number of objects in each node of the grouping process is bounded by $f \cdot(k-1)$. Note that $k$-instantiation is different from bucketing where we only have an upper bound on the number of objects in the node.

Fixed-grids, quadtrees, k-d trees, indexkd tree grid file, EXCELL, as well as other hierarchical representations are good for range searching queries such as finding all cities within 80 miles of St. Louis. In particular, they act as pruning devices on the amount of search that will be performed as many points will not be examined since their containing cells lie outside the query range. These representations are generally very easy to implement and have good expected execution times, although they are quite difficult to analyze from a mathematical standpoint. However, their worst cases, despite being rare, can be quite bad. These worst cases can be avoided by making use of variants of range trees [11] and priority search trees [51]. For more details about these data structures, see Chapter 18.

### 16.3 Bucketing Methods

There are four principal approaches to decomposing the space from which the objects are drawn. The first approach makes use of an object hierarchy and the space decomposition is obtained in an indirect manner as the method propagates the space occupied by the objects up the hierarchy with the identity of the propagated objects being implicit to the hierarchy. In particular, associated with each object is a an object description (e.g., for region data, it is the set of locations in space corresponding to the cells that make up the object). Actually, since this information may be rather voluminous, it is often the case that an approximation of the space occupied by the object is propagated up the hierarchy rather than the collection of individual cells that are spanned by the object. For spatial data, the approximation is usually the minimum bounding rectangle for the object, while for nonspatial data it is simply the hyperrectangle whose sides have lengths equal to the ranges of the values of the attributes. Therefore, associated with each element in the hierarchy is a bounding rectangle corresponding to the union of the bounding rectangles associated with the elements immediately below it.

The R-tree (e.g., $[7,31]$ ) is an example of an object hierarchy which finds use especially in database applications. The number of objects or bounding rectangles that are aggregated in each node is permitted to range between $m \leq\lceil M / 2\rceil$ and $M$. The root node in an R-tree has at least two entries unless it is a leaf node in which case it has just one entry corresponding to the bounding rectangle of an object. The R-tree is usually built as the objects are encountered rather than waiting until all objects have been input. The hierarchy


FIGURE 16.5: Example collection of line segments embedded in a $4 \times 4$ grid.

(a)

(b)

FIGURE 16.6: (a) R-tree for the collection of line segments with $m=2$ and $\mathrm{M}=3$, in Figure 16.5, and (b) the spatial extents of the bounding rectangles. Notice that the leaf nodes in the index also store bounding rectangles although this is only shown for the nonleaf nodes.
is implemented as a tree structure with grouping being based, in part, on proximity of the objects or bounding rectangles.

For example, consider the collection of line segment objects given in Figure 16.5 shown embedded in a $4 \times 4$ grid. Figure 16.6a is an example R-tree for this collection with $m=2$ and $M=3$. Figure 16.6 b shows the spatial extent of the bounding rectangles of the nodes in Figure 16.6a, with heavy lines denoting the bounding rectangles corresponding to the leaf nodes, and broken lines denoting the bounding rectangles corresponding to the subtrees rooted at the nonleaf nodes. Note that the R-tree is not unique. Its structure depends heavily on the order in which the individual objects were inserted into (and possibly deleted from) the tree.

Given that each R-tree node can contain a varying number of objects or bounding rect-
angles, it is not surprising that the R-tree was inspired by the B-tree [6]. Therefore, nodes are viewed as analogous to disk pages. Thus the parameters defining the tree (i.e., $m$ and $M$ ) are chosen so that a small number of nodes is visited during a spatial query (i.e., point and range queries), which means that $m$ and $M$ are usually quite large. The actual implementation of the R -tree is really a $\mathrm{B}^{+}$-tree [18] as the objects are restricted to the leaf nodes.

The efficiency of the R-tree for search operations depends on its ability to distinguish between occupied space and unoccupied space (i.e., coverage), and to prevent a node from being examined needlessly due to a false overlap with other nodes. In other words, we want to minimize coverage and overlap. These goals guide the initial R-tree creation process as well, subject to the previously mentioned constraint that the R-tree is usually built as the objects are encountered rather than waiting until all objects have been input.

The drawback of the R-tree (and any representation based on an object hierarchy) is that it does not result in a disjoint decomposition of space. The problem is that an object is only associated with one bounding rectangle (e.g., line segment i in Figure 16.6 is associated with bounding rectangle R5, yet it passes through R1, R2, R4, and R5, as well as through R0 as do all the line segments). In the worst case, this means that when we wish to determine which object (e.g., an intersecting line in a collection of line segment objects, or a containing rectangle in a collection of rectangle objects) is associated with a particular point in the two-dimensional space from which the objects are drawn, we may have to search the entire collection. For example, in Figure 16.6, when searching for the line segment that passes through point Q, we need to examine bounding rectangles R0, R1, R4, R2, and R5, rather than just R0, R2, and R5.

This drawback can be overcome by using one of three other approaches which are based on a decomposition of space into disjoint cells. Their common property is that the objects are decomposed into disjoint subobjects such that each of the subobjects is associated with a different cell. They differ in the degree of regularity imposed by their underlying decomposition rules, and by the way in which the cells are aggregated into buckets.

The price paid for the disjointness is that in order to determine the area covered by a particular object, we have to retrieve all the cells that it occupies. This price is also paid when we want to delete an object. Fortunately, deletion is not so common in such applications. A related costly consequence of disjointness is that when we wish to determine all the objects that occur in a particular region, we often need to retrieve some of the objects more than once $[1,2,19]$. This is particularly troublesome when the result of the operation serves as input to another operation via composition of functions. For example, suppose we wish to compute the perimeter of all the objects in a given region. Clearly, each object's perimeter should only be computed once. Eliminating the duplicates is a serious issue (see [1] for a discussion of how to deal with this problem for a collection of line segment objects, and [2] for a collection of rectangle objects).

