# Computing and Rendering Point Set Surfaces 

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#### Abstract

We advocate the use of point sets to represent shapes. We provide a definition of a smooth manifold surface from a set of points close to the original surface. The definition is based on local maps from differential geometry, which are approximated by the method of moving least squares (MLS). The computation of points on the surface is local, which results in an out-of-core technique that can handle any point set. We show that the approximation error is bounded and present tools to increase or decrease the density of the points, thus allowing an adjustment of the spacing among the points to control the error. To display the point set surface, we introduce a novel point rendering technique. The idea is to evaluate the local maps according to the image resolution. This results in high quality shading effects and smooth silhouettes at interactive frame rates.


Index Terms-Surface representation and reconstruction, moving least squares, point sample rendering, 3D acquisition.

## 1 Introduction

POINT sets are receiving a growing amount of attention as a representation of models in computer graphics. One reason for this is the emergence of affordable and accurate scanning devices generating a dense point set, which is an initial representation of the physical model [34]. Another reason is that highly detailed surfaces require a large number of small primitives, which contribute to less than a pixel when displayed, so that points become an effective display primitive [41], [46].

A point-based representation should be as small as possible while conveying the shape, in the sense that the point set is neither noisy nor redundant. In [1], we have presented tools to adjust the density of points so that a smooth surface can be well-reconstructed. Fig. 1 shows a point set with varying density. Our approach is motivated by differential geometry and aims at minimizing the geometric error of the approximation. This is done by locally approximating the surface with polynomials using moving least squares (MLS). Here, we include proofs and explanations of the underlying mathematics, detail a more robust way to compute points on the surface, and derive bounds on the approximation error.

We understand the generation of points on the surface of a shape as a sampling process. The number of points is adjusted by either up-sampling or down-sampling the representation. Given a data set of points $P=\left\{p_{i}\right\}$ (possibly

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acquired by a 3D scanning device), we define a smooth surface $S_{P}$ (MLS surface) based on the input points (the definition of the surface is given in Section 3). We suggest replacing the points $P$ defining $S_{P}$ with a reduced set $R=$ $\left\{r_{i}\right\}$ defining an MLS surface $S_{R}$ which approximates $S_{P}$. This general paradigm is illustrated in 2D in Fig. 2: Points $P$, depicted in purple, define a curve $S_{P}$ (also in purple). $S_{P}$ is resampled with points $r_{i} \in S_{P}$ (red points). This typically lighter point set, called the representation points, now defines the red curve $S_{R}$ which approximates $S_{P}$.

Compared to the earlier version [1], we give more and new details on the (computational) properties of the surface definition:
Smooth manifold: The surface defined by the point set is a 2-manifold and expected to be $C^{\infty}$ smooth, given that the points are sufficiently close to the smooth surface being represented.
Bounded sampling error: Let $S_{R}$ be defined by the set of representation points $\left\{r_{i}\right\} \subset S_{P}$. The representation has bounded error $\varepsilon$, if $d\left(S_{P}, S_{R}\right)<\varepsilon$, where $d(\cdot, \cdot)$ is the Hausdorff distance.
Local computation: For computing a point on the surface only a local neighborhood of that point is required. This results in a small memory footprint which depends only on the anticipated feature size and not the number of points (in contrast to several other implicit surface definitions, e.g., those based on radial basis functions).
In addition to giving more implementation details for the rendering method from [1], we give a justification that builds on the error bounds introduced here. This connection to the bounded error substantiates the claimed properties of our rendering scheme:
High quality: Since $S_{R}$ is a smooth surface, proper resampling leads to smooth silhouettes and normals, resulting in superior rendering quality at interactive frame rates.


Fig. 1. A point set representing a statue of an angel. The density of points and, thus, the accuracy of the shape representation are changing (intentionally) along the vertical direction.

Single step procedure: Resampling respects screen space resolution and guarantees sufficient sampling, i.e., no holes have to be filled in a postprocessing step.

## 2 Related Work

### 2.1 Consolidation

Recent technological and algorithmic advances have improved the process of automatic acquisition of 3D models. Acquiring the geometry of an object starts with data acquisition, usually performed with a range scanner. This raw data contains errors (e.g., line-of-sight error cite [21], [47]) mainly due to noise intrinsic to the sensor used and its interaction with the real-world object being acquired. For a nontrivial object, it is necessary to perform multiple scans, each in its own coordinate system, and to register the scans [6]. In general, areas of the objects are likely to be covered by several samples from scans performed from different positions. One can think of the output of the registration as a thick point set.

A common approach is to generate a triangulated surface model over the thick point set. There are several efficient triangulation techniques, such as [2], [3], [5], [7], [17]. One of


Fig. 2. An illustration of the paradigm: The possibly noisy or redundant point set (purple curve). This manifold is sampled with (red) representation points. The representation points define a different manifold (red curve). The spacing of representation points depends on the desired accuracy of the approximation.
the shortcomings of this approach is that the triangulated model is likely to be rough, containing bumps and other kinds of undesirable features, such as holes and tunnels, and be nonmanifold. Further processing of the triangulated models, such as smoothing [51], [14] or manifold conversion cite [20], becomes necessary. The prominent difficulty is that the point set might not actually interpolate a smooth surface. We call consolidation the process of "massaging" the point set into a surface. Some techniques, such as Hoppe et al. [23], Curless and Levoy [12], and Wheeler et al. [55] consolidate their sampled data by using an implicit representation based on a distance function defined on a volumetric grid. In [23], the distances are taken as the signed distance to a locally defined tangent plan. This technique needs further processing [24], [22] to generate a smooth surface. Curless and Levoy [12] use the structure of the range scans and essentially scan convert each range surface into the volume, properly weighting the multiply scanned areas. Their technique is robust to noise and is able to take relative confidence of the samples into account. The work of Wheeler et al. [55] computes the signed distance to a consensus surface defined by weighted averaging of the different scans. One of the nice properties of the volumetric approach is that it is possible to prove, under certain conditions, that the output is a least-square fit of the input points (see [12]).

