Out-of-Core Construction and Visualization of Multiresolution Surfaces

P. Lindstrom


February 3, 2003
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This work was performed under the auspices of the U.S. Department of Energy by the University of California, Lawrence Livermore National Laboratory under Contract No. W-7405-Eng-48.
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Peter Lindstrom
Lawrence Livermore National Laboratory

Abstract

We present a method for end-to-end out-of-core simplification and view-dependent visualization of large surfaces. The method consists of three phases: (1) memory insensitive simplification; (2) memory insensitive construction of a multiresolution hierarchy; and (3) run-time, output-sensitive, view-dependent rendering and navigation of the mesh. The first two off-line phases are performed entirely on disk, and use only a small, constant amount of memory, whereas the run-time system pages in only the rendered parts of the mesh in a cache coherent manner. As a result, we are able to process and visualize arbitrarily large meshes given a sufficient amount of disk space; a constant multiple of the size of the input mesh.

Similar to recent work on out-of-core simplification, our memory insensitive method uses vertex clustering on a rectilinear octree grid to coarsen and create a hierarchy for the mesh, and a quadric error metric to choose vertex positions at all levels of resolution. We show how the quadric information can be used to concisely represent vertex position, surface normal, error, and curvature information for anisotropic view-dependent coarsening and silhouette preservation.

The run-time component of our system uses asynchronous rendering and view-dependent refinement driven by screen-space error and visibility. The system exploits frame-to-frame coherence and has been designed to allow preemptive refinement at the granularity of individual vertices to support refinement on a time budget.

Our results indicate a significant improvement in processing speed over previous methods for out-of-core multiresolution surface construction. Meanwhile, all phases of the method are disk and memory efficient, and are fairly straightforward to implement.


Keywords: surface simplification, large-data visualization, view-dependent refinement, out-of-core algorithms

1 Introduction

Recent advances in scanning technology and the ever increasing size of computer simulations have led to a rapid increase in the availability and size of geometric data sets. Massive polygonal data sets, consisting of hundreds of millions of faces, are becoming quite common [Bernardini et al. 1999; Levoy et al. 2000]. While the performance of graphics hardware has also seen a dramatic rise in the last few years, our ability to produce data sets that overload the capabilities of state-of-the-art graphics chips has led researchers to develop methods for automatic model simplification and run-time level-of-detail (LOD) management [Funkhouser and Séquin 1993].

Whereas many polygonal environments consist of a large collection of moderately complex objects, such as those used in video games, recent trends are for single objects to consist of millions of polygons. Examples of such models include terrain surfaces and high-resolution range scans. The traditional approach of storing a few static levels of detail is not viable for such large objects, which are often viewed in a manner that they vary greatly in screen resolution over the surface. As a result, methods for view-dependent simplification have been proposed, in which a continuous level-of-detail hierarchy is first constructed and then adapted at a fine granularity at run-time [Lindstrom et al. 1996; Xia and Varshney 1996; Duchaineau et al. 1997; Hoppe 1997; Luebke and Erikson 1997; El-Sana and Varshney 1999; Erikson et al. 2001].

The techniques mentioned above have been used successfully for simplifying models up to a few million triangles. In the last few years, however, polygonal models have become so large that they often greatly exceed our ability to perform conventional in-core simplification, and a number of techniques have been devised for simplifying such models out-of-core, e.g. [Bernardini et al. 1999; Lindstrom 2000; Lindstrom and Silva 2001; Shaffer and Garland 2001]. These methods, however, all produce static, single-resolution meshes, which in a sense is inappropriate for such large surfaces, because in order to view them interactively they have to be simplified to such a degree that many important details are lost. Rather, we would like to construct a level-of-detail hierarchy for such large surfaces, and then use view-dependent techniques to render and explore them at varying resolution without significant loss in fidelity. While there has been some recent work on out-of-core

Figure 1: Various view-dependent meshes produced by our system. The mesh in the middle has been coarsened aggressively outside the view frustum (here shown as a rectangle).
construction of LOD hierarchies, most notably by El-Sana and Chiang [2000] and by Prince [2000], these methods have rather long execution times, can be somewhat difficult to implement, and still rely on having a large amount of memory available for in-core processing. Furthermore, the results presented in these papers are for models of rather modest size (just a million or a few million triangles), making it difficult to judge how well they scale to truly large models.

In this paper, we present an alternative approach to out-of-core simplification for view-dependent refinement, by extending the static simplification algorithm by Lindstrom and Silva [2001]. Our end-to-end off-line method is memory insensitive, meaning that it can run successfully with essentially an arbitrarily small amount of memory (in our case, less than 8 MB). We achieve this by storing all intermediate computations in temporary files on disk, and take advantage of fast sequential disk access. In addition to being memory efficient, our method is considerably faster than previously published techniques, running at a triangle reduction rate of up to 60,000 triangles per second. We also present data structures for concisely encoding quadratic matrices and the per-vertex information needed later at run-time during view-dependent rendering. Finally, our method is straightforward to implement and is a simple but significant extension of Lindstrom and Silva’s memory insensitive simplification algorithm. We will present the various steps in our algorithm after covering related work in the area.

2 Previous Work

In this section, we cover previous work on view-dependent and out-of-core surface simplification. Because there has been extensive work on simplification, we here only cover a few of the most related algorithms in the field.

Among the first methods for view-dependent simplification for general polygonal models was the technique by Xia and Varshney [1996]. Their method uses edge collapse to construct a binary tree of possible coarsening operations off-line. At run-time, they use a screen space metric, based on geometric error and proximity to silhouettes and specular highlights, for determining which edges to collapse. Hoppe extended his work on progressive meshes to view-dependent refinement [Hoppe 1997]. Similar to [Xia and Varshney 1996], Hoppe uses edge collapse, but his algorithm provides greater freedom in choosing the order of edge collapses, which generally results in higher quality adaptive meshes. His method also makes use of geomorphing to reduce temporal LOD artifacts. More recently, El-Sana and Varshney [1999] presented a method, based on the more general vertex merge operation, that is able to merge and simplify topologically disjoint parts of an object. This method has been improved by El-Sana and Bachmat [2002], who aggregate parts of the multiresolution hierarchy to speed up LOD selection and rendering.

