Hierarchical Data Structures for Interactive Volume Visualization

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Abstract

In order to interactively investigate large-scale 3D data sets, we propose an improved hierarchical data structure for structured grids and an original hierarchical data structure for unstructured grids. These multi-tiered implementations allow the user to interactively control both the local and global density of the mesh. Therefore, the user can interactively refine areas of interest and decimate peripheral regions. By controlling the density of the mesh throughout the volume, the user controls where computational cycles are spent and gains a deeper insight into the geometric structure of the mesh.

1 Introduction

Computational scientists and engineers are producing ever larger sets of volume data. While the specific application problems encountered by computational scientists vary widely across (as well as within) disciplines, the computational structure of the collected data varies remarkably little and is usually represented in the form of discrete approximations of scalar and vector fields. In general, the scalar and vector field data are measured or computationally simulated at discrete points throughout a volume. The main spatial variance among these volume data sets is typically characterized by whether the grid is structured or unstructured. Interactively visualizing the geometrical structure of underlying models and simulation results for large-scale 3D problems on both structured and unstructured grids remains a significant challenge.

Interactively visualizing models and simulation results presents both an aesthetic and a computational challenge. These challenges are especially important in visualizing data on unstructured grids, when users often wish to view the mesh structure in order to confirm mesh refinement within regions of high gradients. The aesthetic element of the visualization is important for more than merely aesthetic reasons: the mere rendering of myriads of unstructured mesh elements on a 2D screen is generally not informative. On the computational side, the time required to explore such large solution spaces has traditionally been prohibitive to real-time interaction.

The ability to perform mesh and volume visualization interactively affords the user a global, high-level view of the solution space and then allows her/him to probe more deeply in to selected areas of interest. This notion of "closing the loop" gives the user more direct control of the calculations being performed. By employing a "computational steering" paradigm, the user is no longer required to wait for lengthy calculations to complete before seeing results. Instead, the user can guide the computation by feeding back parameter changes to the system based on initial, often partial results from a larger calculation [14]. Integrating the user into the computational visualization process means faster convergence time for the user to achieve the desired results, and, ultimately, increased productivity.

A variety of volume visualization techniques exist for probing 3D fields and vary based on the type of field being queried (i.e. scalar, vector, tensor, or a combination of these), and the type of information being sought. Some examples of these techniques include isosurface extraction, streamline tracing, and volume rendering. In this paper, we focus our attention on the isosurfacing algorithm, comparing performance times of this volume visualization methods implemented with and without our data structures.

In addition to visualizing volumes, our data structures are used for interactively visualizing large-scale, unstructured meshes. For this challenge we focus solely on the node locations and connectivities (i.e. the elements) of unstructured fields, independent of the values stored at those nodes. Scientists are interested in probing the structure of meshes, because scientists have increasingly found that solution accuracy is inextricably tied to mesh discretization. For example, in the finite element (FE) method, the error across a tetrahedral element is often directly proportional to the gradient across that element. In order to reduce the total error in an FE solution, one can refine the mesh in the areas of high gradients, and the system can then be re-solved [1, 2, 3, 4]. As mesh element size decreases, the result is guaranteed to converge to the analytic solution [4]. Unfortunately, the mesh refinement techniques most frequently used by scientists often yield widely varying densities within the support mesh, thereby creating wide discrepancies in the accuracy of the visualization. We have found, however, that by storing the mesh hierarchically, we can allow the user to interactively refine and de-refine the mesh locally as well as through local regions of interest. We will show that this interaction greatly increases the clarity of the mesh and decreases the time associated with understanding the structure of the mesh. Furthermore, we will compare, both quantitatively and qualitatively, our mesh visualization techniques with others currently in use.

2 Background

Historically, a common method for interactively visualizing large volume data sets has been to operate on just a subset of the domain at a time. This reduction serves two primary functions. First, it limits the amount of computation being performed, because the algorithm ignores areas outside of the reduced domain. Second, it reduces the amount of "screen clutter" created when the results are rendered to the display. Examples of these types of domain reducing algorithms range from clipping plane and seeding algorithms [5], to complete automatic refinement in regions containing information of interest [6]. Clipping planes and seeding algorithms localize the region of operation and allow tight user control over the spatial domain. In contrast, completely automatic refinement limits the computational domain, but does not support a means for interactively constraining where work is done spatially.