The volumetric sign-distance techniques described above are related to a new field in computer graphics called Volume Graphics pioneered by Kaufman and colleagues [26], [54], [50], which aims at accurately defining how to deal with volumetric data directly and answering questions related to the proper way to convert between surface and volume representations.

It is also possible to consolidate the point set by performing weighted averaging directly on the data points. In [53], model triangulation is performed first, then averaging is performed in areas which overlap. In [49], the data points are first averaged, weighted by a confidence in each measurement, and then triangulated.

Another approach to defining surfaces from the data points is to perform some type of surface fitting [18], such as fitting a polynomial [31] or an algebraic surface [42] to the data. In general, it is necessary to know the intrinsic topology of the data and (sometimes) have a parametrization before surface fitting can be applied. Since this is a nontrivial task, Krishnamurthy and Levoy [28] have proposed a semiautomatic technique for fitting smooth surfaces to dense polygon meshes created by Curless and Levoy [12]. Another form of surface fitting algorithms couples some form of high-level model recognition with a fitting process [44].

The process of sampling (or resampling) surfaces has been studied in different settings. For instance, surface simplification algorithms [9] sample surfaces in different ways to optimize rendering performance. Related to our work are algorithms which use particle systems for sampling surfaces. Turk [52] proposes a technique for computing level of details of triangular surfaces by first randomly spreading points on a triangular surface, then optimizing their positions by letting each point repel their neighbors. He uses an approximation of surface curvature to weight the number of points which should be placed in a given area of the surface. A related approach is to use physically-based particle systems to sample an implicit surface [56], [13]. Crossno and Angel [11] describe a system for sampling isosurfaces, where they use the curvature to automatically modulate the repulsive forces.

Lee [30] uses a moving-least squares approach to the reconstruction of curves from unorganized and noisy points. He proposes a solution for reconstructing two and three-dimensional curves by thinning the point sets. Although his approach resembles the one used here (and based on theory developed in [33]), his projection procedure is different and requires several iterations to converge to a clean point set (i.e., it is not actually a projection, but more of a converging smoothing step).

An alternative to our point-based modeling mechanism is proposed by Linsen [36]. His work is based on extending the $k$ neighborhood of a point to a "fan" by using an angle criterion (i.e., a neighborhood should cover the full 360 degrees around). Using this simple scheme, Linsen proposes a variety of operations for point clouds, including rendering, smoothing, and some modeling operations.

### 2.2 Point Sample Rendering

Following the pioneering work of Levoy and Whitted [35], several researchers have recently proposed using "points" as the basic rendering primitive, instead of traditional rendering primitives, such as triangulated models. One of the main reasons for this trend is that, in complex models, the triangle size is decreasing to pixel resolution. This is particularly true for real-world objects acquired as "textured" point clouds [39].

Grossman and Dally [19] presented techniques for converting geometric models into point-sampled data sets and algorithms for efficiently rendering the point sets. Their technique addresses several fundamental issues, including the sampling rate of conversion from triangles to points, and several rendering issues, including handling "gaps" in the rendered images and efficient visibility data structures.

The Surfels technique of Pfister et al. [41] builds and improves on this earlier work. They present alternative techniques for the sampling of the triangle mesh, including visibility testing, texture filtering, and shading.

Rusinkiewicz and Levoy [46] introduce a technique which uses a hierarchy of spheres of different radii to model a high-resolution model. Their technique uses small spheres to model the vertices at the highest resolution and a set of bounding spheres to model intermediate levels. Together with each spherical sample, they also save other associated data, such as normals. Their system is capable of time-critical rendering as it adapts the depth of tree traversal to the available time for rendering a given frame. In [45], they show how their system can be used for streaming models over a network.

Kalaiah and Varshney introduced an effective method for rendering point primitives that requires the computation of the principal curvatures and a local coordinate frame for each point [25]. Their approach renders a surface as a collection of local neighborhoods and it is similar to our rendering technique, proposed later, although they do not use dynamic level of detail in their system.

All the above techniques account for local illumination. Schaufler and Jensen [48] propose computing global illumination effects directly on point-sampled geometry by a ray tracing technique. The actual intersection point is computed, based on a local approximation of the surface, assuming a uniform sampling of the surface.

Point-based rendering suffers from the limited resolution of the fixed number of sample points representing the model. At some distance, the screen space resolution is high relative to the point samples, which causes undersampling. Tackling this problem by interpolating the surface points in image space is limited. A better approach is to resample the surface during rendering at the desired resolution in objectspace, guaranteeing that sampling density is sufficient with respect to the image space resolution.

Hybrid polygon-point approaches have been proposed. Cohen et al. [10] introduce a simplification technique which transitions triangles into (possibly multiple) points for faster rendering. Their system uses an extension of Floriani et al.'s Multi-Triangulation data structure [15], [16]. A similar system has been developed by Chen and Nguyen [8] as an extension of QSplat.

