# Efficient Reconstruction of Large Scattered Geometric Datasets using the Partition of Unity and Radial Basis Functions 

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#### Abstract

We present a new scheme for the reconstruction of large geometric data. It is based on the well-known radial basis function model combined with an adaptive spatial and functional subdivision associated with a family of functions forming a partition of unity. This combination offers robust and efficient solution to a great variety of 2D and 3D reconstruction problems, such as the reconstruction of implicit curves or surfaces with attributes starting from unorganized point sets, image or mesh repairing, shape morphing or shape deformation, etc. After having presented the theoretical background, the paper mainly focuses on implementation details and issues, as well as on applications and experimental results.


Keywords: radial basis functions, partition of unity, surface reconstruction, implicit modeling

## 1 MOTIVATION

In many applications fields, real-world datasets are often provided as a non-uniform, unorganized set of a large amount of discrete data. The main (and maybe the most difficult) problem to solve, in order to exploit these scattered data, is to efficiently and precisely reconstruct a continuous function starting from this dataset. In the field of geometric modeling, for instance, this problem has become of major importance due to the rapid development of 3D range scanners that acquire 3D geometries as an unstructured set of points.

Reconstructing a continuous function starting from unorganized data sets has been intensively studied over the last decades. Generally, the techniques can be divided into two major categories. The first category tries to generate a parametric function that interpolates or approximates the initial dataset. Piecewise linear approximation is the easiest and the most popular technique in this category. The second category interpolates or approximates the dataset by building a family of implicit real-valued scalar functions where the reconstructed domain is defined as the zero-set of them.

We will focus in this paper on the reconstruction of implicit surfaces without loss of generality. About 20 years ago, in an extensive survey, Franke [Frank82] identified radial basis functions (RBFs) as one of the most accurate and stable methods to solve scattered data interpolation problems. The pionneering work to interpolating surfaces

[^0]

Figure 1: Example of surface reconstruction from unorganized points.
using RBFs starting from unorganized point sets can be attributed to Savchenko et al. [Savch95] and Turk and O'Brien [Turk98]. Using these techniques, the implicit surface is calculated by solving a linear system. Unfortunately, since the RBFs have global support, the equations lead to a dense linear system. Hence, both techniques fail to reconstruct surfaces from large point sets consisting of more than several thousands of points.

To overcome this problem, by using Wendland's compactly supported RBFs [Wendl95], Morse et al. [Morse01] showed how to reconstruct implicit surfaces from larger datasets since the involved linear system becomes sparse. This algorithm was further improved by Kojekine et al. [Kojek03] by organizing the involved sparse matrix into a band-diagonal sparse matrix which can be solved more efficiently. Unfortunately, the radius of support has to be chosen globally, which means that the method is not robust against non-uniform datasets where the density of the samples may vary significantly over the dataset. A multi-scale approach such as the one proposed by Ohtake et al. [Ohtak03b] overcomes this limitation, but it is not feasible for the approximation of noisy data. Anyway, for all techniques using compactly supported radial basis functions, the number of points that can be processed is still limited since the techniques remain global in nature.

A different approach to interpolate large point sets has
been proposed by Carr et al. [Carr01] based on a fast evaluation of RBF technique by Beatson et al. [Beats97, Beats92] using fast multipole methods. Unfortunately, the far field expansion has to be done for every radial basis function, and is very complex to implement.

