FUNDAMENTAL TECHNIQUES FOR GEOMETRIC AND SOLID MODELING

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I. INTRODUCTION

There are three well-established paradigms for representing solids that are based on the boundary, on spatial subdivision, and on construction from primitives using regularized set operations. Around these paradigms, a substantial literature has grown and many efficient and ingenious algorithms have been devised for working with solids so represented. Yet despite this extensive work, many tasks of interest remain that appear to be cumbersome to implement based on these traditional solid and surface representations. For instance, given a solid, how can we derive a new solid that is the offset of the old one, having

1Supported in part by ONR Contract N00014-90-J-1599, NSF Grant CCR 86-19817, and NSF Grant ECD 86-03017.
2Supported by NSF Grant CCR 86-19817.
a surface that is at constant distance from the old one? To devise a complete algorithm for this task is not simple. In fact, the mathematical difficulty of offsetting a general curved surface is in marked contrast to the simplicity with which this task can be defined and communicated between people. Difficulties of this kind suggest that we remain on the lookout for new solid and surface representations that might facilitate such operations.

In this chapter, we present a number of new paradigms for representing solids and surfaces. These new approaches show potential for pressing practical problems, and, in some cases, have already delivered. Yet one should not conclude that they will therefore displace traditional representation paradigms. Indeed, a representation is intimately linked with algorithmic efficiency and convenience, and so one should expect that there will always be a need to switch to a different representation in response to the algorithmic problem at hand. The new representations we discuss here are intended to supplement the repertoire of geometric and solid modeling, not to supplant it.

In accordance with our outlook that many representations will continue to coexist, we begin by reviewing the three classical representation paradigms and the algorithmic ideas underlying their interrogation. So, we review **spatial subdivision, boundary representation, and constructive solid geometry**. We then discuss in some detail the **B-rep index**, a spatial subdivision structure that integrates boundary-based and spatial-subdivision representations. The B-rep index has proved to be extremely valuable in applications in which a large number of moving objects are queried for collision, and where, upon collision, a geometric analysis must be made of the locale at which a contact has been determined. The favorable performance of the B-rep index in this situation is based on the ease with which lines and points can be classified, with respect to a solid. But the B-rep index also does well in generating rectangular meshes of solid domains, and is also amenable to algorithmic enhancements that significantly increase the robustness of geometric computations.

Technically, constructing the B-rep index from a boundary-based representation (B-rep) is akin to the problem of converting a B-rep to constructive solid geometry. This problem is fully solved in the polyhedral case, but only partially solved in the curved-surface case.

Just as the performance of a solid representation depends on the operation one has in mind, surface representations exhibit a similar relativity. The two major paradigms are the parametric and the implicit surface representations. Again, each has specific strengths and weaknesses, and certain rather intuitive and desirable operations, including offsetting, are difficult to carry out exactly. Here, the **dimensionality paradigm** offers some attractive alternatives. We review the dimensionality paradigm in the section on constrained-surface representations, giving a detailed example illustrating the method, and discussing how some of the standard interrogation algorithms on surfaces so represented can be implemented.
A major strong point of the dimensionality paradigm is its ability to represent exactly the bisectors of curved surfaces. That is, given two surfaces, we can represent exactly those points in three space that have equal distance from the given surfaces. This allows us to represent the skeleton precisely, another alternate solid representation scheme that recently has generated much interest.

Originally proposed as a shape representation in computer vision, the skeleton has been used successfully in a number of algorithms for generating finite-element meshes completely automatically. Furthermore, the skeleton is intimately related to a classical geometric concept, the cyclographic map, which can be thought of as an explicit map of the Euclidean distance of the points in space from a given geometric shape. More than that, the skeleton is the locus of the shocks in the Hamilton-Jacobi equation, and thus can be computed in principle with standard PDE solvers.

We discuss the skeleton in the final section of this chapter, explaining how it relates to cyclographic maps and to the Hamilton-Jacobi equation, and discuss several algorithms for computing the skeleton. We then review two algorithms for finite-element mesh generation, and remark on potential applications in geometric tolerancing.

II. TRADITIONAL PARADIGMS FOR REPRESENTING SOLIDS

Point set topology provides a precise language for describing the basic properties of solids. Let $\mathbb{E}^3$ denote three-dimensional Euclidean space with the usual topology, and let $A$ be a subset of $\mathbb{E}^3$. The interior of $A$ is denoted by $iA$, the boundary of $A$ by $bA$, and the complement of $A$ by $cA$. Note that the interior, the boundary and the complement of $A$ partition $\mathbb{E}^3$. The closure of $A$, $kA$, is obtained by adding to $A$ the limits of convergent point sequences in $iA$. The point set $A$ is regular if $A = k\overline{A}$. If $A$ is not regular, then we regularize it by forming $rA = k\overline{A}$. A set is bounded if it is contained in an open ball. An $r$-set is a regular and bounded set.

Geometrically, the boundaries of $r$-sets may be extremely complex [2]. In solid modeling, only surfaces that are simple in a technical sense are of interest [3]. Roughly speaking, the boundary must be finitely describable, and must be of finite variation; i.e., any line segment is contained or intersects the boundary in finitely many points. Then solids are $r$-sets that are simple in this sense.

A solid can be represented explicitly by its boundary, or by its volume, or implicitly by specifying operations on volumetric primitives that construct it. Accordingly, there are three dominant schemas: boundary representations, cell-decomposition representations, and constructive solid geometry; e.g., [2,4].
A. BOUNDARY REPRESENTATIONS

The boundary of a solid consists of vertices, edges and faces. For each of these entities, the geometric part of the representation fixes the shape and/or location in space, and the topological part records the adjacencies. The combination of the topological and geometrical information is a boundary representation (B-rep) of a solid.

In all, there are nine adjacency relationships between the three types of topological entities. For example, the face-edge adjacency relationship specifies for each face the adjacent edges. A complete solid modeling representation must allow the retrieval of any topological entity and any of the nine adjacency relationships. However, explicitly maintaining all nine adjacency relationships is redundant. Weller has shown that three of the ordered adjacency relationships are sufficient to obtain all others [5]. There is a space/time tradeoff: explicitly maintaining all adjacency relationships requires more space but little time retrieving them. On the other hand, maintaining only sufficient adjacency relationship requires little space but more time to derive the others. A comparison of the space/time tradeoffs of different representation schemas, each consisting of a different subset of the topological adjacency relationships, can be found in [6].

Many different kinds of boundary representations are used for representing solids with a manifold surface. An early representation schema is Baumgart's winged-edge data structure for manifold solids [7]. In the winged-edge data
structure, an edge node records the information about the edge orientation, the face adjacencies, and the clockwise and counterclockwise successor and predecessor edges about the adjacent faces. An example representing the top face of a box is shown in Figure 1. The representation assumes that the faces are simply connected. Multiply connected faces need to be partitioned in this representation.

Braid modified the winged-edge data structure to include multiply connected faces by introducing a fourth topological entity called a loop. In Braid's data structure, each face consists of one or more edge loops, each bounding a hole in the face [8].

Yamaguchi and Tokieda modified the winged-edge data structure differently by introducing bridge edges to allow multiply connected faces [9]. A bridge edge is a double edge that connects two edge cycles of a given face. In their representation, all faces are triangulated using bridge edges. Other manifold representations are Mäntylä's half-edge data structure [4], Guibas and Stolfi's quad-edge representation [10], Hanrahan's face-edge representation [11], and Ansaldi, De Floriani and Falcidieno's hierarchical face adjacency hypergraph [12]. There are others.

The result of set operations can be a solid with a nonmanifold boundary. Manifold representations such as the winged-edge data structure [7,13] can handle nonmanifold solids only in special cases, and this complicates the algorithms for set operations unnecessarily [14]. In response, nonmanifold representations have been investigated; e.g., by Kevin Weiler who introduced the radial-edge data structure [15]. The radial-edge data structure can accommodate general nonmanifold models. More generally, the data structure can model arbitrary solids and structures that are not regularized. For example, a single object
may contain wire-frames, isolated vertices and edges, and arbitrary regions that need not be bounded. Focusing on regularized set operations, others have developed variations of the half-edge data structure to allow nonmanifolds. Two similar such data structures are Karasick's star-edge data structure [18], and Vaněček's edge-based data structure [17,18]. Karasick's representation contains face loop and shell information explicitly. Vaněček's representation uses bridge edges instead, but without triangulating faces. An example of Vaněček's representation is shown in Figure 2.

B. CELL-DECOMPOSITION REPRESENTATIONS

Instead of representing the boundary of an object, the object can be represented explicitly by its volume. The volume is represented as a collection of cells of a partition of space. The various data structures differ in how they organize the cells and what information about the object each cell contains. Some widely used data structures are hierarchical and subdivide space recursively.

The simplest hierarchical data structure is the region octree based on regular decomposition. The method partitions a cuboidal region into eight equal sized octants. The region is represented by a node in the region octree, and the eight octants are its eight children. Each node in the tree is labeled either as a gray node, if it is further decomposed, a white node if it is completely outside the object, or a black node if it is completely inside the object. A simple example is shown in Figure 3.

Region octrees are suitable only for solids with faces that are parallel to the principal axes. Solids with inclined faces are approximated by region octrees. Thus they give only a rough description of the boundary of the object. The accuracy of the approximation depends on the subdivision level. For a thorough
Figure 4: Examples of Face, Edge, and Vertex Nodes of Extended Octree.

discussion of octrees, hierarchical data structures, and related algorithms, see Samet [19].

Region octrees have been generalized to take the object boundary into account, economizing the space requirements of the tree. Samet and Webber developed the PM-octree which added nodes of type vertex, edge and face to the already present black, white and gray nodes. For regions that contain portions of the boundary, the recursive decomposition terminates when the region contains exactly one vertex, one edge or one face. For example, a region with a vertex and its adjacent edges and faces is represented directly and is not further decomposed. Variations of PM-octrees are the polytree by Carlsson, Chakravarty and Vanderschel [20], and the extended octree of Navazo, Ayala and Brunet [21].

The two tree structures differ in minor ways. Extended octrees assume only manifolds, while polytrees allow nonmanifolds. Extended octrees maintain for each leaf node a list of the oriented support planes of the faces intersecting the region, along with information on how to construct the faces. In contrast, in polytrees each leaf node contains a list of polygons representing the parts of the faces that intersect that region. No adjacency information is kept between the edges of two adjacent polygons in one region or between adjacent regions. Thus all regions have independent information.

Both the extended octree and the polytree are allowed to grow only to a prespecified depth. For certain solids, larger trees would be required, for instance, for a solid two vertices that are very close together. A region might then require further decomposition, but is not decomposed. Instead, it is represented by a nasty node that stores a list of pointers to all the planes (or polygons) it contains. The size of region octrees, extended octrees and polytrees is proportional to the solid's surface area and resolution [22,21]. Furthermore, the tree structure is sensitive to rotations other than by 90°.

A generic drawback of the octree and its variants is that the topological structure of the boundary is not readily available. A single face of a solid might
Figure 6: CSG Representation and Corresponding Polyhedron

...
1. SET OPERATIONS IN CSG AND CONVERSION TO B-REP

The constructive representation of a solid is a tree structure in which the internal nodes represent either the regularized set operators or rigid transformations, and the leaf nodes represent primitive solids. Since this is an implicit representation, performing a boolean operation on two solids given by CSG trees is trivial: A new tree is formed by creating a new root labeled with the set operation and with the CSG trees of the two solids as children. However, boundary evaluation which generates the B-rep described by the given CSG, is not trivial.

In their work on CSG, Voelcker and Requicha showed that the regularized operations can be defined as the selective combination of certain sets, called the classification sets. The classification sets are computed by a membership classification function \( M[X, S] \) which partitions a candidate set \( X \) with respect to a reference set \( S \). The reference set \( S \) is regular in a topological space \( W \subset \mathbb{R}^3 \), while the candidate set \( X \) is regular in a subspace \( W' \subset W \), in the relative topology.

**Definition 1** If \( X \) is regular in \( W' \) and \( S \) is regular in \( W \), then the membership classification function, \( M[X, S] \), is defined as follows:

\[
M[X,S] = (X_{\text{in}}S, X_{\text{on}}S, X_{\text{out}}S),
\]

where

\[
X_{\text{in}}S = X \cap iS = \mathcal{H}'(X \cap iS),
\]

\[
X_{\text{on}}S = X \cap bS, \text{ and}
\]

\[
X_{\text{out}}S = X \cap cS.
\]

The classification sets \( X_{\text{in}}S, X_{\text{on}}S, \) and \( X_{\text{out}}S \) correspond to regular subsets of \( X \) that are respectively inside, on the boundary, or outside \( S \). Together they form the regular partition of \( X \):
Theorem 1 [Tilove] If $M[X, S] = (X_{inS}, X_{onS}, X_{outS})$ then

$$X = X_{inS} \cup^{r'} X_{onS} \cup^{r'} X_{outS},$$

and

$$X_{inS} \cap^{r'} X_{onS} = X_{onS} \cap^{r'} X_{outS} = X_{inS} \cap^{r'} X_{outS} = \emptyset.$$  

Through the use of the membership classification function, a classification result of a regularized composition of set operations can be expressed recursively in terms of the classification results. The following theorem states this result for the intersection operation. See Tilove's Master's thesis [31] for similar theorems for the operations of regularized union and set difference.