## 3 Defining the Surface-Projecting

Our approach relies on the idea that the given point set implicitly defines a surface. We build upon recent work by Levin [33]. The main idea is the definition of a projection procedure, which projects any point near the point set onto the surface. Then, the MLS surface is defined as the points projecting onto themselves. In the following, we explain the projection procedure, prove the projection and manifold properties, motivate the smoothness conjecture, and give details on how to efficiently compute the projection.

### 3.1 The Projection Procedure

Let points $p_{i} \in \mathbb{R}^{3}, i \in\{1, \ldots, N\}$, be sampled from a surface $S$ (possibly with a measurement noise). The goal is to project a point $r \in \mathbb{R}^{3}$ near $S$ onto a two-dimensional


Fig. 3. The MLS projection procedure: First, a local reference domain $H$ for the purple point $r$ is generated. The projection of $r$ onto $H$ defines its origin $q$ (the red point). Then, a local polynomial approximation $g$ to the heights $f_{i}$ of points $p_{i}$ over $H$ is computed. In both cases, the weight for each of the $p_{i}$ is a function of the distance to $q$ (the red point). The projection of $r$ onto $g$ (the blue point) is the result of the MLS projection procedure.
surface $S_{P}$ that approximates the $p_{i}$. The following procedure is motivated by differential geometry, namely, that the surface can be locally approximated by a function.

1. Reference domain: Find a local reference domain (plane) for $r$ (see Fig. 3). The local plane $H=$ $\left\{x \mid\langle n, x\rangle-D=0, x \in \mathbb{R}^{3}\right\}, n \in \mathbb{R}^{3},\|n\|=1$ is computed so as to minimize a local weighted sum of squared distances of the points $p_{i}$ to the plane. The weights attached to $p_{i}$ are defined as the function of the distance of $p_{i}$ to the projection of $r$ into the plane $H$, rather than the distance to $r$. Assume $q$ is the projection of $r$ onto $H$, then $H$ is found by locally minimizing

$$
\begin{equation*}
\sum_{i=1}^{N}\left(\left\langle n, p_{i}\right\rangle-D\right)^{2} \theta\left(\left\|p_{i}-q\right\|\right), \tag{1}
\end{equation*}
$$

where $\theta$ is a smooth, monotone decreasing function, which is positive on the whole space. By setting $q=$ $r+t n$ for some $t \in R$, (1) can be rewritten as:

$$
\begin{equation*}
\sum_{i=1}^{N}\left\langle n, p_{i}-r-t n\right\rangle^{2} \theta\left(\left\|p_{i}-r-t n\right\|\right) . \tag{2}
\end{equation*}
$$

We define the operator $\mathcal{Q}(r)=q=r+t n$ as the local minimum of (2) with the smallest $t$ and the local tangent plane $H$ near $r$ accordingly. The local reference domain is then given by an orthonormal coordinate system on $H$ so that $q$ is the origin of this system.
2. Local map: The reference domain for $r$ is used to compute a local bivariate polynomial approximation to the surface in a neighborhood of $r$ (see Fig. 3). Let $q_{i}$ be the projection of $p_{i}$ onto $H$, and $f_{i}$ the height of $p_{i}$ over H, i.e., $f_{i}=n \cdot\left(p_{i}-q\right)$. Compute the coefficients of a polynomial approximation $g$ so that the weighted least squares error

$$
\begin{equation*}
\sum_{i=1}^{N}\left(g\left(x_{i}, y_{i}\right)-f_{i}\right)^{2} \theta\left(\left\|p_{i}-q\right\|\right) \tag{3}
\end{equation*}
$$

is minimized. Here, $\left(x_{i}, y_{i}\right)$ is the representation of $q_{i}$ in a local coordinate system in $H$. Note that, again, the distances used for the weight function are from the projection of $r$ onto $H$. The projection $\mathcal{P}$ of $r$ onto $S_{P}$ is defined by the polynomial value at the origin, i.e., $\mathcal{P}(r)=q+g(0,0) n=r+(t+g(0,0)) n$.

### 3.2 Properties of the Projection Procedure

In our application scenario, the projection property (i.e., $\mathcal{P}(\mathcal{P}(r))=\mathcal{P}(r))$ is extremely important: We want to use the above procedure to compute points exactly on the surface. The implication of using a projection operator is that the set of points we project and the order of the points we project do not change the surface.

From (2), it is clear that if $(t, n)$ is a minimizer for $r$, then $(s, n)$ is a minimizer for $r+(s-t) n$. Assuming the minimization is global in a neighborhood $U \in \mathbb{R}^{3}$ of $q$, then $\mathcal{Q}$ is a projection operation in $U$, because the pair $(0, n)$ is a minimizer for $r-t n=q$. Further, $q+g(0,0) n=$ $r+(t+g(0,0)) n$ is also a minimizer and, thus, $\mathcal{P}$ is also a projection in $U$.

We like to stress that the projection property results from using distances to $q$ rather than $r$. If $\theta$ depended on $r$ the procedure would not be a projection because the values of $\theta$ would change along the normal direction.

The surface $S_{P}$ is formally defined as the subset of all points in $\mathbb{R}^{3}$ that project onto themselves. A simple counting argument shows that this subset is a twoparameter family and, thus, $S_{P}$ a 2-manifold. Equation (2) essentially has six degrees of freedom: three for $r$, two for $n$, and one for $t$. On the other hand, there are four independent necessary conditions: For a local minimum, the partial derivatives of the normal and $t$ have to be zero and, for $r$ to be on the surface, $t=0$ is necessary. This leaves $6-4=2$ free parameters for the surface. It is clear from simple examples that the parameter family includes manifolds which cannot be expressed as functions over $\mathbb{R}^{2}$.