Another fast evaluation method of RBFs, based on the partition of unity method, was proposed by Wendland in a theorical survey [Wend102a] and a more practical sketch [Wendl02b] on which we have based our approach.
Two other methods to reconstruct implicit surfaces without using radial basis functions have drawn much attention recently. In the first one, called Point Set Surfaces, presented by Alexa et al. [Alexa01, Alexa03] an implicit surface is reconstructed using a projection operator based on the method of moving least squares. The resulting implicit surfaces is defined by all points that project on themselves, and the defining function is never calculated explicitly. Unfortunately, the computation of the projection operator is rather expensive since a non-linear optimization problem is involved.
In the second one, called MPU implicits, developped recently by Ohtake et al. [Ohtak03a], the partition of unity method [Frank80] is used to reconstruct implicit surfaces. By using weighted sums of different types of piecewise quadratic functions capturing the local shape of the surface, implicit surfaces from very large point sets can be reconstructed while preserving sharp features.
We combine in this paper two well-known methods in order to obtain a new reconstruction scheme for large datasets. RBFs are used to solve a set of small local problems and the partition of unity (POU) method combines the local solutions together to get the final reconstruction. As we will see, this combination is not only robust and efficient, but also offers a high level of scalability as the data does not need to be completely stored in memory, but can be accessed sequentially from disk.
Note that our presentation focuses more on geometric data, because it is the application field where there are the larger available datasets. But our technique is valid for any dimension discrete dataset. This is not the case for alternative reconstruction techniques such as Point Set Surfaces [Alexa01, Alexa03] or MPU [Ohtak03a], which are totally driven by the specific properties of 3D surface reconstruction. As an example of multidimensional reconstruction we present a technique that combines the reconstruction of geometric and colorimetric discrete data to generate an implicit surface with solid texture.
The paper is oganized as follows: Section 2 provides the theoretical background of our technique. In Section 3, we present our implementation and the practical choices to obtain an easy, efficient, and robust reconstruction algorithm. Section 4 presents some 3D applications and experimentals results, and in Section 5 we conclude and indicate some directions to future work.

## 2 THEORY

### 2.1 Radial basis functions

Given the set of $N$ pairwise distinct points $P=\left\{\mathbf{p}_{1}, \ldots, \mathbf{p}_{N}\right\}$ of dimension $d: \mathbf{p}_{k} \in \mathbb{R}^{d}$, and the set of values $\left\{h_{1}, \ldots h_{N}\right\}$, we want to find a function $f: \mathbb{R}^{d} \rightarrow \mathbb{R}$ with

$$
\begin{equation*}
\forall i \quad f\left(\mathbf{p}_{i}\right)=h_{i} . \tag{1}
\end{equation*}
$$

In order to obtain a radial basis function (RBF) reconstruction of the point set $P$, a function $f$ satisfying the equation

$$
\begin{equation*}
f(\mathbf{p})=\sum_{i=1}^{N} \omega_{i} \phi\left(\left\|\mathbf{p}, \mathbf{p}_{i}\right\|\right)+\pi(\mathbf{p}) \tag{2}
\end{equation*}
$$

has to be found. We denote here $\left\|\mathbf{p}_{i}, \mathbf{p}_{j}\right\|$ the Euclidean distance, $\omega_{i}$ the weights, $\phi: \mathbb{R} \rightarrow \mathbb{R}$ a basis function, and $\pi$ a polynomial of degree $m$ depending on the choise of $\phi . \pi(\mathbf{p})=\sum c_{i} \pi_{i}(\mathbf{p})$ with $\left\{\pi_{\alpha}\right\}_{\alpha=1}^{Q}$ a basis in the $d$ dimensional null space containing all real-valued polynomials in $d$ variables and of order at most $m$, hence $Q=\binom{m+d}{d}$.

The basis function $\phi$ has to be conditionally positive definite [Iske02], and some popular choices proposed in the literature are shown below:

$$
\begin{array}{rll}
\text { biharmonic } & \phi(r)=r & \text { with } \pi \text { of degree } 1 \\
\text { pseudo-cubic } & \phi(r)=r^{3} & \text { with } \pi \text { of degree } 1 \\
\text { triharmonic } & \phi(r)=r^{3} & \text { with } \pi \text { of degree } 2 \tag{5}
\end{array}
$$

As we have an under-determined system with $N+Q$ unknowns ( $\omega$ and $\mathbf{c}$ ) and $N$ equations, so-called natural additional constraints for the coefficients $\omega$ are added, so that

$$
\begin{equation*}
\sum_{i} \omega_{i} c_{1}=\sum_{i} \omega_{i} c_{2}=\ldots=\sum_{i} \omega_{i} c_{Q}=0 \tag{6}
\end{equation*}
$$