A different approach to view-dependent refinement was taken by Luebke and Erikson [1997]. Rather than relying on edge collapse or vertex pair contraction, they use an even more general coarsening operation—vertex clustering—which allows a large collection of vertices to be merged in a single atomic operation. Like us, they make use of an octree decomposition of space, rather than a general binary tree over the set of mesh vertices. Like others, they make use of normal cones to detect when the surface is near a silhouette. Luebke and Hallen [2001] later used this framework for performing view-dependent “imperceptible simplification.”

The off-line processing techniques above all assume that the hierarchy can be constructed in-core. To simplify large meshes out-of-core, Lindstrom [2000] proposed a technique, based on vertex clustering on a uniform grid, that makes use of Garland and Heckbert’s quadratic error metric [Garland and Heckbert 1997]. Lindstrom’s method performs a single sweep over the mesh and constructs an in-core representation of the simplified model. More recently Lindstrom and Silva [2001] extended this method by removing the requirement of having enough RAM to store the simplified model. Their memory insensitive technique uses a constant amount of memory and performs a series of external sorts to allow sequential access to the on-disk data. Our first simplification phase is based upon their algorithm. To provide a higher level of adaptivity, Shaffer and Garland [2001] proposed making two instead of one pass over the mesh. In the first pass, uniform clustering like in [Lindstrom 2000] is performed, after which a binary space partitioning (BSP) tree is constructed from the accumulated quadric information. In the second pass, the BSP tree is used to recluster the mesh. Other recent out-of-core simplification methods for adaptive but static levels of detail are presented in [Choudhury and Watson 2002; Fei et al. 2002; Garland and Shaffer 2002].

There have only been a few published methods for out-of-core simplification for view-dependent refinement. Hoppe applied his view-dependent progressive mesh work to terrain data [Hoppe 1998]. His approach is to partition the terrain into a block hierarchy and simplify the blocks independently. Then, the blocks are merged and the seams between them are simplified further. Prince [2000] later extended Hoppe’s out-of-core simplification method for terrain to arbitrary polygonal surfaces. Prince’s method, like ours, makes use of quadric error metrics, but uses edge collapse as the coarsening primitive. While effective for medium-size models, his out-of-core method still requires much RAM and may be too slow for simplifying very large models. Cignoni et al. [2002] describe efficient data structures for a similar block hierarchy based technique that allows a greater degree of simplification across block boundaries. Their results suggest an improvement over [Prince 2000] in simplification speed and memory efficiency, but they do not report on using their data structures for interactive view-dependent refinement. El-Sana and Chiang [2000] proposed a novel out-of-core technique for view-dependent simplification by segmenting the mesh into independent patches. These patches are such that the edge collapse order inherently preserves the boundaries between them, thus simplifying and stitching the patches together can be done without the need for explicit boundary constraints. Unfortunately, the models used by El-Sana and Chiang are small by out-of-core simplification standards, and it is not clear how well their method scales. For large enough models the average patch size is likely to shrink to the point that each edge must be loaded and collapsed individually, leading to excessive thrashing. More recently, DeCoro and Pajarola [2002] described external data structures and a run-time system for out-of-core visualization. Their framework assumes edge collapse is used to coarsen the mesh, but no out-of-core method is presented for constructing the multiresolution structure. Rusinkiewicz and Levoy [2000] proposed an interesting alternative to polygon-based view-dependent refinement. Their QSplat algorithm clusters the triangle mesh into a vertex hierarchy, and then uses point primitives to render the mesh. While conceptually simple, most current hardware is optimized for triangle rather than point rendering, and the quality afforded by real-time point-based rendering can be rather low. Still, hybrid techniques like Cohen et al.’s point- and triangle-based simplification [Cohen et al. 2001] may prove useful. Finally, there are a number of out-of-core visualization systems based on managing collections of static levels-of-detail, including [Aliaga et al. 1999; Erikson et al. 2001; Varadan and Manocha 2002]. While related to our method, the focus of these works is on the run-time handling of a large number of small objects, as opposed to the off-line construction and run-time fine-grained adaptation of a single large mesh.

To our knowledge, our system is the first to perform both scalable out-of-core simplification and view-dependent visualization of general surfaces from external memory. We will describe our algorithm in the following sections.
3 Algorithm Overview

Our view-dependent algorithm consists of three phases: simplification, level-of-detail hierarchy construction, and run-time view-dependent refinement and rendering. The first two phases are run off-line to produce an on-disk level-of-detail representation of the mesh. The run-time component then traverses this hierarchy, pages in the data needed, and produces an adaptive mesh that can be displayed interactively. Our main approach is to use a sparse octree decomposition of space over a uniform rectilinear grid, similar to Lubeck and Erikson’s view-dependent simplification algorithm [Lubeck and Erikson 1997]. The octree is sparse in the sense that only those nodes that contain at least one vertex from the input mesh are retained. Each node in this octree corresponds to a vertex at some level of resolution, and adaptive mesh simplification and refinement are performed by collapsing and expanding nodes (i.e., removing and creating child nodes, respectively) in the octree. In this section, we give a brief overview of the three phases of our algorithm, and provide further details in the following sections.

The first phase—simplification—is based on the OoCSx memory insensitive mesh simplification algorithm by Lindstrom and Silva [2001]. Their method is a memory efficient variation on the out-of-core simplification algorithm by Lindstrom [2000], which in turn was inspired by Rossignac and Borrel’s [1993] vertex clustering algorithm. Using a uniform grid to partition space, all vertices that fall in the same grid cell are merged (clustered) to a single vertex. In this process, the triangles that collapse to an edge or a point are discarded. Whereas vertex clustering has the potential to introduce non-manifold vertices and edges, this is typically not a concern in interactive visualization, which is our primary application. On the contrary, modest topology simplification is often desired to eliminate the “topological noise” introduced in the data scanning process. For the sake of geometric accuracy, representative vertices for the clusters are chosen based on minimizing the quadric error—a weighted sum of squared distances to the triangle planes of the input mesh—which can be encoded using a symmetric $4 \times 4$ matrix [Garland and Heckbert 1997].

