GPU-Assisted Computation of Centroidal Voronoi Tessellation

Guodong Rong, Yang Liu, Wenping Wang, Xiaotian Yin, Xianfeng Gu, and Xiaohu Guo

Abstract—Centroidal Voronoi tessellations (CVT) are widely used in computational science and engineering. The most commonly used method is Lloyd’s method, and recently the L-BFGS method is shown to be faster than Lloyd’s method for computing the CVT. However, these methods run on the CPU and are still too slow for many practical applications. We present techniques to implement these methods on the GPU for computing the CVT on 2D planes and on surfaces, and demonstrate significant speedup of these GPU-based methods over their CPU counterparts. For CVT computation on a surface, we use a geometry image stored in the GPU to represent the surface for computing the Voronoi diagram on the surface. In our implementation a new technique is proposed for parallel regional reduction on the GPU for evaluating integrals over Voronoi cells.

Index Terms—Centroidal Voronoi Tessellation, Graphics Hardware, Lloyd’s Algorithm, L-BFGS Algorithm, Remeshing.

1 Introduction

Voronoi diagrams are well studied in computational geometry and have many applications in areas like computer graphics, visualization, pattern recognition, etc. [1], [2]. An evenly-space tessellation of a given domain $\Omega$ is produced by a special type of Voronoi diagram, called Centroidal Voronoi Tessellation (CVT); see for example Fig. 1. The uniformity of the cells of an optimal CVT has been conjectured by Gersho [3] and proved in 2D [4], while confirmed empirically in 3D [5]. This property makes the CVT useful in many applications, including graph drawing [6], decorative arts simulation [7], [8], [9], grid generation and optimization [10], vector field visualization [11], [12], surface remeshing [13], [14], [15], [16] and medial axis approximation [17]. In this paper we study how to speed up the computation of the CVT using the GPU.

1.1 Preliminaries

We first present the definition and properties of the CVT and some typical algorithms for computing the CVT. More details about the CVT can be found in the survey [18].

Given $n$ sites $x_1, x_2, \ldots, x_n$ in a domain $\Omega \subset \mathbb{R}^N$, the Voronoi diagram is defined as the collection of the Voronoi cells $\Omega_i$, $i = 1, 2, \ldots, n$, defined as

$$\Omega_i = \{x \in \Omega : \|x - x_i\| < \|x - x_j\|, i \neq j\}.$$  

**Fig. 1.** (a) The CVT of a square 2D domain with 200 sites; (b) The constrained CVT on a torus with 600 sites.

The centroidal Voronoi tessellation is a special Voronoi diagram in which each site $x_i$ coincides with the centroid $c_i$ of its Voronoi cell $\Omega_i$:

$$c_i = \frac{\int_{\Omega_i} \rho(x)x \, d\sigma}{\int_{\Omega_i} \rho(x) \, d\sigma}.
\quad (1)$$

where $\rho(x) > 0$ is a density function. An example of a CVT of 200 sites in a square is shown in Fig. 1(a).

The CVT energy function $F$ is defined for the ordered set of samples (i.e. sites) $X = (x_1, x_2, \ldots, x_n)$ in $\Omega$ as

$$F(X) = \sum_{i=1}^{n} f_i(X) = \sum_{i=1}^{n} \int_{\Omega_i} \rho(x)\|x - x_i\|^2 \, d\sigma,
\quad (2)$$

where the $\Omega_i$ are the Voronoi cells of the sites $x_i$. It can be shown [18] that the regions $\Omega_i$ of the sites $x_i$ form a CVT in the domain $\Omega$ if and only if the gradient of $F(X)$ vanishes, that is, a critical point of $F(X)$. Therefore, a necessary condition for $F$ to be locally minimized is that the regions $\Omega_i$ form a CVT. A CVT that locally minimizes $F$ will be called a stable CVT. In practice one often seeks a stable CVT since it usually produces a more regular and compact tessellation than a CVT that is not a local minimizer of $F$. 

- **G. Rong and X. Guo** are with the Department of Computer Science, University of Texas at Dallas, Richardson, TX, USA. E-mail: {guodongrong, xguo}@utdallas.edu.
- **Y. Liu** is with Project ALICE, INRIA, Villers les Nancy, France. E-mail: liuyang@loria.fr.
- **W. Wang** is with the Department of Computer Science, University of Hong Kong, Hong Kong, China. E-mail: wenping@cs.hku.hk.
- **X. Yin** and **X. Gu** are with the Department of Computer Science, State University of New York at Stony Brook, Stony Brook, NY, USA. E-mail: {xyin, gu}@cs.sunysb.edu.
Furthermore, a CVT that globally minimizes $F$ is called an optimal CVT. Because there are a large number of stable CVTs when the number of sites is large, in general, it is difficult to compute an optimal CVT.

For a non-convex domain, the centroid of a cell $\Omega_i$, computed by (1) may lie outside $\Omega_i$. In this case the centroid is replaced by the constrained centroid inside $\Omega_i$, defined as

$$c_i^* = \arg \min_{p \in \Omega_i} \int_{\Omega_i} \rho(x)||x - p||^2 \, d\sigma.$$  \hfill (3)

When all the sites coincide with the constrained centroids within the domain $\Omega$, the CVT is called a constrained centroidal Voronoi tessellation (CCVT) [19]. The CCVT can also similarly be defined on a surface $S \subset \mathbb{R}^3$, i.e. by constraining all the sites to lie on the surface. Fig. 1(b) shows an example of a CCVT of 600 sites on a torus.