2.1 Structured Grids

The complete automatic refinement algorithm is an isosurfacing algorithm which was introduced by Wilhelm and Van Gelder in 1992 [6]. The strength of this algorithm comes from its efficient use of octrees to store the volume data. The hierarchical nature of octree space decomposition enables the algorithm to trivially reject large portions of the domain, without having to query any part of the subtree within the rejected region. The original octree data structure was proposed independently at approximately the same time by Hunter et.al. [7], and has found application areas in mesh generation [8], modeling [9] and rendering [10], to name just a few. In Wilhelm and Van Gelder's implementation, their Branch-On-Need Octree (BONO) algorithm stores the maximum and minimum scalar values of the region spanned by each subtree in the region's parent node. The algorithm can then recursively traverse the tree, only isosurfacing those subtrees with ranges containing the value being sought. They report overall speed-ups between 132% and 321% when comparing their algorithm to the marching cubes isosurface extraction algorithm [11]. In this paper, we present a further extension to the octree data structure. Our extension allows the user to interactively control the traversal depth within different regions of the tree. With this improved octree, the user can now interactively control how detailed the extracted isosurface will be locally within the domain, and can thus reduce the amount of time spent computing outside the area of interest.

2.2 Unstructured Grids

The second data structure we have implemented is a counterpart to the octree data structure. This new structure applies data hierarchies to unstructured fields. This structure has enabled us to address two difficult problems: unstructured mesh visualization and unstructured volume visualization. The first problem arises solely from the unstructured nature of the geometric mesh, and trying from the need to visualize the structure of the grid elements in a way that is meaningful to the researcher. For small meshes containing less than a couple of dozen elements, this problem does not become an issue. All of the elements can be rendered to the screen, and the user can inspect the mesh with little or no difficulty. But when the mesh expands to hundreds, thousands, or even millions of elements, simply rendering all of these cells no longer conveys practical information. Current research in this area has focused on methods to limit the domain of the mesh, operating on only a subsection of the volume at a time. Examples of this type of method include clipping planes and seed/grow algorithms [5]. In the clipping planes algorithm, elements (or parts of elements) are

displayed only if they fall within the volume created by the planes. The seed/grow algorithm allows the user to choose a seed element from the mesh and a depth parameter, and then renders all elements connected within that depth of the seed element. These algorithms are very useful for exploring small local regions of large meshes, but they fail to convey a sense of the global structure. The second difficult problem associated with unstructured field visualization is that of how to interact with large fields using volume visualization algorithms. Here again it is possible to extract some smaller subregion from the field, and accordingly to operate only on that local partition, but the same "tunnel-vision" problem is encountered here. The user sacrifices any global view of the problem, and is allowed to interact only with small subsections of the field at a time.

In this paper, we propose a new hierarchical data structure for storing unstructured fields. This new structure contains a range of resolutions of the original mesh, with lower levels of the hierarchy containing more refined representations. This implementation allows the user to interact with and visualize the entire domain at varying levels of resolution, and to move seamlessly between those levels for rapid exploration of the field.

3 Methods

3.1 Octrees

We have extended the original octree data structure to contain a "LEAF" bit at each node. This extension provides the user control over how deeply an algorithm will search a path of the octree for data. The construct works in this way: for every path from the top of the octree to the bottom, exactly one of the visited nodes will have a LEAF flag set. This new construct serves many applications. For example, a user isosurfacing a large field would often find it useful to get a general sense of the topology of the surface through the field before generating the surface in full detail. In this way, the user can interactively isosurface even a very large field. Furthermore, the user can select a very fine discretization level (i.e. move the LEAF bits down in the octree) in areas of determined interest, and select a lower discretization level (i.e. move the LEAF bits up in the octree) in areas that are not as critical. This gives the user fine-tuned control over where the field operator (i.e. isosurfacing algorithm) is spending its compute time.

An attractive feature of this data structure is that algorithms such as isosurfacing need not recompute in areas that have not changed as the user locally refines and derefines the structure. We can cache the isosurface portions generated through regions and recompute them only when that region changes or when the threshold value is changed. This allows the user to deeply probe local regions without spending time to recompute old information.

Several convenient methods have been added to the octree data structure to allow the user to control the LEAF bits in the tree. These methods give the user the ability to push all of the bits to the top, to push all of the bits to the bottom, to move all of the bits up or down one level, and to move all of the bits in a specific region up or down one level. All of these operators are implemented in such a way as to maintain the integrity of the structure (i.e. there is always exactly one LEAF bit set along any path in the tree). They are controlled via a simple user interface consisting of a cross-hair widget [12] and a set of Tcl/Tk push-buttons [13]. Fig. 1 shows the SCIRun [14] visual programming map constructed for this algorithm. The cross-hair widget can be interactively moved through the field to identify regions of the octree, and the buttons are linked to method callbacks which move the LEAF bits. All of the interactions are very fast since they involve only testing and setting bits, but formally the global operations are O(n), and the local operations are $O(\log n)$.