The first method based on disjointness partitions the embedding space into disjoint subspaces, and hence the individual objects into subobjects, so that each subspace consists of disjoint subobjects. The subspaces are then aggregated and grouped in another structure, such as a B-tree, so that all subsequent groupings are disjoint at each level of the structure. The result is termed a k -d-B-tree [59]. The $\mathrm{R}^{+}$-tree $[67,70]$ is a modification of the k -d-B-tree where at each level we replace the subspace by the minimum bounding rectangle of the subobjects or subtrees that it contains. The cell tree [30] is based on the same principle as the $\mathrm{R}^{+}$-tree except that the collections of objects are bounded by minimum convex polyhedra instead of minimum bounding rectangles.

The $\mathrm{R}^{+}$-tree (as well as the other related representations) is motivated by a desire to avoid overlap among the bounding rectangles. Each object is associated with all the bounding

(a)

(b)

FIGURE 16.7: (a) $\mathrm{R}^{+}$-tree for the collection of line segments in Figure 16.5 with $\mathrm{m}=2$ and $\mathrm{M}=3$, and (b) the spatial extents of the bounding rectangles. Notice that the leaf nodes in the index also store bounding rectangles although this is only shown for the nonleaf nodes.
rectangles that it intersects. All bounding rectangles in the tree (with the exception of the bounding rectangles for the objects at the leaf nodes) are non-overlapping*. The result is that there may be several paths starting at the root to the same object. This may lead to an increase in the height of the tree. However, retrieval time is sped up.

Figure 16.7 is an example of one possible $\mathrm{R}^{+}$-tree for the collection of line segments in Figure 16.5. This particular tree is of order $(2,3)$ although in general it is not possible to guarantee that all nodes will always have a minimum of 2 entries. In particular, the expected B-tree performance guarantees are not valid (i.e., pages are not guaranteed to be $m / M$ full) unless we are willing to perform very complicated record insertion and deletion procedures. Notice that line segment objects c, h, and i appear in two different nodes. Of course, other variants are possible since the $\mathrm{R}^{+}$-tree is not unique.

The problem with representations such as the k-d-B-tree and the $\mathrm{R}^{+}$-tree is that overflow in a leaf node may cause overflow of nodes at shallower depths in the tree whose subsequent partitioning may cause repartitioning at deeper levels in the tree. There are several ways of overcoming the repartitioning problem. One approach is to use the LSD-tree [32] at the cost of poorer storage utilization. An alternative approach is to use representations such as the hB-tree [49] and the BANG file [27] which remove the requirement that each block be a hyper-rectangle at the cost of multiple postings. This has a similar effect as that obtained when decomposing an object into several subobjects in order to overcome the nondisjoint decomposition problem when using an object hierarchy. The multiple posting problem is overcome by the BV-tree [28] which decouples the partitioning and grouping processes at the cost that the resulting tree is no longer balanced although as in the PK-tree [73] (which we point out in Section 16.2 is also based on decoupling), use of relatively large fanout

[^1]values ensure that the resulting trees are relatively shallow.
Methods such as the $\mathrm{R}^{+}$-tree (as well as the R -tree) also have the drawback that the decomposition is data-dependent. This means that it is difficult to perform tasks that require composition of different operations and data sets (e.g., set-theoretic operations such as overlay). The problem is that although these methods are good are distinguishing between occupied and unoccupied space in the underlying space (termed image in much of the subsequent discussion) under consideration, they re unable to correlate occupied space in two distinct images, and likewise for unoccupied space in the two images.

In contrast, the remaining two approaches to the decomposition of space into disjoint cells have a greater degree of data-independence. They are based on a regular decomposition. The space can be decomposed either into blocks of uniform size (e.g., the uniform grid [24]) or adapt the decomposition to the distribution of the data (e.g., a quadtree-based approach such as [66]). In the former case, all the blocks are congruent (e.g., the $4 \times 4$ grid in Figure 16.5). In the latter case, the widths of the blocks are restricted to be powers of two and their positions are also restricted. Since the positions of the subdivision lines are restricted, and essentially the same for all images of the same size, it is easy to correlate occupied and unoccupied space in different images.

The uniform grid is ideal for uniformly-distributed data, while quadtree-based approaches are suited for arbitrarily-distributed data. In the case of uniformly-distributed data, quadtreebased approaches degenerate to a uniform grid, albeit they have a higher overhead. Both the uniform grid and the quadtree-based approaches lend themselves to set-theoretic operations and thus they are ideal for tasks which require the composition of different operations and data sets. In general, since spatial data is not usually uniformly distributed, the quadtreebased regular decomposition approach is more flexible. The drawback of quadtree-like methods is their sensitivity to positioning in the sense that the placement of the objects relative to the decomposition lines of the space in which they are embedded effects their storage costs and the amount of decomposition that takes place. This is overcome to a large extent by using a bucketing adaptation that decomposes a block only if it contains more than $b$ objects.

### 16.4 Region Data

There are many ways of representing region data. We can represent a region either by its boundary (termed a boundary-based representation) or by its interior (termed an interiorbased representation). In this section, we focus on representations of collections of regions by their interior. In some applications, regions are really objects that are composed of smaller primitive objects by use of geometric transformations and Boolean set operations. Constructive Solid Geometry (CSG) [58] is a term usually used to describe such representations. They are beyond the scope of this chapter. Instead, unless noted otherwise, our discussion is restricted to regions consisting of congruent cells of unit area (volume) with sides (faces) of unit size that are orthogonal to the coordinate axes.