Two commonly used algorithms for computing the CVT are Lloyd’s algorithm [20] and the L-BFGS algorithm [21]. Although converging faster than Lloyd’s algorithm, the L-BFGS algorithm still needs a long time to compute a CVT. In this paper we describe how to leverage the parallel computational power of the programmable graphics processing unit (GPU) to speed up these two methods for computing the CVT in 2D and on surfaces.

Using the GPU for computing the CVT in 2D is a straightforward idea, since a 2D domain can be represented naturally by a 2D texture in the GPU. The idea of computing the CCVT on a surface is similar, but we need to first construct a parametric representation of the surface using the geometry image. With the geometry image represented as a texture in the GPU, we run the jump flooding algorithm [22] to compute the Voronoi diagrams in each iteration of CVT computation. Here the pixels of the geometry image store the 3D coordinates of sampled points on the surface and we use the Euclidean distance 3D for computing the Voronoi diagram. We note that this is different from the method in [15], which computes Voronoi diagrams using distances in a 2D parametric domain.

1.2 Contributions

Our contributions are efficient implementations of Lloyd’s algorithm and the L-BFGS algorithm on the GPU for the computation of the CVT in 2D and on a surface. We propose a new technique for computing Voronoi diagrams on surfaces and a novel way of using vertex programs to perform the regional reduction over Voronoi cells. Significant speedup is achieved by our GPU programs in various cases of CVT computation.

All our GPU programs are implemented with the shader language Cg. Although general purpose languages on the GPU (e.g. CUDA) are more popular now, our tests show that Cg is better suited for implementing the algorithms for CVT computation. We will explain the reasons behind this in more details in Section 5.4.

The remainder of the paper is organized as follows: Section 2 briefly reviews related work. Section 3 explains how to compute the CVT on a 2D plane with the GPU. The idea is then extended in Section 4 to computing the CCVT on surfaces. The experimental results and comparisons are given in Section 5. Section 6 concludes the paper with discussions of future research.

2 PREVIOUS WORK

We will briefly review existing algorithms on the CPU for computing the CVT. We also give a brief survey of previous work on using the GPU to compute the Voronoi diagram.

2.1 CVT Algorithms

MacQueen’s probabilistic method is one of the earliest methods for computing the CVT [23]. Ju et al. integrated MacQueen’s method and Lloyd’s algorithm on a parallel platform [24]. The most commonly used algorithm for computing the CVT in 2D/3D is Lloyd’s algorithm [20] for its simplicity and robust. However this method has linear convergence and is very slow in practice. The multi-grid method has been proposed to accelerate Lloyd’s algorithm [25].

It has recently been shown that the CVT energy function is $C^2$ continuous [21]. Justified by the $C^2$ smoothness of the CVT energy function, Liu et al. applied a quasi-Newton method – the L-BFGS algorithm – to computing the CVT in 2D, 3D and on surfaces, and showed that the algorithm is faster than Lloyd’s algorithm [21].

2.2 GPU Algorithms

With the rapid advance of the GPU, the general-purpose computation on the GPU (GP-GPU) has become an active topic [26]. In the following we will review previous work on using the GPU to compute Voronoi diagrams.

Hoff et al. [27] built a right-angle cone for every site and rendered them from bottom to get a Voronoi diagram of these sites. Denny’s method [28] is similar to Hoff et al.’s but changes the cones to depth textures to get better quality and speed. Fischer and Gotsman [29] lifted the sites to a paraboloid and rendered planes tangent to the paraboloid to obtain the Voronoi diagram, thus avoiding tessellating the cones. Note that the Voronoi diagrams computed by GPU algorithms in 2D are defined by color-coded pixel maps, hence they are only discrete approximations to the true Voronoi diagrams defined by polygons. All these algorithms are designed for computation in 2D, and it is not clear how to extend them to compute the Voronoi diagram on a surface.

The jump flooding algorithm (JFA) [22] is another method for computing the Voronoi diagram in a 2D discrete domain represented in pixels. The JFA propagates information (e.g. coordinates of the sites in the Voronoi diagram problem) from the sites to all other pixels in
parallel, similar to the flood-filling algorithm, but with faster speed due to its use of varying step lengths. We will use the JFA to compute the Voronoi diagram in 2D as well as on a surface. Like [19], [21] we use the Euclidean distance to approximate the geodesic distance on a surface.

To compute the Voronoi diagram on a surface, one may compute a 3D Voronoi diagram and find its intersection with the surface. The GPU has also been used to compute 3D Voronoi diagrams [27], [30], [31], [32], [33]. Weber et al. [34] adopted the raster scan method in the geometry image to compute the Voronoi diagram using the geodesic distance on a surface. This method handles disk-like open surfaces only. The geodesic distance, though more accurate than the Euclidean distance, is much less efficient to compute on a mesh surface, even with GPU acceleration.

Vasconcelos et al.’s work [35] is the only known successful attempt so far using the GPU to compute the CVT in a plane. They implemented the Lloyd’s method on the GPU to compute the CVT on a 2D plane, focusing on the computation of the centroids. A predefined mask is used to conservatively estimate the Voronoi cell for every site. Since the diameter of a Voronoi cell may be very big, to ensure an accurate result, the mask must be as big as the whole texture, which makes the method inefficient.

In contrast, we use the vertex program to perform scattering and use the framebuffer blending function to accumulate the coordinates. Our approach is simpler and works well for computing both the CVT on a 2D plane and the CCVT on a surface.