The OoCSx algorithm performs all these tasks on disk and avoids costly random accesses using a sequence of external sorts, followed by fast sequential accesses. The intermediate output of this algorithm is a set of triangles for the simplified mesh and a list of quadric matrices for its vertices. Note that, unless the cluster grid is fine enough, the mesh will be irreversibly decimated in this first phase, and some fine details of the original full-resolution mesh may be lost. However, in our experience, this potential loss in detail is rarely noticeable. Each triangle that survives the simplification is later assigned to an internal node in the octree hierarchy, as given by an octocode—a bit string that uniquely identifies a grid cell by position and resolution in the octree. Also to facilitate the ensuing octree construction, we augment the per-vertex quadric matrices with a vertex position, computed from the quadric matrix, and an approximate vertex normal, and output these fields to a vertex file. The vertex file is output in octcode order, such that sibling nodes are stored together. This phase of the algorithm is described in Section 4.

The octree construction phase begins by sorting the triangle file by octocode, such that the triangle order follows the order in which nodes are created in the octree. We then build the octree by first processing the vertex file sequentially. Sibling nodes are identified and grouped together, and the quadric and other geometric information is merged to form a new level. Meanwhile, the triangle file is scanned sequentially and triangles are deposited in the nodes where they belong. The generated parent nodes are output sequentially (again in octcode order) to a temporary file for that given level of resolution, and the process is applied iteratively until all levels in the octree are represented, ending with a temporary file containing a single node—the root node. This bottom-up construction is then followed by a final top-down, level-by-level traversal, during which nodes are linked together in a single file and the hierarchical structure is output in the desired order, from coarse to fine resolution. Note that all the I/O accesses used in our simplification and octree construction are sequential in nature, and are therefore very efficient. The algorithm for the octree construction phase is described in Section 5, while the data structures for the output produced are discussed in detail in Section 6.

In the run-time phase, the file containing the LOD hierarchy, which could potentially greatly exceed the available main memory, is memory mapped so that it can be accessed as though it were resident in continuous memory. For simplicity, we let the operating system perform on-demand paging of the external data, although this data could equally well be accessed using an explicit paging scheme. To minimize run-time computation, an in-core “copy” is made of each active node in the octree. As new nodes are created, we extract data from the external LOD hierarchy, compute any per-vertex and per-triangle information not explicitly stored, and write this data to a dynamically allocated data structure. These nodes are maintained in a dynamic subset of the complete external octree, and are expanded and collapsed, as determined by view-dependent error and visibility criteria, to adapt to the current viewpoint. The triangles contained in the active nodes and the currently active vertices they reference are then output to a triangle list and rendered. To avoid stalling the rendering when parts of the mesh need to be paged in, we decouple rendering and refinement/paging as two asynchronous threads. Also, for the sake of interactivity, we impose a time limit on the refinement thread to guarantee frequent updates of the mesh, and generally perform refinement breadth-first to ensure that detail is progressively added evenly over the mesh whenever the allotted refinement time is insufficient. The in-core, dynamic data structures are presented in Section 7, while the steps of the view-dependent refinement are given in Section 8.

4 Simplification

The simplification phase of our algorithm is based on, and is essentially identical to, the beginning stages of the memory insensitive technique OoCSx described in [Lindstrom and Silva 2001]. Therefore, we will only briefly cover this part of our algorithm, and we will focus on the few differences between the two methods. Pseudo-code for the simplification phase is given in Table 1.

Before the simplification begins, the user chooses the resolution of a uniform rectilinear grid that completely contains the input mesh. This grid is constrained to have dimensions $2^n \times 2^n \times 2^n$, for some positive integer $n$. The cells in this grid correspond to the leaf nodes in an $(n+1)$-level octree that ultimately forms a multi-resolution representation of the mesh. In the discussion below, we will use the terms grid cell, cluster, and node interchangeably.

As in [Lindstrom and Silva 2001], we process the input mesh, which is represented as a triangle soup (a sequence of triplets of vertex coordinates), one triangle at a time. For each triangle $t$, we compute a 4-vector

\[ \mathbf{n}_t = \mathbf{w}_t \left( \frac{\mathbf{n}_v}{d_v} \right) \]  

for an implicit plane equation $\mathbf{n}_t \cdot \mathbf{x} + d_v = 0$ that will later be used to construct a quadric matrix. The weight $\mathbf{w}_t$ allows non-uniform weighting of the planes (see below). Then, for each of $t$‘s three vertices, we quantize the vertex coordinates to an integer grid cell location, and then convert this location to an octocode $v$ (cf. [Li and Loew 1987]). These octocodes are represented as follows: The root node has octocode $v = 1$ and the $k^{th}$ child of a node $v$ is computed as $8v + k$, with $k = 0, \ldots, 7$. These octocodes have the property that they are ordered by level, from top to bottom, and sibling nodes...
have consecutive octcodes. Whereas these octcodes are different from the cluster IDs used in OctoCSx, this is of no consequence to the simplification algorithm—any one-to-one mapping between quantized coordinates and cluster IDs will do. The planes and associated octcodes are then output sequentially to a temporary plane file, P. If the triangle’s vertices belong to different clusters, we determine which octree node v_t to assign t to. As in [Luebke and Erikson 1997], we choose v_t to be the lowest common ancestor of all pairs of vertices from t, such that expanding the node v_t would introduce the triangle t (analogous to the introduction of triangles caused by a vertex split in [Hoppe 1996]). Unfortunately, the order given by v_t does not exactly match the bottom-up order in which nodes are visited during the octree construction. Therefore, we compute a modified sort key v_t* for t that orders nodes by increasing octcode within each level, but reverses the inter-level order such that leaf nodes are listed first. Finally, the non-degenerate octree and its reversed octcode are written to a triangle file T_out.