Theorem 2 [Tilove] Let

$$M[X, A] = (X_{inA}, X_{onA}, X_{outA})$$

$$M[X, B] = (X_{inB}, X_{onB}, X_{outB})$$

and let $S = A \cap^{r} B$. Then $M[X, S] = (X_{inS}, X_{onS}, X_{outS})$, where:

$$X_{inS} = X_{inA} \cap^{r'} X_{inB},$$

$$X_{onS} = (X_{inA} \cap^{r'} X_{onB}) \cap^{r'} (X_{onA} \cap^{r'} X_{inA})$$

$$\cap^{r'} \{ p \in X_{onA} \cap^{r'} X_{onB} | N(p; S) = \emptyset \},$$

and

$$X_{outS} = X_{outA} \cup^{r'} X_{outB} \cup^{r'}$$

$$\{ p \in X_{onA} \cap^{r'} X_{onB} | N(p; S) \neq \emptyset \}.$$  

In the theorem, $N(p; S)$ is the neighborhood of the point $p \in S$. The neighborhood is defined as the intersection of $S$ and an open ball, $B(p, r)$, with radius $r$ and center $p$, namely

$$N(p; S) = S \cap B(p, r), \text{ for small } r > 0.$$  

Theorem 2 forms the basis of the membership classification used in regularized intersection. For the regularized union and difference, analogous theorems can be formulated. Given a CSG tree $S$, the following algorithm evaluates the membership classification function for a candidate set $X$:

```
M[Candidate set X, CSG tree S]
    If S is a primitive then
        return(prim-M(X, S))
    else
        return(combine(M[X, left(S)], M[X, right(S)], operation(S)))
```
Prim-M is a classification function based on the known set of primitive objects, and classifies $X$ with respect to the primitive $S$. Combine merges the classification results of the two subtrees according to the set operation \textit{operation}(S), as prescribed by Theorem 2 and its variations for the other boolean operations.

The membership classification algorithm is used to construct the equivalent B-rep of a CSG tree, by a \textit{generate-and-test} paradigm [32]. In the incremental version of the algorithm, the B-reps of the primitives are constructed and are passed up the tree to the root. At each internal node, the B-reps are classified and combined. In a single-phase algorithm, the faces, or the support surfaces of all the primitives are classified against the entire CSG tree to obtain the faces of the final B-rep and are then assembled. Intermediate B-reps are not constructed in this case [2].

2. SET OPERATIONS ON BOUNDARY REPRESENTATIONS

For computing the union, intersection or difference of two solids in B-rep, we need to find the intersection curves of their boundaries on both solids and must classify the faces of both B-reps to determine which ones are inside, outside or on the boundary of the other solid. With the classification, the resulting B-rep can be constructed. We can perform the boolean operations by the following four-step algorithm, where $A$ and $B$ are the input B-reps:

\textbf{Boundary Merging:} Fragment B-rep $A$ so that no face or edge penetrates or grazes $B$, and conversely fragment B-rep $B$ so that no face or edge penetrates or grazes $A$.

\textbf{Boundary Classification:} Classify the faces of both $A$ and $B$ to obtain an eight-way classification. The set of faces of each B-rep is partitioned into four classification sets

\begin{align*}
F(A) &= A_{out}B \cup A_{in}B \cup A_{with}B \cup A_{anti}B, \\
F(B) &= B_{out}A \cup B_{in}A \cup B_{with}A \cup B_{anti}A.
\end{align*}

Corresponding to the faces that are outside, inside, on the boundary with same orientation and on the boundary with an orientation opposite to the other solid.

\textbf{Construction:} Construct the resulting B-rep according to the operation by either merging the two B-reps and removing unused faces, or by copying the needed faces from both B-reps to form a new B-rep. The needed faces are the faces of exactly three of the eight classification set as shown in Table 2.
Table 2: Faces Needed In Set Operation Indicated by $\oplus$ and $\ominus$. Face Orientation Must Be Reversed for $\ominus$

Topological Reduction: Apply topological reduction on the resulting B-rep to form maximally connected faces. Adjacent coplanar faces and adjacent collinear edges are merged.

Exactly how these four steps are implemented varies in the proposed algorithms. Two algorithms follow the four steps exactly, the one by Laidlaw, Trumbore and Hughes [33] and the one by Mäntylä [4]. In Laidlaw's algorithm, the faces of the two objects that penetrate the other object are subdivided. Classification of the faces is performed by casting a ray from a face of one object through the other object. Classifying each face separately is avoided by first grouping together all adjacent faces that do not penetrate the boundary. After classification, faces that do not contribute to the resulting object are removed, and the remaining faces of the two objects are glued to form the new object. The representation assumes convex faces which somewhat simplifies the face/face intersection algorithm. Mäntylä's set operation algorithm allows arbitrary polygonal faces and uses a vertex-neighborhood computation to classify the faces [14].

Robust implementations of the boolean set operators on B-reps are difficult to achieve with finite-precision arithmetic [34]. Typically, the source of the problem is that different numerical computations may imply related geometric facts, such as incidence or coincidence. In this situation, two separate computations may lead to contradictory conclusions, and this entails failure of the algorithm. For a survey of this problem and approaches to solving it see [35], or [2, Chapter 4].

3. SET OPERATIONS ON OCTREES

The union, difference and intersection operations are easy to implement for region octrees, by a coordinated traversal of both trees. Figure 8 shows a two-dimensional example.

Boolean operations on extended octrees and polytrees are straightforward extensions of the region octree algorithms. Suppose we want to compute the
intersection of two region octrees. The two input trees are simultaneously traversed, and a new tree is constructed bottom up, combining corresponding nodes of the input trees. Note that it may be necessary to split nodes in order to obtain compatible levels of detail. For the extended octrees and polytrees, the algorithms must take into account the three additional node types, namely, the vertex, the edge, and the face nodes, as shown in Table 3. The intersection of two face regions can result in a black node, a white node, a face node, an edge node, or a gray node. For example, suppose one of the nodes is a gray node and the other is a vertex node. Since at this level, it is not known how the vertex region interacts with the descendents of the gray node, the vertex node region must be subdivided into eight octants. This changes the problem from a gray/vertex pair to a gray/gray pair, at the cost of processing the eight child pairings recursively. Eventually, nodes representing regions of the same level of detail are reached, and are intersected. The intersection may require additional subdivision in case the geometric structure obtained is too complicated.

It is customary to implement only one binary operation, say regularized intersection, and to obtain the remaining operations by one intersection and several complement operations. The complement of an extended octree is computed by changing all the black nodes into white nodes, changing all the white nodes into black nodes, and changing the orientation of all the plane equations in the other nodes.

If the recursion reaches the limit depth, further recursion is abandoned and a nasty node is created. Nasty nodes are either left empty, or they are associated with a list of all the faces that intersect it. The rationale of this step is that the regions represented are so small that referencing them is unlikely. In the event they are referenced, they can be evaluated further on demand.
Table 3: Possible Node Types For Intersection Using Octree Representation

<table>
<thead>
<tr>
<th>Column</th>
<th>White</th>
<th>Black</th>
<th>Face</th>
<th>Edge</th>
<th>Vertex</th>
<th>Gray</th>
</tr>
</thead>
<tbody>
<tr>
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<td>W</td>
<td>W</td>
<td>W</td>
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</tr>
<tr>
<td>Black</td>
<td>W</td>
<td>B</td>
<td>F</td>
<td>E</td>
<td>V</td>
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</tr>
<tr>
<td>Face</td>
<td>W</td>
<td>F</td>
<td>WFEVG</td>
<td>WFEVG</td>
<td>WFEVG</td>
<td>G</td>
</tr>
<tr>
<td>Edge</td>
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<td>E</td>
<td>WFEVG</td>
<td>WFVG</td>
<td>WVG</td>
<td>G</td>
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<tr>
<td>Vertex</td>
<td>W</td>
<td>V</td>
<td>WFEVG</td>
<td>WVG</td>
<td>WVG</td>
<td>G</td>
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<tr>
<td>Gray</td>
<td>W</td>
<td>G</td>
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<td>G</td>
<td>G</td>
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</tbody>
</table>

III. MULTIDIMENSIONAL SPACE PARTITIONING

We have reviewed three schemas for representing solids: boundary representations, cell decomposition representations, and constructive solid geometry. Each has certain advantages over the others, and many geometric modeling systems improve efficiency of their operations by maintaining solids in several different representations, possibly converting from one representation to another when appropriate. Systems that use explicit representations frequently use a B-rep as the primary representation and add some volume-based information to speed up access to parts of the B-rep. In this section, we present as an alternative a unified representation that combines a B-rep with a volume-based representation, called the B-rep index, and show how to implement some common geometric operations using this unified representation.

The B-rep index is an extension of the BSP tree [23]. A $d$-dimensional BSP tree hierarchically decomposes space into $d$-dimensional convex regions. Given a solid, the root node of a BSP tree represents the entire space while the leaves represent regions that are either completely inside or completely outside the solid. Each internal node contains an oriented cut plane that divides the region represented by the node into two open subregions, one above and one below the cut plane. This recursive decomposition of $d$-dimensional space is sufficient to uniquely represent polyhedral solids. One of the many operations that the BSP tree can support is the classification of a point in relation to a solid, that is, whether the point lies inside, on the boundary or outside the solid. However, since the BSP tree decomposes only the $d$-dimensional space and does not partition the boundary of the solid which is of lower dimension, the BSP tree cannot provide detailed information about the boundary. Yet many geometric modeling problems including Boolean set operations, orthogonal grid generation, and collision detection and analysis need to retrieve this boundary information.
where \( x \) is a face, edge, or vertex. If \( x \) is a face, the second component is a cyclic list of \( n \) vertices bounding the face, and the third component is a list of lines containing consecutive pairs of points in the cyclic list. Note that \( x \) can be a subpolygon of a face of the B-rep. If \( x \) is an edge \( e \), the triple is \([e,(p_0,p_1),(\ell)]\) where \( p_0 \) and \( p_1 \) bound the edge and \( \ell \) is the line containing \( p_0 \) and \( p_1 \). Again, \( e \) could be a segment of an edge of the B-rep. For a vertex \( v \), the triple is \([v,(p),(\ell)]\), where \( p \) gives the coordinates of the vertex. Note that \((p_0,p_1)\) could be two closely-spaced points on a longer edge. Thus explicitly representing the support line increases accuracy when constructing support planes through the two points.

\[
\text{CreateBrepIndex}(\text{Set of triples } X, \text{ Dimension } d, \text{ Set of planes } \mathcal{P})
\]

\[
\text{if } |X| > 1 \text{ then } \\
\quad \text{let } P \leftarrow \text{ChooseCutPlane}(X, d, \mathcal{P}) \\
\quad \{X_a, X_b, X_c\} \leftarrow \text{Cut}(X, P) \\
\quad i_a \leftarrow \text{if } X_a = \emptyset \text{ then OUTSIDE else CreateBrepIndex}(X_a, d, \mathcal{P}) \\
\quad i_b \leftarrow \text{CreateBrepIndex}(X_b, d-1, \mathcal{P} \cup \{P\}) \\
\quad i_c \leftarrow \text{if } X_c = \emptyset \text{ then INSIDE else CreateBrepIndex}(X_c, d, \mathcal{P}) \\
\quad \text{return Node}(P, i_a, i_b, i_c) \\
\} \text{ else return } x \text{ given that } X = \{[x, (p), (\ell)]\}
\]

The function finds an oriented cut plane, \( P \), that contains entity \( x \) that occurs in a triple in \( X \) and is such that \( P \) does not already appear in \( \mathcal{P} \). If \( x \)
is a face, then the cut plane is the support plane of the face. If \( z \) is an edge or a vertex, the cut plane is the support plane of one of the adjacent faces of \( z \). If there is more than one adjacent face, and this may be the case for a vertex or a nonmanifold edge, then a plane is chosen that maximizes transversality with the planes in \( P \). If the support plane of every adjacent face is already in \( P \), then \( P \) is chosen to be perpendicular to the planes in \( P \).

In the left part of Figure 11, for example, the two solids shown are cut by a plane \( P \) that contains the shaded face of the left solid. The cut generates 15 entities contained in the plane. Seven of them are open line segments and these will be further isolated by other cut planes.