There is some overhead associated with using an octree representation of a field as opposed to a generic 3D array. To build the octree takes $O(n \log n)$ time, but this structure needs to be built only once and can then be stored as a persistent object. Accessing the value at a point in the field is slower with the augmented octree, now requiring $O(\log n)$ time, instead of just the constant time required to access a field value from an array or from an intelligently implemented non-augmented octree. Octrees can also be a memory liability if they are not stored correctly. Wilhelm and Van Gelder showed that efficiently implemented octrees can take up less that 15%of the memory required to store the field as a straight 3D array. However, because we store the eight corner values in each node, we fail to show savings in our memory consumption. Furthermore, memory overhead is required to store the LEAF bit at each node, as well as to store all of the pointers to the children nodes. In the end, our structure requires approximately twice as much memory as the generic array structure. The final disadvantage we have found with utilizing the augmented octree data structure is that the isosurfaces created across regions discretized to different grades can be geometrically discontinuous. These discontinuities are generally small jumps across the disparate regions, but they are noticeable and should be mentioned, if for no other reason than to avoid confusion.

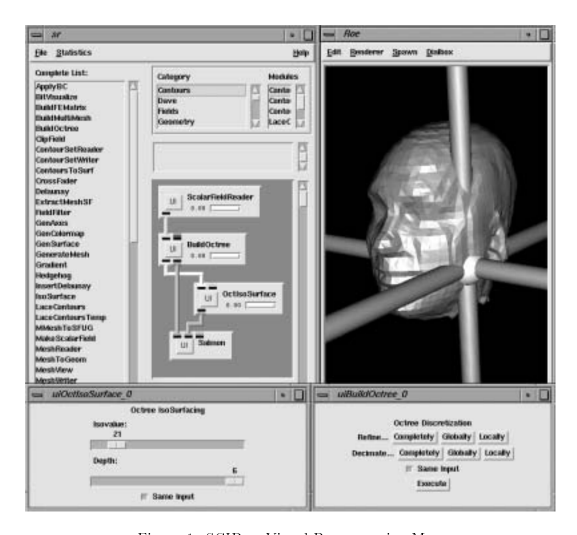


Figure 1: SCIRun Visual Programming Map

3.2 Multimesh

Our solution to the problem of managing large unstructured fields for interactive visualization is to break the original mesh into a hierarchy of sub-meshes which represent some increasing percentage of the original mesh. We call this data structure a "multimesh." A similar technique, detail elision (progressive refinement), has existed for a long time in computer graphics for representing objects in a scene at different levels of detail based on their distance from the viewer. The seminal work on this topic was conducted by Clark [15] and continues to be used today in progressive refinement for architectural walk-throughs and other interactive applications [16]. For our multimesh data structure, the bottom level of the multimesh hierarchy contains the original mesh, with all of its nodes and connectivities. Each level up contains a subset of the nodes on the level below and a new set of connections for those nodes. At the highest level there are just a few nodes left in the mesh. Once this data structure has been generated from the original mesh, the user can interactively move between the levels of the hierarchy to visualize the mesh structure and field data. Effectively, the user now has multiple representations of the original mesh, each with a different number of nodes and elements. The coarsest representations might be used for gaining a global sense of the field, whereas more detailed levels would be useful for viewing the exact behavior of the mesh. Since the multimesh is stored as an array of independent meshes, it is simple to extract a mesh from the hierarchy. Furthermore, it is possible to then pass the extracted mesh into a filter if the user wants to interact with a localized region of the field.

The major issues associated with constructing a multimesh data structure involve deciding which subset of nodes will be contained in each higher level and determining the connections of those nodes. A simple solution is to decide on a "survival percentage" that indicates what percentage of the nodes on a given level will exist at the level above, and then randomly selecting enough nodes to fill that percentage. This method has the attractive property that the mesh is evenly de-refined, or decimated, throughout. Areas that were very finely meshed to begin with maintain a proportionately high percentage of the nodes at every level, and the global structure of the mesh, though coarser, is maintained at all levels. For example, imagine we are given an unstructured mesh with 100,000 nodes, and are asked to create a multimesh hierarchy with a 50% survival percentage and 6 hierarchical levels. The resulting data structure would contain an array of meshes, the sixth mesh containing all 100,000 nodes, the fifth containing approximately 50,000 nodes, the fourth approximately 25,000 nodes, the third approximately 12,500 nodes, the second approximately 6,250 nodes, and the first approximately 3,125 nodes. All of these meshes will have similar node densities, and all will approximate the original mesh, but the variance in the number of nodes (and correspondingly the number of elements) from one level to another makes the lower levels of the mesh interactive for volume visualization, whereas the original mesh may not have been.