3 CVT on 2D Plane

There are two main steps in both Lloyd’s algorithm and the L-BFGS algorithm: 1) computing the Voronoi diagram of the current sites, and 2) finding new positions of the sites for the next iteration. For step 1, we compute a discrete approximation of the Voronoi diagram using the jump flooding algorithm (JFA) [22] on the GPU. We propose a new regional reduction method for efficiently computing various integrals needed in step 2. In the following we will briefly review the JFA and present the regional reduction technique.

3.1 Jump Flooding Algorithm

Suppose that there is a site in an $n \times n$ texture in the GPU and we want to propagate some information (e.g. the coordinates of the site) from the site to all the other pixels. The JFA performs this in several passes. For an initial step length $k$ that is a power of 2 (e.g. $2^\lceil \log n \rceil$), a pixel at $(x, y)$ passes its information to its neighbors (eight at most) at $(x + i, y + j)$, where $i, j \in \{-k, 0, k\}$ (Fig. 2). Then in the subsequent passes, the same propagation is performed for a pixel using the step length that is half of the previous step length $k$. The iteration is stopped when the step length reaches 1. Fig. 2 illustrates how the JFA fills up an $8 \times 8$ texture using three passes with the single initial site at the bottom left corner.

When using the JFA to compute the Voronoi diagram in a 2D texture, there are multiple sites and the information to be propagated from each site is its coordinates. Upon receiving the coordinates of different sites, each pixel compares its distances to these sites to find the nearest site whose Voronoi cell it belongs to. Thus all the pixels are classified to form a Voronoi diagram.

Despite its fast speed, the JFA may misclassify a small number of pixels [22]. We use 1+JFA [33], an improved version of JFA, to compute the Voronoi diagram in our GPU implementations. On average, the error rate of 1+JFA is less than 0.25 pixels in a texture with the resolution of $512 \times 512$ for less than 10,000 sites, which is accurate enough for most practical graphics applications utilizing the CVT. To be brief, we will refer 1+JFA as JFA as well.

A sufficiently large initial step length is needed to ensure that each pixel is reached by its nearest site. On the other hand, one should try to use a small but “safe” initial step length to reduce the computation time incurred by unnecessary JFA passes. A safe choice is $2^\lceil \log n \rceil$, which is necessary for computing a Voronoi diagram when the sites are distributed in such a way that each Voronoi cell is narrow and long, as the cells generated by a sequence of collinear sites. However, this initial step length is, overall, very conservative because after a few iterations of CVT computation (with either Lloyd’s algorithm or the L-BFGS algorithm) the sites are normally already distributed rather evenly, therefore a much smaller initial step length would be sufficient for each pixel to be reached by its nearest site.

This consideration leads us to use the following scheme for selecting the initial step length. In the VD computation of the first CVT iteration, the initial step length of the JFA is set to be $2^\lceil \log n \rceil$. For the VD computation in the next $m$ CVT iterations, we compute the average distance from each site to all the pixels in its Voronoi cell using the regional reduction (to be introduced in next subsection), and set the initial step length of the JFA to be the double of the maximum of all these average distances. Our experiments show that $m = 5$ gives satisfactory performance. Then in the remaining CVT iterations (i.e. after 5 iterations) the initial step length of the JFA is set to be that used in the 5th iteration.
3.2 Regional Reduction

Several different integrals over the Voronoi cells of all the sites need to be evaluated in CVT computation, that is, for computing the value of the CVT energy function or computing the centroid as needed by the Lloyd algorithm. To approximate these integrals, we need to perform summations over the pixels of all the Voronoi cells in parallel, which calls for solving the so-called regional reduction problem.

In the reduction problem, one needs to reduce a number of values to a single one, such as sum, maximum, minimum, etc. More precisely, the reduction problem takes as input a set of values \( v_1, \ldots, v_n \) and outputs a single value \( v = v_1 \oplus \ldots \oplus v_n \), where \( \oplus \) is an associative and commutative operator. Thus, summation is a special case of the reduction problem.

The reduction problem can be solved in \( O(\log n) \) passes on the GPU using a fragment program [36]. Existing algorithms can only perform global reductions, that is, reducing values of all the pixels in a texture into one single value. However, in CVT computation the domain \( \Omega \) is decomposed into multiple regions (that is, Voronoi cells) \( \Omega_i \) and we need to compute the sums of values of the pixels of different regions in parallel. Therefore, we face a regional reduction problem rather than a global one.

We propose to use the method of rendering points for regional reduction. A single point is rendered for every pixel, and its position is changed in the vertex program. All the points in the Voronoi cell of site \( x_i \) are translated to the same position decided by its ID \( i \). For example, the site \( x_i \) corresponds to the position \( (i/w,i\%w) \), where \( w \) is the width of the texture used and \( '/' \) and \( \% \) are division and modulus operators for integers, respectively. The result is a texture containing the reduction values for all the Voronoi cells, one pixel for each site, which are packed in the order of the site’s ID. Fig. 3 shows an illustration of this operation.

First, every point is rasterized into one fragment. The fragment program processing these fragments then writes the values to a texture recording the results. The depth test or framebuffer blending operations can be used to reduce all the values corresponding to the same pixel to a single value which is stored in a result texture.

For example, if we want to compute the maximum of the values, we can write the values into the depth channel as well as the color channels for every fragment, and set the depth test function so that only the fragment with the maximum value is stored into the result texture (e.g., using \texttt{glDepthFunc(GL\_GREATER)}). If we want to compute the sum of the values, we can write the values into color channels for every fragment, and set the blending function so that the values of all the fragments are added and the result is stored into the result texture (e.g., using \texttt{glBlendFunc(GL\_ONE, GL\_ONE)}).