All entities of \( X \) that cross the cut plane \( P \) are split by the intersection with \( P \). The resulting subentities are then put into the sets \( X_a \), \( X_b \), and \( X_c \) according to whether they lie above, on or below \( P \). A face crossing \( P \) is split into faces lying above and below, and edges lying on \( P \). An edge crossing \( P \) is split into the edges above and below and the vertex on \( P \). Figure 12 illustrates the fragmentation with plane \( P \) of a face, its 16 edges and its 16 vertices. After the cut, the region above \( P \) consists of two faces, six edges and four vertices. The region on \( P \) consists of seven vertices and six edges. The region below \( P \) consists of three faces, nine edges, and six vertices.

Figure 12: Fragmentation of a Face, Bordering Edges and Vertices by Cut Plane \( P \).
B. COMPRESSING THE B-REP INDEX

After the B-rep index has been so created, it can be used directly. However, the B-rep index can be compressed considerably, and doing so speeds up accessing it. Compression proceeds in three steps:

1. Locate and eliminate all redundant cut planes.

2. Reorder some cuts to uncover and eliminate additional redundant cuts.
   By restructuring the tree so that the cut planes are queried in a different order, other cut planes may become redundant and the corresponding nodes eliminated.

3. If the tree is badly unbalanced (for instance in the case of convex objects) rebalance the tree by introducing a few well-chosen redundant cuts.

We give tree rewrite rules that compress the tree and illustrate the rules by compressing the tree of Figure 10. In prefix notation, the subtree rooted in \( n \) is written as \( P(a, b, c) \) where \( P \) is the cut plane at \( n \), and \( a, b, \) and \( c \) denote the above, on, and below subtrees of \( n \), respectively.

The first rule states that any extended region can be cut redundantly by a plane that is transversal with that region. For \( P \cap R(a) \subset R(a) \) and \( \dim(a) > 0 \),

\[
a \Rightarrow P(a, a, a). \quad (R1)
\]

The converse rule states that this redundant cut can be removed

\[
P(a, a, a) \Rightarrow a. \quad (R2)
\]

If the cut plane orientation is reversed, then the order of the children must be reversed also:

\[
P(a, b, c) \Leftrightarrow -P(c, b, a). \quad (R3)
\]

Two transversal cuts of a given region commute as follows:

\[
P_1(P_2(a, b, c), P_2(d, e, f), P_2(g, h, i)) \Leftrightarrow P_2(P_1(a, d, g), P_1(b, e, h), P_1(c, f, i)). \quad (R4)
\]

The above rules do not need any geometric information. In contrast, the next rule requires knowing the orientation of the cut planes and generalizes AVL tree rotation [36]. Given that the cut plane of a child node does not intersect the cut plane of the parent node within the parent region, the order of the cuts can be interchanged in a way that depends on the relative orientation of the cut planes.

Given that \( P_1 \cap P_2 \cap R(n) = \emptyset \),

\[
P_1(P_2(a, b, c), d, c) \Rightarrow \begin{cases} P_2(a, b, P_1(c, d, e)) & \text{if } R(c) \subset P_2^- \\ P_2(P_1(a, d, e), b, c) & \text{if } R(c) \subset P_2^+ \end{cases}, \quad (R5a)
\]
Figure 13: Elimination of Redundant Cut $P_1$ by Rule 7.

and

$$P_1(a, b, P_2(c, d, e)) \Rightarrow \begin{cases} P_2(P_1(a, b, c), d, e) & \text{if } R(a) \subset P_1^+ \\ P_2(c, d, P_1(a, b, e)) & \text{if } R(a) \subset P_1^- \end{cases} \quad (R5b)$$

From the given rules additional rules can be derived. For instance,

$$P_1(P_2(a, b, c, c, P_3(c, d, e)) \Rightarrow P_2(a, b, P_3(c, d, e))$$

$$P_1(P_2(a, b, c, c, P_3(c, d, e)) \Rightarrow P_3(P_2(a, b, c), d, e)) \quad (R6)$$

and

$$P_1(P_2(a, b, c), c, c) \Rightarrow P_2(a, b, c). \quad (R7)$$

Both rules have additional variations that are derived using Rule (R3).

To illustrate the use of the rewrite rules, we compress the B-rep index of Figure 10. When a subtree is changed, the rule that applies is indicated in the superscript. We begin the compression by noting that $P_5$ is redundant, and remove it by propagating it to the bottom of the tree where it is removed by Rule (R2).

$$P_1(P_3(\text{out}, П_2(\text{out}, v_3, e_5),$$

$$\quad P_3(P_2(\text{out}, e_2, f), P_2(\text{out}, e_2, f), P_2(\text{out}, e_2, f)))^{R4}),$$

$$\quad P_3(P_2(\text{out}, v_1, e_1), v_2, P_3(P_2(\text{out}, e_2, f), f, f)^{R5}),$$

$$\quad P_3(P_2(\text{out}, v_4, e_2), P_4(\text{out}, e_4, f))^{R4},$$

$$\quad P_4(\text{out}, e_4, f), P_3(\text{out}, e_4, f)))$$

$$= P_1(P_3(\text{out}, П_2(\text{out}, v_3, e_5),$$

$$\quad P_3(P_2(\text{out}, \text{out}, \text{out})^{R2}, P_5(e_2, e_2, e_2)^{R2}, P_5(f, f, f)^{R2}),$$

$$\quad P_3(P_2(\text{out}, v_1, e_1), v_2, P_2(\text{out}, e_2, f)),$$

$$\quad \ldots)$$

$$= P_1(P_3(\text{out}, П_2(\text{out}, v_3, e_5), П_2(\text{out}, e_2, f)),$$

$$\quad P_3(\text{out}, v_1, e_1), v_2, P_2(\text{out}, e_2, f),$$

$$\quad P_4(\text{out}, П_2(\text{out}, v_4, e_4), П_2(\text{out}, e_2, f)))$$
Figure 14: The Compressed B-rep Index of B-rep Index of Figure 10.

Next we observe that the subtree $P_2(out, e_2, f)$ appears multiply as a cut of $P_1$, and so we exchange the order of the cuts by moving $P_2$ up and exchanging with Rule R4.

$P_1(P_3(out^{R1}, P_2(out, v_3, e_3), P_2(out, e_2, f))^{R4}$,
$P_3(P_4(out, v_1, e_1), v_2, P_2(out, e_2, f))^{R5}$,
$P_4(out^{R1}, P_2(out, v_4, e_4), P_2(out, e_2, f))^{R4}$

$= P_1(P_3(out, P_3(out, v_3, e_2), P_3(out, e_3, f))$,
$P_2(out, e_2, P_3(P_4(out, v_1, e_1), v_2, f))$,
$P_2(out, P_4(out, v_4, e_2), P_4(out, e_4, f))^{R4}$

$= P_2(P_1(out, out, out))^{R2}$,
$P_1(P_3(out, v_3, e_2), e_2, P_4(out, v_4, e_2))^{R6}$,
$P_1(P_3(out, e_3, f), P_3(P_4(out, v_1, e_1), v_2, f))$,
$P_4(out, e_4, f))^{R4}$

$= P_2(out, P_3(out, v_3, P_4(out, v_4, e_2))$,
$P_1(P_3(out, e_3, f), P_3(P_4(out, v_1, e_1), v_2, f))$,
$P_4(out, e_4, f))^{R4}$.

The original B-rep index tree consisted of 17 internal nodes; the final tree has only eight nodes, as shown in Figure 14. In this simple example, we achieved a reduction in the size of the tree by more than fifty percent. Vaněček has shown that for a B-rep with $v$ vertices, $e$ edges and $f$ faces its B-rep index has at least $v + e + f$ internal nodes, and $1 + 2(v + e + f)$ leaf regions [37]. In 2D, the number of internal nodes is at least $v + e$, and the number of leaf regions is at least $1 + 2(v + e)$. Our 2D example has four vertices and four edges, so the compressed tree with eight nodes is minimal.
C. POINT AND LINE CLASSIFICATION

Most geometric computations reduce to point/solid and line/solid classification. This includes boundary merging, collision detection, grid generation and ray casting. The point/solid classification determines whether a point is inside, outside or on the boundary of a solid. With the B-rep index, if the point lies on the boundary of the solid, the boundary entity on which it lies is also determined. The line/solid classification partitions the line where it penetrates or touches the boundary.

Classifying a point with the B-rep index is simple. Starting from the root, check the position of the point relative to the cut planes of the internal nodes, and proceed through the corresponding subtrees until a leaf is reached. At the leaf the point classification is known.

```
classify(Point p, Node n)
  while n is an internal node do {
    Let d be the signed distance of p to P(n)
    n ← \{
      ABOVE(n) if d > 0
      BELOW(n) if d < -1
      ON(n) otherwise
    }
  }
  return n
```

Since an exact answer to "a lies on b" cannot be given in floating point arithmetic, we compute an approximate answer that depends on a tolerance \( \epsilon > 0 \).

The tolerance can be viewed as a thickness of the boundary. Since using exact arithmetic is not practical, the tolerance will affect the classification. Recall that a given B-rep does not have a unique M3P tree. The tree structure depends on the order and choice of the cutting planes. The example of Figure 15
Figure 16: Line-Segment/Solid Classification.

shows a boundary fragment with two different MSP subtrees. Figure 15(a) is

\[ P_1(..., P_2(\text{out}, v_1, P_3(\text{out}, v_2, e)), ...) \]

and Figure 15(b) is

\[ P_1(..., P_2(\text{out}, v_2, P_3(\text{out}, v_1, e)), ...) \]

and these are equivalent by Rule R5b. Now consider the point indicated by
the circle in the figures. Using tree (a), the point is found to be coincident
with \( v_1 \), but using tree (b), the point is found to be coincident with \( v_2 \). This
shows that equivalent MSP trees for a given solid do not classify points the
same way. This discrepancy is caused by a tolerance that is too large relative
to the separation between two entities. In particular, as the tolerance changes,
the classification result may change. It is possible that a point classified as
coincident with a vertex, at one tolerance, is found coincident with another
vertex at a different tolerance.

Consider now the line/solid classification problem and refer to Figure 16
for an example. The classification of the line segment \( pq \) should have the result

\[ [\text{out}, (f_1, p), \text{in}, (c_1, q), f_2, (v_4, p_4), c_2, (v_5, p_5), \text{out}] \]

(1)

where \( p, q, p_4, p_5 \) are the points at which the line penetrates the corresponding
entities, and the \( v \)'s, \( c \)'s and \( f \)'s are the vertices, edges and faces of the B-rep.

An intuitive way to classify the line is to pass the line segment down the
tree, split the line segment whenever it crosses \( P(u) \) at a node \( u \), and classify
the portions above, on, and below recursively. This approach, however, is
problematic when the line segment forms a very small angle with a cutting
plane. Figure 17 shows a line segment, \( \ell \), close to vertex \( v \). Although \( \ell \) can be
arbitrarily close to \( v \), the line segment crosses the plane \( P_1 \) far away from \( v \),
because of the small angle between \( P_1 \) and the line. So the point of intersection
on \( P_1 \) and the two subsegments fall into the regions outside the solid. This is
not intuitive, partly because the thickness of the \( \pm \epsilon \) region around the planes
Figure 17: Part of Face With Line Segment $\ell$ Crossing at $v$ Within Tolerance.

has not been well accounted for in the above approach. A better classification is obtained when the interval of $\ell$ passing through the $\pm \epsilon$ regions of $P_1$ and $P_2$ is computed. The closest point $r$ on $\ell$ to $v$ is then computed by projecting $v$ onto $\ell$.

The line classification thus proceeds by passing the original interval $[t_1, t_2]$ from the top to the bottom of the B-rep index in phase one, followed by a second phase in which the final line partition is determined. In phase one, the interval $[t_1, t_2]$ reaches the node $n$ and is partitioned as either $[t_1, t_2]$ or $[t_1, t_4, t_5, t_2]$ depending on the orientation of the plane $P(n)$. The three subintervals are processed in the corresponding subtrees. An interval is given a preliminary classification at the leaf node it reaches.

Next, the sequence of intervals is reduced by merging adjacent intervals with common classifications and by isolating the boundary penetrations. In the example of Figure 17, after step 1, we have the classification

$$[\text{out}, e_1, v, \text{out}, \text{out}]$$

After merging adjacent equal classifications, this sequence is reduced to

$$[\text{out}, e_1, v, \text{out}]$$

The classification corresponds to the sequence of dimensions $d_i$, $[3, 1, 0, 3]$. We replace each maximal bitonic sequence $d_i > \cdots > d_j < \cdots < d_k$ with the triple $d_i, d_j, d_k$. That is, all intervening intervals and their classification are dropped. Then the sequence is reduced to $[3, 0, 3]$ with the associated classification $[\text{out}, v, \text{out}]$. The reduced classification is used to compute the final interval partition, by computing the intersection of the line segment with the remaining faces, edges, and vertices. Note that this may require projecting say a vertex onto the line segment.