We have improved on this method to allow the user to interactively select which areas of the mesh are comparatively more important, and to skew the node survival percentages such that important nodes have a greater chance of surviving to the next level. We have accomplished this by building an "importance weighting field" in 3-space, where each node can look up its importance based on its geometric location. The user interactively places point sources into the scene and defines the "charge" and "fall-off rate" associated with each of these sources. If a source has a large positive charge and a high fall-off rate, the nodes in its immediate vicinity have very high chances of surviving to lower levels. Alternatively, a source with a small negative charge and low fall-off rate can be placed in a region of the mesh which is less important, and the nodes in that region will be less likely to survive to lower levels. In this way the user gains local, as well as global control of the mesh decimation process, and the constructed multimesh will enable the user to more quickly understand the unstructured field. However, it turns out that it is desirable to make the system somewhat random, in order to remove a uniform sampling of the nodes. To accomplish this we associate a random weighting value with each node. In a non-random system, the nodes with the lowest weightings would be removed at each higher level in the structure. In contrast, a completely random system would ignore the node weightings altogether and randomly choose which nodes to remove. The compromise we implemented between these extremes allows us to remove nodes based on a linear combination of a node's actual weighting and it's random value. The linear combination is based on the chaos bias value b between 0 and 1, such that n = b * r + (1 - b) * w. When the chaos bias is 0 the node levels are completely dependent on the source charges in the scene, and when the chaos bias is 1 the node levels are determined completely randomly.

There are generally two ways to build the mesh connectivities for each level of the hierarchy – top down, or bottom up. Bottom up construction requires first copying the existing mesh into the bottom level of the hierarchy, and then iteratively removing the nodes and re-meshing as we construct higher levels. The difficulty raised is that of how to re-mesh areas of the mesh when nodes are removed. Yu implemented an algorithm for local mesh derefinement [17], and his methods are very successful if boundary nodes are never removed, the domain is convex, and the original mesh is Delaunay. However, as this is not always the case (especially with regard to the removal of boundary nodes), we have implemented a top-down construction. For this construction, we ignore the original mesh connectivities, and, adding nodes in order of importance, construct the levels of the mesh from the top down. Each node is inserted via a Watson's style algorithm [18], and as a result remains Delaunay at all

times. The top-down method is considerably faster than the bottom-up approach, requiring on the average $O(n \log n)$ time to build all levels of the multimesh.

The multimesh data structure exploits the redundancy of nodes in multiple levels of the hierarchy in two ways: by storing only one copy of each node, and by storing pointers rather than full node structures in each mesh. Unfortunately, there is not a corresponding feature for connectivities, and each mesh in the hierarchy must maintain its own list of element connectivities. The space requirements for a multimesh depend very much on the depth of the hierarchy and the "global survival rate" parameters.

The primary weakness of this method, as we have implemented it, is that within the use of the method non-Delaunay meshes are currently destroyed. We have no mechanism to account for or respect the topology of non-convex meshes. If a fast, reliable version of such a mechanism were found or developed, the above multimesh creation algorithm could be easily rewritten in a bottom-up manner, and the resulting meshes throughout the hierarchy would maintain the original non-Delaunay structure. This is an area for future work.

4 Results

We have applied our improved octree data structure to isosurface extraction of a magnetic resonance image (MRI) data set. The original scan is of a patient's head and contains 32 axial slices with 256×256 pixels per slice. We resampled this data to construct a $32 \times 32 \times 32$ grid. We used this data set to explore our isosurfacing algorithm. All of our timings were done on an Indigo2 Extreme with an R4400 processor and 128MB of memory.