The equations (1), (2), and (6) determine the following linear system:

$$
\begin{align*}
A \mathbf{x} & =\mathbf{b}  \tag{7}\\
A & =\left[\begin{array}{cc}
\Phi & P^{T} \\
P & 0
\end{array}\right]  \tag{8}\\
\Phi & =\left[\left.\phi\left(\left\|\mathbf{p}_{\mathbf{i}}, \mathbf{p}_{\mathbf{j}}\right\|\right)\right|_{\substack{i=1 \ldots N \\
j=1 . \ldots N}}\right] \\
P & =\left[\left.\pi_{\alpha}\left(\mathbf{p}_{\mathbf{i}}\right)\right|_{\substack{i=1 \ldots N \\
\alpha=1 \ldots Q}}\right] \\
\mathbf{x} & =\left[\omega_{1}, \omega_{2}, \ldots, \omega_{N}, c_{1}, c_{2}, \ldots, c_{Q}\right]^{T} \\
\mathbf{b} & =[h_{1}, h_{2}, \ldots, h_{N}, \underbrace{0,0, \ldots, 0}_{Q \text { times }}]^{T} \tag{9}
\end{align*}
$$

The solution vector $\mathbf{x}$ is composed of the weights $\omega_{i}$ and the polynomial coefficients $c_{i}$ for equation (2) and represent a solution of the interpolation problem given by (1).
If an approximation rather than an interpolation is required, one solution is to modify the diagonal of the matrix $\Phi \leftarrow \Phi-8 k \pi \rho \mathbb{I}$ (see Carr [Carr01, Carr03]), where the parameter $\rho$ controls the fitting tolerance (i.e. the result is getting smoother when $\rho$ is increased).

### 2.2 Partition of unity method

The main idea of the partition of unity (POU) approach is to divide the global domain of interest into smaller domains where the problem can be solved locally. More formally, the global difficult problem $P$ is decomposed into several smaller local problems $P_{i}$ and their local solutions $S_{i}$ are combined together using the weighting coefficients of $S_{i}$ that act as smooth "glueing" functions to obtain a global solution $S$.

Consider a global domain $\Omega$ and divide it into $M$ "slightly" overlapping subdomains $\left\{\Omega_{i}\right\}_{i=1}^{M}$ with $\Omega \subseteq$ $\bigcup_{i} \Omega_{i}$. On this set of subdomains $\left\{\Omega_{i}\right\}_{i=1}^{M}$, we construct a partition of unity, i.e. a collection of non-negative functions $\left\{w_{i}\right\}_{i=1}^{M}$ with limited support $\operatorname{supp}\left(w_{i}\right) \subseteq \Omega_{i}$ and with $\sum w_{i}=1$ in the entire domain $\Omega$.

For each $\Omega_{i}$, a set $P_{i}=\left\{\mathbf{p} \in P \mid \mathbf{p} \in \Omega_{i}\right\}$ is constructed, and a local reconstruction function $f_{i}$ is computed. The global solution is then defined as a combination of the local functions weighted by the partition functions $w_{i}$.

$$
\begin{equation*}
F(\mathbf{p})=\sum_{i=1}^{M} f_{i}(\mathbf{p}) w_{i}(\mathbf{p}) \tag{10}
\end{equation*}
$$

The condition $\sum w_{i}=1$ is obtained from any other set of smooth functions $W_{i}$ by a normalization procedure

$$
\begin{equation*}
w_{i}(\mathbf{p})=\frac{W_{i}(\mathbf{p})}{\sum_{j} W_{j}(\mathbf{p})} . \tag{11}
\end{equation*}
$$

Any function $W_{i}$ is appropriated, but to guarantee the continuity of the global interpolation function $F$, it has to be continuous at the boundary of the regions $\Omega_{i}$.