Regions with arbitrary boundaries are usually represented by either using approximating bounding rectangles or more general boundary-based representations that are applicable to collections of line segments that do not necessarily form regions. In that case, we do not restrict the line segments to be perpendicular to the coordinate axes. Such representations are discussed in Section 16.6. It should be clear that although our presentation and examples in this section deal primarily with two-dimensional data, they are valid for regions of any dimensionality.

The region data is assumed to be uniform in the sense that all the cells that comprise
each region are of the same type. In other words, each region is homogeneous. Of course, an image may consist of several distinct regions. Perhaps the best definition of a region is as a set of four-connected cells (i.e., in two dimensions, the cells are adjacent along an edge rather than a vertex) each of which is of the same type. For example, we may have a crop map where the regions correspond to the four-connected cells on which the same crop is grown. Each region is represented by the collection of cells that comprise it. The set of collections of cells that make up all of the regions is often termed an image array because of the nature in which they are accessed when performing operations on them. In particular, the array serves as an access structure in determining the region associated with a location of a cell as well as all remaining cells that comprise the region.

When the region is represented by its interior, then often we can reduce the storage requirements by aggregating identically-valued cells into blocks. In the rest of this section we discuss different methods of aggregating the cells that comprise each region into blocks as well as the methods used to represent the collections of blocks that comprise each region in the image.

The collection of blocks is usually a result of a space decomposition process with a set of rules that guide it. There are many possible decompositions. When the decomposition is recursive, we have the situation that the decomposition occurs in stages and often, although not always, the results of the stages form a containment hierarchy. This means that a block $b$ obtained in stage $i$ is decomposed into a set of blocks $b_{j}$ that span the same space. Blocks $b_{j}$ are, in turn, decomposed in stage $i+1$ using the same decomposition rule. Some decomposition rules restrict the possible sizes and shapes of the blocks as well as their placement in space. Some examples include:

- congruent blocks at each stage
- similar blocks at all stages
- all sides of a block are of equal size
- all sides of each block are powers of two
- etc.

Other decomposition rules dispense with the requirement that the blocks be rectangular (i.e., there exist decompositions using other shapes such as triangles, etc.), while still others do not require that they be orthogonal, although, as stated before, we do make these assumptions here. In addition, the blocks may be disjoint or be allowed to overlap. Clearly, the choice is large. In the following, we briefly explore some of these decomposition processes. We restrict ourselves to disjoint decompositions, although this need not be the case (e.g., the field tree [23]).

The most general decomposition permits aggregation along all dimensions. In other words, the decomposition is arbitrary. The blocks need not be uniform or similar. The only requirement is that the blocks span the space of the environment. The drawback of arbitrary decompositions is that there is little structure associated with them. This means that it is difficult to answer queries such as determining the region associated with a given point, besides exhaustive search through the blocks. Thus we need an additional data structure known as an index or an access structure. A very simple decomposition rule that lends itself to such an index in the form of an array is one that partitions a $d$-dimensional space having coordinate axes $x_{i}$ into $d$-dimensional blocks by use of $h_{i}$ hyperplanes that are parallel to the hyperplane formed by $x_{i}=0(1 \leq i \leq d)$. The result is a collection of $\prod_{i=1}^{d}\left(h_{i}+1\right)$ blocks. These blocks form a grid of irregular-sized blocks rather than congruent blocks. There is no recursion involved in the decomposition process. We term the resulting decomposition an irregular grid as the partition lines are at arbitrary positions in contrast to a uniform
grid [24] where the partition lines are positioned so that all of the resulting grid cells are congruent.

Although the blocks in the irregular grid are not congruent, we can still impose an array access structure by adding $d$ access structures termed linear scales. The linear scales indicate the position of the partitioning hyperplanes that are parallel to the hyperplane formed by $x_{i}=0(1 \leq i \leq d)$. Thus given a location $l$ in space, say $(a, b)$ in two-dimensional space, the linear scales for the $x$ and $y$ coordinate values indicate the column and row, respectively, of the array access structure entry which corresponds to the block that contains $l$. The linear scales are usually represented as one-dimensional arrays although they can be implemented using tree access structures such as binary search trees, range trees, segment trees, etc.

Perhaps the most widely known decompositions into blocks are those referred to by the general terms quadtree and octree [60,63]. They are usually used to describe a class of representations for two and three-dimensional data (and higher as well), respectively, that are the result of a recursive decomposition of the environment (i.e., space) containing the regions into blocks (not necessarily rectangular) until the data in each block satisfies some condition (e.g., with respect to its size, the nature of the regions that comprise it, the number of regions in it, etc.). The positions and/or sizes of the blocks may be restricted or arbitrary. It is interesting to note that quadtrees and octrees may be used with both interior-based and boundary-based representations although only the former are discussed in this section.

There are many variants of quadtrees and octrees (see also Sections 16.2, 16.5, and 16.6), and they are used in numerous application areas including high energy physics, VLSI, finite element analysis, and many others. Below, we focus on region quadtrees [43] and to a lesser extent on region octrees [39,53] They are specific examples of interior-based representations for two and three-dimensional region data (variants for data of higher dimension also exist), respectively, that permit further aggregation of identically-valued cells.

Region quadtrees and region octrees are instances of a restricted-decomposition rule where the environment containing the regions is recursively decomposed into four or eight, respectively, rectangular congruent blocks until each block is either completely occupied by a region or is empty (such a decomposition process is termed regular). For example, Figure 16.8 a is the block decomposition for the region quadtree corresponding to three regions A, B, and C. Notice that in this case, all the blocks are square, have sides whose size is a power of 2, and are located at specific positions. In particular, assuming an origin at the lower-left corner of the image containing the regions, then the coordinate values of the lower-left corner of each block (e.g., $(a, b)$ in two dimensions) of size $2^{i} \times 2^{i}$ satisfy the property that $a \bmod 2^{i}=0$ and $b \bmod 2^{i}=0$.