Our regional reduction method works for a connected region as well as a set consisting of disconnected regions. This is important because the Voronoi diagram of a surface may contain disconnected Voronoi cells near the boundaries of the geometry image (see Section 4). Furthermore, a Voronoi cell in a pixel plane may contain disconnected parts [37]; such a case can be handled properly by our regional reduction method.

3.3 Lloyd’s Algorithm in 2D

Every iteration in Lloyd’s algorithm contains two steps: 1) computing the Voronoi diagram of the current sites; and 2) computing the centroids for Voronoi cells as the new sites in the next iteration. These two steps are iterated until certain termination condition is met.

On a 2D plane the centroids are computed using (1). Using regional reduction the integrations in (1) are approximated by summations, so we have

\[
 c_i = \frac{\sum_{x \in \Omega_i} \rho(x) \Delta \sigma}{\sum_{x \in \Omega_i} \rho(x) \Delta \sigma} = \frac{\sum_{x \in \Omega_i} \rho(x) x}{\sum_{x \in \Omega_i} \rho(x)},
\]

where the \( x \) is the pixel position and \( \Delta \sigma \) the constant area occupied by each pixel.

The numerator and denominator in (4) are computed using the regional reduction in the same pass since they share the same domain. The numerator is a 2D vector stored in red and green channels of the texture, and the denominator is a scalar value stored in the blue channel.

We stress that the coordinates of each site of the Voronoi diagram are floating point numbers, albeit they are stored in the discrete pixel closest to it. Since the Voronoi cells are composed of pixels, the Voronoi boundaries have only pixel precision. The algorithm will stop when the Voronoi diagram does not change. Alternatively, a different termination condition can be used by checking if the value of CVT energy function \( F \) or its gradient has reached a threshold.

3.4 L-BFGS Algorithm in 2D

The L-BFGS algorithm is a quasi-Newton algorithm that is more efficient than Lloyd’s method for CVT computation [21]. In every iteration of the L-BFGS algorithm, we need to compute the CVT energy \( F \) and its partial derivatives with respect to all the sites. These values are accumulated with those values of the \( m \) preceding iterations to approximate the inverse Hessian matrix.
More specifically, define \( s_k = X_k - X_{k-1} \) and \( y_k = \nabla F_k - \nabla F_{k-1} \), where \( X_k \) and \( \nabla F_k \) are the ordered set of sites and the gradient of the CVT energy function \( F \) at the \( k \)-th iteration, respectively. Then the approximated inverse Hessian matrix \( H_k \) is updated by

\[
H_k = V_k^T H_{k-1} V_k + \rho_k s_k s_k^T
\]

where \( \rho_k = 1/y_k^T s_k \), and \( V_k = I - \rho_k y_k s_k^T \). The new sites \( X_{k+1} \) for the next iteration is given by \( X_{k+1} = X_k - H_k \nabla F_k \).

The first-order partial derivative of \( F \) with respect to the site \( x_i \) is [2]:

\[
\frac{\partial F}{\partial x_i} = 2 \int_{\Omega_i} \rho(x)(x_i - x) \, d\sigma.
\]

This integral is approximated by summation in the GPU using the regional reduction, performed in the same pass as computing the energy function \( F \). These values are written into a texture and read back to the CPU, where they are used to compute the approximated inverse Hessian matrix \( H_k \) using (5) and then the new sites \( X_{k+1} \). Hence, our implementation of the L-BFGS method is not entirely done on GPU.

4 CCVT ON SURFACES

In this section we will explain the pipeline of computing the CCVT on a surface, following the flow in Fig. 4.

4.1 Geometry Image for Surface Representation

For a given surface we first compute a conformal parameterization [38] of the surface over a 2D rectangular domain and use the parameterization to construct a regular quad mesh to represent the surface. Then the surface can be represented by a 2D texture called the geometry image [39] whose pixels store the 3D coordinates of the corresponding mesh vertices. The red, green and blue channels of a geometry image store the 3D coordinates of the mesh vertices, respectively. The surface normal vectors at the vertices are also stored in a 2D texture of the same size, called normal vector image, which will be used in the L-BFGS algorithm for computing the CCVT on a surface.

Due to parameterization distortion, each pixel is assigned a weight equal to the area it covers on the surface. This weight is computed as follows. Suppose that the grid is subdivided to give a regular triangulation as shown in Fig. 5. Every pixel is supposed to cover one-third of each triangle incident to the corresponding vertex of the pixel. Thus the weight of the pixel is set to the 1/3 of the areas of all the triangles incident to the corresponding vertex. The weight will be called the distortion factor.

The pipeline of computing the CCVT is shown in Fig. 4. The left most part of Fig. 4 shows the geometry image and the normal vector image of the surface of David Head in the top row, and the surface itself and a checkerboard texture showing the parameterization in the bottom row. The computed distortion factors are shown in the left middle part of Fig. 4.

Like the 2D case, computing the CCVT on a surface
using either Lloyd’s algorithm or the L-BFGS algorithm takes two main steps in each iteration: 1) computing the Voronoi diagram of the current sites on the surface, and 2) computing the new sites for the next iteration. We will explain these two steps in the following subsections.

4.2 Computing Voronoi Diagram on Surfaces
Suppose that a given surface is represented as a geometry image. We first store the 3D coordinates of the initial sites in the nearest corresponding pixels of the geometry image, and then perform the JFA to compute a Voronoi diagram on the surface. Note that the Euclidean distance between points in 3D space is used as approximation to the geodesic distance when computing the Voronoi diagram on the surface.

