Adaptive enumeration of implicit surfaces with affine arithmetic

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Abstract

We discuss adaptive enumeration and rendering methods for implicit surfaces, using octrees computed with affine arithmetic, a new tool for range analysis. Affine arithmetic is similar to standard interval arithmetic, but takes into account correlations between operands and sub-formulas, generally providing much tighter bounds for the computed quantities. The resulting octrees are accordingly much smaller, and the rendering faster.

We also describe applications of affine arithmetic to intersection and ray tracing of implicit surfaces.

KEYWORDS: cellular models, interval analysis, rendering, implicit surfaces.

1 Introduction

Implicit surfaces have recently become popular in computer graphics and solid modeling. In order to exploit existing hardware and algorithms, it is often necessary to approximate such surfaces by models with simpler geometry, such as polygonal meshes or voxel arrays. Therefore, it is important to find efficient approximation methods for implicit surfaces.

Let \( S \) be a surface defined implicitly by the equation \( h(x, y, z) = 0 \). A simple and general technique for computing an approximation of \( S \) in a region \( \Omega \) is:

1. decompose \( \Omega \) into small cells;
2. identify which cells intersect \( S \);
3. approximate \( S \) within each intersecting cell.

The enumeration of the intersecting cells is usually the most expensive step in this method. In the simplest schema, the cells that intersect \( S \) are identified by sampling. The function \( h \) is evaluated at the vertices of each cell; if the signs of those values are not identical, then the cell necessarily intersects \( S \). Obviously, the converse does not hold: if

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the values have all the same sign, we cannot conclude that the cell does not intersect $S$. This is a form of aliasing in the sampling related to the size of the cell. Therefore, cell sizes must be carefully chosen to avoid missing features due to undersampling.

Typically, the cellular decomposition of $\Omega$ is a regular grid of cubes. If there are $n$ cubes along each main direction, then there are $n^3$ cubes to be scanned in a full enumeration, but only $O(n^2)$ cubes will intersect $S$. Thus, choosing a smaller cell size to avoid aliasing will greatly increase the number of cells to be scanned, and also increase the fraction of “useless” tests. Therefore, this approximation method is simple but not efficient.

Figure 1 shows a full enumeration of the curve defined implicitly by $y^2 - x^3 + x = 0$, in the square $\Omega = [-2 .. 2] \times [-2 .. 2]$, using a $16 \times 16$ grid. Intersecting cells were identified by sampling and appear in grey (note how few they are: only 44 out of 256). Black dots mark the points where the curve crosses cell edges; they can be joined to form a polygonal approximation to the curve.

![Figure 1: full enumeration of the curve given implicitly by $y^2 - x^3 + x = 0$](image)

In Section 2, we review some general methods that try to find the cells that intersect $S$ without enumerating all cells in $\Omega$. The most reliable of those seems to be recursive subdivision of space based on range analysis, i.e., on estimates for the range of values taken by $h$ on subsets of $\Omega$. Interval arithmetic is the natural technique for range analysis. However, as we point out in Section 3, the excessive conservatism of interval arithmetic may greatly reduce the efficiency of the search. In Section 4, we describe affine arithmetic, a computation technique that generally provides much tighter bounds than interval arithmetic. In Section 5, we describe some experiments that exploit the properties of affine arithmetic to efficiently enumerate implicit surfaces. In Section 6, we show how to render implicit surfaces directly, during enumeration, without explicitly creating a cellular model. In Section 7, we discuss other applications of affine arithmetic for computing implicit objects. Finally, Section 8 contains some conclusions and outlines directions for future work.
2 Efficient enumeration methods

Several methods exist for finding the cells intersecting a surface $S$ without visiting all cells in a cellular decomposition of a region $\Omega$. The goal is to avoid testing all $O(n^3)$ cells, since only $O(n^2)$ cells intersect $S$.

The most popular efficient enumeration methods are still based on sampling. However, hierarchical decomposition methods are more efficient, more robust, and also suitable for parallel implementation. In this section, we review some of these methods.

2.1 Sampling-based methods

Continuation methods sample the surface only in the immediate neighbourhood of known intersecting cells [3, 4, 1, 6]. These methods only work for cellular subdivisions of $\Omega$ whose topology and geometry are well understood [6], so that it is simple to advance from one intersecting cell to an adjacent one, using “pivoting” techniques [1].