The traditional, and most natural, access structure for a region quadtree corresponding to a $d$-dimensional image is a tree with a fanout of $2^{d}$ (e.g., Figure 16.8b). Each leaf node in the tree corresponds to a different block $b$ and contains the identity of the region associated with $b$. Each nonleaf node $f$ corresponds to a block whose volume is the union of the blocks corresponding to the $2^{d}$ sons of $f$. In this case, the tree is a containment hierarchy and closely parallels the decomposition in the sense that they are both recursive processes and the blocks corresponding to nodes at different depths of the tree are similar in shape. Of course, the region quadtree could also be represented by using a mapping from the domain of the blocks to a subset of the integers and then imposing a tree access structure such as a $\mathrm{B}^{+}$-tree on the result of the mapping as was described in Section 16.2 for point data stored in a PR quadtree.

As the dimensionality of the space (i.e., $d$ ) increases, each level of decomposition in the region quadtree results in many new blocks as the fanout value $2^{d}$ is high. In particular, it is too large for a practical implementation of the tree access structure. In this case, an


FIGURE 16.8: (a) Block decomposition and (b) its tree representation for the region quadtree corresponding to a collection of three regions $\mathrm{A}, \mathrm{B}$, and C .
access structure termed a bintree $[44,65,72]$ with a fanout value of 2 is used. The bintree is defined in a manner analogous to the region quadtree except that at each subdivision stage, the space is decomposed into two equal-sized parts. In two dimensions, at odd stages we partition along the $y$ axis and at even stages we partition along the $x$ axis. In general, in the case of $d$ dimensions, we cycle through the different axes every $d$ levels in the bintree.

The region quadtree, as well as the bintree, is a regular decomposition. This means that the blocks are congruent - that is, at each level of decomposition, all of the resulting blocks are of the same shape and size. We can also use decompositions where the sizes of the blocks are not restricted in the sense that the only restriction is that they be rectangular and be a result of a recursive decomposition process. In this case, the representations that we described must be modified so that the sizes of the individual blocks can be obtained. An example of such a structure is an adaptation of the point quadtree [22] to regions. Although the point quadtree was designed to represent points in a higher dimensional space, the blocks resulting from its use to decompose space do correspond to regions. The difference from the region quadtree is that in the point quadtree, the positions of the partitions are arbitrary, whereas they are a result of a partitioning process into $2^{d}$ congruent blocks (e.g., quartering in two dimensions) in the case of the region quadtree.

As in the case of the region quadtree, as the dimensionality $d$ of the space increases, each level of decomposition in the point quadtree results in many new blocks since the fanout value $2^{d}$ is high. In particular, it is too large for a practical implementation of the tree access structure. In this case, we can adapt the k-d tree [8], which has a fanout value of 2, to regions. As in the point quadtree, although the k -d tree was designed to represent points in a higher dimensional space, the blocks resulting from its use to decompose space do correspond to regions. Thus the relationship of the k -d tree to the point quadtree is the same as the relationship of the bintree to the region quadtree. In fact, the k-d tree is the precursor of the bintree and its adaptation to regions is defined in a similar manner in the sense that for $d$-dimensional data we cycle through the $d$ axes every $d$ levels in the k-d tree. The difference is that in the k-d tree, the positions of the partitions are arbitrary, whereas they are a result of a halving process in the case of the bintree.

The k-d tree can be further generalized so that the partitions take place on the various axes at an arbitrary order, and, in fact, the partitions need not be made on every coordinate axis. The k-d tree is a special case of the BSP tree (denoting Binary Space Partitioning) [29] where the partitioning hyperplanes are restricted to be parallel to the axes, whereas in the


FIGURE 16.9: (a) An arbitrary space decomposition and (b) its BSP tree. The arrows indicate the direction of the positive halfspaces.

BSP tree they have an arbitrary orientation. The BSP tree is a binary tree. In order to be able to assign regions to the left and right subtrees, we need to associate a direction with each subdivision line. In particular, the subdivision lines are treated as separators between two halfspaces ${ }^{\dagger}$. Let the subdivision line have the equation $a \cdot x+b \cdot y+c=0$. We say that the right subtree is the 'positive' side and contains all subdivision lines formed by separators that satisfy $a \cdot x+b \cdot y+c \geq 0$. Similarly, we say that the left subtree is 'negative' and contains all subdivision lines formed by separators that satisfy $a \cdot x+b \cdot y+c<0$. As an example, consider Figure 16.9a which is an arbitrary space decomposition whose BSP tree is given in Figure 16.9b. Notice the use of arrows to indicate the direction of the positive halfspaces. The BSP tree is used in computer graphics to facilitate viewing. It is discussed in greater detail in Chapter 20.

As mentioned before, the various hierarchical data structures that we described can also be used to represent regions in three dimensions and higher. As an example, we briefly describe the region octree which is the three-dimensional analog of the region quadtree. It is constructed in the following manner. We start with an image in the form of a cubical volume and recursively subdivide it into eight congruent disjoint cubes (called octants) until blocks are obtained of a uniform color or a predetermined level of decomposition is reached. Figure 16.10a is an example of a simple three-dimensional object whose region octree block decomposition is given in Figure 16.10b and whose tree representation is given in Figure 16.10c.