The reduction above does not address how to reduce classification sequences that contain adjacent, different entities of the same dimension. Consider line
Figure 18: Ambiguous Line Classification With Oversized Tolerance.

$t$ of Figure 18. Its classification after step one is

$$[\text{out}, e_1, e_2, e_3, e_4, \text{out}].$$  \hspace{1cm} (2)

The correct classification is either

$$[\text{out}, v, \text{out}],$$

or

$$[\text{out}, e_1, f_1, e_2, \text{out}, e_3, f_2, e_4, \text{out}].$$

From Sequence (2) we cannot derive the two classifications nor can we determine which one is correct. In this situation, the line has to be reclassified with a different tolerance. In our example, a larger tolerance yields the first classification, whereas a smaller tolerance yields the second.

In many applications, both point/solid and line/solid classification are done. Consider the following example, and refer to Figure 17. The segment $t$ enters the tolerance region of edge $e_1$ from the left, at a point $q$. There is a part of $t$ adjacent to $q$ that is in the tolerance region of $e_1$, but not in the tolerance region of $v$. Let $p$ be a point on that part of $t$. Classified as a point, $p$ is found to be on the edge $e_1$. When $t$ is classified, however, the final segment classification will be $[\text{out}, v, \text{out}]$, thus $p$ will be determined to be out, a contradictory classification, even though both classifications used the same tolerance. In problems such as grid generation, we account for this possibility.

D. ORTHOGONAL GRID GENERATION

We generate orthogonal grids using line/solid classification. Orthogonal grids are used by finite-differencing solvers for partial differential equations (PDE), for example in Ellpack [38], a system for solving elliptic PDEs. The grids can be regular, with uniform distance between grid points, or adaptive, with grid points that are denser in some parts of the domain.

A regular three-dimensional grid can be specified by two extreme points of the grid's bounding cuboid, and by the number of grid planes $n_x$, $n_y$, and
$n_z$ along each major axis. The intersection of the grid planes defines $n{x}_z n_y + n_y n_z + n_z n_x$ grid lines and $n_x n_y n_z$ grid points. Our task is to classify all grid points and determine where the grid lines cross the boundary between grid points. It is clear from the previous section that grid points and grid lines should not be classified independently. Instead, we classify only the grid lines and process local inconsistencies, thus obtaining a consistent grid point and grid line classification.

Consider the two-dimensional example of Figure 19(a) in which the two grid lines $\ell_1$ and $\ell_2$ were classified independently. The intersections of $\ell_1$ and $\ell_2$ are the grid point $p$ that is near to the vertex $v$. On $\ell_1$, $p$ is classified as in; on $\ell_2$, it is classified as out, because of the projection computation when segmenting the lines. Classified as a point, $p$ would be $v$. By shifting them to the grid point, the two vertex projections are made coincident, giving $p$ a consistent classification; Figure 19(b). This approach requires that the tolerance regions are substantially smaller than the minimum grid point separation. Otherwise, there could be several grid points to which to shift the two projections of $v$. The classification of grid points that lie within tolerance of an edge or a face is corrected in the same way.

The cost of classifying the grid lines is $O((n_x n_y + n_y n_z + n_z n_x) L(m))$, where $L(m)$ is the cost of the line/solid classification in a MSP tree with $m$ nodes. The cost of correcting the grid point classifications by shifting the vertex projections is $O(n_x n_y n_z)$. The method has been implemented; [26].

E. COLLISION DETECTION

Our approach to orthogonal grid generation is also used for collision detection and analysis of moving solid objects, [27]. The collision problem arises in
Figure 20: (a) Solids $s_1$ and $s_2$ With Respect to Local Coordinates. b) $s_1$ and $s_2$ Placed in Global Frame of Reference. c) $s_1$ Mapped To Local $s_2$-Frame of Reference. d) $s_2$ Mapped To Local $s_1$-Frame of Reference.

computer simulations of physical systems, based on rigid-body dynamics; for example, in the Newton system [39,40]. Here, objects of arbitrary shape are permitted, and their motion cannot be predicted in advance, so that collisions can happen anywhere and at any time. Moreover, objects may be in contact with each other for some time, and such contacts begin and end in ways that depend entirely on the dynamics of the system, and again cannot be predicted beforehand. So, the positional relationships of all objects must be analyzed.

In the Newton system, the dynamics evaluation inquires at certain time instances whether two bodies are in contact, and if so, a geometric analysis of contact locality is required. Collision detection and analysis can be reduced to a static problem in Newton as follows: All solids have a local frame of reference centered at the mass center. Objects are placed in relation to each other by a $4 \times 4$ transformation matrix $M$ that positions and orients the local frame with respect to a global frame of reference. As objects move, the matrix $M$ is changed accordingly.

Consider two solids $s_1$ and $s_2$ and their associated transforms $M_1$ and $M_2$. Then solid $s_1$ can be related to the local frame of reference of $s_2$ by the transformation

$$M_{12} = M_2^{-1} \cdot M_1$$

See also Figure 20. Similarly, $M_1^{-1} \cdot M_2$ relates $s_2$ to the local frame of $s_1$. We classify each edge of $s_1$ against $s_2$ after transforming the edges with $M_{12}$, and classify each edge of $s_2$, after transformation by $M_{21}$, against $s_1$. Note that this strategy does not require changing the B-rep index of either solid, irrespective of their motion over time. The method has been implemented in the Newton system.

As in the case of orthogonal grid generation, floating-point arithmetic causes robustness problems. Consider an edge of $s_1$ touching an edge of $s_2$ at an angle. Projecting one edge onto the other determines a pair of contact
points, one point on each edge. When classifying the edges of \( s_1 \) we obtain one pair, and when classifying the edges of \( s_2 \) we obtain another pair. The points in the two pairs should coincide, but often they do not. We resolve this discrepancy by averaging corresponding points. It is also possible that the first classification reports two touching edges, whereas the second classification does not find this crossing. In this case, we give priority to the contact detection. These strategies have been very successful in the Newton system. Clearly, such heuristics ameliorate robustness problems but do not completely eliminate them. Indeed, solving the robustness problem completely is extremely costly, [2, Chapter 4]. Thus, in practice it is often acceptable to use incomplete solutions as long as they eliminate most robustness problems that arise.

F. BOOLEAN SET OPERATIONS WITH THE B-REP INDEX

We implement the boolean set operations using the B-rep index with the four-step algorithm of Section II.D.1. That is, we merge the boundaries, classify the faces, construct the result solid, and then compress its representation. Let \( A \) and \( B \) be the input solids. We could use the algorithm of Section III.A to construct a B-rep index of both B-reps together, obtaining an MSP tree in which the leaves have two classification fields, one for the region with respect to solid \( A \), the other with respect to solid \( B \). Since the B-reps of \( A \) and \( B \) are not explicitly subdivided, it is difficult to construct the B-rep index of the result. In particular, extracting the curves of intersection from the joint index would have to be done without explicit adjacency information.

A better approach is to construct (implicitly) a joint B-rep index of both input solids that explicitly subdivides the boundaries of \( A \) and \( B \), so that the intersection curves on the surfaces can be extracted easier. Algorithm \texttt{MergeBreps} does this by maintaining separate entity sets, with \( X \) initially all boundary entities of \( A \) and \( Y \) all boundary entities of \( B \). We intend to cross-classify entities, so that, ultimately, the boundary entities of \( A \) are classified as in or out with respect to \( B \), or are classified by the coincident entity of \( B \). Likewise, the entities of \( B \) should be cross-classified with respect to \( A \). We associate with each entity \( x \) a classification field, \( C(x) \), initially set to unknown. At the end, the classification will be known for coincident and intersection entities, but not for entities that are outside or inside the other solid.

Figure 21 shows two objects before and after boundary merging, with the dotted lines indicating coincident entities. Note that the merging step, constructing the joint MSP tree, subdivides both boundaries, but only in the vicinity of the intersection curves. In particular, the right edge of the box in the figure is not separated from its bounding vertices.
Figure 21: Objects Before and After Boundary Merging.

```plaintext
MergeBrepSet\{Set of entities \(\mathcal{X}, \mathcal{Y}\); Dimension \(d\); Set of planes \(\mathcal{P}\)\)
if \(|\mathcal{X}| = 1\) and \(|\mathcal{Y}| = 1\) then
  \(C(\mathcal{X}) \leftarrow y\), and \(C(\mathcal{Y}) \leftarrow x\), where \(\mathcal{X} = \{x\}\), and \(\mathcal{Y} = \{y\}\)
else {
  let \(P \leftarrow \text{ChooseCutPlane}(\mathcal{X}, \mathcal{Y}, d, \mathcal{P})\)
  \((\mathcal{X}_0, \mathcal{X}_a, \mathcal{X}_b) \leftarrow \text{Cut}(\mathcal{X}, P)\)
  \((\mathcal{Y}_0, \mathcal{Y}_a, \mathcal{Y}_b) \leftarrow \text{Cut}(\mathcal{Y}, P)\)
  if \(\mathcal{X}_a \neq \emptyset\) and \(\mathcal{X}_b \neq \emptyset\) then
    MergeBreps(\(\mathcal{X}_a, \mathcal{X}_0, d, \mathcal{P}\))
  if \(\mathcal{Y}_a \neq \emptyset\) and \(\mathcal{Y}_b \neq \emptyset\) then
    MergeBreps(\(\mathcal{Y}_a, \mathcal{Y}_0, d, \mathcal{P}\))
  if \(\mathcal{X}_a \neq \emptyset\) and \(\mathcal{Y}_b \neq \emptyset\) then
    MergeBreps(\(\mathcal{X}_b, \mathcal{Y}_0, d-1, \mathcal{P} \cup \{P\}\))
}
```

After the joint MSP tree \(T\) has been constructed, we must cross-classify the remaining entities. We do this by casting a ray as follows. Let \(x\) be an entity, say of \(A\), whose classification is unknown, and which therefore is either in or out \(B\). Note that \(x\) is attached to a leaf of \(T\) that represents a convex region, and that the parent's region is also convex. The parent's region must contain part of the boundary of \(B\), say some entities including \(y\). We cast a ray from an interior point of \(x\) to an interior point of \(y\). Then only ray intersections with boundary elements in the parent's region are considered, because the region is convex, and these entities are indexed by the subtree rooted in the parent. Thus, classifying by ray casting is very efficient, since only a small part of the boundary must be considered.

After this second step, every entity \(x\) of either B-rep has the correct classification \(C(x)\), which is either the labels \(\text{in}\), or \(\text{out}\), or a pointer to a vertex, edge,
paradigms. Nevertheless, implicit surfaces are useful in geometric modeling, and [2] gives many details of their properties and describes techniques for interrogating them. Implicit algebraic surfaces include in principle all parametric surfaces, but not vice versa. Current techniques for converting from parametric to implicit form, and methods for parameterizing implicit algebraic surfaces that possess a rational parametric form, can be demanding symbolic computations [45]. Simple algorithms are known for monoids, a special class of surfaces that includes all quadratic surfaces. See [2] for details.

In certain applications, one would like to construct curved surfaces that satisfy prescribed constraints. In many cases, these surfaces are readily explained and informally specified in intuitive geometric terms. For instance, given two surfaces \( f \) and \( g \), consider all points in space that have equal minimum distance from the given surfaces. Such points form the \textit{equal-distance} or \textit{Voronoi surface} of \( f \) and \( g \). Another example is the \textit{rolling-ball blend}, a blending surface between given surfaces \( f \) and \( g \) obtained as follows: Roll a sphere such that it maintains contact with both \( f \) and \( g \), at all times. Considering the volume in space traversed by the sphere, the rolling ball blend is defined by the surface of that volume.

The basic property of these and similar other surface definitions is that the new surface to be defined can be expressed in terms of one or more base surfaces and a number of intuitive geometric constraints. Despite this conceptual simplicity, it is by no means a trivial undertaking to represent such surfaces in exact mathematical terms, in a manner suitable for computation. Often, neither implicit nor parametric surface representations can be found, and so we seek an alternative to these two representation schemata. The \textit{dimensionality paradigm} provides a straightforward and mathematically rigorous method for exactly representing surfaces defined through constraints. The dimensionality paradigm simplifies representing complicated surfaces, and it permits generic algorithms for manipulating them. The method is also of interest because it is an enabling technology for constructing promising curved-solid representations such as the skeleton discussed later in Section V.

B. THE DIMENSIONALITY PARADIGM

We observe that the definition of complex constrained surfaces often simplifies when we consider the surface as the natural projection of a manifold in higher-dimensional space. The manifold is defined by a system of nonlinear equations in \( n \) variables, where \( n > 3 \). The extra dimensions may be point coordinates on the base surface(s), distances, or other quantities used to express the constraints that must be obeyed. The surface we want is then the natural projection of this manifold into a three-dimensional subspace.