### 2.3 Complexity analysis

The solution of the linear system (7) of size $N$ requires $O\left(N^{3}\right)$ floating point operations and $O\left(N^{2}\right)$ core-memory cells. Thus it is clear that direct methods are not suitable for a number of constraints greater than several thousands.
In the partition of unity approach, $\Omega$ is divided into $M$ subdomains. With a "good", quasi-uniform distribution, every $\Omega_{i}$ contains $N / M$ constraints in average. The new solution requires $O\left(M(N / M)^{3}\right)$ operations and $O\left((N / M)^{2}\right)$ cells. As $N / M$ can be considered as a constant, the reconstruction complexity is in $O(N)$, hence it is linear with respect to the number of constraints.

Another benefit using the POU method is in the evaluation of the interpolating function (2). The global RBF approach requires $O(N)$ operations for one single evaluation. Using the POU approach, two steps are required: first, finding all regions containing the point to evaluate, and second, evaluating the radial basis functions of the small local regions. Thus, the number of operations required is $O(M+N / M)$. Using an appropriate partitioning and data structure for fast neighbor searching like octree or kd-tree, we can reduce the first step to $O(\log N)$ or even $O(1)$, and with $N / M$ considered as a constant, the evaluation complexity is $O(1)$.

## 3 RECONSTRUCTION SCHEME

### 3.1 Partitioning and local reconstruction

According to the equations (10) and (11) two families of functions have to be built: the weighting functions $W_{i}$ and the local reconstruction functions $f_{i}$. Our reconstruction algorithm requires two steps: a space partition step that determines a set of overlapping domains $\left\{\Omega_{i}\right\}$ with associated weighting functions $\left\{W_{i}\right\}$, and a reconstruction step that computes the set of local functions $\left\{f_{i}\right\}$.
As we strive for an optimal reconstruction time, we have to obtain a quasi-uniform repartition of the points in the

```
Algorithm 1 Partition \(\left(P, \Omega_{i}\right)\)
Require: points \(P\), domain \(\Omega_{i}\)
Ensure: set of domains \(\left\{\Omega_{j}\right\}\)
    compute \(n\) number of points \(P\)
    if \(n>T_{\text {max }}\) then
        subdivide \(\Omega_{i}\) into overlapping \(\Omega_{i}^{1}, \ldots, \Omega_{i}^{k}\)
        Partition \(\left(P, \Omega_{i}^{1}\right)\)
        \(\dddot{\text { Partition }}\left(P, \Omega_{i}^{m}\right)\)
    else if \(n<T_{\text {min }}\) then
        while \(n \notin\left[T_{\text {min }}, T_{\text {max }}\right]\) do
            if \(n<T_{\text {min }}\) then
                enlarge \(\Omega_{i}\)
            else if \(n>T_{\text {max }}\) then
                reduce \(\Omega_{i}\)
            end if
        end while
        domain OK , add \(\Omega_{i}\) to \(\left\{\Omega_{j}\right\}\)
    else
        domain OK , add \(\Omega_{i}\) to \(\left\{\Omega_{j}\right\}\)
    end if
```

domains $\Omega_{i}$. The recursive algorithm 1 describes an adaptive subdivision of the domain $\Omega_{i}$, so that each subdomain contains between $T_{\text {min }}$ and $T_{\max }$ points.

If we call this function with the bounding domain $\Omega$ of the point set $P$, the result is a set $\left\{\Omega_{1}, \ldots, \Omega_{M}\right\}$ where $\forall \Omega_{i} T_{\text {min }} \leq \operatorname{Card}\left(P_{i}\right) \leq T_{\text {max }}$.
Starting from the domain $\Omega_{j}$ we divide it recursively into $m$ overlapping domains $\Omega_{j}^{1} \ldots \Omega_{j}^{m}$. When the number of points $P_{j}^{n}$ in $\Omega_{j}^{n}$ domain is higher than $T_{\max }$ we continue with the recursive subdivision. If the number of points is smaller than $T_{\max }$, the recursion is terminated and the current domain is added to the $\left\{\Omega_{i}\right\}$ set. However, if a domain $\Omega_{i}$ does not contain enough points, the local interpolation $f_{i}$ can lead to unexpected results. Hence, when the number of points is smaller than $T_{\text {min }}$, we have to adjust $\Omega_{i}$. The adjusting scheme is an iterative process when $\Omega_{i}$ is enlarged and reduced until the number of points $P_{i}$ is between $T_{\min }$ and $T_{\max }$ as explained in the following subsection.

