# Best Quadratic Spline Approximation for Hierarchical Visualization 

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#### Abstract

We present a method for hierarchical data approximation using quadratic simplicial elements for domain decomposition and field approximation. Higher-order simplicial elements can approximate data better than linear elements. Thus, fewer quadratic elements are required to achieve similar approximation quality. We use quadratic basis functions and compute best quadratic simplicial spline approximations that are $C^{0}$-continuous everywhere. We adaptively refine a simplicial approximation by identifying and bisecting simplicial elements with largest errors. It is possible to store multiple approximation levels of increasing quality. We have tested the suitability and efficiency of our hierarchical data approximation scheme by applying it to several data sets.


Categories and Subject Descriptors (according to ACM CCS): I.4.10 [Image Processing and Computer Vision]: Hierarchical; I.4.2 [Image Processing and Computer Vision]: Approximate Methods

## 1. Introduction

The trend in science and engineering applications has been to produce larger data sets since computers and imaging technology are getting faster and storage space is increasing. Large amounts of data are difficult to visualize and it is impossible to directly visualize on inexpensive computers. Many visualization techniques exist that visualize certain types of large data, however, a general solution does not exist. A hierarchical method provides the foundation for a solution. Linear and quadratic decomposition elements can be used to form an approximation hierarchy representing large data; a user can then visualize this hierarchy on an inexpensive machine.

We only consider quadratic simplicial elements. In the 2D case, we use quadratic triangles whose edges are straight line segments; in the 3D case, we use quadratic tetrahedral elements whose edges are straight line segments and faces are planar triangles. We use a linear transformation to map
the so-called standard simplex to the corresponding simplicial region in 2D/3D space. Furthermore, we use a quadratic polynomial defined over each simplicial element to locally approximate the dependent variable(s).

Our overall goal is the construction of a hierarchical data approximation over 2D or 3D domains using a bestapproximation approach based on quadratic polynomials defined over the simplices defining the domain. Our approach belongs to the class of refinement methods. These methods are based on the principle of refining intermediate data approximations by inserting additional points or elements until a certain termination (error) criterion is satisfied. We have developed our method with a focus on massive scientific data visualization, see ${ }^{13}$. To enable interactive frame rates for massive data visualization, it is possible to either use lowresolution best approximations everywhere or to adaptively "insert" high-resolution approximations locally into an oth-
erwise relatively coarse approximation. The overall approximation algorithm is based on these steps:

- Initial simplicial domain decomposition. We construct a coarse simplicial decomposition of the domain. (The linear transformations, mapping the standard simplex defined in parameter space to simplices in physical space, are defined by specifying corresponding point pairs in the two spaces such that one obtains a one-to-one, bijective mapping.)
- Best approximation. In the 2D case, each simplicial element has six associated knots, one knot per corner and one knot per edge. Six knots in parameter space are associated with six points in physical space, and this defines the needed mapping for a simplex. (Accordingly, the number of knots is ten in the 3D case.) For simplicity, we consider only knots that are uniformly distributed along the edges of the standard simplex. We associate a quadratic polynomial with each simplicial element that approximates the dependent variable(s) over the corresponding region in space. We represent each quadratic basis polynomial in Bernstein-Bézier form, see ${ }^{6}$. Assuming that the function to be approximated, typically a scalar- or vector-valued function, is known in analytical form, it is possible to compute the unique best quadratic spline approximation defined as a linear combination of a set of quadratic basis functions. The best approximation, understood in a least squares sense, is the result of solving the normal equations, see ${ }^{5}$.
- Adaptive bisection. We compute a local error value for each simplicial element once a best approximation is computed. We use the $L_{2}$ norm to compute simplex-specific error values. The set of simplices is ordered according to these simplex-specific, local error values. To compute a "next-level" best quadratic approximation, we determine a certain percentage of simplices with largest error values and bisect them by splitting them at the midpoint of their longest edge. If a simplex's longest edge is not unique, we choose the edge randomly. Splitting a simplex into two simplices induces additional splits for all those simplices that share the split edge. We update a simplicial domain decomposition by considering all edge bisections and computing a new best approximation. We repeat the process of identifying simplices with largest errors, bisecting these simplices, and computing a new best approximation until we obtain an approximation for which a global approximation error is below a user-specified error threshold, or until a user-specified maximal number of simplices is reached.
- Hierarchical data representation. To support level-ofdetail visualization we can store multiple best approximations of different resolutions. For each best approximation, we need to store the polynomial coefficients of each simplicial element. We store a fixed number of best approximations such that either the number of simplices increases in a specified fashion or the maximal simplex-