After the input has been exhausted, the plane file is sorted on the octcode field using an external sort. As in [Lindstrom and Silva 2001], we use rsort [Linderman 1996] for this task. We then process all planes, one cluster at a time, and construct a 4 × 4 quadric matrix Q_v for each cluster v:

\[ Q_v = \sum_t \hat{n}_t \hat{n}_t^T = \sum_t w_t^2 \left( \frac{\hat{n}_t \hat{n}_t^T}{d_t^2} \right) \]  

We also construct an approximate normal \( \hat{n}_v \) for the cluster, e.g. as the sum of the cluster’s face normals, which will later be used to resolve an ambiguity in sign of the actual normal extracted from the quadric matrix. If space is at a premium, we can avoid storing \( \hat{n}_v \) by performing additional computation and encoding this sign directly in the quadric matrix (see Section 6.1.2). Given a quadric matrix Q_v, we proceed by computing a representative vertex for the cluster that minimizes the quadric error [Lindstrom and Silva 2001]. Finally, we output the quadric matrices, along with their octcodes, vertex positions, and normals, to an intermediate vertex file V_n for level n in the octree (the bottommost level), and we remove the temporary plane file P. For each leaf node that contains at least one vertex from the original mesh, we now have a single vertex for the full-resolution simplified mesh. The original OctoCSx simplification algorithm would at this point perform multiple external sorts on the triangle file to replace the cluster IDs with vertex indices. Because

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\( \hat{n}_v \) is a proprietary software developed at AT&T Labs.

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### 5 Octree Construction

The second phase of our algorithm takes the simplified mesh, represented as a list of vertices V_n, augmented with quadric error information, and a list of triangles T, and constructs a coarse-to-fine level-of-detail representation H of the mesh. For each node in H, we store vertex information, such as position, normal, error, etc., as well as a (potentially empty) list of triangles. The triangles of a node are the ones that are eliminated when the node is collapsed. We will focus later on the particular data structures used for the nodes in H, and spend this section describing the steps of the octree construction algorithm.

Table 2 lists pseudo-code for the octree construction. We begin by externally sorting the triangle file on its reversed octcode field v*. This is done to group the triangles by octree node and to place them in the order in which the nodes will be processed. Following this sort, we begin constructing the internal nodes of the octree (lines 2–9). Recall that the simplification has already produced vertex positions and normals for the leaf nodes V_n on level n. Because V_n is sorted on octcode, and because sibling nodes have consecutive octcodes, we can easily scan V_n and fetch the per-vertex data for each group of siblings. From these we compute a quadric matrix and approximate normal for the parent using simple addition. We additionally compute the optimal vertex position \( p \), the minimum
For the on-disk leaf node, we write only the vertex position and normal vector. The triangles are represented as triplets of octcodes corresponding to leaf nodes in the hierarchy. The vertex fields are derived entirely from the quadric matrices, and will be discussed in the following sections.

### 6 External Data Structures

We now turn our attention to the data structures used for our external on-disk representation of the level-of-detail hierarchy $H$. Each internal node in $H$ consists of vertex information (position, normal, etc.) and a list of triangles and pointers to its children. These data structures are given by Table 3. Because leaf nodes have no triangles associated with them, and because error evaluation and view culling are done only for internal nodes, we store only position and normal information with leaf nodes. The triangles are represented as triplets of octcodes corresponding to leaf nodes in the hierarchy. The vertex fields are derived entirely from the quadric matrices, and will be discussed in the following sections.

#### 6.1 Vertex Data

In this section, we describe how to compute and store the per-vertex information needed in our view-dependent renderer from the $4 \times 4$ quadric matrix $Q$. The data we are primarily concerned with are the vertex position $p$, the surface normal $n$, the quadric error $e$ at $p$, and a matrix $K$ that encodes the normal curvature and is used to measure how large the error appears from different view directions. We could compute and store $(K, n, p, e)$ directly, however this information would require $6 + 3 + 3 + 1 = 13$ scalar values, whereas the original quadric matrix requires only 10 values (assuming we take advantage of the fact that $Q$ and $K$ are symmetric). Instead, we will make use of an alternative representation $(r, \lambda, p, e)$—three 3-vectors and a scalar—that allows $K$ and $n$ to be computed quickly. In addition to these fields, we also store the radius $r$ of a bounding sphere, centered on $p$, used for view frustum culling. We will describe how to compute this radius later.

#### 6.1.1 Vertex Position and Quadric Error

In Section 4, we explained how to compute the quadric matrix $Q$ for a vertex, or more generally a node in the LOD hierarchy. As in [Lindstrom 2000], we decompose the quadric matrix as

$$Q = \begin{pmatrix} \mathbf{A} & -\mathbf{b} \\ -\mathbf{b}^T & c \end{pmatrix}$$

(4)

We can then write the quadric error $\epsilon(x)$ as

$$\epsilon(x) = x^T \mathbf{A} x - 2\mathbf{b}^T x + c$$

(5)

Let $p$ be the position of the vertex associated with the node and let $\nabla \epsilon(p) = 2(\mathbf{A} p - \mathbf{b})$ be the gradient of $\epsilon$ at $p$. We now rewrite $\epsilon$:

$$\epsilon(x) = (x - p)^T \mathbf{A} (x - p) + 2(\mathbf{A} p - \mathbf{b})^T (x - p)$$

$$+ (p^T \mathbf{A} p - 2 \mathbf{b}^T p + c)$$

(6)

$$= (x - p)^T \mathbf{A} (x - p) + \nabla \epsilon(p)^T (x - p) + \epsilon(p)$$

Because we want to minimize the quadric error, we generally choose the vertex position $p$ to be a minimizer for $\epsilon$, which implies $\nabla \epsilon(p) = 0$ and

$$\epsilon(x) = (x - p)^T \mathbf{A} (x - p) + \epsilon(p)$$

$$\epsilon(p) = c - p^T \mathbf{A} p$$

(7)

That is, if $p = \arg \min \epsilon$, we can parameterize the quadric error as $(\mathbf{A}, p, \epsilon)$. This parameterization is valid whether $\mathbf{A}$ is singular or not. If $\mathbf{A}$ is singular, $\epsilon$ has infinitely many minima, in which case we choose the one closest to the grid cell center. In rare circumstances our chosen optimum $p$ falls outside its associated grid cell. When this happens, we constrain $p$ using the procedure outlined in [Lindstrom and Silva 2001], which may result in a $p$ that does not minimize $\epsilon$. Regardless of this special case, if we do not have to compute $\epsilon$ at points other than the vertex position $p$ (irrespective of our choice of $p$), then the parameterization $(\mathbf{A}, p, \epsilon)$ is useful since it directly gives us the vertex position $p$ and the error $\epsilon$ at $p$. Still, we are left with determining the normal $\hat{n}$ and the curvature matrix $K$. As shown below, these two quantities can both be derived from the matrix $\mathbf{A}$.