We use a single geometry image for parameterizing a surface of arbitrary topology, with necessary topological cutting to facilitate parameterization. Although only surfaces with simple topology (open surface, cylinder and torus) can be represented by a geometry image without singularity. The singular points arising from a surface of complex topology do not pose any difficulty to our GPU programs. This is because the JFA only uses the 3D coordinates in the Voronoi diagram computation, and does not use the texture coordinates, which are not unique for singular points.

Due to topological cut, a Voronoi region on a surface may be split into disconnected regions in the geometry image, as shown by the example in Fig. 6. Here, for example, because the surface under consideration has a cylinder topology, we need to identify the two opposite boundaries of the geometry image when running the JFA to ensure that a correct Voronoi diagram is computed on the surface. Two disconnected parts in the geometry image of a connected Voronoi region on the surface are marked by ellipses in Fig. 6(b). In this case we compute the Voronoi diagram by identifying boundaries produced by topological cut; the result for this example is shown in Fig. 6(a).

Equipped with the above routine for computing the Voronoi diagram on a surface, we now explain how to implement Lloyd’s algorithm and the L-BFGS algorithm on the GPU to compute the CCVT on a surface.

4.3 Lloyd’s Algorithm on Surfaces
Using the geometry image to compute the CCVT on a surface with Lloyd’s algorithm follows the same procedure as for computing the CVT in 2D, except that we need to compute the constrained centroids using (3). The following property about the constrained centroid will be useful [19]: if \( \mathbf{c}_i \) and \( \mathbf{c}_i' \) are the centroid and the constrained centroid of the Voronoi cell \( \Omega_i \), respectively, then \( \mathbf{c}_i \mathbf{c}_i' \) is parallel to the surface normal vector at \( \mathbf{c}_i' \). On the other hand, we observe that if \( \mathbf{c}_i' \) is the nearest point in \( \Omega_i \) to \( \mathbf{c}_i \), and it is not on the boundary of \( \Omega_i \), then \( \mathbf{c}_i \mathbf{c}_i' \) is parallel to the surface normal vector at \( \mathbf{c}_i' \). Based on these observations we find it an effective heuristic to use the nearest point \( \mathbf{c}_i' \) as the constrained centroid \( \mathbf{c}_i \) in Lloyd’s algorithm, although they are not always identical, since \( \mathbf{c}_i \mathbf{c}_i' \) being parallel to the surface normal vector at \( \mathbf{c}_i' \) is not a sufficient condition for \( \mathbf{c}_i' \) to be the constrained centroid.

The nearest point \( \mathbf{c}_i' \) is computed as follows. For the corresponding mesh vertex of every pixel in \( \Omega_i \), we compute the nearest point within its six incident triangles to the centroid \( \mathbf{c}_i \). To avoid redundant computations, we only need to check two incident triangles for every vertex (e.g. the two shaded triangles in Fig. 5 for the center vertex), and the other four incident triangles will be dealt by other neighboring vertices. In this way we find a nearest point to the centroid \( \mathbf{c}_i \) for every pixel in \( \Omega_i \). Then by computing the minimum of the distances from these points to \( \mathbf{c}_i \) using the regional reduction, we find the nearest point \( \mathbf{c}_i' \) within \( \Omega_i \) to \( \mathbf{c}_i \).

The number of iterations needed by both Lloyd’s and the L-BFGS algorithms to obtain the CCVT can be reduced significantly if the initial sites are roughly evenly distributed on the surface. To obtain such initial sites, we use the distortion factors as a probability density function to sample the initial sites in the geometry image. Therefore, a pixel with a larger distortion factor is more likely to be selected as the initial site. The right middle part of Fig. 4 shows 1,000 initial sites sampled according to the distortion factors and the Voronoi diagram generated in the parameter domain (top row), as well as on the surface (bottom row). The final sites and the CCVT are shown in the right most part of Fig. 4.

4.4 L-BFGS Algorithm on Surfaces
The initial sites for the L-BFGS algorithm are also sampled according to distortion factors. In every iteration, to update the sites we need to evaluate the CVT energy function \( F \) and its partial derivatives. Because the sites are constrained to be on the surface, we only use the tangential components of the partial derivatives as

\[
\frac{\partial F}{\partial x_i} \bigg|_\Omega = \frac{\partial F}{\partial x_i} - \left( \frac{\partial F}{\partial x_i} \cdot \mathbf{N}(x_i) \right) \mathbf{N}(x_i),
\]

(7)
Fig. 7. The Voronoi diagram of 1,000 initial sites in the 2D domain of $[-1, 1] \times [-1, 1]$ (a), and CVT results generated by the L-BFGS algorithm using (b) GPU and (c) CPU. (Unshaded cells are hexagons.)

where $N(x_i)$ is the surface normal vector at $x_i$ stored in the normal vector image. Therefore we can use shaders to evaluate $F$ and its partial derivatives on the GPU efficiently, in the same way as in the 2D case.

The updated sites computed in the L-BFGS algorithm may not lie exactly on the surface. We compute their nearest points in their respective Voronoi cells on the surface as the new sites on the surface.

5 Experimental Results and Discussion

We implement our programs using Microsoft Visual C++.net 2005 and NVIDIA Cg 2.0. The hardware platform is Intel Core 2 Duo 2.93GHz with 2GB DDR2 RAM, and NVIDIA GeForce GTX 280 with 1GB DDR3 VRAM. For the L-BFGS algorithm, we use our HLBFGS library [40]. We have compared our results with the CPU version of Lloyd’s algorithm and the GPU version of the L-BFGS algorithm in [21]. We use $m = 7$ for all L-BFGS algorithms in our experiments; that is, we use the gradients of the 7 previous iterations to estimate the inverse Hessian in our implementation.