Figure 1: Correspondence between 2D basis functions $B_{i, j}^{2}$ and knots (indicated by bullets and circles) in uv-parameter space.
specific error decreases in a certain way from one resolution to the next.

We discuss these steps in more detail in the following sections.

## 2. Previous Work

Related work in the areas of hierarchical data representation and visualization is discussed in ${ }^{12,17,20,27}$. Simplification methods (methods that begin with a high resolution of data and then simplify by removing data) are described in 10, 14, 18, 29, 30 . Wavelet methods, in general, work well for rectilinear 2D and 3D grids and are described in 2,3,24. Refinement methods (methods that begin with few data and then refine by adding more data), similar to our method, are described in ${ }^{15,16}$. Data-dependent triangulation schemes, i.e., schemes concerned with the construction of piecewise linear approximations using near-optimally shaped and placed simplicial elements, are described in ${ }^{22}$. From a more general perspective, our work is also related to grid generation, and references for this area are ${ }^{8,19,28}$. Finite element methods are discussed in ${ }^{31}$.

## 3. Mapping the Standard Simplex

In the 2D case, the standard simplex in parameter space is the triangle with corners $(0,0),(1,0)$, and $(0,1)$. We associate a 2D quadratic Bernstein-Bézier polynomial $B_{i, j}^{2}(u, v)$ (abbreviated by $B_{i, j}^{2}$ ), defined as

$$
\begin{align*}
& B_{i, j}^{2}(u, v)= \\
& \quad \frac{2!}{(2-i-j)!i!j!}(1-u-v)^{2-i-j} u^{i} v^{j},  \tag{1}\\
& i, j \geq 0, i+j \leq 2
\end{align*}
$$

with each corner and midpoint of each edge. The six basis polynomials correspond to the six knots $\mathbf{u}_{i, j}=\left(u_{i, j}, v_{i, j}\right)=$ $\left(\frac{i}{2}, \frac{j}{2}\right), i, j \geq 0, i+j \leq 2$ in parameter space, see Figure 1.

In the 3D case, the standard simplex is the tetrahedron with corners $(0,0,0),(1,0,0),(0,1,0)$, and $(0,0,1)$.

We associate a 3D quadratic Bernstein-Bézier polynomial $B_{i, j, k}^{2}(u, v, w)$ (abbreviated by $B_{i, j, k}^{2}$ ), defined as

$$
\begin{align*}
& B_{i, j, k}^{2}(u, v, w)= \\
& \frac{2!}{(2-i-j-k)!i!j!k!}(1-u-v-w)^{2-i-j-k} u^{i} v^{j} w^{k},  \tag{2}\\
& i, j, k \geq 0, \quad i+j+k \leq 2
\end{align*}
$$

with each corner and edge. The ten basis polynomials correspond to the ten knots $\mathbf{u}_{i, j, k}=\left(u_{i, j, k}, v_{i, j, k}, w_{i, j, k}\right)=$ $\left(\frac{i}{2}, \frac{j}{2}, \frac{k}{2}\right), i, j, k \geq 0, i+j+k \leq 2$ in parameter space.

## 4. Initial Decomposition

The main objective driving the development of our method is the hierarchical representation of very large scientific data, where real-time and adaptive data visualization are crucial. Data sets resulting from computational simulations are typically defined on a grid, and the dependent variables are associated with either the vertices, also called nodes in the finite element literature, or the elements defining the grid. Either of these types of data can be approximated. Triangulating the convex hull of the original set of data sites with a coarse triangulation defines an initial decomposition of the domain. The 3D case requires us to construct a tetrahedralization of the convex hull. We decompose quadrilaterals with two triangles (2D case) hexahedra with five tetrahedra (3D case). The elements we use are geometrically linear. However, there are quadratic polynomials defined "over" them that approximate the dependent field variable(s).