We extracted an isosurface from the volume corresponding to the value of skin. Our algorithm required a set-up time of 3.54 seconds to construct the octree data structure. This set-up time is somewhat negligible though, since the octree only needs to be computed once. We then applied the isosurfacing algorithm for different levels of mesh refinement, and compared our results to those generated via a marching cubes style algorithm [11]. Marching cubes required 0.64 seconds to extract the isosurface. Wilhelm and Van Gelder's BONO algorithm [6] was somewhat faster, requiring only 0.43 seconds to execute. The execution time for our algorithm varied linearly, depending on how deeply the mesh was discretized. The isosurface in Fig. 2 was generated from the completely refined octree, and also required 0.43 seconds. As an example

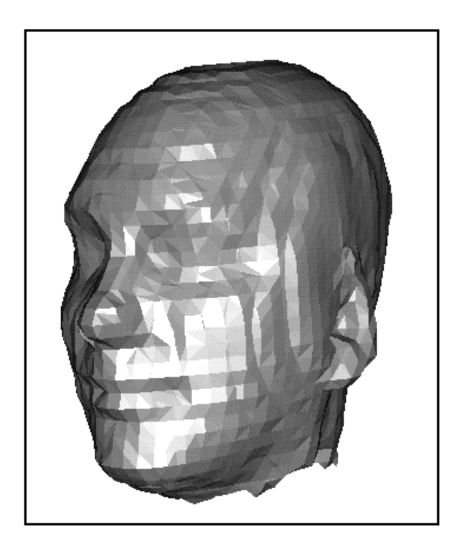


Figure 2: Fully Refined Octree Isosurface

of locally de-refining peripheral regions of the mesh, we focus our attention on the side, front side of the head, and de-refine peripheral regions. We were able to reduce the isosurfacing time by nearly half, to 0.22 seconds, after locally de-refining the back half of the head as seen in Fig. 3. Further reducing the domain, we de-refine the left side of the head, and are now able to isosurface the field in 0.12 seconds.

Next, we applied the multimesh isosurfacing and mesh visualization algorithms to a simulated voltage field computed through the same cranial domain described above. We constructed this field by simulating the effect of a dipole source located in the right frontal lobe. The charge and orientation of the dipole were consistent with experimental neural activity recorded from epilepsy patients during seizures. To construct

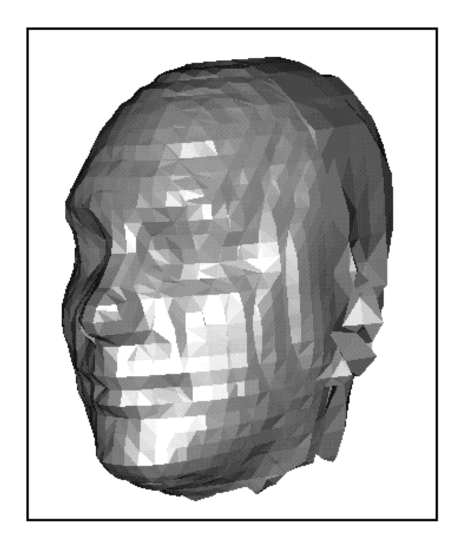


Figure 3: Half-Refined Octree Isosurface

the anatomical domain for the simulation, the MRI data set was first segmented into distinct material regions. These regions were then discretized into 137,923 nodes and 884,343 tetrahedral elements, which form the support structure for the field. After the mesh was constructed, Poisson's equation for electrical conduction was approximated using the FE method. This simulation results in scalar and vector values, which are subsampled at 1000 nodes. We used this simulation data to test our multimesh visualization methods. The right, front side of the head is again chosen as the region of interest, and we generate 7 levels of mesh refinement for our multimesh hierarchy. Isosurfacing the fully refined field required 0.82 seconds of CPU time, and the mesh at this level is too dense to yield any insight to the viewer. Switching to a less refined version of the mesh, we are able to gain real intuition to the mesh structure, and to reduce our isosurfacing time to 0.12 seconds, with little change in the quality of the surface.

5 Conclusions

Through original and improved hierarchical data structures, we have been able to substantially improve the interactivity of mesh and volume visualization. These data structures allow the user to identify regions of interest and then interact with the visualization algorithms to constrain work to be done only in those regions.

There are many opportunities to further improve the data structures that we have implemented. Specifically, we intend to implement Wilhelm and Van Gelder's octree architecture, which relies on maintaining full nodes at the bottom of the octree and partial nodes at the top. This restructuring provides for fuller trees with ultimately fewer nodes. In addition, we would like to incorporate their hashing scheme for visiting nodes during isosurfacing to further improve our performance.

Based on the success we found applying our hierarchical data structures to isosurface extraction, we plan to extend these data structures to more volume visualization techniques. We intend to integrate both hierarchical data structures into streamline tracing and volume rendering algorithms.

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