Adaptive refinement techniques start with a coarse cellular subdivision, and further subdivide only those cells that do intersect $S$ [3, 10, 21]. Such techniques frequently use estimates of the curvature of $S$ in a cell as an additional subdivision criterion, thus building a cellular subdivision that is adapted not only to the position of $S$ inside $\Omega$, but also to the intrinsic geometry of $S$. Polygonal approximations based on such subdivisions are thus quite efficient.

All these methods based on sampling have one fundamental difficulty, besides aliasing: finding a set of initial “seed” cells intersecting each connected component of $S$ in $\Omega$. (Unlike parametric surfaces, implicit surfaces can have several connected components; see Figure 1.) Obviously, if $S$ is a general implicit surface, and we are restricted to evaluating $h$ at individual points, then this problem has no satisfactory solution. Even if we assume that $S$ has no extended features thinner than the minimum cell size (so that there are no aliasing problems), we still must sample the whole region $\Omega$ uniformly at that resolution; otherwise, we may miss some component of $S$.

2.2 Hierarchical decomposition methods

In order to discard large portions of $\Omega$ quickly and reliably, we need a more powerful test than point sampling. The latter can only prove the presence of $S$ in some region of $\Omega$; to reduce the number of cells scanned, we need a test procedure that can also prove the absence of $S$ in a region.

The hierarchical decomposition methods rely on such a test to explore $\Omega$ recursively, starting with $\Omega$ itself as the initial cell. If a cell is proved to be empty, it is ignored; otherwise, it is subdivided into smaller cells, which are then explored recursively, until the cells are small enough to approximate $S$ [18, 19, 7, 17, 20].

The meaning of “small enough” depends on the application. For rendering, it might mean “smaller than a pixel”. For other applications, such as modeling, it may depend on some other numerical criterion. For instance, testing how closely the surface can be approximated by a linear function inside the cell allows polygonal approximations to adapt to the curvature of the surface.
Note that the test procedure is not required to be complete, in the sense that it may fail to prove either the presence or the absence of \( S \) in a given cell. In particular, a cell that is declared “small enough” may still have unknown status. Each application must decide what to do with those “indeterminate” cells: discard them, treat them just like the cells that do intersect \( S \), or handle them in some special way. Point sampling may be useful at this stage to help identify some intersecting cells.

The subdivision of \( \Omega \) resulting from a hierarchical decomposition is not a regular grid, but rather a tree of nested cells. The leaves in this tree form a cellular model for \( S \). There are many variants of this method; they differ in the shape of cells and the method of subdivision. In particular, if the cells are cubical and divided into eight equal parts, then the resulting subdivision is called an octree [16]. Other choices give binary space partitions [8], 3-d trees [2], hierarchical triangulations [15], and many more.

### 3 Interval arithmetic

In order to prove that a given cell \( C \) does not intersect \( S \), we must prove that the function \( h \) does not vanish inside \( C \). For some classes of surfaces, this fact can usually be established by specific tests. For instance, if \( h \) is a polynomial, we can compute its Bézier-Bernstein coefficients for that cell: if they are all positive, or all negative, then the same will be true of \( h \). Unfortunately, this method cannot be used for general (non-polynomial) functions; or even for polynomials of high degree presented in factored form, such as \( (x^2 + y^2 + z^4)^{20} - 1 \).

Range analysis can be used to provide general test procedures for hierarchical decompositions. In range analysis, an estimate is computed for the whole range of values taken by the function \( h \) on the points of a cell \( C \). This estimate is an interval guaranteed to contain \( h(C) \). If this interval does not contain zero, then the function \( h \) cannot vanish inside \( C \) and the test has proved that the cell does not intersect \( S \). However, if the interval does contain zero, we cannot conclude that the cell intersects \( S \) because estimates are not required to be exact.

The classical technique of interval arithmetic (IA), also known as interval analysis, provides a natural tool for range analysis. In IA, each quantity is represented by an interval of floating-point numbers. Those intervals are added, subtracted, multiplied, etc., in such a way that each computed interval is guaranteed to contain the (unknown) value of the quantity it represents [14].

To test whether a cell \( C \) intersects the surface \( S \), we can evaluate \( h(x, y, z) \) with IA, letting \( x \), \( y \) and \( z \) be the projections of the cell onto the coordinate axes. The interval thus computed will contain all values of \( h \) for points inside the cell. If this interval is entirely positive, or entirely negative, then we have proved that \( C \) does not intersect \( S \).