All the programs in this paper use IEEE standard float point numbers (32-bit).

5.1 Results of CVT in 2D

The first test example is the computation of the CVT in the 2D domain of $[-1, 1] \times [-1, 1]$ mapped to a $512 \times 512$ texture, with 1,000 random initial sites. The CVTs computed by the GPU program and CPU program are shown in Fig 7. It is clear that the uniformity of the sites is greatly improved in both results. We plot the CVT energy values and their gradients versus the number of iterations for Lloyd’s algorithm and the L-BFGS algorithm in Fig. 8. The red and green curves are for our GPU programs and the black and blue curves for CPU ones. For clarity, we show zoom-in views of the shaded regions in Fig. 8(b) and Fig. 8(d).

The experiments show that the GPU result is very close to the CPU one, although there are fewer hexagon cells in the GPU result and the final CVT energy value generated by the GPU is slightly higher. We may evaluate the quality of an approximate CVT by considering its relative difference from CVT computed with the CPU, defined as

$$\frac{CVT_{energy} - final\_energy\_CPU}{final\_energy\_CPU} \times 100\%,$$

where $CVT_{energy}$ is the CVT energy of the approximate CVT. For the tests shown in Fig. 8 the relative difference is 0.69% for the result of Lloyd’s algorithm on GPU and 0.58% for that of the L-BFGS algorithm on GPU. These differences can be attributed to two factors. First, a different local minimum with higher energy is produced by the GPU program. Second, the quantization errors in the GPU implementation are responsible.

Table 1 compares the total running times of different programs for this example. The GPU program and CPU program of the same algorithm use the same number of iterations. Because of the line search used in the L-BFGS minimizer, the function evaluating gradients and energy value may be called more than once in every iteration. Within every function call, we need to rebuild the Voronoi diagram and compute the gradients of the CVT energy, and this is the most time-consuming part of the L-BFGS algorithm and dominates the running time. For this reason, #Iter. for the L-BFGS method in Table 1 is the number of the function calls.

We see that the GPU program of Lloyd’s method is several time faster than the CPU program, because all the computations for Lloyd’s algorithm are executed on the GPU. However, the GPU program of the L-BFGS method has only about 30% speedup over its CPU counterpart. That is because, although the most time-consuming parts for the L-BFGS algorithm are executed on the GPU, some of its computations are done on the CPU which takes about 23% of the total time and is not accelerated by the
GPU; and the communication between the CPU and the GPU also takes about 5% of the total time.

We have also compared our GPU program of Lloyd’s method to the algorithm proposed by Vasconcelos et al. [35]. For 1,000 sites, even with a very small mask (32 × 32), their program needs 1.416 seconds for the same number of iterations as in Table 1. If the size of the mask is set to be same as the screen resolution (512), the running time becomes 44.521 seconds.

Our GPU programs can also compute the CVT with a non-constant density function, as the density can be sampled and stored as floating point numbers in a texture. An example is shown in Fig 9, comparing the CVTs computed by the GPU program and the CPU program of the L-BFGS method. The final CVT energy values and the total running time are listed in Table 2.

### Table 1: Comparison of energy F and running time (in seconds) of different programs using 1,000 initial sites in the 2D domain of [−1, 1] × [−1, 1].

<table>
<thead>
<tr>
<th>Program</th>
<th>#Iter</th>
<th>GPU Time</th>
<th>CPU Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lloyd</td>
<td>216</td>
<td>2.628 × 10^3 0.550</td>
<td>2.610 × 10^3 1.535</td>
</tr>
<tr>
<td>L-BFGS</td>
<td>92</td>
<td>2.612 × 10^3 0.512</td>
<td>2.597 × 10^3 0.703</td>
</tr>
</tbody>
</table>

Fig. 9. CVT results of 1,000 initial sites generated by the L-BFGS algorithm in the 2D domain of [−1, 1] × [−1, 1] with the density function \( \rho(x) = e^{-20x^2 - 20y^2} + 0.05\sin^2(\pi x)\sin^2(\pi y) \) using (a) GPU and (b) CPU.

### Table 2: Comparison of energy F and running time (in seconds) of different programs using 1,000 initial sites in the 2D domain of [−1, 1] × [−1, 1] with non-constant densities.

<table>
<thead>
<tr>
<th>Program</th>
<th>#Iter</th>
<th>GPU Time</th>
<th>CPU Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lloyd</td>
<td>627</td>
<td>6.524 × 10^-5 2.529</td>
<td>6.163 × 10^-5 28.530</td>
</tr>
<tr>
<td>L-BFGS</td>
<td>376</td>
<td>6.185 × 10^-5 2.301</td>
<td>6.116 × 10^-5 16.889</td>
</tr>
</tbody>
</table>

### Table 3: Information of the five models used in this paper. The last two columns are for the number of boundaries and the resolution of geometry images.