The particular charm of this surface definition is that it avoids piecewise parameterizations. No subset of the surface is parameterized over a planar piece, but every single point has its own support plane. This avoids the common problems of piecewise parameterizations for shapes, e.g., parameterization dependence, distortions in the parameterization, and continuity issues along the boundaries of pieces.

However, in this approach, we have the problem of proving continuity for any point on the surface because its neighbors have different support planes in general. The intuition for $S_{P} \in C^{\infty}$ is, of course, that (1) interpreted as a function $\mathbb{R}^{6} \rightarrow \mathbb{R}$ is $C^{\infty}$ so that a particular kernel of its gradient is also $C^{\infty}$.

The approximation of single points is mainly dictated by the radial weight function $\theta$. The weight function suggested in [33] is a Gaussian

$$
\begin{equation*}
\theta(d)=e^{-\frac{d^{2}}{h^{2}}}, \tag{4}
\end{equation*}
$$

where $h$ is a fixed parameter reflecting the anticipated spacing between neighboring points. By changing $h$, the surface can be tuned to smooth out features of size $<h$ in $S$. More specifically, a small value for $h$ causes the Gaussian to


Fig. 4. The effect of different values for parameter $h$. A point set representing an Aphrodite statue defines an MLS surface. (a) shows an MLS surface resulting from a small value and reveals a surface micro-structure. (b) shows a larger value for $h$, smoothing out small features or noise.
decay faster and the approximation is more local. Conversely, large values for $h$ result in a more global approximation, smoothing out sharp or oscillatory features of the surface. Fig. 4 illustrates the effect of different $h$ values.

### 3.3 Computing the Projection

We explain how to efficiently compute the projection and what values should be chosen for the polynomial degree and $h$. Furthermore, we discuss tradeoffs between accuracy and speed.

Step 1 of the projection procedure is a nonlinear optimization problem. Usually, (2) will have more than one local minimum. By definition, the local minimum with the smallest $t$ has to be chosen, which means the plane should be close to $r$. For minimizing (2), we have to use some iterative scheme which descends toward the next local minimum. Without any additional information, we start with $t=0$ and first approximate the normal. Note that, in this case, the weights $\theta_{i}=\theta\left(\left\|p_{i}-r\right\|\right)$ are fixed. Let $B=$ $\left\{b_{j k}\right\}, B \in \mathbb{R}^{3 \times 3}$ be the matrix of weighted covariances defined by

$$
b_{j k}=\sum_{i} \theta_{i}\left(p_{i_{j}}-r_{j}\right)\left(p_{i_{k}}-r_{k}\right) .
$$

Then, the minimization problem (2) can be rewritten in bilinear form

$$
\begin{equation*}
\min _{\|n\|=1} n^{T} B n \tag{5}
\end{equation*}
$$

and the solution of the minimization problem is given as the Eigenvector of $B$ that corresponds to the smallest Eigenvalue.

If a normal is computed (or known in advance), it is fixed and the function is minimized with respect to $t$. This is a nonlinear minimization problem in one dimension. In general, it is not solvable with deterministic numerical methods because the number of minima is only bounded by the number $N$ of points $p_{i}$. However, in all practical cases, we have found that, for $t \in[-h / 2, h / 2]$, there is only one local minimum. This is intuitively clear as $h$ is connected to the feature size and features smaller than $h$ are smoothed out, i.e., features with distance smaller than $h$ do not provide several minima. For this reason, we make the assumption that any point $r$ to be projected is at most $h / 2$ away from its projection. In all our practical applications, this is naturally satisfied.

Using these prerequisites, the local minimum is bracketed with $\pm h / 2$. The partial derivative

$$
\begin{equation*}
2 \sum_{i=1}^{N}\left\langle n, p_{i}-r-t n\right\rangle\left(1+\frac{\left\langle n, p_{i}-r-t n\right\rangle^{2}}{h^{2}}\right) e^{\left\|p_{i}-r-t n\right\|^{2} / h^{2}} \tag{6}
\end{equation*}
$$

can be easily evaluated together with the function itself. The derivative is exploited in a simple iterative minimization scheme, as explained in [43, Chapter 10.3].

Once $t \neq 0$, fixing $t$ and minimization with respect to the normal direction is also a nonlinear problem because $q=$ $r+t n$ changes and, thus, the weights change also. The search space can be visualized as the tangent planes of a sphere with
center point $r$ and radius $t$. However, in practice, we have found the normal (or $q$ ) to change only slightly so that we approximate the sphere locally around the current value of $r+t n$ as the current plane defined $t$ and $n$. On this plane, a conjugate gradient scheme [43] is used to minimize among all $q$ on the plane. The main idea is to fix a subspace for minimization in which $n$ cannot vanish so that the constraint $\|n\|=1$ is always satisfied. The use of a simple linear subspace makes the computation of partial derivatives efficient and, thus, conjugate gradient methods applicable. Clearly, this search space effectively changes $t$, resulting in a theoretically worse convergence behavior. In practice, the difference between the sphere and the plane is small for the region of interest and the effect is not noticeable.