## 5. Best Approximation

We assume that the field to be approximated over a domain is known analytically. Should this not be the case, e.g., in the case of scattered data (when one is given a set of randomly distributed points with associated function values without connectivity information), it is possible to construct an analytical representation by performing a prior data interpolation or approximation step, see ${ }^{7,23}$. In the case that a data set is defined on a grid, the required analytical definition is given by a piecewise linear function for a simplicial (triangular or tetrahedral) grid and a piecewise bilinear/trilinear function in the case of quadrilateral/hexahedral grid cells. We denote the analytical function to be approximated over the domain by $F(\mathbf{x})$ (abbreviated by $F$ ). Based on an initial simplicial domain decomposition, we compute the corresponding best piecewise quadratic approximation of $F(\mathbf{x})$ by solving the normal equations, see ${ }^{5}$. The normal equations determine the set of coefficients for the desired quadratic spline representation-a best approximation in the least squares sense.

Corner vertices of simplicial elements may be shared by any number of simplices, and we denote the basis function associated with a corner vertex $\mathbf{v}_{i}$ by $f_{i}(\mathbf{x})$. An edge of a simplicial element may be shared by no more than two simplices


Figure 2: Types of basis functions: basis function associated with corner (left) and edge (right).


Figure 3: Basis functions associated with the platelet of vertex $\mathbf{v}_{i}$ and the edge neighbors of edge $\mathbf{e}_{j}$.
in the 2D case and by an arbitrary number of simplices in the 3D case. We denote a basis function associated with the midpoint of a simplex edge $e_{j}$ by $g_{j}(\mathbf{x})$. We refer to the set of simplices sharing a common corner vertex as the platelet of this corner, and we call the set of simplices sharing a common edge edge neighbors. Thus, a set of platelet simplices defines the region in space over which a basis function associated with the corresponding corner vertex is non-zero. Edge neighbors, associated with a particular edge, define the region in space over which a basis function associated with this edge is non-zero. Figure 2 shows the two types of basis functions for the bivariate case. Figure 3 shows the basis functions associated with the platelet of a vertex and the edge neighbors of an edge.

We denote a best approximation as $a(\mathbf{x})$, and we write it as a linear combination of the basis functions associated with all distinct simplex corners ("corner basis functions" $f_{i}$ ) and simplex edges ("edge basis functions" $g_{i}$ ). Assuming that there are $m$ distinct corners and $n$ distinct edges, we can write a best approximation as

$$
\begin{equation*}
a(\mathbf{x})=\sum_{i=1}^{m} c_{i} f_{i}(\mathbf{x})+\sum_{j=1}^{n} d_{j} g_{j}(\mathbf{x}) \tag{3}
\end{equation*}
$$

We must solve the normal equations to obtain the unknown coefficients $c_{i}$ and $d_{j}$. In matrix form, the normal equations are

$$
\begin{align*}
& \left(\begin{array}{cccccc}
\left\langle f_{1}, f_{1}\right\rangle & \ldots & \left\langle f_{1}, f_{m}\right\rangle & \left\langle f_{1}, g_{1}\right\rangle & \ldots & \left\langle f_{1}, g_{n}\right\rangle \\
\vdots & & & & & \vdots \\
\left\langle f_{m}, f_{1}\right\rangle & \ldots & \left\langle f_{m}, f_{m}\right\rangle & \left\langle f_{m}, g_{1}\right\rangle & \ldots & \left\langle f_{m}, g_{n}\right\rangle \\
\left\langle g_{1}, f_{1}\right\rangle & \ldots & \left\langle g_{1}, f_{m}\right\rangle & \left\langle g_{1}, g_{1}\right\rangle & \ldots & \left\langle g_{1}, g_{n}\right\rangle \\
\vdots & & & & & \vdots \\
\left\langle g_{n}, f_{1}\right\rangle & \ldots & \left\langle g_{n}, f_{m}\right\rangle & \left\langle g_{n}, g_{1}\right\rangle & \ldots & \left\langle g_{n}, g_{n}\right\rangle
\end{array}\right) \\
& \left(\begin{array}{c}
c_{1} \\
\vdots \\
c_{m} \\
d_{1} \\
\vdots \\
d_{n}
\end{array}\right)=\left(\begin{array}{c}
\left\langle F, f_{1}\right\rangle \\
\vdots \\
\left\langle F, f_{m}\right\rangle \\
\left\langle F, g_{1}\right\rangle \\
\vdots \\
\left\langle F, g_{n}\right\rangle
\end{array}\right), \tag{4}
\end{align*}
$$