Hierarchical decomposition methods based on IA have recently been proposed for the enumeration of implicit surfaces in computer graphics applications [19, 7, 17]. Those methods have become quite popular, due to their ability to handle arbitrarily complex non-polynomial surfaces, and their immunity to round-off errors.

Figure 2 shows a hierarchical enumeration based on IA of the curve shown in Figure 1, but now using a \( 32 \times 32 \) grid. Note that large portions of \( \Omega \) were discarded at early stages.
Section 5 contains other examples.

Figure 2: hierarchical enumeration of $y^2 - x^3 + x = 0$

3.1 The error explosion problem

The main weakness of IA is that it tends to be too conservative: the computed interval for a quantity may be much wider than the true range of that quantity, often to the point of uselessness. This over-conservatism is mainly due to the assumption that the (unknown) values of the arguments to primitive operations may vary independently over the given interval. If there are any mathematical constraints between these arguments, then not all combinations of values in the corresponding intervals will be valid. In that case, the result interval computed by IA may be much wider than the true range of the result quantity.

For example, consider evaluating $x(10 - x)$, where $x$ is known to lie in the interval $[4, 6]$. Applying the IA formulas blindly, we get

\[
10 - x = [10, 10] - [4, 6] = [4, 6]
\]
\[
x(10 - x) = [4, 6] \cdot [4, 6] = [16, 36],
\]
which is 20 times wider than the true range of the expression $x(10 - x)$ over $[4, 6]$, namely $[24, 25]$. The large discrepancy between the two intervals is due to the inverse relation between the quantities $x$ and $10 - x$, which is not known to the IA multiplication algorithm.

The over-conservatism of IA is particularly bad in long computation chains, where the intervals computed by one stage of the chain are the inputs to the following stage. In such cases, one often observes an "error explosion": as the evaluation advances down the chain, the relative accuracy of the computed intervals decreases exponentially, and they soon become too wide to be useful, by many orders of magnitude. Unfortunately, long computations chains are not uncommon in computer graphics applications.
4 Affine arithmetic

To address the “error explosion” problem in IA, Comba and Stolfi [5] proposed a new model for numerical computation, called affine arithmetic (AA). Like standard IA, affine arithmetic keeps track automatically of the round-off and truncation errors affecting each computed quantity. Unlike IA, however, affine arithmetic keeps track of correlations between those quantities. Thanks to this extra information, AA is able to provide much tighter intervals than IA, especially in long computation chains.

The key feature of AA is an extended encoding of quantities from which one can determine, in addition to their ranges, also certain relationships to other quantities — such as the ones existing between $x$ and $10 - x$ in the example. Specifically, a partially unknown quantity $x$ is represented in AA by an affine form $\hat{x}$, which is a first-degree polynomial:

$$\hat{x} = x_0 + x_1 \varepsilon_1 + x_2 \varepsilon_2 + \cdots + x_n \varepsilon_n.$$ 

Here, the $x_i$ are known real coefficients (stored as floating-point numbers), and the $\varepsilon_i$ are symbolic variables whose values are unknown but assumed to lie in the interval $U = [-1 .. +1]$.

Each $\varepsilon_i$ is called a noise symbol: it stands for an independent source of error or uncertainty that contributes to the total uncertainty of the quantity $x$; the corresponding coefficient $x_i$ gives the magnitude of that contribution. The source of error may be external (due to original uncertainty in some input quantity) or internal (due to round-off and truncation errors committed in the computation of $\hat{x}$).

A somewhat similar approach has been proposed by Hansen [11], in which quantities are represented instead by affine combinations of a fixed number of intervals. In AA, new noise symbols are dynamically created during a long computation (Section 4.3).

As one may expect, affine arithmetic is more complex and expensive than ordinary interval arithmetic. However, its higher accuracy is worth the extra cost in many applications, including adaptive enumeration of implicit objects, as we show in Section 5.

4.1 Conversions between IA and AA

If $\hat{x} = x_0 + x_1 \varepsilon_1 + \cdots + x_n \varepsilon_n$ is an affine form for a quantity $x$, then the value of $x$ is guaranteed to be in the range of $\hat{x}$, which is the interval

$$[\hat{x}] = [x_0 - \xi .. x_0 + \xi], \quad \xi = \sum_{i=1}^{n} |x_i|.$$ 

Note that $[\hat{x}]$ is the smallest interval that contains all possible values of $\hat{x}$, assuming that each $\varepsilon_i$ ranges independently over the interval $U = [-1 .. +1]$.