Using these pieces, the overall implementation of computing the support plane looks like this:
Initial normal estimate in $r$ : The normal in a point might be given as part of the input (e.g., estimated from range images) or could be computed when refining a point set surface (e.g., from close points in the already processed point set). If no normal is available, it is computed using the Eigenvector of the matrix of weighted covariances $B$.
Iterative nonlinear minimization: The following two steps are repeated as long as any of the parameters changes more than a predefined $\varepsilon$ :

1. Minimize along $t$, where the minimum is initially bracketed by $t= \pm h / 2$.
2. Minimize $q$ on the current plane $H:(t, n)$ using conjugate gradients. The new value for $q=r+t n$ leads to new values for the pair $(t, n)$.
The last pair $(t, n)$ defines the resulting support plane $H$.
The second step of the projection procedure is a standard linear least squares problem: Once the plane $H$ is computed, the weights $\theta_{i}=\theta\left(\left\|p_{i}-q\right\|\right)$ are known. The gradient of (3) over the unknown coefficients of the polynomial leads to a system of linear equations of size equal to the number of coefficients, e.g., 10 for a third degree polynomial.

Through experimentation, we found that high degree polynomials are likely to oscillate. Polynomials of degree 3 to 4 have proven to be very useful as they produce good fits of the neighborhood, do not oscillate, and are quickly computed.

### 3.4 Data Structures and Tradeoffs

The most time-consuming step in computing the projection of a point $r$ is collecting the coefficients from each of the $p_{i}$ (i.e., computing the sum). Implemented naively, this process takes $O(N)$ time, where $N$ is the number of points. We exploit the effect of the quickly decreasing weight function in two ways:

1. In a certain distance from $r$, the weight function is effectively zero. We call this the neglect distance $d_{n}$, which depends solely on $h$. A regular grid with cell size $2 d_{n}$ is used to partition the point set. For each projection operation, a maximum of eight cells is needed. This results in a very small memory footprint and yields a simple and effective out-of-core
implementation, which makes the storage requirements of this approach independent of the total size of the point set.
2. Using only the selected cells, the terms are collected using a hierarchical method inspired by solutions to the N-body problem [4]. The basic observation is that a cluster of points far from $r$ can be combined into one point. To exploit this idea, each cell is organized as an Octree. Leaf nodes contain the $p_{i}$; inner nodes contain information about the number of points in the subtree and their centroid. Then, terms are collected from the nodes of the Octree. If the node's dimension is much smaller than its distance to $r$, the centroid is used for computing the coefficients; otherwise, the subtree is traversed. In addition, whole nodes can be neglected if their distance to $r$ is larger than $d_{n}$.
The idea of neglecting points could be also made independent of numerical issues by using a compactly supported weight function. However, the weight function has to be smooth. An example for such an alternative is $\theta(x)=2 x^{3}-3 x^{2}+1$.

A simple way to trade accuracy for speed is to assume that the plane $H$ passes through the point to be projected. This assumption is reasonable for input points, which are expected to be close to the surface they define (e.g., input that has been smoothed). This simplification saves the cost of the iterative minimization scheme.

### 3.5 Results

Actual timings for the projection procedure depend heavily on the feature size $h$. On a standard Pentium PC, the points of the bunny were projected at a rate of 1,500-3,500 points per second. This means the 36 K points of the bunny are projected onto the surface they define themselves in 1030 seconds. Smaller values for $h$ lead to faster projection since the neighborhoods and, thus, the number of points taken into account are smaller.

As has been stressed before, the memory requirements mainly depend on the local feature size $h$. As long as $h$ is small with respect to the total diameter of the model the use of main memory of the projection procedure is negligible.

## 4 Approximation Error

Consider again the setting depicted in Fig. 2. Input points $\left\{p_{i}\right\}$ define a surface $S_{P}$, which are then represented by a set of points $\left\{r_{i}\right\} \in S_{P}$. However, the set $\left\{r_{i}\right\}$ defines a surface $S_{R}$ which approximates $S_{P}$. Naturally, we would like to have an upper bound on the distance between $S_{R}$ and $S_{P}$.

From differential geometry, we know that a smooth surface can be locally represented as a function over a local coordinate system. We also know that, in approximating a bivariate function, $f$ by a polynomial $g$ of total degree $m$, the approximation error is $\|g-f\| \leq M \cdot h^{m+1}$ [32]. The constant $M$ involves the $(m+1)$ th derivatives of $f$, i.e., $M \in O\left(\left\|f^{(m+1)}\right\|\right)$. In the case of surface approximation, since $S_{P}$ is infinitely smooth, there exists a constant $M_{m+1}$, involving the $(m+1)$ th derivatives of $S_{P}$ such that

$$
\begin{equation*}
\left\|S_{P}-S_{R}\right\| \leq M_{m+1} h^{m+1} \tag{7}
\end{equation*}
$$

where $S_{R}$ is computed using polynomials of degree $m$.
Note that this error bound holds for the MLS surface $S_{R}$, which is obtained by the projection procedure applied to the data set $R$, using the same parameter $h$ as for $S_{P}$. Moreover, the same type of approximation order holds for piecewise approximation: Assume each point $r_{i}$ defines a support plane so that $S_{P}$ is a function over a patch $[-h, h]^{2}$ around $r_{i}$ on the plane and, further, that the points $\left\{r_{i}\right\}$ leave no hole of radius more than $h$ on the surface. Then, the above arguments hold and the approximation error of the union of local, nonconforming, polynomial patches $G_{i}$ around points $r_{i}$ approximates $S_{P}$ as

$$
\begin{equation*}
\left\|S_{P}-\bigcup G_{i}\right\| \leq M_{m+1} h^{m+1} \tag{8}
\end{equation*}
$$