The aggregation of cells into blocks in region quadtrees and region octrees is motivated, in part, by a desire to save space. Some of the decompositions have quite a bit of structure thereby leading to inflexibility in choosing partition lines, etc. In fact, at times, maintaining the original image with an array access structure may be more effective from the standpoint of storage requirements. In the following, we point out some important implications of the use of these aggregations. In particular, we focus on the region quadtree and region octree. Similar results could also be obtained for the remaining block decompositions.

[^2]

FIGURE 16.10: (a) Example three-dimensional object; (b) its region octree block decomposition; and (c) its tree representation.

The aggregation of similarly-valued cells into blocks has an important effect on the execution time of the algorithms that make use of the region quadtree. In particular, most algorithms that operate on images represented by a region quadtree are implemented by a preorder traversal of the quadtree and, thus, their execution time is generally a linear function of the number of nodes in the quadtree. A key to the analysis of the execution time of quadtree algorithms is the Quadtree Complexity Theorem [39] which states that the number of nodes in a region quadtree representation for a simple polygon (i.e., with non-intersecting edges and without holes) is $O(p+q)$ for a $2^{q} \times 2^{q}$ image with perimeter $p$ measured in terms of the width of unit-sized cells (i.e., pixels). In all but the most pathological cases (e.g., a small square of unit width centered in a large image), the $q$ factor is negligible and thus the number of nodes is $O(p)$.

The Quadtree Complexity Theorem also holds for three-dimensional data [52] (i.e., represented by a region octree) where perimeter is replaced by surface area, as well as for objects of higher dimensions $d$ for which it is proportional to the size of the $(d-1)$-dimensional interfaces between these objects. The most important consequence of the Quadtree Complexity Theorem is that it means that most algorithms that execute on a region quadtree representation of an image, instead of one that simply imposes an array access structure on the original collection of cells, usually have an execution time that is proportional to the number of blocks in the image rather than the number of unit-sized cells. In its most general case, this means that the use of the region quadtree, with an appropriate access structure, in solving a problem in $d$-dimensional space will lead to a solution whose execution time is proportional to the $(d-1)$-dimensional space of the surface of the original $d$-dimensional image. On the other hand, use of the array access structure on the original collection of cells results in a solution whose execution time is proportional to the number of cells that comprise the image. Therefore, region quadtrees and region octrees act like dimension-reducing devices.

### 16.5 Rectangle Data

The rectangle data type lies somewhere between the point and region data types. It can also be viewed as a special case of the region data type in the sense that it is a region with only four sides. Rectangles are often used to approximate other objects in an image for which they serve as the minimum rectilinear enclosing object. For example, bounding rectangles are used in cartographic applications to approximate objects such as lakes, forests, hills,
etc. In such a case, the approximation gives an indication of the existence of an object. Of course, the exact boundaries of the object are also stored; but they are only accessed if greater precision is needed. For such applications, the number of elements in the collection is usually small, and most often the sizes of the rectangles are of the same order of magnitude as the space from which they are drawn.

Rectangles are also used in VLSI design rule checking as a model of chip components for the analysis of their proper placement. Again, the rectangles serve as minimum enclosing objects. In this application, the size of the collection is quite large (e.g., millions of components) and the sizes of the rectangles are several orders of magnitude smaller than the space from which they are drawn.

It should be clear that the actual representation that is used depends heavily on the problem environment. At times, the rectangle is treated as the Cartesian product of two onedimensional intervals with the horizontal intervals being treated in a different manner than the vertical intervals. In fact, the representation issue is often reduced to one of representing intervals. For example, this is the case in the use of the plane-sweep paradigm [57] in the solution of rectangle problems such as determining all pairs of intersecting rectangles. In this case, each interval is represented by its left and right endpoints. The solution makes use of two passes.

The first pass sorts the rectangles in ascending order on the basis of their left and right sides (i.e., $x$ coordinate values) and forms a list. The second pass sweeps a vertical scan line through the sorted list from left to right halting at each one of these points, say $p$. At any instant, all rectangles that intersect the scan line are considered active and are the only ones whose intersection needs to be checked with the rectangle associated with $p$. This means that each time the sweep line halts, a rectangle either becomes active (causing it to be inserted in the set of active rectangles) or ceases to be active (causing it to be deleted from the set of active rectangles). Thus the key to the algorithm is its ability to keep track of the active rectangles (actually just their vertical sides) as well as to perform the actual one-dimensional intersection test.

Data structures such as the segment tree [9], interval tree [20], and the priority search tree [51] can be used to organize the vertical sides of the active rectangles so that, for $N$ rectangles and $F$ intersecting pairs of rectangles, the problem can be solved in $O(N$. $\left.\log _{2} N+F\right)$ time. All three data structures enable intersection detection, insertion, and deletion to be executed in $O\left(\log _{2} N\right)$ time. The difference between them is that the segment tree requires $O\left(N \cdot \log _{2} N\right)$ space while the interval tree and the priority search tree only need $O(N)$ space. These algorithms require that the set of rectangles be known in advance. However, they work even when the size of the set of active rectangles exceeds the amount of available memory, in which case multiple passes are made over the data [41]. For more details about these data structures, see Chapter 18.

In this chapter, we are primarily interested in dynamic problems (i.e., the set of rectangles is constantly changing). The data structures that are chosen for the collection of the rectangles are differentiated by the way in which each rectangle is represented. One representation discussed in Section 16.1 reduces each rectangle to a point in a higher dimensional space, and then treats the problem as if we have a collection of points [33]. Again, each rectangle is a Cartesian product of two one-dimensional intervals where the difference from its use with the plane-sweep paradigm is that each interval is represented by its centroid and extent. Each set of intervals in a particular dimension is, in turn, represented by a grid file [55] which is described in Section 16.2.