The second step is quite simply: for every domain $\Omega_{i}$ in the $\left\{\Omega_{i}\right\}$ set, a RBF reconstruction $f_{i}$ is computed from the point set $P_{i}$ as shown in section 2.1.


Figure 2: Space subdivision ( $\Omega_{i}$ and $\Omega_{i}^{4}$ ) and domain enlarging $\left(\Omega_{i}^{3}\right)$ in order to adaptively balance the number of points per domain.

Figure 2, shows a hierarchy of domains, created by our partitioning algorithm. The final $\left\{\Omega_{i}\right\}$ set contains all the leafs of the tree.

The $T_{\text {max }}$ parameter controls time and stability. Increasing its value improves the local reconstruction at the cost of evaluation time. For the surface reconstruction, $T_{\max } \in[120,200]$ is a reasonable threshold. We use also $T_{\text {min }}>30$ that leads to a reconstruction without visible artifacts.

### 3.2 Operations on domains

The domains $\Omega_{i}$ can be of any shape, in practice, we use two simple and convex objects: axis-aligned ellipsoids of center $C$ and axis $A$ (Figure 3(a)), and axis-aligned bounding boxes defined from the two opposite corners $S$ and $T$ (Figure 3(a))
In order to enlarge or reduce a domain we simply scale it by a factor while keeping the domain center as shown Figure 3(b).

Figure 3(c) shows a subdivision step. In this stage a domain $\Omega$ is divided in eight (in the 3D case) equal-sized, overlapping subdomains based on a classical octree decomposition.
The table formalizes such operations according to $S, T, C$ and $A$ parameters. We denote the new parameters of the transformed domain with $S^{\prime}, T^{\prime}, C^{\prime}$ and $A^{\prime}$.

|  | ellipsoid $\Omega_{e}$ | box $\Omega_{b}$ |
| :--- | :--- | :--- |
| enlarge or | $A^{\prime}=k A$ | $S^{\prime}=S-\frac{(k-1)(T-S)}{2}$ |
| reduce | $C^{\prime}=C$ | $T^{\prime}=T+\frac{(k-1)(T-S)}{2}$ |
| subdivide | $A^{\prime}=\frac{\sqrt{3}}{2} d A$ | $S^{\prime}=S-\frac{(d-1)(T-S)}{4}$ |
|  | $C^{\prime}=C-\frac{A}{2}$ | $T^{\prime}=\frac{S+T}{2}+\frac{(d-1)(T-S)}{4}$ |

The cofficient $k$ denotes a scale factor in the iterative process of the enlarge/reduce scheme in order to adapt the domain size to the number of points. In practice, we take $k=1.05$ for enlarge and $k=0.98$ for reduce. The coefficient $d$ is an overlap factor. A larger value for $d$ yields larger overlapping zones and larger subdomains and hence a more stable reconstruction, but also increases the total number of domains and the reconstruction time ( $d=1.1$ is a good compromise).

### 3.3 Weighting function

The choice of the weighting functions $W_{i}$ determines the continuity between the local solutions $f_{i}$ and the continuity of the global reconstruction function $F$. Our weighting functions $W_{i}$ are defined as the composition of a distance function $D_{i}: \mathbb{R}^{n} \rightarrow[0,1]$, where $D_{i}(\mathbf{p})=1$ at the boundary of $\Omega_{i}$ and a decay function $V:[0,1] \rightarrow[0,1]$ : $W_{i}(\mathbf{p})=V \circ D_{i}(\mathbf{p})$.
We propose two formulations for the distance function $D_{i} ; D_{i}^{b}$ for a box and $D_{i}^{e}$ for an ellipsoid:

$$
\begin{aligned}
& D_{i}^{b}(\mathbf{p})=1-\prod_{r \in x, y, z} \frac{4\left(\mathbf{p}^{(r)}-S^{(r)}\right)\left(T^{(r)}-\mathbf{p}^{(r)}\right)}{\left(T^{(r)}-S^{(r)}\right)^{2}} \\
& D_{i}^{e}(\mathbf{p})=\sum_{r \in x, y, z} \frac{\left(\mathbf{p}^{(r)}-C^{(r)}\right)^{2}}{\left(A^{(r)}\right)^{2}}
\end{aligned}
$$