where $\langle G, H\rangle$ denotes the inner product of the functions $G(\mathbf{x})$ and $H(\mathbf{x})$, i.e.,

$$
\begin{equation*}
\langle G, H\rangle=\int_{\text {Common Domain of } G \text { and } H} G(\mathbf{x}) H(\mathbf{x}) d \mathbf{x} \tag{5}
\end{equation*}
$$

In our construction, we must compute inner products over the simplices. Since all simplicial elements in physical space are defined by linear mappings of the standard simplex, we can simplify integration by making use of the change-ofvariables theorem, see ${ }^{21}$, which relates integration in physical space to integration in parameter space. In the 2D case, integrals are computed according to the formula

$$
\begin{align*}
& \int_{\text {Physical Simplex }} G(x, y) d x d y= \\
& \int_{\text {Standard Simplex }} G(x(u, v), y(u, v)) J(u, v) d u d v \tag{6}
\end{align*}
$$

where $J(u, v)$ denotes the Jacobian associated with the mapping of the standard simplex to the corresponding simplex in physical space. The Jacobian is the determinant

$$
J(u, v)=\left|\begin{array}{cc}
\frac{\partial}{\partial u} x(u, v) & \frac{\partial}{\partial v} x(u, v)  \tag{7}\\
\frac{\partial}{\partial u} y(u, v) & \frac{\partial}{\partial v} y(u, v)
\end{array}\right| .
$$

Thus, to effectively compute integrals of functions over triangles we only need to consider the linear transformation

$$
\begin{align*}
& {\left[\begin{array}{l}
x(u, v) \\
y(u, v)
\end{array}\right]=}  \tag{8}\\
& \quad\left[\begin{array}{ll}
x_{1,0}-x_{0,0} & x_{0,1}-x_{0,0} \\
y_{1,0}-y_{0,0} & y_{0,1}-y_{0,0}
\end{array}\right]\left[\begin{array}{l}
u \\
v
\end{array}\right]+\left[\begin{array}{l}
x_{0} \\
y_{0}
\end{array}\right] .
\end{align*}
$$

This transformation maps the standard triangle with vertices $\mathbf{u}_{0}=[0,0]^{T}, \mathbf{u}_{1}=[1,0]^{T}$, and $\mathbf{u}_{2}=[0,1]^{T}$ in the $u v$ plane to the arbitrary simplex $S$ with corner vertices $\mathbf{v}_{0}=$
$\left[x_{0,0}, y_{0,0}\right]^{T}, \mathbf{v}_{1}=\left[x_{1,0}, y_{1,0}\right]^{T}$, and $\mathbf{v}_{2}=\left[x_{0,1}, y_{0,1}\right]^{T}$ in the $x y$ plane. (Both triangles must be oriented counterclockwise). For this linear mapping, the change-of-variables theorem yields

$$
\begin{equation*}
\int_{S} G(x, y) d x d y=J \int_{v=0}^{1} \int_{u=0}^{1-v} G(x(u, v), y(u, v)) d u d v \tag{9}
\end{equation*}
$$

where the Jacobian $J$ is given by

$$
J=\operatorname{det}\left[\begin{array}{ll}
x_{1,0}-x_{0,0} & x_{0,1}-x_{0,0}  \tag{10}\\
y_{1,0}-y_{0,0} & y_{0,1}-y_{0,0}
\end{array}\right] .
$$