When the base surfaces are algebraic, and this is normally the case in geometric and solid modeling, then the resulting system of equations can be
processed by a number of symbolic computation algorithms that eliminate all additional variables and arrive at a single implicit equation for the surface. Such an approach is normally intractable, for it routinely arrives at elimination problems that are well beyond what hardware and software can deliver in the foreseeable future. Worse yet, a number of elimination algorithms require exact arithmetic, further exacerbating the magnitude of the computation. Therefore, we will work directly with the system of equations.

We will demonstrate the definitional approach with an example. We have developed a number of algorithms for surfaces defined using the dimensionality paradigm. The algorithms are general and do not require knowledge of the geometric nature of the surfaces. Therefore, they will work unchanged on offsets, on blends, on equal-distance surfaces, and so on. Some algorithms are local in nature. That is, given a point on the surface, the algorithms will explore the surface in the vicinity of the point. Such algorithms can and have been globalized, by suitably embedding the exploration into a spatial grid that coordinates the computation. We will sketch the salient features of many of these algorithms also.

C. AN EXAMPLE DEFINITION

We consider the definition of an equal-distance surface as an example of the dimensionality paradigm. Assume that we are given two implicit surfaces \( f(x, y, z) = 0 \) and \( g(x, y, z) = 0 \). Using a declarative style, we can then describe the equal-distance surface as follows:

(i) Let \( p = (x, y, z) \) be a point on the equal-distance surface. Moreover, let \( p_f = (u_1, v_1, w_1) \) be a point at minimum distance from \( p \) on \( f \), and let \( p_g = (u_2, v_2, w_2) \) be a point at minimum distance from \( p \) on the surface \( g \). Then:

(ii) The point \( p_f \) satisfies the equation of \( f \), and the point \( p_g \) satisfies the equation of \( g \).

(iii) The distance \( (p, p_f) \) is \( d \) and is equal to the distance \( (p, p_g) \).

(iv) The line \( \overline{p_f p_g} \) is normal to \( f \) at \( p_f \).

(v) The line \( \overline{p_f p_g} \) is normal to \( g \) at \( p_g \).

Note that Assertion (i) declares the names of nine variables, the coordinates of three points, whereas Assertions (ii)–(v) simply state the geometric relationships that these points must satisfy.

In order to obtain an equational representation of the equal-distance surface, we translate the Assertions (ii)–(v), using the variable names of (i). We obtain in sequence:

\[
\begin{align*}
    f(u_1, v_1, w_1) &= 0 \quad (3) \\
    g(u_2, v_2, w_2) &= 0 \quad (4)
\end{align*}
\]
\[(x - u_1)^2 + (y - v_1)^2 + (z - w_1)^2 = d^2\]  
\[(x - u_2)^2 + (y - v_2)^2 + (z - w_2)^2 = d^2\]  
\[\{x - u_1, y - v_1, z - w_1\} \cdot [-f_{u_1}, f_{u_1}, 0] = 0\]  
\[\{x - u_1, y - v_1, z - w_1\} \cdot [f_{w_1}, 0, -f_{u_1}] = 0\]  
\[\{x - u_1, y - v_1, z - w_1\} \cdot [0, -f_{w_1}, f_{u_1}] = 0\]  
\[\{x - u_2, y - v_2, z - w_2\} \cdot [f_{u_2}, f_{u_2}, 0] = 0\]  
\[\{x - u_2, y - v_2, z - w_2\} \cdot [f_{u_2}, 0, -f_{u_2}] = 0\]  
\[\{x - u_2, y - v_2, z - w_2\} \cdot [0, -f_{u_2}, f_{u_2}] = 0\]

Subscripting, as in \(f_{u_1}\), denotes partial differentiation.

Equations (3–6) are quite clear and express Assertions (ii) and (iii). Equations (7–9) together express Assertion (iv), since the three vectors

\[-f_{u_1}, f_{u_1}, 0\]
\[f_{w_1}, 0, -f_{u_1}\]
\[0, -f_{w_1}, f_{u_1}\]

are tangent to \(f\) and span the tangent space as long as \(p_f\) is not a singular point on the surface. Similarly, Eqs. (10–12) together express Assertion (v).

Note that if \(f\) is given in parametric form, then Eq. (3) and Eqs. (7–9) have to be adjusted. This is routine and shows that the methodology is independent of the representation of the base surfaces. In principle, an implicit equation could be derived by eliminating the variables \(\{u_1, v_1, \ldots, u_2, d\}\) from the system, but in almost all cases this computation is not tractable.

The entire system of equations defines a manifold in 10-dimensional space.\(^3\) The projection of that manifold into the \((x, y, z)\)-subspace is the equal-distance surface. A number of papers discuss other examples of surface definitions using the dimensionality paradigm, including

- offset surfaces, [46,45];
- constant-radius blends, [46];
- variable-radius blends, [46,47];
- ruled surfaces in parametric blending, [48];
- trimming surfaces in skeleton computations, [49].

We do not repeat these definitions here.

\(^3\)Note that \(d\) is a variable
D. FAITHFUL DEFINITION SYSTEMS

The equation systems formulated by the dimensionality paradigm entail certain additional point sets that are unwanted because they do not reflect the geometric intent. The origin of these additional solutions is found in possible interdependence of the individual equations at certain points in space. For example, if $p_f$ is a singular point, then Eqs. (7–9) vanish. In consequence, the system also defines a manifold that projects to the equal-distance surface of $g$ and the singular point.

Extraneous solutions present in the system can be excluded by introducing certain additional equations, [50]. These new equations use one or more additional variables that no longer have a direct geometric meaning, but are used to express inequalities through equations. This “trick” originates with refutational approaches in automated geometry theorem proving, [51,52]. Note that all extraneous solutions can be so eliminated. The equations added reflect a generic method, but must account for the geometry of the original system.

A technical subtlety of this work is to define precisely what is meant by “extraneous” solution. Following [50], we define the extraneous solutions in the example of the Voronoi surface, Eqs. (3–12). Let $p = (x, y, z)$ be a point of the equal-distance surface, $p_f = (u_1, v_1, w_1)$ a point on $f$ at minimum distance from $p$, and $p_g = (u_2, v_2, w_2)$ a point on $g$ also at minimum distance from $p$.

The points $p_f$ and $p_g$ are footpoints of $p$ on $f$ and $g$, respectively. Footpoints and the associated surface point(s) are said to correspond. Then a solution is extraneous if it corresponds to a footpoint that, in turn, corresponds to infinitely many solutions. It can be shown that all real extraneous solutions to Eqs. (3–12) arise as follows, [50]:

1. Footpoints $p_f$ or $p_g$ are singular. In this case, we obtain as extraneous solutions points at equal distance from the singular point and the other surface. In case both footpoints are singular but not coincident, there is an additional plane. If both footpoints are singular and coincident, then every point in $\mathbb{R}^3$ is an extraneous solution.

2. The footpoints coincide, are regular, and the surfaces intersect tangentially. In this case, the common surface normal is extraneous.

Note that we must assume for the proof that $f$ and $g$ are algebraic surfaces, because Bezout's Theorem is used to show that all other footpoints correspond to finitely many points of the equal-distance surface.

We explain how to modify the system so as to exclude all extraneous solutions enumerated. Consider the equation

$$\alpha x - 1 = 0$$

It is solved iff both $\alpha$ and $x$ are not zero. Thus, if $x$ is a quantity occurring in some system and $\alpha$ is a new variable, then adjoining this equation to the
system effectively expresses the constraint

\[ z \neq 0 \]

Now consider the equation

\[(\alpha x - 1)(\alpha y - 1) = 0 \]

It expresses effectively

\[(z \neq 0) \lor (y \neq 0)\]

We use this idea to exclude all singular footpoints from the definition of the equal-distance surface by adjoining

\[
(\alpha f_{u_1} - 1)(\alpha f_{v_1} - 1)(\alpha f_{w_1} - 1) = 0
\]

\[
(\beta g_{u_2} - 1)(\beta g_{v_2} - 1)(\beta g_{w_2} - 1) = 0
\]

where \( \alpha \) and \( \beta \) are new variables. The two equations express that not all partial derivatives of \( f \) and of \( g \) vanish simultaneously at footpoints. Likewise, adjoining

\[
(\gamma(u_1 - u_2) - 1)(\gamma(v_1 - v_2) - 1)(\gamma(w_1 - w_2) - 1)(\gamma D - 1) = 0
\]

where

\[
D = \begin{vmatrix}
1 & 1 & 1 \\
1 & f_{u_1} & f_{v_1} \\
1 & g_{v_2} & g_{w_2}
\end{vmatrix}
\]

expresses that the footpoints \( p_f \) and \( p_g \) are distinct, or else that the surface normals through them do not coincide. In consequence, the system of equations

\[
\begin{align*}
 f(u_1, v_1, w_1) &= 0 \\
 g(u_2, v_2, w_2) &= 0 \\
 (x - u_1)^2 + (y - v_1)^2 + (z - w_1)^2 &= d^2 \\
 (x - u_2)^2 + (y - v_2)^2 + (z - w_2)^2 &= d^2 \\
 [x - u_1, y - v_1, z - w_1] \cdot [-f_{u_1}, f_{v_1}, 0] &= 0 \\
 [x - u_1, y - v_1, z - w_1] \cdot [f_{u_1}, 0, -f_{v_1}] &= 0 \\
 [x - u_1, y - v_1, z - w_1] \cdot [0, -f_{u_1}, f_{v_1}] &= 0 \\
 [x - u_2, y - v_2, z - w_2] \cdot [-g_{u_2}, g_{v_2}, 0] &= 0 \\
 [x - u_2, y - v_2, z - w_2] \cdot [g_{u_2}, 0, -g_{v_2}] &= 0 \\
 [x - u_2, y - v_2, z - w_2] \cdot [0, -g_{u_2}, g_{v_2}] &= 0 \\
 (\alpha f_{u_1} - 1)(\alpha f_{v_1} - 1)(\alpha f_{w_1} - 1) &= 0 \\
 (\beta g_{u_2} - 1)(\beta g_{v_2} - 1)(\beta g_{w_2} - 1) &= 0 \\
 (\gamma(u_1 - u_2) - 1)(\gamma(v_1 - v_2) - 1)(\gamma(w_1 - w_2) - 1)(\gamma D - 1) &= 0
\end{align*}
\]
defines the equal-distance surface of \( f \) and \( g \) without extraneous solutions. The same approach has been used in \([50]\) to exclude extraneous solutions from offsets, constant-radius and variable-radius surface definitions.

In this approach to faithful surface definition, the exclusion of all extraneous solutions also removes finitely many solutions that, by continuity, ought to be included in the final surface. For example, associated with two coincident regular footpoints is always the solution \( p = p_l = p_y \), now excluded in case of coincident normals. If the projection of the system into the \((x, y, z)\) subspace is done using variable elimination, then these "missing points" are reintroduced by closure; \([53]\). If the system is interrogated by the methods to be explained next, then the missing points can be reintroduced by compensatory steps in the algorithms. However, in almost all cases the missing points will not be noticed because the numerical processing evaluates only a discrete sample of points, and the missing points are isolated.

E. INTERROGATION OF SURFACES DEFINED WITH THE DIMENSIONALITY PARADIGM

There is a considerable body of algorithmic infrastructure to deal with surfaces defined by systems of nonlinear equations. The following algorithms are now available:

(i) Given two surfaces and an initial point on both, evaluate their intersection; see \([54,2,46]\). The algorithm is robust and can evaluate very high-degree surface intersections without significant precision problems.

(ii) Given a surface and an initial point, evaluate locally the curvatures, \([48]\), and give a local parametric or local explicit surface approximant of arbitrary contact order, \([55,45]\).

(iii) Given a surface and an initial point, globally approximate the surface; \([55]\).

These algorithms are not confined to algebraic equation systems, and can be used as long as the nonlinear equations of the system are continuously differentiable. They are very efficient.