Conversely, given an interval $\bar{x} = [a .. b]$ representing some quantity $x$ in IA, an equivalent affine form for the same quantity is given by $\hat{x} = x_0 + x_k \varepsilon_k$, where

$$x_0 = \frac{b + a}{2} \quad \text{and} \quad x_k = \frac{b - a}{2}.$$
The noise symbol $\varepsilon_k$ symbolizes the uncertainty in the value of $x$. Since the interval $\bar{x}$ tells us nothing about possible constraints between the value of $x$ and that of other variables, $\varepsilon_k$ must be distinct from all other noise symbols previously used in the same computation.

### 4.2 Exploiting correlations in operands

The key feature of AA is that the same noise symbol $\varepsilon_i$ may contribute to the uncertainty of two or more quantities (inputs, outputs, or intermediate results) arising in the evaluation of an expression. The sharing of a noise symbol $\varepsilon_i$ by two affine forms $\hat{x}, \hat{y}$ indicates some partial dependency between the underlying quantities $x, y$. The magnitude and sign of the dependency is determined by the corresponding coefficients $x_i, y_i$.

For example, suppose that the quantities $x, y$ are represented by the affine forms

\[
\begin{align*}
\hat{x} & = 10 + 2\varepsilon_1 + 1\varepsilon_2 - 1\varepsilon_4 \\
\hat{y} & = 20 - 3\varepsilon_1 + 1\varepsilon_3 + 4\varepsilon_4.
\end{align*}
\]

From this data, we can tell that $x$ lies in the interval $[6 .. 14]$ and $y$ lies in $[12 .. 28]$. However, since they both include the same noise variables $\varepsilon_1$ and $\varepsilon_4$ with non-zero coefficients, they are not entirely independent of each other. In fact, the pair $(x, y)$ is constrained to lie in the region of $\mathbb{R}^2$ depicted in Figure 3 (dark grey), which is substantially smaller than the rectangle $[6 .. 14] \times [12 .. 28]$ (light grey). Obviously, this dependency information would be lost if we were to replace $\hat{x}$ and $\hat{y}$ by the ordinary intervals $[\hat{x}]$ and $[\hat{y}]$, even though the latter encode precisely the same ranges of values as the former.

![Figure 3: joint range of two quantities $x$ and $y$, where $\hat{x} = 10 + 2\varepsilon_1 + 1\varepsilon_2 - 1\varepsilon_4$ and $\hat{y} = 20 - 3\varepsilon_1 + 1\varepsilon_3 + 4\varepsilon_4$](image-url)
4.3 Computing with affine arithmetic

To evaluate a formula in AA, we must replace each of its elementary operations $z \leftarrow f(x, y)$ on real numbers by an equivalent operation $\hat{z} \leftarrow \hat{f}(\hat{x}, \hat{y})$ on affine forms, where $\hat{f}$ is a procedure that computes an affine form for $z = f(x, y)$ that is consistent with $\hat{x}, \hat{y}$.

When $f$ is an affine function of $x, y$, the value $\hat{z}$ can be expressed exactly as an affine combination of the noise symbols $\varepsilon_i$. More precisely, if

$$\hat{x} = x_0 + x_1 \varepsilon_1 + \cdots + x_n \varepsilon_n$$
$$\hat{y} = y_0 + y_1 \varepsilon_1 + \cdots + y_n \varepsilon_n,$$

and $a \in \mathbb{R}$, then

$$\hat{x} \pm \hat{y} = (x_0 \pm y_0) + (x_1 \pm y_1) \varepsilon_1 + \cdots + (x_n \pm y_n) \varepsilon_n$$
$$a \hat{x} = (ax_0) + (ax_1) \varepsilon_1 + \cdots + (ax_n) \varepsilon_n$$
$$\hat{x} \pm a = (x_0 \pm a) + x_1 \varepsilon_1 + \cdots + x_n \varepsilon_n.$$

Note that, according to those formulas, the difference $\hat{x} - \hat{z}$ between an affine form and itself is identically zero. In this case, the fact that the two operands share the same noise symbols with the same coefficients reveals that they are actually the same quantity, and not just two quantities that happen to have the same range of possible values. Thanks to this feature, in AA we also have $(\hat{x} + \hat{y}) - \hat{x} = \hat{y}$, $(3\hat{x}) - \hat{x} = 2\hat{x}$, and so on. Such properties are not valid in IA, and are one source of error explosion.