Here, $G_{i}$ is the polynomial patch defined by an MLS polynomial $g_{i}$ for the point $r_{i}$, with corresponding reference plane $H_{i}$ with normal $n_{i}$, a corresponding orthonormal system $\left\{u_{i}, v_{i}, n_{i}\right\}$, and an origin at $q_{i}$.

$$
\begin{equation*}
G_{i}=\left\{q_{i}+x \cdot u_{i}+y \cdot v_{i}+g_{i}(x, y) \cdot n_{i} \mid(x, y) \in[-h, h]^{2}\right\} . \tag{9}
\end{equation*}
$$

These error bounds explain nicely why curvature (see, e.g., [17]) is an important criterion for piecewise linear approximations (meshes): The error of piecewise linear functions depends linearly on the second derivatives and the spacing of points. This means, more points are needed where the curvature is higher.

However, when the surface is approximated with higher order polynomials, curvature is irrelevant to the approximation error. Using cubic polynomials, the approximation error depends on the fourth order derivatives of the surface. Note that our visual system cannot sense smoothness beyond second order [38]. From that point of view, the sampling density locally depends on an "invisible" criterion. We have found it to be sufficient to fix a spacing $h$ of points $r_{i}$ on $S_{P}$.

## 5 Generating the Representation Point Set

A given point set might have erroneous point locations (i.e., is noisy), may contain too many points (i.e., is redundant), or may not have enough points (i.e., is undersampled).

The problem of noise is handled by projecting the points onto the MLS surface they define. The result of the projection procedure is a thin point set. Redundancy is avoided by decimating the point set, taking care that it persists to be a good approximation of the MLS surface defined by the original point set. In the case of undersampling, the input point set needs to be upsampled. In the following sections, we show techniques to remove and add points.

Fig. 5 illustrates the idea of resampling in the example of the Buddha statue. Using the techniques presented below, it is possible to resample the geometry to be evenly sampled on the surface.

### 5.1 Downsampling

Given a point set, the decimation process repeatedly removes the point that contributes the smallest amount of information to the shape. The contribution of a point to the


Fig. 5. Points acquired by range scanning devices or vertices from a processed mesh typically have uneven sampling density on the surface (a). The sampling techniques discussed here allow us to evenly resample the object (b) to ensure a sufficient density for further processing steps, for example rendering (c).
shape or the error of the sampling is dictated by the definition of the shape. If the point set is reconstructed by means of a triangulation, criteria from the specific triangulation algorithm should control the resampling. Criteria include the distance of points on the surface [23], curvature [17], or distance from the medial axis of the shape [2]. For a direct display of the point set on a screen, homogeneous distribution of the points over the surface is required [19], [41]. Here, we derive a criterion motivated by our definition of the surface.

The contribution of a projected point $p_{i}$ to the surface $S_{P}$ can be estimated by comparing $S_{P}$ with $S_{P-\left\{p_{i}\right\}}$. Computing the Hausdorff distance between both surfaces is expensive and not suitable for an iterative removal of points of a large point set. Instead, we approximate the contribution of $p_{i}$ by its distance from its projection onto the surface $S_{P-\left\{p_{i}\right\}}$. Thus, we estimate the difference of $S_{P}$ and $S_{P-\left\{p_{i}\right\}}$ by projecting $p_{i}$ onto $S_{P-\left\{p_{i}\right\}}$ (projecting $p_{i}$ under the assumption it was not part of $P$ ).

The contribution values of all points are inserted into a priority queue. At each step of the decimation process, the point with the smallest error is removed from the point set and from the priority queue. After the removal of a point, the error values of nearby points have to be recalculated since they might have been affected by the removal. This process is repeated until the desired number of points is reached or the contributions of all points exceeds some prespecified bound.


Fig. 6. Noisy input points (green points) are projected onto their smooth MLS curve (orange points). The figures in (a) show the point sets and a close up view. The decimation process is shown in (b). Points are colorcoded as in Fig. 7.

Fig. 6 illustrates our decimation process applied on the set of red points depicted in Fig. 6a. First, the red points are projected on the MLS surface to yield the blue points. A close up view over a part of the points shows the relation between the input (red) points and the projected points. In Fig. 6b, we show three snapshots of the decimation process, where points are colored according to their error value; blue represents low error and red represents high error. Note that, in the sparsest set, all of the points have a high error, that is, their removal will cause a large error. As the decimation proceeds, fewer points remain and their importance grows and the error associated with them is larger. Fig. 7 shows the decimation process in 3D with corresponding renderings of the point sets.

### 5.2 Upsampling

In some cases, the density of the point set might not be sufficient for the intended usage (e.g., direct point rendering or piecewise reconstructions). Motivated by the error bounds presented in Section 4, we try to decrease the spacing among points. Additional points should be placed (and then projected to the MLS surface) where the spacing among points is larger then a specified bound.

The basic idea of our approach is to compute Voronoi diagrams on the MLS surface and add points at vertices of this diagram. Note that the vertices of the Voronoi diagram are exactly those points on the surface with maximum distance to several of the existing points. This idea is related to Lloyd's method [37], i.e., techniques using Voronoi diagrams to achieve a certain distribution of points [40].