The second representation is region-based in the sense that the subdivision of the space from which the rectangles are drawn depends on the physical extent of the rectangle not just one point. Representing the collection of rectangles, in turn, with a tree-like data


FIGURE 16.11: (a) Collection of rectangles and the block decomposition induced by the MX-CIF quadtree; (b) the tree representation of (a); the binary trees for the y axes passing through the root of the tree in (b), and (d) the NE son of the root of the tree in (b).
structure has the advantage that there is a relation between the depth of node in the tree and the size of the rectangle(s) that is (are) associated with it. Interestingly, some of the region-based solutions make use of the same data structures that are used in the solutions based on the plane-sweep paradigm.

There are three types of region-based solutions currently in use. The first two solutions use the R-tree and the $\mathrm{R}^{+}$-tree (discussed in Section 16.3) to store rectangle data (in this case the objects are rectangles instead of arbitrary objects). The third is a quadtree-based approach and uses the MX-CIF quadtree [42] (see also [47] for a related variant).

In the $M X$-CIF quadtree, each rectangle is associated with the quadtree node corresponding to the smallest block which contains it in its entirety. Subdivision ceases whenever a node's block contains no rectangles. Alternatively, subdivision can also cease once a quadtree block is smaller than a predetermined threshold size. This threshold is often chosen to be equal to the expected size of the rectangle [42]. For example, Figure 16.11b is the MX-CIF quadtree for a collection of rectangles given in Figure 16.11a. Rectangles can be associated with both leaf and nonleaf nodes.

It should be clear that more than one rectangle can be associated with a given enclosing block and, thus, often we find it useful to be able to differentiate between them. This is done in the following manner [42]. Let $P$ be a quadtree node with centroid ( $C X, C Y$ ), and let $S$ be the set of rectangles that are associated with $P$. Members of $S$ are organized into two sets according to their intersection (or collinearity of their sides) with the lines passing through the centroid of $P$ 's block - that is, all members of $S$ that intersect the line $x=C X$ form one set and all members of $S$ that intersect the line $y=C Y$ form the other set.

If a rectangle intersects both lines (i.e., it contains the centroid of $P$ 's block), then we adopt the convention that it is stored with the set associated with the line through $x=C X$. These subsets are implemented as binary trees (really tries), which in actuality are onedimensional analogs of the MX-CIF quadtree. For example, Figure 16.11c and Figure 16.11d illustrate the binary trees associated with the $y$ axes passing through the root and the NE son of the root, respectively, of the MX-CIF quadtree of Figure 16.11b. Interestingly, the MX-CIF quadtree is a two-dimensional analog of the interval tree. described above. More precisely, the MX-CIF quadtree is a a two-dimensional analog of the tile tree [50] which is a regular decomposition version of the interval tree. In fact, the tile tree and the onedimensional MX-CIF quadtree are identical when rectangles are not allowed to overlap.

### 16.6 Line Data and Boundaries of Regions

Section 16.4 was devoted to variations on hierarchical decompositions of regions into blocks, an approach to region representation that is based on a description of the region's interior. In this section, we focus on representations that enable the specification of the boundaries of regions, as well as curvilinear data and collections of line segments. The representations are usually based on a series of approximations which provide successively closer fits to the data, often with the aid of bounding rectangles. When the boundaries or line segments have a constant slope (i.e., linear and termed line segments in the rest of this discussion), then an exact representation is possible.

There are several ways of approximating a curvilinear line segment. The first is by digitizing it and then marking the unit-sized cells (i.e., pixels) through which it passes. The second is to approximate it by a set of straight line segments termed a polyline. Assuming a boundary consisting of straight lines (or polylines after the first stage of approximation), the simplest representation of the boundary of a region is the polygon. It consists of vectors which are usually specified in the form of lists of pairs of $x$ and $y$ coordinate values corresponding to their start and end points. The vectors are usually ordered according to their connectivity. One of the most common representations is the chain code [26] which is an approximation of a polygon's boundary by use of a sequence of unit vectors in the four (and sometimes eight) principal directions.

Chain codes, and other polygon representations, break down for data in three dimensions and higher. This is primarily due to the difficulty in ordering their boundaries by connectivity. The problem is that in two dimensions connectivity is determined by ordering the boundary elements $e_{i, j}$ of boundary $b_{i}$ of object $o$ so that the end vertex of the vector $v_{j}$ corresponding to $e_{i, j}$ is the start vertex of the vector $v_{j+1}$ corresponding to $e_{i, j+1}$. Unfortunately, such an implicit ordering does not exist in higher dimensions as the relationship between the boundary elements associated with a particular object are more complex.

Instead, we must make use of data structures which capture the topology of the object in terms of its faces, edges, and vertices. The winged-edge data structure is one such representation which serves as the basis of the boundary model (also known as BRep [5]). For more details about these data structures, see Chapter 17.

Polygon representations are very local. In particular, if we are at one position on the boundary, we don't know anything about the rest of the boundary without traversing it element-by-element. Thus, using such representations, given a random point in space, it is very difficult to find the nearest line to it as the lines are not sorted. This is in contrast to hierarchical representations which are global in nature. They are primarily based on rectangular approximations to the data as well as on a regular decomposition in two dimensions. In the rest of this section, we discuss a number of such representations.

In Section 16.3 we already examined two hierarchical representations (i.e., the R-tree and the $\mathrm{R}^{+}$-tree) that propagate object approximations in the form of bounding rectangles. In this case, the sides of the bounding rectangles had to be parallel to the coordinate axes of the space from which the objects are drawn. In contrast, the strip tree [4] is a hierarchical representation of a single curve that successively approximates segments of it with bounding rectangles that does not require that the sides be parallel to the coordinate axes. The only requirement is that the curve be continuous; it need not be differentiable.