Depending on the choice of the decay function $V$, a more or less smooth weighting function $W_{i}$ is created with the desired continuity over the domain $\Omega_{i}$. We suggest to use one of the following formulations for $V$ that were chosen


Figure 3: Operations on the ellipsoid and box regions.


Figure 4: Distance function and weighting function for a rectangular region.
by including some simple constraints similar to the construction of base spline functions $(V(0)=1, V(1)=0$, $V^{\prime}(0)=V^{\prime}(1)=0$, etc.).

$$
\begin{array}{ll}
\mathbb{C}^{0}: & V^{0}(d)=1-d \\
\mathbb{C}^{1}: & V^{1}(d)=2 d^{3}-3 d^{2}+1 \\
\mathbb{C}^{2}: & V^{2}(d)=-6 d^{5}+15 d^{4}-10 d^{3}+1
\end{array}
$$

Figure 4 shows the geometric interpretation of these formules for a 2 D rectangular domain.

### 3.4 Reconstruction constraints

When the set of values $\left\{h_{1}, \ldots h_{k}\right\}$ of the conditions (1) are distinct, we can solve various reconstruction problems using the technique described so far. However, when reconstructing implicit surfaces from a set of $N$ distinct points $P=\left\{\mathbf{p}_{1}, \ldots, \mathbf{p}_{N}\right\}$, we want to find a function satisfying

$$
\begin{equation*}
\forall i F\left(\mathbf{p}_{i}\right)=0 \tag{12}
\end{equation*}
$$



Figure 5: Reconstruction of the Stanford Bunny and Dragon from different initial point sets.
and the conditions (1) are not sufficient because $h_{1}=h_{2}=\ldots=h_{N}=0$.
Unfortunately, with these conditions, the system (7) has the trivial solution, the constant function $F(\mathbf{p})=0$, which is not useful. One common solution [Turk98, Carr01] is to use off-surface points: the constraints (12) are completed by an additional set of points $P^{\prime}=\left\{\mathbf{p}_{1}^{\prime}, \ldots, \mathbf{p}_{l}^{\prime}\right\}$ where $F\left(\mathbf{p}_{i}^{\prime}\right)=h_{i} \neq 0$. These off-surface points $\mathbf{p}_{i}^{\prime}$ can be computed starting from the initial points $\mathbf{p}_{j}$ and moving them along the normal vector: $\mathbf{p}_{i}^{\prime}=\mathbf{p}_{j}+k_{j} \mathbf{n}_{j}$. The normal is usually obtained during data acquisition, however, when the normal is not available, it can be estimated from neighboring points [Hoppe92].
Based on the common convention that $F(\mathbf{p})>0$ inside and $F(\mathbf{p})<0$ outside the surface, Turk and O'Brien [Turk98] add one off-surface point at the exterior of the surface $\left(k_{j}>0, h_{i}=-1\right)$ on every initial point. Carr et al. [Carr01] propose to add two new points on both sides of the surface $\left(k_{j}>0, h_{i}=-1\right.$ or $\left.k_{j}<0, h_{i}=-1\right)$ for a subset of the initial points.

In practice, we found that taking a translation value $k_{i}$ as $1 \%$ of the length of the bounding box is often, but not always, sufficient. Carr et al. [Carr01] give a simple condition to reconstruct surfaces without auto-intersections: for every off-surface point $\mathbf{p}_{i}^{\prime}=\mathbf{p}_{j}+k_{j} \mathbf{n}_{j}$, the nearest surface point has to be the point $\mathbf{p}_{j}$ that it is derived of.