#### 6.1.2 Surface Normal

To compute the surface normal $\hat{n}$, note that the matrix $\mathbf{A}$ is the covariance matrix (with zero mean) for the set of (weighted) normals of the triangles in the cluster [Garland 1999, §4.2.2]. Thus, the eigenvector for the largest eigenvalue $\lambda_1$ of $\mathbf{A}$ corresponds to...
the dominant normal direction \( \hat{n} \). Note that if \( \hat{n} \) is an eigenvector, then so is \(-\hat{n}\). Because the sign of the normal matters for correct rendering, we will show later how to resolve this ambiguity. Using an eigen decomposition of \( A \), we have

\[
A = \mathbf{P} \Lambda \mathbf{P}^T
\]

\[
P = (x_1, x_2, x_3) \quad \mathbf{P}_n \parallel \hat{n}
\]

\[
\Lambda = \text{diag}(\lambda_1, \lambda_2, \lambda_3) \quad \lambda_1 \geq \lambda_2 \geq \lambda_3 \geq 0
\]

where \( \Lambda \) is a diagonal matrix of (non-negative) eigenvalues, and \( \mathbf{P} \) is orthogonal with determinant \( \text{det}(\mathbf{P}) = 1 \). That is, \( \mathbf{P} \) is a rotation matrix, which can be represented using as little as three parameters. We have chosen to use a 3-parameter axis-angle representation that is similar to the standard unit quaternion representation. Let \( \mathbf{P} \) correspond to a rotation around a unit vector \( \mathbf{r} \) by an angle \( \theta \). Then the vector

\[
\mathbf{r} = (r_x, r_y, r_z) = \sqrt{2} \sin \frac{\theta}{2} \hat{\mathbf{r}}
\]

completely represents \( \mathbf{P} \). We will not go into detail of how to compute \( \mathbf{P} \) from \( \mathbf{P} \), but refer the reader to any tutorial on quaternions, e.g. [Shoemake 1991]. We recover \( \mathbf{P} \) from \( \mathbf{r} \) as follows:

\[
\mathbf{P} = \left(\begin{array}{ccc}
1 - r_x^2 - r_y^2 & r_x r_y - \alpha r_z & r_x r_z + \alpha y \\
 r_x r_y + \alpha z & 1 - r_y^2 - r_z^2 & r_y r_z - \alpha x \\
r_x r_z - \alpha y & r_y r_z + \alpha x & 1 - r_z^2 - r_x^2
\end{array}\right)
\]

\[
\alpha = \sqrt{2 - r_x^2 - r_y^2 - r_z^2}
\]

Thus, by storing \( \mathbf{r} \), we can quickly compute \( \mathbf{P} \) and the normal \( \hat{n} \) from the first column of \( \mathbf{P} \). To recover \( A \), we also store the eigenvalues \( \lambda \).

As noted above, the canonical decomposition \( A = \mathbf{P} \Lambda \mathbf{P}^T \) does not necessarily lead to a matrix \( \mathbf{P} \) whose first column equals the surface normal in sign. However, if we already know the (approximate) normal, which is the case in our simplification algorithm (see Section 4), then we can test whether the normal obtained from \( \mathbf{P} \) matches the given normal. If the two vectors point in opposite directions, then we encode this fact by negating \( \lambda_1 \). Because \( A \) is non-zero and positive semi-definite, we must have \( \lambda_1 > 0 \), and we can therefore safely use the sign bit of \( \lambda_1 \) to encode the sign of \( n \).

As mentioned in Section 4, we explicitly store approximate normals in the vertex files. To save disk space, however, we could use the technique just described for extracting normals from quadric matrices. We would then compute, but not store, a surface normal for each leaf node during simplification, compare it against the normal obtained from the quadric matrix \( Q \), and encode its sign difference in \( Q \). Similar to the argument above, because \( Q \) is non-zero and positive semi-definite, \( \text{tr}(Q) > 0 \), and the need to flip the extracted normal can be encoded by negating the diagonal of \( Q \).

### 6.1.3 Curvature Matrix

Our view-dependent error metric takes advantage of the fact that geometric displacements parallel to the view direction are less perceptible than those perpendicular to the view direction. Thus geometry viewed straight-on can often be coarsened significantly more than geometry near silhouettes—a fact that has been exploited by other view-dependent methods, e.g. [Lindstrom et al. 1996; Xia and Varshney 1996; Hoppe 1997; Luebke and Erikson 1997]. This is illustrated in Figure 2 for a smoothed and slightly curved octahedron. Rather than using a cone to bound the normals [Hoppe 1997; Luebke and Erikson 1997], we account for this directional by analyzing the normal spread given by the quadric matrix.

Let \( \hat{n}_t \) be the unit normal, and consequently the direction of geometric error, associated with triangle \( t \). Furthermore, let \( \hat{\mathbf{v}} \) be the unit vector from the cluster’s representative vertex that we are computing the error for with the viewpoint. In our anisotropic error projection, we modulate the error associated with \( t \) by the sine of the angle \( \gamma_t \) between \( \hat{n}_t \) and \( \hat{\mathbf{v}} \), i.e. by the factor \( \gamma_t = \| \hat{n}_t \times \hat{\mathbf{v}} \| \). (This modulation factor was also used in [Lindstrom et al. 1996; Hoppe 1997].) Thus, when \( \gamma_t = 0^0 \) implies that we are near a silhouette and the projected error is at a maximum. We can rewrite (the square of) \( \gamma_t \) as follows:

\[
\gamma_t^2 = \| \hat{n}_t \times \hat{\mathbf{v}} \|^2 = \hat{\mathbf{v}}^T \hat{n}_t \hat{n}_t - (\hat{\mathbf{v}}^T \hat{n}_t)^2 = \hat{\mathbf{v}}^T (I - \hat{n}_t \hat{n}_t^T) \hat{\mathbf{v}}
\]