The 3D case is a straightforward extension; here, the Jacobian is given by

$$
\begin{align*}
& J= \\
& \operatorname{det}\left[\begin{array}{lll}
x_{1,0,0}-x_{0,0,0} & x_{0,1,0}-x_{0,0,0} & x_{0,0,1}-x_{0,0,0} \\
y_{1,0,0}-y_{0,0,0} & y_{0,1,0}-y_{0,0,0} & y_{0,0,1}-y_{0,0,0} \\
z_{1,0,0}-z_{0,0,0} & z_{0,1,0}-z_{0,0,0} & z_{0,0,1}-z_{0,0,0}
\end{array}\right] \tag{11}
\end{align*}
$$

The matrices involved in the best-approximation step are sparse because all basis functions have local support. Several methods exist for bandwidth reduction, efficient factorization, and inversion of such sparse matrices, see ${ }^{4,9,11,25,26}$. We use an efficient sparse matrix representation and system solver to compute the coefficients in linear time.

The computation of the inner products appearing in the normal equations requires multi-dimensional integration over simplicial elements. While the change-of-variables theorem reduces this integration to integration over the standard simplex, we still need to perform relatively expensive numerical integration for the calculation of the inner products appearing on the right-hand side of the normal equations, i.e., the integrals of the types $\left\langle F, f_{i}\right\rangle$ and $\left\langle F, g_{j}\right\rangle$. Since $F(\mathbf{x})$ can, in general, be any analytically defined function, numerical integration can potentially become expensive. We use Romberg integration for the computation of these right-hand-side inner products, see ${ }^{1,16}$.

Once we have computed a best approximation for a particular simplicial domain decomposition, we analyze the local approximation quality to identify simplices that should be refined (bisected) to further improve approximation quality. In the following section, we discuss the principle we use for adaptive bisection.

## 6. Adaptive Bisection

For each simplicial element $S_{i}$ in a particular domain decomposition, we compute a local approximation error $e_{i}$. We define this error as

$$
\begin{equation*}
e_{i}=\left(\int_{S_{i}}(F(\mathbf{x})-a(\mathbf{x}))^{2} d \mathbf{x}\right)^{\frac{1}{2}} \tag{12}
\end{equation*}
$$



Figure 4: Bisection of simplices in bivariate and trivariate cases. Darker simplex is the one selected for bisection.

Selecting and bisecting simplices of maximal error are the steps used to refine the mesh. In general, we choose a certain percentage of the simplices to be refined.

We refine a simplicial element by bisecting at the midpoint of its longest edge. All simplices sharing the split edge are bisected to avoid "hanging nodes" and, therefore, to preserve a conforming mesh. The bisection step is shown in Figure 4. Bisection steps lead to new simplicial domain decompositions, and we must compute new best quadratic spline approximations for each one.

We continue to bisect a certain percentage of simplices in the intermediate approximations until either the number of simplices in a decomposition exceeds some userspecified maximal number or until an approximation is obtained whose global error is less than a user-specified tolerance. (We defined the global error of an approximation as the sum of all local simplex errors).

The final result of our method is a set of independent best quadratic spline approximations that can be used for the purposes of interactive and/or adaptive level-of-detail visualization.

## 7. Results

We have tested our method for several test data. In general, we compare a piecewise quadratic approximation to a piecewise linear approximation. We visualize quadratic simplices by tessellating them using many linear elements.

Figure 5 shows a quadratic and linear spline approximation for comparison. The quadratic approximation can, in theory, approximate this function exactly. (Numerical floating-point error is introduced in practice. For the shown example, this numerical floating-point error is on the order of $10^{-14}$ ). The linear approximation must use a relatively large number of elements to represent this quadratic function with small error. The global error for the quadratic spline is $3.6 \times 10^{-14}$ and the global error for the linear approximation error is $1.6 \times 10^{-6}$. The linear approximation was computed using the method described in ${ }^{16}$.