When $f$ is not an affine operation, $\hat{z}$ cannot be expressed exactly as an affine combination of the $\varepsilon_i$. In that case, we pick the best affine approximation (best in the Chebyshev sense of minimizing the maximum error), and then append an extra term $\varepsilon_k z_k$ to represent the error introduced by this approximation:

$$\hat{z} = z_0 + z_1 \varepsilon_1 + \cdots + z_n \varepsilon_n + \varepsilon_k z_k.$$

Here, $\varepsilon_k$ must be a brand new noise symbol (distinct from all other noise symbols in the same computation) and $z_k$ must be an upper bound for the approximation error. Note that, unlike Hansen’s generalized interval arithmetic, new noise symbols are created during a long AA computation.

Using this approach, the multiplication of two affine forms $\hat{x}, \hat{y}$ is given by

$$z_0 = x_0 y_0$$
$$z_i = x_0 y_i + y_0 x_i \quad (i = 1, \ldots, n)$$
$$z_k = u v,$$

where

$$u = \sum_{i=1}^{n} |x_i|, \quad v = \sum_{i=1}^{n} |y_i|.$$  

Similar formulas can be given for the other elementary operations and functions. For instance, it turns out [5] that the square root of an affine form $\hat{x} = x_0 + x_1 \varepsilon_1 + \cdots + x_n \varepsilon_n$
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is given by

\[
\begin{align*}
  z_0 &= \alpha x_0 + \beta \\
  z_i &= \alpha x_i \\
  z_k &= \delta,
\end{align*}
\]

where

\[
\begin{align*}
  \alpha &= \frac{1}{\sqrt{a} + \sqrt{b}} \\
  \beta &= \frac{\sqrt{a} + \sqrt{b}}{8} + \frac{1}{2} \frac{\sqrt{a} \sqrt{b}}{\sqrt{a} + \sqrt{b}} \\
  \delta &= \frac{1}{8} \frac{(\sqrt{b} - \sqrt{a})^2}{\sqrt{a} + \sqrt{b}}
\end{align*}
\]

and \([a .. b]\) is the interval \([\hat{x}]\).

4.4 Example revisited

Consider again evaluating \(z = x(10 - x)\), for \(x\) in the interval \([4 .. 6]\), but now using AA instead of IA:

\[
\begin{align*}
  \hat{x} &= 5 + 1\epsilon_1 \\
  10 - \hat{x} &= 5 - 1\epsilon_1 \\
  \hat{z} = \hat{x}(10 - \hat{x}) &= 25 + 0\epsilon_1 - 1\epsilon_2.
\end{align*}
\]

Observe that the influence of the noise symbol \(\epsilon_1\) in the factors happened to cancel out (to first order) in the product.

The range of \(z\) implied by the affine form \(\hat{z}\) is

\([\hat{z}] = [25 - 1 .. 25 + 1] = [24 .. 26]\),

which is much closer to \([24 .. 25]\), the true range of \(z\). Recall that IA gave \([16 .. 36]\) for this expression.

5 Examples of adaptive enumeration

In this section, we show examples that compare the performance of IA and AA in adaptive enumeration. Because pictures of three dimensional decompositions are not easy to understand, we only give pictures of the enumeration of two-dimensional implicit curves.

In all these examples, the goal is to build a cellular model of a curve given implicitly by \(h(x, y) = 0\) inside some rectangle \(\Omega\). The model consists of a set of cells from some fixed regular grid, whose union is guaranteed to contain all zeros of \(h\) in \(\Omega\). These cells are shown in grey in the pictures.
5.1 A quartic

Take the quartic curve defined by

\[ h(x, y) = x^2 + y^2 + xy - (xy)^2/2 - 1/4 \]

in the square \( \Omega = [-2 .. 2] \times [-2 .. 2] \), using a \( 32 \times 32 \) grid of cells. In a full enumeration, \( 1024 = 32 \times 32 \) cells have to be scanned, but the curve actually enters only 66 of these cells.

Figure 4 illustrates an adaptive enumeration with interval arithmetic and affine arithmetic, using a 2-d tree. With IA, the range of \( h \) was evaluated 847 times and 246 cells remained in the model (i.e., could not be shown to be disjoint from the curve). With AA, the range of \( h \) was evaluated 451 times and only 70 cells remained in the model. Thus, IA generated a model with 180 useless cells whereas AA generated a model with only 4 useless cells, even at such a relatively low resolution.