However, computing the Voronoi diagram on the MLS surface is excessive and local approximations are used instead. More specifically, our technique works as follows: In each step, one of the existing points is selected randomly. A local linear approximation is built and nearby points are projected onto this plane. The Voronoi diagram of these points is computed. Each Voronoi vertex is the center of a circle that touches three or more of the points without including any point. The circle with largest radius is chosen and its center is projected to the MLS surface. The process is repeated iteratively until the radius of the largest circle is less than a user-specified threshold (see Fig. 8). At the end of the process, the density of points is locally near-uniform on the surface. Fig. 8 shows the original sparse point set containing 800 points, and the same object after adding 20 K points over its MLS surface.

## 6 Rendering

The challenge of our interactive point rendering approach is to use the representation points and (when necessary) create new points by sampling the implicitly defined surface at a resolution sufficient to conform to the screen space resolution (see Fig. 9 for an illustration of that approach).

Usually, the representation points are not sufficient to render the object in screen space. In some regions, it is not necessary to render all points as they are occluded, backfacing, or have higher density than needed. However, typically, points are not dense enough to be projected directly as a single pixel and more points need to be generated by interpolation in object space.


Fig. 7. The point set representing an Aphrodite statue is projected onto a smooth MLS-surface. (a) After removing redundant points, a set of 37 K points represents the statue. (b) The corresponding rendering is shown. The point set is decimated using point removal. (c) An intermediate stage of the reduction process is shown. Note that the points are color-coded with respect to their importance. Blue points do not contribute much to the shape and might be removed; red points are important for the definition of the shape. The final point set in (e) contains 20K points. The corresponding rendering is depicted in (d) and is visually close to the one in (b).


Fig. 8. The upsampling process: Points are added at vertices of the Voronoi diagram. In each step, the vertex with the largest empty circle is chosen. The process is repeated until the radius of the largest circle is smaller than a specified bound. The wavy torus originally consisting of 800 points has been upsampled to 20 K points.

### 6.1 Culling and View Dependency

The structure of our rendering system is similar to QSplat [46]. The input points are arranged into a bounding sphere hierarchy. For each node, we store a position, a radius, a normal coverage, and, optionally, a color. The leaf nodes additionally store the orientation of the support plane and the coefficients of the associated polynomial. The hierarchy is used to cull the nodes with the view frustum and to apply a hierarchical backface culling [29]. Note that culling is important for our approach since the cost of rendering the leaf nodes (evaluating the polynomials) is relatively high compared to simpler primitives. Moreover, if the traversal reaches a node with an extent that projects to a size of less than a pixel, this node is simply projected to the frame-buffer
without traversing its subtree. When the traversal reaches a leaf node and the extent of its bounding sphere projects to more than one pixel in screen space, additional points have to be generated.

The radius of a bounding sphere in leaf nodes is simply set to the anticipated feature size $h$. The bounding spheres for inner nodes naturally bound all of their subtree's bounding spheres. To compute the size of a bounding sphere in pixel space, the radius is scaled by the model-view transform and then divided by the distance of center and eye point in $z$ direction to accommodate the projective transform.

### 6.2 Sampling Additional Points

The basic idea is to generate a grid of points sufficient to cover the extent of a leaf node. However, projecting the grid points using the method described in Section 3 is too slow in interactive applications. The main idea of our interactive rendering approach is to sample the polynomials associated with the representation points rather than really projecting the points.

It is clear that the union of these polynomial patches is not a continous surface. However, the Hausdorff-error of this approximation is not worse than the error of the surface computed by projecting every point using the operator $\mathcal{P}$ (see Section 4). Since the Hausdorff-error in object space limits the screen space error, the error bound can be used to make the approximation error invisble in the resulting image.

However, to conform with the requirements formulated in Section 4, the point set and the associated polyonmials are required to be near-uniform on the surface. It might be necessary to first process a given point set with the upsampling methods presented in Section 5. This way, we ensure that the local, nonconforming (i.e., overlapping or intersecting) polynomials are a good approximation to the surface inside a patch $[-h, h]^{2}$ around a point and, thus, the resulting image shows a smooth surface. However, most


Fig. 9. Upsampling could be used generate enough points on the surface to conform with the resolution of the image to be rendered. The right image shows a close up rendered with splats.


Fig. 10. The patch size of a polynomial: Points inside a ball of radius $h$ around the red point $r$ are projected onto the support plane of the red point. The patch size is defined as the bounding disk (in local coordinates) of the projections. Note that using a disk of radius $h$ would lead to unpleasant effects in some cases, as the polynomial might leave the ball of radius $h$.
dense point sets can be readily displayed with the approach presented here. For example, Fig. 11 shows several renderings of the original Stanford Bunny data.

It is critical to properly define the extent of a polynomial patch on the supporting plane such that neighboring patches are guaranteed to overlap (to avoid holes), but do not overlap more than necessary. Since no interpoint connectivity information is available, it is unclear which points are immediate neighbors of a given point on the surface.

To compute the extent of a polynomial patch associated to the point $p_{i}$ on the support plane $H$, all points inside a ball of radius $h$ around the projection $q=\mathcal{Q}\left(p_{i}\right)$ are collected. These points are projected to the support plane $H$, which leads to local $(u, v)$ coordinates for each projected point. The extent is defined by a circle around $q$ that encloses all projected points. More specifically, assume $q$ has the local coordinate ( 0,0 ), the radius of the circle is given by the largest 2-norm of all local $(u, v)$ coordinates of projected points. Since the spacing of points is expected to be less than $h$, patches of neighboring points are guaranteed to overlap.

Note that using a constant extent (e.g., a disk of radius $h$ on the support plane) can lead to errors as the polynomial $g$ over $H$ might leave the ball of radius $h$, in which a good approximation of the point set is expected. Fig. 10 illustrates the computation of the patch sizes.