This modulation factor for a single triangle \( t \) can then be extended to a set of triangles \( T \) in a cluster as a weighted sum:

\[
\gamma_T^2 = \frac{\sum_{t \in T} \gamma_t^2}{\sum_{t \in T} \gamma_t^2} = \hat{\mathbf{v}}^T \sum_{t \in T} \left( I - \hat{n}_t \hat{n}_t^T \right) \hat{\mathbf{v}}
\]

\[
\hat{\mathbf{v}}^T \left( I - \frac{\sum_{t \in T} \hat{n}_t \hat{n}_t^T}{\sum_{t \in T} \gamma_t^2} \right) \hat{\mathbf{v}} = \hat{\mathbf{v}}^T \left( I - \frac{\mathbf{P} \Lambda \mathbf{P}^T}{\sum_{t \in T} \gamma_t^2} \right) \hat{\mathbf{v}}
\]

This weight \( \tilde{w}_c \) is the same as the one used to weight triangles in the quadric matrix construction (see Section 4). The expression for the trace of \( A \), \( \text{tr}(A) = \sum_{t \in T} \gamma_t^2 \), follows from the fact that \( \text{tr}(A) \) is linear and \( \text{tr}(\hat{n}_t \hat{n}_t^T) = \hat{n}_t \hat{n}_t^T = 1 \).

We call \( K \) the “curvature matrix,” because it encodes the amount that the surface curves in different directions. From the definition of \( K \), we see that as the normals spread out the curvature grows, the eigenvalues approach a common value, \( K \) approaches \( \frac{1}{3} I \), and as a result \( \eta_T \) becomes more and more isotropic. If on the other hand \( \lambda_1 \) is much larger than the other eigenvalues, then the normals are tightly clustered, the curvature is small, and \( \eta_T \) allows aggressive coarsening when \( \hat{\mathbf{v}} \) is near the dominant normal direction.

### 6.1.4 Bounding Sphere

In order to support fast hierarchical view frustum culling, we compute the tightest possible hierarchy of nested bounding spheres over the vertices in \( H \) (cf. [Lindstrom and Pascucci 2001]). The radius for a node \( p \) located at \( p \), is given by

\[
r_p = \max_{c \in C_p} (\| p - p_c \| + r_c)
\]

where \( C_p \) are the children of \( p \). Using these bounding spheres for view frustum culling, we are guaranteed that a vertex is culled only if and its descendants are not visible. Note that for conservative view culling the bounding sphere should also enclose all edge connected neighbors of a vertex [Hoppe 1997]. However, because the adjacent vertices of a node depend nontrivially on run-time level-of-detail decisions, we cannot compute useful conservative bounding spheres off-line. In practice, our bounding sphere hierarchy is loose enough to not cause any visible artifacts. See Figure 1 for an example of view frustum culling.
7 In-Core Data Structures

In this section we describe the in-core data structures used in our run-time view-dependent renderer. These data structures closely resemble the ones used by Luebke and Erikson [1997], but have been modified to work in an out-of-core setting, where only part of the octree is assumed to be memory resident. That is, we maintain an in-core subset $H'$ of the LOD hierarchy $H$, such that the leaf nodes of $H'$ correspond to the vertices of the adaptively refined mesh. We refer to the subset $H'$ of $H$ as the set of active nodes. The triangles of the refined mesh are those stored in the internal (non-leaf) nodes of $H'$, Each triangle vertex, represented on disk as an octcode $v$ for a leaf node in $H$, is mapped to a proxy vertex—either $v$ itself or its lowest active ancestor. Thus, the actual vertices used for a triangle change dynamically as the mesh is adaptively refined and simplified.

Table 4 lists the octree node data structures that are dynamically allocated at run-time. We have already covered the fields for the vertex data in Section 6, and will here discuss the remaining fields for the node. The triangle data consists of a list of triangles and a list of pointers to triangle vertices that currently reference this node as their proxy. Thus the vertex data type consists of an octcode for a leaf node and an in-core octree node pointer to the vertex’s proxy. Whenever a node is expanded, we need to modify all the triangle vertices that have the node as a proxy, which is accomplished by maintaining a list of back references to those vertices. Similarly, when a node is collapsed, its children’s references are first accessed, and the associated proxies are modified to point to the collapsed node.

In contrast to [Luebke and Erikson 1997], where the entire hierarchy is assumed to be memory resident, we cannot pre-compute the list of back references (or “tris” using their terminology), because these references might point to triangles in nodes that are not active and therefore have not been dynamically allocated. We could of course store these references as non-memory-specific octocodes, but have opted instead to maintain only references to triangles in active nodes, and to update the reference lists on-the-fly as triangles are added and removed. Similarly, we do not construct the triangle list $T$ for a node until it is expanded. We will further discuss the operations on the octree $H'$ in the following section.

The octree node pointers $\text{prev}$ and $\text{next}$ are part of a doubly linked list of all active nodes in $H'$, except for leaf nodes in $H$, which cannot be expanded and for which collapsing has no effect. This list is traversed linearly during run-time refinement, as explained in the next section.

8 View-Dependent Refinement

We are now ready to describe the steps pertaining to the final phase of our out-of-core view-dependent renderer: the run-time component. Because the level-of-detail hierarchy stored on disk may exceed the amount of available memory, we must be careful to page in only the active nodes of the hierarchy. Rather than making use of an explicit paging system, we rely on the use of read-only memory mapping to associate the on-disk hierarchy with a logically contiguous address space, and let the operating system fetch the data from disk when it is first accessed. Similar strategies for out-of-core rendering of large terrain have been employed, for example, by Hoppe [1998] and by Lindstrom and Pascucci [2001]. This approach to data paging is particularly attractive when the refinement and rendering tasks are decoupled and run asynchronously, as is the case in our system. Also, as demonstrated in [Lindstrom and Pascucci 2001], by arranging the data and the accesses to it in a cache coherent manner, it is possible to substantially improve the paging performance. Indeed, our choice of arranging the octree in a coarse-to-fine, breadth-first layout on disk was made intentionally after having been inspired by the quadtree layout in [Lindstrom and Pascucci 2001]. We point out that, since we do not access the memory-mapped hierarchy continuously, but copy and “decompress” the compact on-disk data to a more readily usable in-core structure when a node becomes active, it is entirely possible to use an explicit paging scheme in place of memory mapping. The benefits of doing so include improved scalability (many operating systems limit memory mapping to a 32-bit addressing space), more general geometry cache replacement policies (although the malloc system call, when implemented, provides this functionality), as well as potentially better memory usage if only small pieces of each memory page is needed (which is generally not the case in our system). The downsides of explicit paging include higher paging overhead, the need to keep an explicit “page table” for resident data, and implementing a robust data replacement policy that adapts well to the dynamically changing memory resources in a multiuser/multiprocess environment.