<table>
<thead>
<tr>
<th>Model</th>
<th>#Vertices</th>
<th>#Faces</th>
<th>Genus</th>
<th>#B</th>
<th>GI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Torus</td>
<td>4,096</td>
<td>7,938</td>
<td>1</td>
<td>0</td>
<td>512×512</td>
</tr>
<tr>
<td>Lion</td>
<td>5,321</td>
<td>10,200</td>
<td>0</td>
<td>0</td>
<td>512×190</td>
</tr>
<tr>
<td>Sculpture</td>
<td>5,471</td>
<td>10,364</td>
<td>3</td>
<td>0</td>
<td>512×372</td>
</tr>
<tr>
<td>Body</td>
<td>13,978</td>
<td>27,295</td>
<td>0</td>
<td>2</td>
<td>512×456</td>
</tr>
<tr>
<td>David Head</td>
<td>51,038</td>
<td>101,144</td>
<td>0</td>
<td>1</td>
<td>512×416</td>
</tr>
</tbody>
</table>

5.2 Results of CCVT on Surfaces

We will compare our GPU programs and CPU programs of Lloyd’s algorithm and the L-BFGS algorithm using five surface models: Torus (Fig. 1), Lion (Fig. 10), Sculpture (Fig. 11), Body (Fig. 6), and David Head (Fig. 4). Table 3 lists information about these five models. All of our experiments use 1,000 initial sites on the surface sampled according to distortion factors. For each model, the same set of initial sites are used as input for all programs. The CPU programs utilize the fast algorithm introduced in [41] which can greatly accelerate the computation of the intersection between the surface and the 3D Voronoi diagram. For every model the geometry image is pre-computed with user interaction in less than 10 seconds. This time is not included in the total running time reported below.

The final energy values F and the total running time of the entire iteration procedure are listed in Table 4 and Table 5. Like the 2D cases, for the L-BFGS algorithm, the number of the function calls for VD computation is listed, rather than the number of iterations. The GPU program and the CPU program have the same number of iterations for Lloyd’s algorithm or the same number of function calls for the L-BFGS algorithm.

It is observed that the GPU programs perform about one order of magnitude faster than their CPU counterparts. This speedup is more than that of the 2D case because the CPU programs for computing the CCVT of a surface need to compute a 3D Voronoi diagram and find its intersection with the surface, which is a very time consuming task compared with computing a Voronoi diagram in a 2D domain. Although the GPU programs also compute distances in 3D, the whole algorithms are still efficiently performed in a 2D domain.

The relative differences between the GPU and CPU results range from 0.88% to 8.36% due to different distortions of surface parameterizations. Fig. 10 and Fig. 11 show two sets of results with the largest relative differences of Lion (genus 0) and Sculpture (genus 3): the Voronoi diagram of the initial sites, and the CCVTs generated using the GPU and the CPU of Lloyd’s algorithm and the L-BFGS algorithm, with the same number of iterations for each algorithm. We note that the GPU results have larger energy values than their CPU counterparts due its approximation nature.

In addition to the comparison in terms of visual inspection and energy values, we may also measure the geometric uniformity of the sites and their Voronoi cells.
in the CCVT results. For every site $x_i$, we define the radius $r_i$ of its Voronoi cell $\Omega_i$, the distance $d_i$ to its nearest neighboring site, and the area $a_i$ of its Voronoi cell $\Omega_i$ as follows:

$$r_i = \max_{x \in \Omega_i} ||x - x_i||, d_i = \min_{j \neq i} ||x_i - x_j||, a_i = Area(\Omega_i).$$

The standard deviations of $r_i$, $d_i$, and $a_i$ are used to measure the uniformity of a set of sites. Table 6 lists the standard deviations for initial sites and the sites in the CCVT results on Lion. The uniformity of the initial sites is greatly improved in all the results. Again, the GPU results are overall still not as good as the CPU ones.
TABLE 6
Comparison of the uniformity of 1,000 initial sites and the sites in CCVT results generated by different programs on the surface of Lion.

<table>
<thead>
<tr>
<th>Program</th>
<th>STDEV(ri)</th>
<th>STDEV(di)</th>
<th>STDEV(ai)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Sites</td>
<td>$1.450 \times 10^{-2}$</td>
<td>$1.809 \times 10^{-2}$</td>
<td>$3.717 \times 10^{-3}$</td>
</tr>
<tr>
<td>Lloyd - GPU</td>
<td>$7.422 \times 10^{-3}$</td>
<td>$9.246 \times 10^{-3}$</td>
<td>$1.902 \times 10^{-3}$</td>
</tr>
<tr>
<td>Lloyd - CPU</td>
<td>$3.228 \times 10^{-3}$</td>
<td>$6.090 \times 10^{-3}$</td>
<td>$1.002 \times 10^{-3}$</td>
</tr>
<tr>
<td>L-BFGS - GPU</td>
<td>$6.992 \times 10^{-3}$</td>
<td>$9.657 \times 10^{-3}$</td>
<td>$1.805 \times 10^{-3}$</td>
</tr>
<tr>
<td>L-BFGS - CPU</td>
<td>$2.868 \times 10^{-3}$</td>
<td>$5.824 \times 10^{-3}$</td>
<td>$9.046 \times 10^{-4}$</td>
</tr>
</tbody>
</table>

The quality of the CCVT we computed with the GPU is heavily affected by the distortion factors. If the distortion factors are very large in a certain part of a surface, the Voronoi cells in this part are mapped to a very small region in the geometry image. Then the resolution of the geometry image will not be adequate for accurate computation in this part of the surface, resulting in larger errors in the computation of centroids (for Lloyd’s algorithm) or energy value and gradients (for the L-BFGS algorithm). To illustrate this, we compare the results on David Head using two different parameterizations (see Fig. 12). Clearly, the second parameterization has larger distortion, which leads to more artifacts than the first parameterization. This problem could be alleviated by using a geometry image of higher resolution (see the example in Fig. 12(e) and (h)), or using multiple geometry images based a multi-chart surface parameterization [42].