The strip tree data structure consists of a binary tree whose root represents the bounding rectangle of the entire curve. The rectangle associated with the root corresponds to a rectangular strip, that encloses the curve, whose sides are parallel to the line joining the endpoints of the curve. The curve is then partitioned in two at one of the locations where it touches the bounding rectangle (these are not tangent points as the curve only needs to be


FIGURE 16.12: (a) MX quadtree and (b) edge quadtree for the collection of line segments of Figure 16.5.
continuous; it need not be differentiable). Each subcurve is then surrounded by a bounding rectangle and the partitioning process is applied recursively. This process stops when the width of each strip is less than a predetermined value.

In order to be able to cope with more complex curves such as those that arise in the case of object boundaries, the notion of a strip tree must be extended. In particular, closed curves and curves that extend past their endpoints require some special treatment. The general idea is that these curves are enclosed by rectangles which are split into two rectangular strips, and from now on the strip tree is used as before.

The strip tree is similar to the point quadtree in the sense that the points at which the curve is decomposed depend on the data. In contrast, a representation based on the region quadtree has fixed decomposition points. Similarly, strip tree methods approximate curvilinear data with rectangles of arbitrary orientation, while methods based on the region quadtree achieve analogous results by use of a collection of disjoint squares having sides of length power of two. In the following we discuss a number of adaptations of the region quadtree for representing curvilinear data.

The simplest adaptation of the region quadtree is the MX quadtree [39, 40]. It is built by digitizing the line segments and labeling each unit-sized cell (i.e., pixel) through which it passes as of type boundary. The remaining pixels are marked WHITE and are merged, if possible, into larger and larger quadtree blocks. Figure 16.12a is the MX quadtree for the collection of line segment objects in Figure 16.5. A drawback of the MX quadtree is that it associates a thickness with a line. Also, it is difficult to detect the presence of a vertex whenever five or more line segments meet.

The edge quadtree $[68,74]$ is a refinement of the MX quadtree based on the observation that the number of squares in the decomposition can be reduced by terminating the subdivision whenever the square contains a single curve that can be approximated by a single straight line. For example, Figure 16.12b is the edge quadtree for the collection of line segment objects in Figure 16.5. Applying this process leads to quadtrees in which long edges are represented by large blocks or a sequence of large blocks. However, small blocks are required in the vicinity of corners or intersecting edges. Of course, many blocks will contain no edge information at all.

The PM quadtree family $[54,66]$ (see also edge-EXCELL [71]) represents an attempt


FIGURE 16.13: (a) $\mathrm{PM}_{1}$ quadtree and (b) PMR quadtree for the collection of line segments of Figure 16.5.
to overcome some of the problems associated with the edge quadtree in the representation of collections of polygons (termed polygonal maps). In particular, the edge quadtree is an approximation because vertices are represented by pixels. There are a number of variants of the PM quadtree. These variants are either vertex-based or edge-based. They are all built by applying the principle of repeatedly breaking up the collection of vertices and edges (forming the polygonal map) until obtaining a subset that is sufficiently simple so that it can be organized by some other data structure.

The $\mathrm{PM}_{1}$ quadtree [66] is an example of a vertex-based PM quadtree. Its decomposition rule stipulates that partitioning occurs as long as a block contains more than one line segment unless the line segments are all incident at the same vertex which is also in the same block (e.g., Figure 16.13a). Given a polygonal map whose vertices are drawn from a grid (say $2^{m} \times 2^{m}$ ), and where edges are not permitted to intersect at points other than the grid points (i.e., vertices), it can be shown that the maximum depth of any leaf node in the $\mathrm{PM}_{1}$ quadtree is bounded from above by $4 m+1$ [64]. This enables a determination of the maximum amount of storage that will be necessary for each node.

A similar representation has been devised for three-dimensional images (e.g., [3] and the references cited in [63]). The decomposition criteria are such that no node contains more than one face, edge, or vertex unless the faces all meet at the same vertex or are adjacent to the same edge. This representation is quite useful since its space requirements for polyhedral objects are significantly smaller than those of a region octree.

The PMR quadtree [54] is an edge-based variant of the PM quadtree. It makes use of a probabilistic splitting rule. A node is permitted to contain a variable number of line segments. A line segment is stored in a PMR quadtree by inserting it into the nodes corresponding to all the blocks that it intersects. During this process, the occupancy of each node that is intersected by the line segment is checked to see if the insertion causes it to exceed a predetermined splitting threshold. If the splitting threshold is exceeded, then the node's block is split once, and only once, into four equal quadrants.

For example, Figure 16.13b is the PMR quadtree for the collection of line segment objects in Figure 16.5 with a splitting threshold value of 2 . The line segments are inserted in alphabetic order (i.e., a-i). It should be clear that the shape of the PMR quadtree depends on the order in which the line segments are inserted. Note the difference from the $\mathrm{PM}_{1}$
quadtree in Figure 16.13a - that is, the NE block of the SW quadrant is decomposed in the $\mathrm{PM}_{1}$ quadtree while the SE block of the SW quadrant is not decomposed in the $\mathrm{PM}_{1}$ quadtree.

On the other hand, a line segment is deleted from a PMR quadtree by removing it from the nodes corresponding to all the blocks that it intersects. During this process, the occupancy of the node and its siblings is checked to see if the deletion causes the total number of line segments in them to be less than the predetermined splitting threshold. If the splitting threshold exceeds the occupancy of the node and its siblings, then they are merged and the merging process is reapplied to the resulting node and its siblings. Notice the asymmetry between the splitting and merging rules.