Given the general framework above, we now describe the two tasks of adaptive refinement and rendering.

8.1 Refinement Algorithm

Similar to several other components of our algorithm, our adaptive octree refinement closely follows the strategy employed by Luebke and Erikson [1997]. We begin by creating a single node for the root of the dynamic octree $H'$. During refinement, we make use of two complementary operations; node expansion and collapse. When expanding a node we add its children; when collapsing a node we remove its descendants. Pseudo-code for these steps is listed in Table 5. Note that node expansion is valid only if a node is on the front, i.e., if it is a leaf node in the dynamic octree. Expanding a node requires dynamic memory allocation and a minor processing step to initialize the fields in Table 4. In addition to the implicit caching of on-disk data performed by the operating system, the overhead of this step can be reduced by making use of an explicit caching scheme in place of memory mapping. The benefits of doing so include improved scalability (many operating systems limit memory mapping to a 32-bit addressing space), more general geometry cache replacement policies (although the malloc system call, when implemented, provides this functionality), as well as potentially better memory usage if only small pieces of each memory page is needed (which is generally not the case in our system). The downsides of explicit paging include higher paging overhead, the need to keep an explicit “page table” for resident data, and implementing a robust data replacement policy that adapts well to the dynamically changing memory resources in a multiuser/multiprocess environment.

Given the general framework above, we now describe the two tasks of adaptive refinement and rendering.
8.2 Screen Space Metric

The decision whether to collapse or expand a node is governed by a screen space error metric and an error threshold. In our screen space metric, we make use of the quadric error $\varepsilon$ and position $p$ of a node's representative vertex, as well as the curvature matrix $K$. We saw earlier that the quadric error measures (squared) volumetric errors. For view-dependent refinement, we need to express $\varepsilon$ in terms of a screen space geometric deviation, i.e., in units of length. While there are several ways of doing this, our approach is to assume that the volumetric errors correspond to some hypotheti-
cal volume, e.g., a sphere, and we simply compute the radius of the corresponding sphere and use it as the error term. One nice feature of this mapping is that it is monotonic, i.e., the error of a parent node remains at least as large as its children's. This computation is done once when the node is constructed, and the $\varepsilon$ field in the dynamic
node is assumed to have units of length.

As a base metric, we set the screen space error to be proportional to the ratio of the object space error and the distance from the view-
point to the node. As suggested in Section 6.1.3, to incorporate directionality into our metric for silhouette preservation, we modu-
late the base metric by the factor $\eta$. Thus, our screen space metric $\rho$ can be written as

$$\rho = \lambda \eta \frac{\varepsilon}{d} = \lambda \varepsilon \frac{\sqrt{v^T K v}}{v^T v}$$

where $d = ||v||$, $v = e - p$ is the vector from the node to the viewpoint $e$, and $\lambda$ is the screen resolution in pixels per radian.

The metric $\rho$ has a nice geometrical interpretation, due in part to our definition of $\eta$. Using the law of sines, one can show that

$$\eta \varepsilon \frac{v}{d} = \sin \theta \frac{v}{d} = \sin \phi_i \frac{||v + c_i n||}{||v||} \approx \phi_i$$

where $\phi_i$ is the visual angle (in radians) subtended by the point-
to-surface displacement $c_i n$ between $p$ and $t$. The relative error in
this approximation is bounded by $\frac{1}{\sqrt{1 - \varepsilon^2}}$, and is thus low when $\varepsilon \ll d$, which is generally the case. As a result, $\xi$ is (to a good approxima-
tion) a weighted average of the angular extents of the surface
errors.

To determine whether a node is active or not, we compare the screen space error $\rho$ against a user-specified threshold $\tau$. For effi-
cyency reasons, we square and rearrange some terms, and obtain the
following expression:

$$\text{active} \iff \rho > \tau$$
$$\iff \lambda^2 \frac{\varepsilon^2 (v^T K v)}{||v||^2} > \tau^2 (v^T v)^2$$
$$\iff \varepsilon^2 (v^T K v) > \kappa^2 (v^T v)^2$$

where $\kappa = \frac{\tau}{\lambda}$ is the angular error threshold in radians.

Because the octree is pruned whenever a node is found to be
inactive, we should ideally ensure that a node's projected error is always larger than those of its children. Note that the quadric
errors by their additive nature already satisfy this nesting property.

Similarly, the amount of curvature encoded in the matrix $K$ can
only increase when quadric matrices are combined. However, the
direction and length of the vector $v$ vary from node to node, mak-
ing it possible to violate the nesting condition. A general technique
for handling this view-dependent problem was presented in [Lind-
strom and Pascucci 2001], in which a nested sphere hierarchy is
computed and used in place of the positions of individual vertices.
This sphere hierarchy is the same as the one we compute for view
frustum culling purposes. While not implemented here, we believe
that it would be rather straightforward to incorporate the sphere hi-
ernarchy into our screen space metric.
9 Results

In this section we present experimental results of running our algorithms. We used a number of polygonal test models, including a massive 373 million triangle model of Michelangelo’s St. Matthew statue [Levoy et al. 2000]. All models, except the Buddha, were simplified on one processor of a 250 MHz R10000 SGI Onyx2 with 40.5 GB of main memory. The Buddha model was simplified on a Linux PC with two 800 MHz Pentium III processors, 880 MB of RAM, and a GeForce3 graphics card. We used this PC for the accompanying video of the Lucy data set. To stress the out-of-core aspect of our system, we used a lower-end dual-processor PC, with 256 MB of RAM and GeForce2 graphics, for the Isosurface video.