5.2 Quadratic electrostatic potential

Consider now the “quadratic” electrostatic potential defined by \( n \) charges \( q_i \) located at points \((x_i, y_i)\):

\[ h(x, y) = \sum_{i=1}^{n} \frac{q_i}{(x - x_i)^2 + (y - y_i)^2} . \]

(True electrostatic potential depends on distances, not on square of distances.) In this case, the interesting curves are the equipotential curves \( h(x, y) = h_0 \). Such “potential” functions and their variations are useful for modeling with implicit objects [22].

Figure 5 illustrates an adaptive quadtree enumeration with interval arithmetic and affine arithmetic of an equipotential curve for the quadratic electrostatic potential defined by four
equal charges. The grid is 64 × 64; a full enumeration would have to scan 4096 cells. In both cases, the range of \( h \) was evaluated 1449 times (a coincidence). However, 346 cells remained in the model generated by IA, whereas only 299 remained in the model generated by AA.

![Figure 5: adaptive enumeration of quadratic electrostatic potential with IA (left) and AA (right)](image)

5.3 The quartic in 3d

As a three-dimensional example, consider an octree enumeration of the quartic surface defined by

\[
h(x, y, z) = x^2 + y^2 + xy - (xy)^2/2 - 1/4 - z,
\]

in the cube \( \Omega = [-2..2] \times [-2..2] \times [-2..2] \).

We give two tables showing how the size of the model (number of leaves in the octree) and the time (in seconds) taken to compute it depend on the resolution of the underlying grid (the resolution is \( n = 2^L \) at level \( L \)). Table 1 shows this data for IA; Table 2 shows this data for AA. Times were measured in an otherwise idle SPARCstation 10.

We observe that AA evaluates \( h \) half as many times as IA, and generates a model half the size of the model generated by IA; the size of the IA model is approximately 5\( n^2 \), whereas the size of the AA model is approximately 2.4\( n^2 \). However, AA is 4.3 times slower than IA; the time for the IA model is approximately 86\( n^2 \mu \text{sec} \) whereas the time for the AA model is approximately 370\( n^2 \mu \text{sec} \) (note that the time is linear in the size of the model.)
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<td>6738841</td>
<td>2516356</td>
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Table 1: performance of adaptive enumeration of 3d quartic with IA

<table>
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Table 2: performance of adaptive enumeration of 3d quartic with AA
6 Rendering

The octrees built during enumeration can be used to speed up ray tracing algorithms [9, 23]. However, a "lower" quality rendering can be done directly, concurrently with the enumeration; there is no need to explicitly build an octree.

By composing the function $h$ with the appropriate projective map, we can take the domain $\Omega$ to be a rectangular box, aligned with the coordinate axes, and compute the image of the orthogonal projection of the surface $S$ defined by $h$ onto a horizontal plane located above the box $\Omega$. For simplicity, we assume that the resolution of the image is $n \times n$, where $n = 2^L$. The enumeration proceeds down the octree to level $L$. When we reach a leaf of the octree (i.e., a cell at level $L$), we compute the colour of the corresponding pixel in the image. Each cell has an integer address $(x, y, z)$; all cells with the same $x, y$ project onto the same pixel $(x, y)$. To remove hidden parts, we use a "painter's algorithm", carefully traversing the octree from bottom to top (in the $z$ direction).

In order to compute the colour of a pixel in shaded images of $S$, we need to compute also the mean normal direction to $S$ in the cell corresponding to the pixel. The normal direction is the direction of the gradient of $h$. Full enumeration and adaptive enumeration with interval arithmetic do not directly provide an estimate for $\nabla h$. For full enumeration, one typically uses an estimate based on central differences [12]. For adaptive enumeration with interval arithmetic, techniques such as formal differentiation (evaluated in either floating point or IA) or Bauer-Strassen differentiation [13] would be required to estimate $\nabla h$ in each visible cell. On the other hand, affine arithmetic directly provides an estimate for $\nabla h$. Recall that, to compute $h$ in a cell represented by the intervals $\bar{x}$, $\bar{y}$, $\bar{z}$, we take the corresponding affine forms $\hat{x} = x_0 + x_1\varepsilon_1$, $\hat{y} = y_0 + y_2\varepsilon_2$, $\hat{z} = z_0 + z_3\varepsilon_3$, and compute $\hat{h} = h_0 + h_1\varepsilon_1 + h_2\varepsilon_2 + h_3\varepsilon_3 + \ldots$ Then we can take $(h_1/x_1, h_2/y_2, h_3/z_3)$ as an estimate for the mean value of $\nabla h$ in this cell, and from this estimate compute an appropriate colour for the corresponding pixel in the image.