The PMR quadtree is very good for answering queries such as finding the nearest line to a given point [34-37] (see [38] for an empirical comparison with hierarchical object representations such as the R -tree and $\mathrm{R}^{+}$-tree). It is preferred over the $\mathrm{PM}_{1}$ quadtree (as well as the MX and edge quadtrees) as it results in far fewer subdivisions. In particular, in the PMR quadtree there is no need to subdivide in order to separate line segments that are very "close" or whose vertices are very "close," which is the case for the $\mathrm{PM}_{1}$ quadtree. This is important since four blocks are created at each subdivision step. Thus when many subdivision steps that occur in the $\mathrm{PM}_{1}$ quadtree result in creating many empty blocks, the storage requirements of the $\mathrm{PM}_{1}$ quadtree will be considerably higher than those of the PMR quadtree. Generally, as the splitting threshold is increased, the storage requirements of the PMR quadtree decrease while the time necessary to perform operations on it will increase.

Using a random image model and geometric probability, it has been shown [48], theoretically and empirically using both random and real map data, that for sufficiently high values of the splitting threshold (i.e., $\geq 4$ ), the number of nodes in a PMR quadtree is asymptotically proportional to the number of line segments and is independent of the maximum depth of the tree. In contrast, using the same model, the number of nodes in the $\mathrm{PM}_{1}$ quadtree is a product of the number of lines and the maximal depth of the tree (i.e., $n$ for a $2^{n} \times 2^{n}$ image). The same experiments and analysis for the MX quadtree confirmed the results predicted by the Quadtree Complexity Theorem (see Section 16.4) which is that the number of nodes is proportional to the total length of the line segments.

Observe that although a bucket in the PMR quadtree can contain more line segments than the splitting threshold, this is not a problem. In fact, it can be shown [63] that the maximum number of line segments in a bucket is bounded by the sum of the splitting threshold and the depth of the block (i.e., the number of times the original space has been decomposed to yield this block).

### 16.7 Research Issues and Summary

A review has been presented of a number of representations of multidimensional data. Our focus has been on multidimensional spatial data with extent rather than just multidimensional point data. There has been a particular emphasis on hierarchical representations. Such representations are based on the "divide-and-conquer" problem-solving paradigm. They are of interest because they enable focusing computational resources on the interesting subsets of data. Thus, there is no need to expend work where the payoff is small. Although many of the operations for which they are used can often be performed equally as efficiently, or more so, with other data structures, hierarchical data structures are attractive because of their conceptual clarity and ease of implementation.

When the hierarchical data structures are based on the principle of regular decomposition,
we have the added benefit that different data sets (often of differing types) are in registration. This means that they are partitioned in known positions which are often the same or subsets of one another for the different data sets. This is true for all the features including regions, points, rectangles, lines, volumes, etc. The result is that a query such as "finding all cities with more than 20,000 inhabitants in wheat growing regions within 30 miles of the Mississippi River" can be executed by simply overlaying the region (crops), point (i.e., cities), and river maps even though they represent data of different types. Alternatively, we may extract regions such as those within 30 miles of the Mississippi River. Such operations find use in applications involving spatial data such as geographic information systems.

Current research in multidimensional representations is highly application-dependent in the sense that the work is driven by the application. Many of the recent developments have been motivated by the interaction with databases. The choice of a proper representation plays a key role in the speed with which responses are provided to queries. Knowledge of the underlying data distribution is also a factor and research is ongoing to make use of this information in the process of making a choice. Most of the initial applications in which the representation of multidimensional data has been important have involved spatial data of the kind described in this chapter. Such data is intrinsically of low dimensionality (i.e., two and three).

Future applications involve higher dimensional data for applications such as image databases where the data are often points in feature space. Unfortunately, for such applications, the performance of most indexing methods that rely on a decomposition of the underlying space is often unsatisfactory when compared with not using an index at all (e.g., [16]). The problem is that for uniformly-distributed data, most of the data is found to be at or near the boundary of the space in which it lies [13]. The result means that the query region usually overlaps all of the leaf node regions that are created by the decomposition process and thus a sequential scan is preferable. This has led to a number of alternative representations that try to speed up the scan (e.g., VA-file [75], VA ${ }^{+}$-file [21], IQ-tree [15], etc.). Nevertheless, representations such as the pyramid technique [14] are based on the principle that most of the data lies near the surface and therefore subdivide the data space as if it is an onion by peeling off hypervolumes that are close to its boundary. This is achieved by first dividing the hypercube corresponding to the $d$-dimensional data space into $2 d$ pyramids having the center of the data space as their top point and one of the faces of the hypercube as its base. These pyramids are subsequently cut into slices that are parallel to their base. Of course, the high-dimensional data is not necessarily uniformly-distributed which has led to other data structures with good performance (e.g., the hybrid tree [17]). Clearly, more work needs to be done in this area.

## Acknowledgment

This work was supported, in part, by the National Science Foundation under Grants EIA-99-00268, IIS-00-86162, and EIA-00-91474 is gratefully acknowledged.

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[^0]:    *All figures (c)2003 by Hanan Samet.

[^1]:    *From a theoretical viewpoint, the bounding rectangles for the objects at the leaf nodes should also be disjoint However, this may be impossible (e.g., when the objects are line segments and if many of the line segments intersect at a point).

[^2]:    ${ }^{\dagger}$ A (linear) halfspace in $d$-dimensional space is defined by the inequality $\sum_{i=0}^{d} a_{i} \cdot x_{i} \geq 0$ on the $d+1$ homogeneous coordinates $\left(x_{0}=1\right)$. The halfspace is represented by a column vector $a$. In vector notation, the inequality is written as $a \cdot x \geq 0$. In the case of equality, it defines a hyperplane with $a$ as its normal. It is important to note that halfspaces are volume elements; they are not boundary elements.