## 4 APPLICATIONS AND RESULTS

All results presented in this section were performed on an Intel Pentium 1.7 GHz with 512 MB of RAM running Linux. To solve the linear systems, we used the linear solver from the GNU Scientific Library package [GNU] based on LU-decomposition.
To visualise the resulting implicit surfaces, we used a polygonizer such as [Loren87, Bloom94] to create a polygonal mesh. We can even imagine a more topologically stable algorithm based on knowlegde about the initial points [Cresp02]. As other possible solutions,
we can mention the oriented particles method [Witki94, Lomba95] and a mixed forward and backward warping method [Reute03].

Our first application is the reconstruction of implicit surfaces starting from unorganized point sets. Table 1 presents the processing times depending on the initial number of constraints.
We denote $\# P$ the number of points (with two additional off-surface points the total number of constraints is $3 \# P$ ), $M$ the number of regions, $t_{\text {rec }}$ the geometry reconstruction time in seconds, $t_{p o l y}$ the polygonization time in seconds using Bloomenthal polygonizer [Bloom94] with a resolution of $2 \%$ of object's bounding box size, and $e_{R M S}$ the RMS error of the reconstructed surface.

| Model | $\# \mathrm{P}$ | M | $t_{\text {rec }}$ | $e_{R M S}$ | $t_{\text {poly }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Bunny | 1000 | 50 | 6 | 0.19 | 126 |
|  | 2000 | 157 | 17 | 0.17 | 182 |
|  | 4000 | 223 | 24 | 0.13 | 134 |
|  | 8000 | 676 | 73 | 0.10 | 218 |
|  | 16000 | 946 | 106 | 0.07 | 170 |
|  | 32000 | 2577 | 329 | 0.04 | 288 |
| Buddha | 25000 | 1814 | 143 | 0.15 | 445 |
|  | 50000 | 3553 | 369 | 0.12 | 701 |
|  | 100000 | 7431 | 717 | 0.10 | 1020 |
|  | 150000 | 10126 | 1216 | 0.08 | 1400 |
|  | 300000 | 22285 | 2652 | 0.02 | 2354 |
|  | 500000 | 33745 | 5254 | 0.005 | 4064 |

Table 1: Processing time and error of different models.
The initial models Bunny, Happy Buddha and Dragon borrowed from Stanford 3D Scanning Repository [Lar] were downsampled and the complete process consisting of reconstruction and polygonizing was applied. At the same time we computed the RMS error of the downsampled constraints set compared to the initial point set. We can confirm the linear complexity of our reconstruction process, whereas usual polygonal reconstruction (based on Delaunay triangulation for instance) is $O(N \log N)$ at best. Note that the global error decreases proportionally to the number of points as it is expected. Figures 1 and 5


Figure 6: Reconstruction time in function of number of points. Bunny and Buddha. See also Table 1


Figure 7: Linear morphing between Max Planck head and Igea.
show the examples of the reconstruction quality.
The influence of the two thresholds $T_{\min }$ and $T_{\max }$ is shown in Table 2. The same 8000 points model of the Dragon [Lar] was processed with different parameters. Visual results have shown that increasing $T_{\max }$ above a threshold does not influence the reconstruction quality but only increases the processing time, whereas a too small $T_{\text {min }}$ can lead to visible artefacts.

| $T_{\min }$ | $T_{\max }$ | M | $t_{\text {rec }}$ | $t_{\text {poly }}$ |
| :---: | :---: | :---: | :---: | :---: |
| 20 | 50 | 1474 | 23 | 93 |
| 50 | 100 | 638 | 82 | 135 |
| 50 | 200 | 434 | 104 | 143 |
| 50 | 300 | 251 | 226 | 187 |
| 50 | 400 | 195 | 364 | 237 |
| 100 | 300 | 251 | 283 | 253 |
| 100 | 400 | 195 | 451 | 309 |

Table 2: Influence of $T_{\min }$ and $T_{\max }$.
All characteristics of implicit surfaces that are reconstructed using our implementation still apply, such as function-based shape modelling [Pasko95]. As an example, we show in Figure 7 the linear morphing from the Max Planck head to the Igea head.