Figure 6 shows a $256 \times 256$ image of the union of two spheres rendered using this technique.

7 Other applications

Affine arithmetic is also useful for other applications related to implicit objects. We describe two such applications briefly: surface intersection, and rendering with ray tracing.

7.1 Intersection of implicit surfaces

The intersection of two implicit surfaces defined by functions $f$ and $g$ can be given implicitly by a single function $h = f^2 + g^2$. This expression is not useful for enumeration based on point sampling because $h$ is always positive and enumeration based on sampling relies on changes of sign in $h$ to identify intersecting cells. On the other hand, enumeration using range analysis is perfectly feasible in this case. Again, affine arithmetic will be able to exploit the correlations present in the expression of $h$ to compute tighter bounds. Alternatively, the
Figure 6: the union of two spheres enumerated and shaded with AA

equation for the intersection can be done by combining the simultaneous enumeration of the two surfaces, rejecting cells that are rejected in at least one enumeration.

7.2 Ray tracing

A fundamental step in ray tracing a surface is finding the first intersection of a ray with the surface. If the surface is given implicitly by \( h(x, y, z) = 0 \), and the ray is the line segment \( \overline{pq} \), then one needs to find the smallest root of the univariate function

\[
f(t) = h((1 - t)x_p + tx_q, (1 - t)y_p + ty_q, (1 - t)z_p + t z_q),
\]

in the interval \( t \in [0..1] \).

A simple algorithm for finding all roots of \( f \) is the following. Evaluate \( \overline{u} = f(t) \) using IA, for the whole interval \( \overline{t} = [0..1] \). If the resulting interval \( \overline{u} \) is strictly positive or strictly negative, then the ray does not intersect the surface. Otherwise, split \( \overline{t} \) in two equal parts, and recursively search each half. Stop the recursion when the interval \( \overline{t} \) is small enough for the application. A simple variant of this algorithm finds the smallest root: only search the right half if the left half has been shown to contain no roots.

One drawback of this algorithm is that evaluating \( \overline{f}(\overline{t}) \) with IA is equivalent to evaluating \( \overline{h}(\overline{x}, \overline{y}, \overline{z}) \) on the intervals \( \overline{x} = [x_p .. x_q], \overline{y} = [y_p .. y_q], \overline{z} = [z_p .. z_q] \) — that is, evaluating \( h \) on the axis-aligned bounding box of the segment \( \overline{pq} \), instead of only along the segment itself. Again, the problem arises because IA does not know that the arguments \( \overline{x}, \overline{y}, \) and \( \overline{z} \) of \( \overline{h} \) are highly correlated. Obviously, the bounding box of the segment \( \overline{pq} \) may intersect the surface even when the segment itself does not.

Replacing standard IA by affine arithmetic will generally improve the performance of this algorithm. Even without any algebraic manipulation, AA will automatically notice that the affine forms \( \overline{x}, \overline{y}, \) and \( \overline{z} \) are strongly correlated, and thus will usually produce tighter bounds for \( f(\overline{t}) \). Moreover, as we have seen above, AA also automatically provides
estimates for the derivative of $f$, allowing us to replace the binary search used in the IA method by Newton’s method, without explicitly computing derivatives.

8 Conclusion

Numerical experiments show that affine arithmetic is indeed more accurate than standard interval arithmetic. On the other hand, AA is more complex and expensive than IA. However, its higher accuracy is worth the extra cost for adaptive enumeration of implicit objects because the models resulting from enumeration with AA are smaller. Although the relative speeds of IA and AA depend on the particular function being evaluated, in our implementation, AA is typically 4–5 times slower than IA. Work is in progress to improve the performance of AA.

An interesting extension of the techniques described here would be to use the extra information provided by AA to estimate surface curvature, so that the enumeration is adapted not only to the location of the surface in the ambient space, but also to its intrinsic geometry, as adaptive refinement techniques do.

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