Less efficient are the known techniques for finding initial points. When nothing is known about the system, then generic techniques such as \([56,57,58, 59,60]\) can be applied. Usually, however, the geometric intent of the system is known and leads to good initial estimates for starting points that can be refined using Newton iteration. Moreover, some of the search techniques from CAGD are in principle applicable. That is, by raising the dimension of the ambient space by \( l \), all equations can be rewritten in Bernstein–Bézier form after which domain reduction is applied using the convex hull property. To implement this, one has to exercise care not to obtain an exponential growth in the number of control points.
1. LOCAL PARAMETRIC APPROXIMATIONS

We assume given a surface definition using the dimensionality paradigm, in the form

\[ f_1(x_1, x_2, \ldots, x_n) = 0 \]
\[ f_2(x_1, x_2, \ldots, x_n) = 0 \]
\[ \vdots \]
\[ f_m(x_1, x_2, \ldots, x_n) = 0 \]

(14)

and an initial point \( p = (u_1, u_2, \ldots, u_n) \) solving the system. At \( p \), we assume that every hypersurface \( f_i \) is regular and twice continuously differentiable, and that the Jacobian matrix

\[ \left( \frac{\partial f_i}{\partial x_j} \right)_{i,j} \]

has rank \( n - 2 \). Then the manifold \( \mathcal{M} \) defined by Eq. (14) is of dimension 2 in the neighborhood of \( p \). Its natural projection into the \((x_1, x_2, x_3)\)-subspace will be denoted by \( \pi(\mathcal{M}) \). We describe how to compute a local approximant to the manifold at \( p \), using the approach of [55,45]. So, we will derive coordinate functions

\[ x_1 = h_1(s, t) \]
\[ x_2 = h_2(s, t) \]
\[ \vdots \]
\[ x_n = h_n(s, t) \]

(15)

such that

\[ p = (h_1(0,0), \ldots, h_n(0,0)) \]

and

\[ f_i(h_1(s,t), h_2(s,t), \ldots, h_n(s,t)) \equiv 0 \]

for \( i = 1, \ldots, m \). Note that the natural projection \( \pi(\mathcal{M}) \) of the manifold \( \mathcal{M} \) into the \((x_1, x_2, x_3)\)-subspace is then locally approximated by

\[ x_1 = h_1(s, t) \]
\[ x_2 = h_2(s, t) \]
\[ x_3 = h_3(s, t) \]

in the vicinity of the projection \( \pi(p) \) of \( p \).

Now the Taylor expansion of the hypersurface \( f_i \) at the point \( p = (u_1, \ldots, u_n) \) is given by
\[ f_i(x_1, \ldots, x_n) = f_i(u_1 + \delta_1, u_2 + \delta_2, \ldots, u_n + \delta_n) = f_i(u_1, \ldots, u_n) + f_i^{(1)} \delta_1 + \cdots + f_i^{(n)} \delta_n + \left[ f_i^{(1,1)} \delta_1^2 + \cdots + f_i^{(n,n)} \delta_n^2 \right]/2 + \cdots + f_i^{(n-1,n)} \delta_{n-1} \delta_n + \text{higher order terms} \] 

(16)

Here, \( f_i^{(j,k)} \) denotes the partial derivative of \( f_i \) by \( x_j \) and \( x_k \).

Next, consider the Taylor expansion of the local coordinate functions, having chosen \( s \) and \( t \) such that \( p = (h_1(0,0), h_2(0,0), \ldots, h_n(0,0)) \):

\[ h_k(s,t) = h_k(0,0) + h_k^{(s)} s + h_k^{(t)} t + [h_k^{(ss)} s^2 + 2 h_k^{(st)} st + h_k^{(tt)} t^2]/2 + \text{higher order terms} \]

where \( h_k^{(s)} \) denotes the partial derivative of \( h_k \) by \( s \), and so on. By assumption, there is a neighborhood of \( p \) in which

\[ f_i(h_1(s,t), h_2(s,t), \ldots, h_n(s,t)) \equiv 0 \]

We set

\[ \delta_1 = h_1^{(s)} s + h_1^{(t)} t + [h_1^{(ss)} s^2 + 2 h_1^{(st)} st + h_1^{(tt)} t^2]/2 + \cdots \]

\[ \delta_2 = h_2^{(s)} s + h_2^{(t)} t + [h_2^{(ss)} s^2 + 2 h_2^{(st)} st + h_2^{(tt)} t^2]/2 + \cdots \]

\[ \vdots \]

\[ \delta_n = h_n^{(s)} s + h_n^{(t)} t + [h_n^{(ss)} s^2 + 2 h_n^{(st)} st + h_n^{(tt)} t^2]/2 + \cdots \]

and note that

\[ \delta_k^2 = (h_k^{(s)})^2 s^2 + 2 h_k^{(ss)} h_k^{(t)} st + (h_k^{(t)})^2 t^2 + \cdots \]

\[ \delta_k \delta_j = h_k^{(s)} h_j^{(s)} s^2 + (h_k^{(s)})^2 h_j^{(t)} st + h_k^{(t)} h_j^{(t)} t^2 + \cdots \]

Now we substitute these quantities into Eq. (16), obtaining a power series in \( s \) and \( t \). The coefficient of each term \( s^a t^b \) in the power series must be zero. In consequence, the following systems of equations are implied:

\[ f_i^{(1)} h_1^{(s)} + f_i^{(2)} h_2^{(s)} + \cdots + f_i^{(n)} h_n^{(s)} = 0 \]  

(17)

\[ f_i^{(1)} h_1^{(t)} + f_i^{(2)} h_2^{(t)} + \cdots + f_i^{(n)} h_n^{(t)} = 0 \]  

(18)
\[ f_i^{(1)} h_1^{(s)2} + \cdots + f_i^{(n)} h_n^{(s)2} = -c_i \] (19)
\[ f_i^{(1)} h_1^{(t)2} + \cdots + f_i^{(n)} h_n^{(t)2} = -d_i \] (20)
\[ f_i^{(1)} h_1^{(s,t)} + \cdots + f_i^{(n)} h_n^{(s,t)} = -e_i \] (21)

where \( i = 1, \ldots, m \). The right hand sides are, respectively,

\[
\begin{align*}
c_i &= f_i^{(1,1)} (h_1^{(s)})^2 + \cdots + f_i^{(n,n)} (h_n^{(s)})^2 \\
&+ 2[f_i^{(1,2)} h_1^{(s)} h_2^{(s)} + \cdots + f_i^{(1,n)} h_1^{(s)} h_n^{(s)}] \\
&+ \cdots + [f_i^{(n-1,n)} h_{n-1}^{(s)} h_n^{(s)}]
\end{align*}
\]

\[
\begin{align*}
d_i &= f_i^{(1,1)} h_1^{(t)} h_1^{(t)} + \cdots + f_i^{(n,n)} h_n^{(t)} h_n^{(t)} \\
&+ f_i^{(1,2)} h_1^{(t)} h_2^{(t)} + \cdots + f_i^{(1,n)} h_1^{(t)} h_n^{(t)} \\
&+ \cdots + f_i^{(n-1,n)} h_{n-1}^{(t)} h_n^{(t)}
\end{align*}
\]

\[
\begin{align*}
e_i &= f_i^{(1,1)} (h_1^{(t)})^2 + \cdots + f_i^{(n,n)} (h_n^{(t)})^2 \\
&+ 2[f_i^{(1,2)} h_1^{(t)} h_2^{(t)} + \cdots + f_i^{(1,n)} h_1^{(t)} h_n^{(t)}] \\
&+ \cdots + [f_i^{(n-1,n)} h_{n-1}^{(t)} h_n^{(t)}]
\end{align*}
\]

The partial derivatives of the coordinate functions \( h_i \) are computed from these systems of linear equations, and define an approximate local parameterization of the manifold given by Eq. (14).

The linear systems of Eqs. (17–21) have rank deficiency 2. Their solutions, therefore, take the form

\[ \alpha_1 \nabla f_1 + \cdots + \alpha_{n-2} \nabla f_{n-2} + \beta t_1 + \gamma t_2 \]

where \( t_1 \) and \( t_2 \) are two linearly independent tangent directions to the surface at the point \( p \). These tangent directions are determined by the method chosen to solve the linear systems.

To choose suitable values for the \( \beta \) and \( \gamma \) coefficients, we proceed as follows. For Eq. (17) and Eq. (18), we let

\[ \alpha_1 = \alpha_2 = \cdots = \alpha_{n-2} = 0 \]

For Eq. (17), we choose \((\beta, \gamma) = (1, 0)\); and for Eq. (18), we choose \((\beta, \gamma) = (0, 1)\). Then the isoparametric curves

\[ (h_1(s,0), \ldots, h_n(s,0)) \quad \text{and} \quad (h_1(0,t), \ldots, h_n(0,t)) \]
intersect transversely at \( p \). Moreover, if \( t_1 \) and \( t_2 \) are orthogonal, then so are the isoparametric curves through \( p \). For Eqs. (19–21) we choose \( \beta = \gamma = 0 \).

We have shown how to construct local linear and quadratic approximants to the manifold \( M \) at \( p \), which also give local approximants to the surface \( \pi(M) \) at \( \pi(p) \). Assuming that the \( f_i \) are sufficiently differentiable, higher-order local approximants are constructed in the same way.

2. CURVATURE DETERMINATION

As shown in [48], it is possible to determine the curvature of the surface \( \pi(M) \) at \( p \). We recall the following

Lemma 3. Let \( n_i \) be the normal vector to the hypersurface \( f_i \) at the point \( p \), for \( 1 \leq i \leq m \). Let \( \alpha \) be such that the last \( n-3 \) components of

\[
\mathbf{n}_0 = \sum_{i=1}^{m} \alpha_i n_i = (a, b, c, 0, ..., 0)
\]

are zero. Then the natural projection of \( \mathbf{n}_0 \) is normal to \( \pi(M) \) at the projection of \( p \).

The normal curvature of \( \pi(M) \) at \( p \) will be determined in a direction that is the natural projection of a tangent \( \mathbf{v} \) to \( M \) at \( p \). The actual computation requires forming a linear combination of the Hessians \( H_i \) of the \( f_i \). It also requires that the vectors \( \mathbf{v} \) and \( \mathbf{n}_0 \) project to vectors of unit length, that is, the sum of squares of the first three components of each vector must be 1. We state how to compute the normal curvature.

Theorem 4. Let \( n_i \) be the normal vector to the hypersurface \( f_i \) at the point \( p \), for \( 1 \leq i \leq m \). Let \( \alpha \) be such that the last \( n-3 \) components of

\[
\mathbf{n}_0 = \sum_{i=1}^{m} \alpha_i n_i = (a, b, c, 0, ..., 0)
\]

are zero, and such that \( a^2 + b^2 + c^2 = 1 \). Let \( \mathbf{v} = (v_1, v_2, ..., v_m) \) be a tangent vector to \( M \) at \( p \) where \( v_1^2 + v_2^2 + v_3^2 = 1 \). Let \( H_i \) be the Hessian of \( f_i \). Then the normal curvature of \( \pi(M) \) at \( p \) in the (projected) direction \( \pi(\mathbf{v}) \) is given by

\[
\kappa = -\mathbf{v}^T (\sum_{i=1}^{m} \alpha_i H_i) \mathbf{v}
\]

It is well-known that the principle curvatures and their directions can be recovered from the normal curvatures in three different directions; e.g., [61]. Thus, the principal curvatures, mean curvature, and Gauss curvature of \( \pi(M) \) can all be determined with help of the theorem.

3. SURFACE INTERSECTION

The derivation of local approximants can be used to construct local approximants to surface intersections as well. As discussed in [54,2], at the point \( p \) we
construct an approximant of the form

\[
  \begin{align*}
  z_1 &= h_1(s) \\
  z_2 &= h_2(s) \\
  &\vdots \\
  z_n &= h_n(s)
  \end{align*}
\]

such that

\[
  p = (h_1(0), \ldots, h_n(0))
\]

The derivation is exactly as in the surface approximations, except that the right hand sides are simpler. Once the approximant has been obtained, to some degree, it is used to derive a new point estimate

\[
  (h_1(s_1), \ldots, h_n(s_1))
\]

which is refined with Newton iteration and becomes the new point at which to construct a local approximant.

This approach to evaluating surface intersections by marching can be very effective. In [40] we show the evaluation of intersections with a variable-radius blend that have an estimated degree in excess of 500. The system defining this curve consists of equations of low to moderate algebraic degree, which is an important aspect of the dimensionality paradigm.

4. GLOBAL APPROXIMATIONS

For curves such as surface intersections it is easy to derive a marching scheme like the one summarized before. In order to obtain a similar scheme for evaluating surfaces we need a device that orients the exploration in space and prevents inadvertently exploring the same neighborhood several times. In [55] this problem has been addressed in the context of the dimensionality paradigm. The technique competes well with other approaches such as Allgower’s simplicial continuation method, [57], or the moving-frame method of Rheinboldt, [62]. All methods are based on the following idea. Given a manifold \( M \) by the system of Eq. (14) and on it a point \( p = (u_1, \ldots, u_n) \), construct a piecewise approximation of \( M \), beginning at \( p \) and extending in all directions.

Rheinboldt’s moving-frame method proceeds by triangulating the tangent space at \( p \), and transferring the triangulation to \( M \) with Newton iteration. Each vertex of the triangulation, after projection to a point \( q \) on \( M \), becomes the center of a new triangulation, of the tangent space at \( q \). The algorithm locally resolves any overlap, but cannot do so globally.