A comparison of a quadratic and a linear spline approx-


Figure 5: Comparison between quadratic spline approximation (left) and linear spline approximation (right). Under each approximation, we show the corresponding domain decomposition. The quadratic approximation uses 9 knots and 2 simplices. The linear approximation uses 111 knots and 187 simplices. The function being approximated is $F(x, y)=x^{2}+y^{2}, x, y \in\left[-\frac{1}{2}, \frac{1}{2}\right]$.
imation is shown in Figure 6. The original image consists of $1536 \times 1024$ pixels. The quadratic spline approximationconsisting of 2989 quadratic simplices-required 158 seconds of computation time while the linear approximationconsisting of 11482 linear simplices-required 536 seconds.

A sample hierarchy of 2D quadratic spline approximations is shown in Figure 7. The original image consists of $121 \times 121$ pixels. Global errors for the six approximations are 329.11, 106.22, 45.53, 12.85, 3.08, and 0.40. Computations times ranged from two to 200 seconds for the six approximations.

A sample hierarchy of 2D quadratic spline approximations is shown in Figure 8. The original image consists of $211 x 144$ pixels. Global errors for the four approximations are $37.05,9.70,1.86$, and 0.45 . Computations times ranged from six to 200 seconds for the six approximations.

A comparison of a quadratic and a linear spline approximation of a 3D skull data set is shown in Figure 9. The original data set consists of 278528 data sites. We visualize the quadratic spline approximation by tessellating each quadratic simplex with 512 linear elements and then extracting an isosurface from the linear elements. The same isosurface for the linear spline approximation was extracted directly from the linear simplices. The quadratic spline approximation has a global error of $2.15 \times 10^{-6}$, and the linear


Figure 6: Comparison between quadratic approximation (left) and linear approximation (right). Original image is shown at the top. The quadratic approximation uses 6076 knots and 2989 simplices. The linear approximation uses 5816 knots and 11482 simplices.


Figure 7: Hierarchical approximations of digital image data set. Original image is shown at the top. Six approximations are shown, $8,20,38,90,225$, and 633 simplices, respectively.


Figure 8: Hierarchical approximations of digital image data set. Original image is shown at the top. Four approximations are shown, 16, 48, 191, and 790 simplices, respectively.


Figure 9: Comparison between quadratic approximation (left) and linear approximation (right). The quadratic approximation uses 7487 knots and 5348 simplices. The linear approximation uses 14667 knots and 78530 simplices.
spline approximation has a global error of $1.65 \times 10^{-2}$. The quadratic spline approximation required about 20 hours of computation time while the linear approximation required less than three.

A sample hierarchy of 3D quadratic spline approximations for a 3D skull data set is shown in Figure 10. Global errors for the six approximations are $1.0 \times 10^{-3}, 4.7 \times 10^{-4}$, $3.9 \times 10^{-5}$, and $2.1 \times 10^{-6}$.

All of the approximations were computed on a 1.8 GHz


Figure 10: Hierarchical approximations of skull data set. Four approximations are shown, 62, 125, 741, and 5348 simplices, respectively.

Pentium IV graphics workstation with 512 MB of main memory.

Linear 2D and 3D approximations were rendered at interactive frame rates. Quadratic 2D approximations required just a few seconds to render per frame. Tessellation of 3D quadratic approximations required several seconds for the highest resolutions. Once tessellated, computing and rendering a contour was at interactive frame rates.

## 8. Conclusions

Quadratic simplicial elements can be used to more compactly approximate data than linear simplicial elements. In general, the use of higher-order simplices should be considered as they can produce better-quality approximations, using a smaller number of simplices.

Additional research incorporating geometrically curved simplices can further improve the quality of approximations by allowing the simplices to decompose more complicatedshaped domains. A decomposition of a domain having curved boundaries would require fewer curved simplices to represent the domain well.

The generated meshes are $C^{0}$ continuous, it is also possible to produce $C^{1}$ approximations. We plan to investigate this enhancement in the future.

Higher-order simplices are growing in importance in visualization as researchers are also using them more frequently for domain decomposition in numerical simulations. Thus, visualization of these simplices is also important because
of their increasing popularity. Direct higher-order visualization techniques such as contouring and volume visualization techniques, must be developed to take advantage of higher-order elements. We are currently working on such techniques.

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