The grid spacing $d$ should be so computed that neighboring points have a screen space distance of less than a pixel. Thus, the grid spacing depends on the orientation of the polyonmial patch with respect to the screen. Since the normals change on each polyonmial patch, we rather use a simple heuristic to conservatively estimate $d$ : The grid spacing $d$ is computed so that a grid perpendicular to the viewing direction is sufficiently sampled in image space.

If the grid is, indeed, perpendicular to the viewing direction, the sampling is also correct on the polynomial. If the grid is not perpendicular to the viewing direction, the projected area might be oversampled or undersampled, depending on the orientation of the support plane and the derivatives of the polyonmial. Note, however, that sufficient sampling is guaranteed if the derivatives are less than 1. In practice, we have rarely encountered higher derivatives, so
we decided not to evaluate the maximum derivatives of all polynomial patches. However, this could be done in a preprocess and the density could be adjusted accordingly.

Upon the view-dependent grid spacing $d$, the polynomials are evaluated by a forward difference approach, where the polynomial is scanned across its extent in its local $u, v$ parametric space. The affine map transforming from support plane coordinates to world coordinates is factored into polynomial evaluation, thus generating points in world coordinates. These points are then fed into the graphics pipeline to be projected to the screen.

Surprisingly, we have found that quad meshes are processed faster by current graphics hardware than a set of points. For this reason we use quad meshes to represent the polynomial patches. This has the additional advantage that, during lazy evaluation (see below), no holes occur.

### 6.3 Grid Pyramids

The time-critical factor is the view-dependent evaluation of the points on the polynomial. Optimally, these are recomputed whenever the projected screen space size changes. To accelerate the rendering process, we store a grid pyramid with various resolutions per representation point. Initially, the pyramid levels are created, but no grid is actually evaluated. When a specific grid resolution is needed, the system creates and stores the level that slightly oversamples the polynomial for a specific resolution such that small changes in the viewing position do not result in new evaluations.

To enhance the interactivity of our approach, we use lazy evaluation during rotating or zooming. Once the viewer stops moving, a proper grid is chosen from the pyramid.

### 6.4 Results

We have tested our approach on a variety of point sets. Fig. 11 shows the renderings of the Stanford Bunny. In Fig. 11a, the original point set is shown. Splatting, Fig. 11b, does not lead to good results because the model is not sampled densely enough. The traditional Gouraud-shaded mesh in Fig. 11c and Fig. 11g is compared to our approach in Fig. 11d and Fig. 11h. Note the accuracy of the highlights. The nonconforming local polynomial patches are shown color-coded in Fig. 11e and Fig. 11f. An example of an environment mapping to demonstrate the normal continuity is given in Fig. 12. Note that silhouettes and normals are smooth, which leads to fewer distortions on the boundary and in the reflections.

The frame rates we achieve are mainly dictated by the number of visible representation points (i.e., graph traversal time) and the number of pixels to be filled. All models depicted in the paper are displayed at more than five frames per second in a $512^{2}$ screen window (see the accompanying video for more information). The number of representation points ranges from 1,000 (for the torus) to 900 K (for the angel statue). Tests are performed on a PC with GeForce2 graphics board.

## 7 Conclusion

In differential geometry, a smooth surface is characterized by the existence of smooth local maps at any point. In this work, we use this as a framework to approximate a smooth


Fig. 11. The Stanford Bunny: The points defining the bunny are depicted in (a) (some points are culled). Points are splatted in (b) to satisfy screen space resolution. Note the difference of a piecewise linear mesh over the points (c) and close up in (g) to the rendering of nonconforming polynomial patches (d) and (h). The patches are color-coded in (e) and (f).
surface defined by a set of points and we introduced new techniques to resample the surface to generate an adequate representation of the surface.

To render such surfaces, the surface is covered by a finite number, as small as possible, of nonconforming, over-


Fig. 12. Comparison of mesh rendering with our technique with environment mapping. The left column shows renderings of a mesh consisting of 1,000 vertices. The right column shows our technique using the vertices as input points. The environment maps emphasize the improved normal and boundary continuity.
lapping, polynomial patches. We showed that the error of these approximations is bounded and dependent on the spacing among points. Thus, it is possible to provide a point set representation that conforms with a specified tolerance.

Our paradigm for representing surfaces advocates the use of a point set (without connectivity) as a representation of shapes. This representation is universal in the sense that it is used from the beginning (i.e., acquisition) to the end (i.e., rendering) of a graphical representation's life cycle. Moreover, we believe that this work is a step toward rendering with higher order polynomials. Note that we have used simple primitives (points) to achieve this goal. This admits to the current trend of integrating high quality rendering features into graphics hardware.

It would be interesting to integrate our approach with combinatorial methods such as the one of Amenta et al. [2]. This would combine topological guarantees with the additional precision of higher order approximations and the possibility of smoothing out noise or small features.

Using different values for $h$, it is easy to generate more smooth or more detailed versions of a surface from one point set (see, for example, Fig. 4). A set of different versions could be used as a smooth-to-detailed hierarchy and would allow for multiresolution modeling [27]. Of course, $h$ is not necessarily a global parameter and could be adapted to the local feature size. Varying $h$ has several implications and utility in handling point sets (see [30] for a nice introduction to the issues in two dimensions), such as properly accounting for differences in sampling rate and levels of noise during the acquisition process. Also,
nonradial functions might be necessary to properly account for sharp features in the models.

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