Allgower’s method is based on a triangulation of ambient space and only evaluates the system of equations. At each vertex of a simplex, Eq. (14) is
evaluated and from the \( m \)-vector of function values a linear approximant is constructed. Moreover, it is then deduced through which faces the linear approximant passes, and based on this information adjacent simplices are considered in the same way. Note that adjacent simplices can be found efficiently by a triangulation schema based on an elegant enumeration. In Allgower's method the ambient \( n \)-dimensional space is triangulated. In consequence, the number of simplices in an elementary volume grows exponentially with the number of variables. However, the method does not need to assume differentiability of the \( f_i \).

In the case of surface definitions with the dimensionality paradigm, the surface that is ultimately of interest is the projection \( \pi(\mathcal{M}) \). Since this surface is in a three-dimensional space, it is advantageous to construct an approximant only for the projection, and Chuang's algorithm \[55\] does this as follows, using a grid in 3-space to detect whether a volume of space has already been explored.

1. At \( p \), construct a local approximant to \( \mathcal{M} \) as described before.

2. Determine how the projected approximant, \((h_1(s,t), h_2(s,t), h_3(s,t))\), intersects the faces of the cube, as a function of \( s \) and \( t \).

3. From the intersection curves, determine the coordinates \((s_1, t_1)\) of a point on the approximant that lies in an adjacent cube.

4. Refine the estimated point with Newton iteration.

There is a tradeoff between the degree of the approximant, the mesh size of the grid, and the difficulty of determining face intersections and adjacent points in Steps 2 and 3. With increasing degree of the approximant a coarser mesh can be tolerated, so that fewer approximant calculations are needed. However, determining the intersection with the faces of the current cube becomes more difficult. The advantage of this method is the fact that the dimension of the meshed space does not depend on the number of variables used to define \( \mathcal{M} \). Yet, by determining the \((s,t)\) curves, each estimate \((s_1, t_1)\) can be pulled back into the \( n \)-space in which \( \mathcal{M} \) is given.

We illustrate the method for linear approximants. Assume that we are at a point \( p = (u_1, u_2, \ldots, u_n) \) on \( \mathcal{M} \) that projects to \( \pi(p) \) in a cube, and we have constructed the linear approximant

\[
L: \quad z_1 = u_1 + u_1 s + w_1 t \\
z_2 = u_2 + u_2 s + w_2 t \\
\vdots \\
z_n = u_n + v_n s + w_n t
\]

\( L \) intersects the faces of the cube containing \( \pi(p) \), say as shown in Figure 22. Each intersection is easily determined: If the face plane is \( z_1 = a \), then the
line in \((s, t)\)-space corresponding to the intersection with \(L\) is simply

\[v_1s + w_1t = a - u_1\]

Intersections with the planes \(y = b\) and \(z = c\) are found analogously. Thus, the intersections define a polygon in \((s, t)\)-space that corresponds to the area of \(L\) contained in the cube, as shown in Figure 23. Possibly with help of additional lines corresponding to the intersection of \(L\) with the faces of adjacent cubes, we can find good estimates \((s_1, t_1)\) for new points in neighboring cells, as illustrated in Figure 21.

Note that the linear approximants found at each point do not have \(C^0\) continuity. Assuming that the mesh is sufficiently fine, a simple solution is to triangulate the points found on \(\pi(M)\). Since we have good adjacency information from the construction, no ambiguities have to be resolved. Other possibilities include determining the principal curvatures and directions at each point on \(\pi(M)\). Thereupon, interpolation algorithms such as [63] could be used to provide a surface approximant that has \(C^1\) continuity.
V. THE SKELETON

The interior skeleton of a three-dimensional solid is the locus of the centers of all inscribed maximal spheres. A sphere is maximal if there is no other inscribed sphere that contains it completely. Other names for the skeleton include medial-axis transform and symmetric-axis transform.

Computer vision has proposed the skeleton as a shape representation; e.g., [64,65]. Applications of the skeleton in pattern recognition are discussed; e.g., in [64,66]. In the context of geometric modeling applications, it has been argued that the skeleton in 2D is useful for generating finite-element meshes [67,68,69], because it allows identifying areas in which the 2D domain is constricted or extended, and so provides the basis for quantifying shape parameters of relevance to the mesh density.

If we know for each skeleton point its distance from the boundary, then the skeleton is an informationally-complete representation in the sense of Requicha [70]. We discuss this representation now, and then consider some of the applications that might be facilitated by it. For the sake of a clear presentation, we begin with a discussion of the skeleton of two-dimensional objects. We then point out how to generalize the techniques to three-dimensional solids.

There is an interesting connection between skeletons and certain concepts from classical geometry and from the theory of differential equations, with consequences on how to compute the skeleton, [71]. We will discuss these connections as well.

A. THE SKELETON REPRESENTATION

Consider a bounded solid $T$, in $\mathbb{R}^2$ or $\mathbb{R}^3$, with compact boundary $\partial T$. For every point $p$, we define its distance from the boundary of $T$ as the minimum
Euclidean distance $d(p, q)$ where $q$ is on $\partial T$. For every $p$, there is always at least one point $q$ on the boundary of $T$ such that $d(p, q)$ is minimum, and such a point $q$ will be called a foot point of $p$ on $T$. Then the interior skeleton of $T$ consists of all points $p$ that are interior points of $T$ and have more than one footpoint on $\partial T$, as well as the limits of point sequences in this set. That is, the skeleton is the relative closure of the set of all interior points that do not have unique footpoints. An example in two dimensions is shown in Figure 25. Similarly, the exterior skeleton of $T$ is the closure of all points $p$ that are exterior to $T$ and have more than one footpoint.

We associate with each skeleton point its distance from $\partial T$, as an additional coordinate. For two-dimensional solids, therefore, the skeleton point $(u, v)$ at distance $r$ from $\partial T$ will be associated with $(u, v, r)$ in 3-space. The skeleton thus becomes a three-dimensional object, as shown in Figure 26. We recover the boundary $\partial T$ from this skeleton representation as follows. Associate with the point $(u, v, r)$ of the skeleton the circle $(x - u)^2 + (y - v)^2 - r^2 = 0$. Then $\partial T$ is the envelope of all such circles. We will see later that the envelope points of the circle can be determined without constructing the envelope.

Figure 25: Interior Skeletons of a Simple Domain

Figure 26: The Skeleton as 3D Object
Likewise, the skeleton point \((u, v, w)\) at distance \(r\) of the three-dimensional solid \(T\) is represented by the point \((u, v, w, r)\) in 4-space, thus considering the skeleton a four-dimensional structure. Then \(\partial T\) is the envelope of the spheres 
\[(x - u)^2 + (y - v)^2 + (z - w)^2 - r^2 = 0.\]

Since the \(r\)-coordinate gives the distance from the boundary, we expect that a translation of the skeleton in the \(r\)-direction represents an interior or exterior offset of \(T\). More precisely, with \(d\) a signed translation distance, let the \(d\)-translate of the skeleton \(S\) consist of the points 
\[\{(u, v, w, r + d) \mid (u, v, w, r) \in S, r + d \geq 0\}\]

Then a negative \(d\)-translate represents the interior \(d\)-offset of \(T\), whereas, in the case of positive \(d\)-translates, self-intersections are possible when \(T\) is not convex. See also Figure 27.

The possibility that the boundaries of positive translates may contain self-intersections suggests that the interior skeleton does not contain sufficient information to generate offsets in a simple way. In fact, exterior offsets will require the exterior skeleton as well. Briefly, a positive translate of the interior skeleton is combined with a negative translate of the exterior skeleton, by the same distance. Then the exterior skeleton points that are clipped by the rule \(r + d \geq 0\) identify self-intersections.

B. CYCLOGRAPHIC MAPS AND IMAGES

The concept of cyclographic maps is due to Müller [72], and has uses in descriptive geometry. Cyclographic maps provide a different conceptualization for the skeleton, and we explore this now in detail.
We consider oriented circles in the \((x, y)\)-plane, calling them *cycles* so as to stress that they are oriented. The plane is considered in \((x, y, z)\)-space. A counter-clockwise orientation is *positive*, as seen from points with positive \(z\) coordinates, and a *negative* orientation is clockwise. If a cycle has the center \((u, v)\) and radius \(r\), we associate with it the point \((u, v, r)\) in 3-space, where \(r\) is signed according to the cycle orientation. This association defines a bijection

\[ \Phi : \mathbb{R}^3 \leftrightarrow C \]

between points in 3-space and cycles in the \((x, y)\)-plane. Note that the points in the plane correspond to cycles of zero radius.

Let \(C\) be an oriented curve in the plane. We investigate which cycles are tangent to \(C\), requiring that the cycles and the curve are consistently oriented, as shown in Figure 28. Let \(p\) be any point on \(C\). Then all cycles tangent to \(C\) at \(p\) have their centers on the normal to \(C\) and are at distance equal to the cycle radius. Hence, the points in 3-space to which those cycles correspond are on a line that is inclined to the \((x, y)\)-plane by 45° and orthographically projects onto the normal of \(C\) at \(p\). The set of all oriented circles tangent to the curve \(C\) therefore corresponds to a ruled surface in 3-space all of whose generators intersect \(C\), project onto the normals of \(C\), and have slope 1 against the \((x, y)\)-plane. This surface is the *cyclographic map of \(C\).*

In general, since all generators of the cyclographic map have equal slope, the ruled surface is developable [72]. In particular, if \(C\) is a line, then the cyclographic map is an inclined plane, and if \(C\) is a circle, then the cyclographic map is a right circular cone with a right angle at the vertex.

Now consider the reverse process, of investigating the cycle geometry that corresponds to a given curve in space. Intuitively, the family of cycles corresponding to the points in space will have an envelope, although the envelope could be imaginary. We call this envelope the *cyclographic image* of the space curve.

In the simplest case, the space curve is a line \(\ell\), and each point on this line maps to a cycle. If \(\ell\) is parallel to the \((x, y)\)-plane, then the cycles have equal radius and their common envelope is a pair of parallel lines in the \((x, y)\)-plane. If \(\ell\) is inclined, with a slope less than 45°, then the cycles are enveloped by a
pair of intersecting lines. The intersection is the point at which \( \ell \) intersects the \((x, y)\)-plane. If the line \( \ell \) is at an angle of 45°, then the envelope is a single line perpendicular to the projection of \( \ell \) onto the \((x, y)\)-plane. For greater inclination angles, the cycles have no real envelope. See also Figure 29.

If we choose a space curve \( C \) other than a line, then to each point \( p \) of \( C \) the corresponding cycle contributes two points to the cyclographic image which are real and distinct if the tangent at \( p \) has slope less than 45°, coincident if the slope is 45°, and conjugate complex otherwise. Moreover, the orthogonal projection of the space curve \( C \) onto the \((x, y)\)-plane bisects the envelope curve, and so \( C \) is the skeleton of the cyclographic image. Furthermore, it can be proved that the tangent to \( C \) at \( p \), and the tangents to the cyclographic image of \( C \), at the points contributed to the envelope by the cycle corresponding to \( p \), are concurrent, as shown in Figure 30. Note that this fact implies a direct construction of the footpoints on the boundary from each skeleton point.

There are a number of interesting relationships between the space curve \( C \) and its cyclographic image that are discussed in [72]. For example, the tangents to \( C \) form the cyclographic map of the cyclographic image. Furthermore, the edge of regression of the cyclographic map projects unto the evolute of the cyclographic image.

Consider the cyclographic map of the boundary \( \partial T \) of a 2D solid \( T \). We will trim this map as follows: For every point \( q = (u, v) \) in the \((x, y)\)-plane, consider the normal of the plane through \( q \). This line intersects the cyclographic map in a number of points. If \( q \) is interior of \( T \), then we choose among the intersection
Figure 31: The Distance Function $S$ Defined by the Cyclographic Map

points the one that lies above the $(x, y)$-plane and has the smallest $z$ coordinate. If $q$ is exterior to $T$, then we choose the intersection point below the $(x, y)$-plane with the largest $z$-coordinate. In either case, we have associated a point that lies on that line of the cyclographic map that goes through the nearest foot point of $q$ on $\partial T$. With this convention, we have defined a single-valued function $S$ whose domain is the $(x, y)$-plane and whose range is a subset of the cyclographic map of $\partial T$. Note that $S$ is zero only on the boundary of $T$. We will see later that this function is characterized by the Hamilton-Jacobi equation. Note that $S$ defines the minimum Euclidean distance of every point of the $(x, y)$-plane to the boundary of $\partial T$. Clearly, its singular curves are the skeleton of $T$. See Figure 31 for an example.

Since the $S$ function is the Euclidean distance function, we observe that it represents all interior and exterior offsets as follows: In order to obtain the boundary of the interior $d$-offset of $T$, intersect $S$ with the plane $z = d$. The boundary of the exterior $d$-offset is given similarly by the intersection of $S$ with the plane $z = -d$. See also Figure 31.