The CCVT can be used for surface remeshing by computing a well-shaped triangulation of the surface as the dual mesh of a CCVT. Fig. 13 shows an example of remeshing the Body surface with 1,000 sites.

5.3 JFA Errors

Ideally, the CVT energy should decrease monotonically during the iterations in both Lloyd’s and the L-BFGS algorithm, if implemented accurately. However, since some pixels may be misclassified by the JFA into other Voronoi cells, the partial CVT energy values computed for those Voronoi cells are slightly different to the accurate values. Despite this, the sites move greatly and the total CVT energy keeps decreasing in early iterations. However, in late iterations, when most of sites are no longer moving, the error of JFA may cause the CVT energy to fluctuate. When this happens, most sites would remain unchanged but a small number of sites may oscillate between some positions.

To evaluate the consequence of this oscillation, we compared the JFA with an implementation on the GPU which computes the distance from every pixel to every site accurately by brute force, and show the results in Fig. 14. We see that the energy values only begins to oscillate in very late iterations due to the errors of the JFA.
In practice, this oscillation is very small and so does not affect the quality of the CVT for most applications in graphics. One may handle this nonconvergent behavior by terminating the computation when the CVT energy is found to increase.

5.4 Shader Language VS. CUDA

CUDA is a relatively new and popular C-like language for general purpose computation on the GPU. Since CUDA has some features not available in traditional shader languages (such as accessing the shared memory), it is much faster for applications which can benefit from the new features. However, Cg is better suited than CUDA for implementing CVT algorithms. To compare Cg and CUDA programs, we list detailed time breakdown for each step in Lloyd’s algorithm and the L-BFGS algorithm for a 2D case in Table 7 and Table 8 (the time is the total time for all iterations). It is clear that the JFA and the regional reduction (computing centroid for Lloyd’s algorithm, and computing CVT energy and its gradient for the L-BFGS algorithm) are the two steps dominating the total running time. The Cg version is faster for both steps than the CUDA version. In total, the CUDA versions are more than 40% slower than the Cg versions.

Table 7
Comparison of Cg and CUDA running time (in seconds) of Lloyd’s algorithm for 1,000 sites in the 2D domain of \([-1, 1] \times [-1, 1]\). All steps are executed 243 times.

<table>
<thead>
<tr>
<th>Step</th>
<th>Cg Time</th>
<th>CUDA Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>JFA</td>
<td>0.637</td>
<td>0.775</td>
</tr>
<tr>
<td>Compute Centroids</td>
<td>0.237</td>
<td>0.413</td>
</tr>
<tr>
<td>Test Stop Condition</td>
<td>0.015</td>
<td>0.107</td>
</tr>
<tr>
<td>Draw New Sites</td>
<td>0.041</td>
<td>0.032</td>
</tr>
<tr>
<td>Total Time</td>
<td>0.931</td>
<td>1.327</td>
</tr>
</tbody>
</table>

The better efficiency of the Cg implementation can be explained as follows. The JFA spends most of its time on accessing memory rather than computation. For every pixel, it needs to read information from nine pixels and write to one pixel (itself). The read addresses required by the JFA are non-coalesced [43] and change in every pass according to different step lengths. So it is very difficult, if not impossible, to make this step efficient with CUDA.

Computing the centroids in Lloyd’s algorithm (as well as the energy value and gradients computation in the L-BFGS algorithm) is essentially a regional reduction problem. The regional reduction is known to be difficult for CUDA, since it requires many threads writing to a same memory address. This is usually implemented using atomic operations [43]. Currently, CUDA only supports atomic operations on integers. So for floating point numbers, we have to mimic the atomic operations by tagging the five least significant bits of the thread ID (see [44], [45] for details). This is inefficient and hardware-dependent, since the warp size must be known in advance.

In conclusion, our algorithms for computing the CVT fit the traditional shader languages better than CUDA at this moment. However, as CUDA is fast evolving, we believe that efficient atomic operations on floating point numbers will be available soon. That will be likely to make CUDA faster than Cg for implementing our GPU algorithms in the near future.

6 Conclusion and Future Work

We have presented new techniques that use the GPU to compute the centroidal Voronoi tessellation both in 2D and on a surface. We proposed a novel algorithm to directly compute the Voronoi diagram on a surface, and also presented a new method using the vertex program to perform the regional reduction. Equipped with these two tools, we implemented two algorithms on the GPU – Lloyd’s algorithm and the L-BFGS algorithm. For Lloyd’s algorithm, the entire procedure is performed on the GPU; and for the L-BFGS algorithm, the major computational work is performed on the GPU. Our GPU implementations of the two methods have shown significant speedup over their CPU counterparts. Although our results are discrete approximations to the true CVTs, we believe that many applications can benefit from the fast computation of the CVT made possible by our GPU-based algorithms.

In our current implementation of the L-BFGS algorithm, only the computation of CVT energy values and gradients, which is the most time-consuming part, is
performed on the GPU. Currently, these values still need to be read back to the CPU for computing the new sites, thus slowing down the overall computation. If we could migrate this task to the GPU, the speedup would be even more significant.

The number of sites of the CVT is currently limited by the size of the 2D texture in the GPU (e.g. for a $512 \times 512$ texture, the number of sites should not exceed 10,000); otherwise, the results would be very poor due to the large approximation errors). Furthermore, when using the geometric image to represent a surface of complex shape, the surface parameterization often has a large distortion and that leads to large discretization error in computation. In the future we will consider applying our GPU-base method to a multiple-chart representation of a surface of complex shape in order to compute the CVT with a much larger number of sites or on a surface of arbitrary shape.

References