Animations We have prepared animations to illustrate the runtime performance of our system.1 Frames from these videos and other images are shown in Figures 3–5. As is evident in the Lucy sequence, we obtain a throughput of up to three million rendered triangles per second using a GeForce3 graphics card and immediate mode rendering. Notice the ability to adapt the mesh to the view frustum, the shape of the surface, and the presence of silhouettes (see, for example, the neck of the statue). The second animation shows a very dense and complex isosurface of the turbulent boundary between two mixing fluids. Because we used backface culling, it is possible to see through much of the back side of the surface (the majority of the video sequence). This video illustrates the progressive nature of the refinement, as mesh updates are made at least once per second, and the panning of data from disk as the mesh is navigated at various scales. The slower rendering speed can be attributed to a lower performance graphics card, as well as memory contention due to the limited amount of memory available (less than one sixth the on-disk size of the data set). Note that we were careful to invalidate the operating system’s disk cache before creating these animations to ensure that no data was memory-resident at startup.

Disk and Memory Usage Table 6 lists numerical results for our off-line method, including the number of triangles, grid size \( (2^n) \), execution time, effective triangle processing rate, and disk usage. Not included in this table is the maximum memory usage. Our method uses only a constant \( O(1) \) amount of memory: 5 MB for the PC and 8 MB for the SGI (the difference is due to different size executable files). Note, however, that the operating system may use additional memory to cache parts of the intermediate files generated by our system, e.g., during sorting. The temporary disk usage of our method can be shown to be a constant multiple (roughly a factor of 5) of the size of the input mesh. While the theoretical usage is linear in both the size of the input and the output, the simplification phase often reduces the mesh to the extent that the overall temporary disk space is entirely dominated by the plane file (which is removed before the octree construction begins) and any disk space used while sorting this file. As can be seen from the last column, the size of the octree output file is on average only a few percent larger (in bytes per triangle) than the size of the input, and ranges from 35–45 bytes per triangle. For the sake of fairness, our triangle soup input format is not as efficient as the more common indexed mesh representation, which requires only half as much storage. The indexed mesh, on the other hand, is impractical for external memory algorithms since it requires random access.

Execution Time As can be seen from the table, for small octrees the total time is dominated by the external sort phase of the simplification. As the octree (and thus size of the output) grows larger, the output phase of the octree construction becomes more significant, but generally requires only a small additional fraction of the simplification time. Even though the off-line construction is done entirely on disk, our memory insensitive algorithm is remarkably fast, and yields an effective triangle processing rate (measured as the size of the input over the total time) of roughly 20,000–60,000 triangles per second (tps). As a point of reference, El-Sana and Chiang [2000] report a reduction rate of roughly 5,300 tps for their largest model, consisting of 1.2 million triangles. This model is considerably smaller than some of those simplified here, and the nature of their method suggests that it would steadily decline in performance as the input grows larger. The method proposed by Prince [2000], while producing high quality meshes, yields about 1,000 tps for the largest model used (11.4 million triangles). Meanwhile, his method requires more than 512 MB of RAM to simplify this model, whereas we use a constant 8 MB. Closest in performance to our method is the one by Cignoni et al. [2002], with up to 13 thousand tps.4 However, the output of their method is not in a form that is directly usable for view-dependent refinement.

The theoretical execution time of our off-line processing is \( O(T_{simp} + T_{out}), \) assuming the external sort is implemented as a radix sort,5 which suggests that our method scales well.

10 Summary and Future Work

We have described a method for constructing a level-of-detail hierarchy for large polygonal meshes. This method performs all computations on disk, and requires only a small, constant amount of RAM. The method uses temporary disk space linear in the size of the input and output, and runs in linear time. Even though virtually no RAM is used, our method executes one to two orders of magnitude faster than previous methods, and achieves a peak simplification rate of over 60,000 triangles per second. We have also presented compact data structures and a suitable error metric for performing out-of-core view-dependent refinement of the resulting mesh hierarchy. Our results show that we obtain interactive frame rates and a throughput of up to three million triangles per second using immediate mode rendering.

4 The performance of their method drops to around 6 thousand tps when their one-time external mesh construction phase is accounted for.

5 Our external sort is a combination of radix and merge sort.
We see several avenues for future work. One limitation of our approach is that the multi-resolution surface constructed does not retain the fidelity of the original mesh. One possible solution to this problem is to keep the triangles that degenerate during simplification and store them with their corresponding leaf nodes. Another limitation is that our 32-bit octocodes support a maximum octree height of 10. Using 64-bit codes would allow a finer grid and less coarsening of the input mesh, although at the expense of larger files. Our run-time system could be improved by using geomorphing to smooth out popping artifacts and prefetching to increase the paging performance, e.g. using separate threads that visit nodes below the active front. We will also investigate how to extend our metric to guarantee nested screen space errors, and how to account for the visual impact of simplification on the shading of the surface.

Finally, we envision that our off-line algorithms can be further enhanced. As noted in [Lindstrom and Silva 2001], the greatest potential for reducing the disk usage lies in using a compressed representation for the rather large file, which by far dominates the temporary storage needed. We also see untapped potential in increasing the speed of our out-of-core method through parallelization. The locality of our data accesses and the octree partitioning of space and work naturally lend themselves to parallel execution.

Acknowledgements

This work was performed under the auspices of the U.S. DOE by LLNL under contract no. W-7405-Eng-48. We thank Stanford University and the Digital Michelangelo Project for providing the Buddha, Lucy, and St. Matthew data sets.

References


Out-of-Core Construction and Visualization of Multiresolution Surfaces: Lindstrom

Figure 3: Frames from the Lucy video. View frustum culling is done against the violet rectangle. (c) The hand is moving into the view, but has not been fully refined yet.

Figure 4: Frames from the Isosurface video. The full multiresolution mesh contains 47 million triangles. (a) The relatively smooth top layer of the dense surface. (b), (c) Close-ups of fine structure on the back side.

Figure 5: View-dependent renderings of the St. Matthew data set for various error thresholds and viewpoints. The original model contains 373 million triangles. (a), (b) Gouraud shaded meshes. (c), (d), (e) Flat shaded meshes and close-ups of the face.