This construction of the offsets interprets why the interior offset can be obtained as the positive $d$-translate of the interior skeleton. The interior skeleton characterizes $S$ in the interior of $T$, but does not reflect the exterior structure of $S$ completely. Thus, the exterior skeleton must be translated to obtain exterior offsets without self-intersection.

It has been pointed out in [73] that a solid can be blended by a succession of interior and exterior offsets, by the rounding radius. The exterior offset of the interior offset obtains all rounds, and the interior offset of the exterior offset obtains all fillets. It is clear, however, that the translation of the $(x, y)$-plane first by $d$ and then by $-d$ recovers the original boundary $\partial T$. It follows that the trimmed cyclographic map of an offset is not equal to the translation of the cyclographic map of $\partial T$. In the construction of interior offsets as negative $d$-translates of the skeleton, this fact finds its counterpart in the clipping effected by the condition $r + d \geq 0$. 
In analogy to the two-dimensional case, we can define cyclographic maps that establish a correspondence between oriented spheres in 3-space, and the points of a four-dimensional space. The map of a solid boundary $\partial T$ is now a curved space that is ruled, and the ruling consists of lines, through every point of $\partial T$, that are inclined against $(x,y,z)$-space by $45^\circ$ in the containing 4-space. Furthermore, the lines project onto the normals in $(x,y,z)$-space of the boundary of $T$. From differential geometry we know that the normals in 3-space envelope the focal surfaces of $\partial T$, [74], and the various constructions and properties outlined above for the two-dimensional case generalize virtually unchanged to the three-dimensional case [75]. In particular, the skeleton of three-dimensional domains is the singular set of the trimmed cyclographic map.

C. THE HAMILTON-JACOBI EQUATION

As we discussed before, the function $S$ is a subset of the cyclographic map, which, in turn, is ruled with each generator inclined $45^\circ$ against the $(x,y)$-plane, or against the $(x,y,z)$-space in the case of 3-dimensional solids $T$. Denoting the distance variable with $t$ for the moment, we can express the geometry of the ruling by the equation

$$S_t^2 = S_x^2 + S_y^2$$

for two-dimensional solids, and by

$$S_t^2 = S_x^2 + S_y^2 + S_z^2$$

for three-dimensional solids. Since we can locally express $t$ as an explicit function of the other variables, the equations become

$$S_x^2 + S_y^2 = 1$$

and

$$S_x^2 + S_y^2 + S_z^2 = 1$$

respectively, but this is just the well-known Hamilton-Jacobi equation. The function $S$, therefore, satisfies this differential equation with the boundary condition

$$S = 0 \text{ on } \partial T$$

With proper sign conventions, it is clear that the Hamilton-Jacobi equation defines the cyclographic map $S$, and, because of the distance interpretation of $t$, $S$ defines the Euclidean minimum distance from $\partial T$ of any point in $(x,y)$- and $(x,y,z)$-space. This fact has been noted before; [76].

We have identified the skeleton as the locus of points at which the cyclographic map is derivative discontinuous. In terms of the Hamilton-Jacobi equation, this set of points is called a shock wave; e.g., [77]. Thus, if we have a PDE solver for integrating the Hamilton-Jacobi equation that locates the
shocks in the solution, it follows that such a solver also finds the skeleton of \( T \). Integration inward from \( \partial T \) evaluates the interior skeleton, whereas integration outward from the boundary evaluates the exterior skeleton.

At the time of writing, we have only limited experience with this approach towards computing the skeleton. In particular, we have implemented the scheme described in [78]. Early experiments seem to indicate that the solver is more general than necessary. More specialized solvers should be faster and more accurate. We are experimenting with alternatives that integrate the equation as an initial value problem, augmented by a correcting step that refines the solution based on the dimensionality paradigm.

D. GEOMETRIC APPROACHES TO COMPUTING THE SKELETON

Early approaches to compute the skeleton in two dimensions have evaluated an approximate distance from the boundary, by one or two passes over a discretization of \( T \). Examples include [66,79].

Computational geometers have investigated the skeleton of polygonal domains. The algorithm by Preparata [80] evaluates first the branching points of the skeleton. Considering certain triples of edges, the associated branch point is the intersection of the bisectors of the edge pairs. Careful sequencing of triples yields an \( O(n^2) \) algorithm for convex polygons. The sequencing can be conceptualized as constructing the interior offsets that contain a branching point. Then a branch point signals that certain edges of the boundary no longer contribute to the offset. An algorithm by Lee [81] constructs the skeleton using a divide-and-conquer approach, and achieves \( O(n\log(n)) \). Preparata’s algorithm was extended by Patrikalakis and Gürsoy to domains bounded by line segments and circular arcs, [68]. As in the polygonal algorithm, branch points are determined but are now the intersection of conics.

Using some conceptual elements of Preparata’s and the Patrikalakis-Gürsoy approach, [82] gives an algorithm for constructing the skeleton of CSG solids in 3-space. The algorithm computes nearest approach points between pairs of elements of the boundary, but not between triples, for that would entail algebraic computations that are of unattractive complexity. The closest approach points found are sorted by their distance from the boundary. In this phase a proximity computation also rejects those closest approach points that are nearer to a third boundary element.

Beginning at the closest approach points, the skeleton is evaluated, by increasing distance from the boundary. At each moment throughout the computation, the skeleton is known up to a current distance. Closest approach points at a greater distance are not processed until their respective distance has been reached. Each face, edge, and vertex of the skeleton is formulated as a suitable set of nonlinear equations, using the dimensionality paradigm, and is
evaluated with help of the approximation algorithm described in Section IV.G. In particular, the grid used to orient the exploration of faces and edges also signals that different skeleton faces are about to intersect.

Intersection of two faces of the skeleton are easily formulated and evaluated. Each face involves two boundary elements from which its points are equidistant. Their intersection is thus obtained by seeking all points equidistant from all boundary elements involved in the definition of the intersecting faces, and only requires combining the equation systems that describe the faces. A proximity computation determines how to trim faces in the vicinity of an edge or a vertex.

The algorithm does not attempt to exploit special geometric configurations of boundary elements. As discussed in [49], many such configurations exist and result in skeleton faces that have a special geometry that could be utilized in the construction of the face.

Bowyer et al. [83] describes an approximate skeleton algorithm for three-dimensional solids constructed from algebraic half spaces of arbitrary degree. The algorithm begins by superimposing a grid whose elements are classified as one of inside, outside, or boundary. Boundary elements intersect the solid boundary, inside elements are in the solid interior, and outside elements are exterior to the solid. Next, for all nonboundary elements, a distance computation determines the approximate minimum distance from the boundary. Those elements that are closest to more than one boundary element constitute the approximate skeleton. A number of heuristics are employed that narrow down the search for closest boundary elements. That is, for each boundary element, the cone of normals to the contained boundary area is determined, since interior elements that do not fall into this cone cannot be nearest to this boundary element. The algorithm has been implemented and incorporated into the Bath solids modeler by David Lavender. Note that the approach generalizes to solids of any dimension.

Price et al. [84] gives an algorithm for determining the branching points of the skeleton of arbitrary planar domains. First, it is observed that a Delaunay triangulation can be used to approximate the skeleton as follows: Select a large number of points from the domain boundary, and construct their Delaunay triangulation.\footnote{See, e.g., [85] for an algorithm to construct Delaunay triangulations.} If the points are sufficiently dense, the triangulation will respect the domain boundary. Moreover, since in a Delaunay triangulation the circumscribed circle of every triangle contains no other triangle vertices in the interior, the skeleton is approximated by the centers of the circumscribing circles.

Price et al. consider how to select a small number of boundary points such that the resulting Delaunay triangulation respects the domain boundary and such that the centers of the circumscribed circles contain all branch points of the skeleton. Several heuristics are described that make the compatibility
test more efficient. In particular, by classifying the types of triangles, it is shown that many triangles need not be tested if certain critical ones respect the boundary. Exact branch points are found later using Newton iteration. The algorithm has been implemented. Note that the approach is relatively insensitive to the types of curves bounding the domain. The ideas and heuristics should generalize to the three-dimensional case. The method is used as basis of a finite-element mesh generation algorithm [67].

E. FINITE-ELEMENT MESH GENERATION

In contrast to grid generation methods like the one we discussed earlier in Section III.D, finite-element mesh generation discretizes a domain using an irregular mesh of simple shapes, often triangles or tetrahedra. Automating finite-element mesh generation is of interest because the meshing requires much effort and considerable manual labor at present. The suitability of the mesh depends on both the geometry of the domain as well as on the nature of the physical problem and its solution. It is therefore important that the mesh not only satisfy geometric criteria, but also that it can be refined iteratively in response to the analysis results.

Systematic finite-element mesh generation has been based on three fundamental approaches:

1. Impose a spatial subdivision, say using octrees. Subdivide the interior cells in a standard way into elements, and do special processing near the boundary; see, e.g., [66].

2. Sample the domain boundary computing a set of points. Construct a Delaunay triangulation of those points, and refine it such that the elements respect the boundary; see, e.g., [87,88].

3. Partition the domain by its skeleton and some additional edges or faces into simple subdomains. Triangulate the subdomains in a way that is compatible at the boundaries between them; e.g., [67,68,69].

It is argued in [69] that Approach (1) does not address the essence of the geometric problem and merely postpones it to those spatial cells that intersect the boundary. However, when suitably subdivided, the octree cells containing parts of the boundary have a relatively simple structure.

Approach (2) is attractive in 2D where favorable aspect ratios of the triangles can be guaranteed. However, no similar guarantees exist in the 3D case, as pointed out in [67].

We will sketch first the algorithm of [69], for generating meshes of polygonal domains. The algorithm uses Approach (3), and recognizes automatically where the domain is constricted. In those areas it is usually the case that the mesh needs to be finer than elsewhere.
In the first stage, the algorithm determines the skeleton of the polygonal domain, and augments it with the shortest paths from certain skeleton points to the boundary. More precisely, at every point of the skeleton where the associated boundary entities change, the shortest paths to the boundary are added. Moreover, the parabolic arcs of the skeleton are replaced with one or two chords according to whether the angle subtended is smaller or greater than 90°. If the arc is replaced with two chords, then the shortest paths from the common chord vertex to the boundary are also added. Figure 32 shows a domain with its skeleton and the result of the first stage on the left. Note that the first stage constructs a domain partition into convex subdomains.

In the second stage, the initial partition is changed by eliminating regions that are very narrow (sinks), and by subdividing extended regions bounded by relatively long skeleton edges. Note that these types of region are quantitatively defined in terms of the types and relative lengths of bounding edges [69]. Figure 32 shows the result of this stage on the right.

Next, the partition of the second stage is meshed. Depending on the local geometry and the type of finite-element analysis, some edges bounding subdomains may be subdivided by additional nodes. The resulting mesh optionally may be subjected to Laplace smoothing or other heuristics, producing a final result such as shown in Figure 33.

Armstrong et al. [67] describe a different algorithm for generating meshes of planar domains. The approach is also based on the skeleton, but quadrilateral elements are generated rather than triangular ones. The skeleton is determined by the algorithm of [84]. After the skeleton has been determined, the induced partition of the domain is refined as follows. A concave boundary corner is subdivided by an additional line so as to obtain angles close to 90°, except when the corner is close to 180°. Skeleton edges that lie between complex
branch points are also subdivided in an effort to obtain four-sided regions in the partition. The regions are then meshed with quadrilateral elements. See also Figure 34.

F. ON GEOMETRIC TOLERANCING

In many applications, a solid designed for manufacture need not have the exact shape defined by the geometric model. Rather, shapes are allowed to vary within certain tolerances. Expressing these tolerances can be extremely complex, because they may depend on the function of the solid part, and its interrelationship to other parts. Geometric tolerancing studies the subproblem of how to express the geometric variation allowed; e.g., [89].

In the simplest case, one could view the surface of the solid design as a nominal shape, and accept all solids with a shape whose surface is within a
zone of constant thickness surrounding the surface of the nominal shape. See Figure 35 for a two-dimensional example. This appealingly simple idea entails some difficulties in traditional solid representations in that the measurement of distance from a curved boundary is not necessarily a simple computation.

By using a skeleton representation, however, such a distance computation simplifies dramatically. For, since dilations and contractions of shapes correspond to translates of the skeleton in the distance dimension (Section V.A), the point-boundary distance can be measured by comparing corresponding skeleton points in their last coordinate. Also, the function S discussed before gives a precise distance measurement, so that evaluating this function on the boundary of the variational shape at once reveals by how much the surfaces deviate.

ACKNOWLEDGEMENTS

We are indebted to H. Pottmann for bringing the work on cyclographic maps to our attention, and to R. Lynch, who noted the connection between offsets and the Hamilton-Jacobi equations. The discussions we have had with both on this subject have been most helpful.

References