The second application is the possibility to combine geometric reconstruction with the reconstruction of additional surface attributes such as color channels, reflectance, and others. Such attributes can be considered as an intersection of the surface and a 3D procedural solid texture, where each attribute is reconstructed separately. Note that the number of constraints to reconstruct the implicit surface can be different from the number of constraints to reconstruct the attributes (Figure 8). As said, this is a unique feature of our RBF/POU reconstruction technique compared to alternative technique such as

Point Set Surfaces or MPU that can only handle geometric constraints.
A nice characteristic is, that the same deformation function applied to the implicit surface and the attributes conserves the geometry and texture coherence. As an example Figure 9 shows a Chameleon model twisted along the z axis.


Figure 8: Independent reconstruction of geometry and texture. First row: surface reconstruction from 10000 points, second row: from 100000 points. First column: texture reconstruction from 10000 points, second column: from 100000 points.

Table 3 presents the reconstruction time for the geometry $\left(t_{\text {geo }}\right)$ and the texture $\left(t_{\text {tex }}\right)$ in seconds, and also shows the memory usage (in MB) during reconstruction. As we can see, the memory peak has a linear complexity what confirms a theoretically limited memory due to local reconstruction stages.

| Model | $\# \mathrm{P}$ | M | $t_{\text {geo }}$ | $t_{\text {tex }}$ | Mem |
| :---: | :---: | :---: | :---: | :---: | :---: |
| King | 2500 | 271 | 9 | 2 | 7 |
|  | 10000 | 809 | 49 | 10 | 10 |
|  | 40000 | 2595 | 222 | 55 | 22 |
| Chameleon | 25000 | 2228 | 109 | 39 | 17 |
|  | 50000 | 3671 | 272 | 77 | 28 |
|  | 75000 | 7203 | 352 | 148 | 42 |
|  | 100000 | 9129 | 586 | 183 | 58 |

Table 3: Geometry and texture reconstruction. Memory peak.

Mesh repairing is another common use of 3D reconstruction. As partition of unity is a purely local method, meshes with very large holes may be incorrectly repaired, so we propose a simple semi-automatic process where the user has to specify additional subdomains that include the hole boundary. Figure 10 shows that it is sufficient to manually add one single region containing the boundary of the hole to repair the mesh as expected.

The following results show additional features of the surface reconstruction. The difference between interpolation and approximation is shown at Figure 11. The same 10000 points model of the Stanford Bunny was reconstructed with a different fitting tolerance $\rho$. Finally, Figure 12 shows that non-uniform, scattered data can be very


Figure 10: Mesh repairing.


Figure 9: Twisted chameleon - simultaneous transformation of reconstructed geometry and texture.


Figure 11: Interpolation and approximation of point set.
robustly reconstructed whether the density variation is smooth (Figure 12(a)) or sharp (Figure 12(b)).

Another interesting application domain is volume reconstruction for medical images. Medical images are often acquired as a set of slices distanced of some millimeters. A difficult problem is to connect the slices by reconstructing the missing information between. We obtained convincing results (not presented here) simply by considering the raw data as a sampling of a 3D signal function and then applying the reconstruction scheme on it.

(a) Non-uniform dataset with smooth density variation

(b) Non-uniform dataset with sharp density variation

Figure 12: Variable point density test.

## 5 CONCLUSION AND FUTURE wORKS

We described a new approach to reconstruct large geometric datasets by dividing the global reconstruction domain into smaller local subdomains, solving the reconstruction problems in the local subdomains using radial basis functions with global support, and combining the solutions together using the partition of unity method. Our approach has a nice behaviour with respect to the size of the dataset. Furthermore, the local reconstruction problems can be solved by various, non-communicating entities due to the independence of the local subdomains. Moreover, the stability of the reconstruction using radial basis functions makes our approach robust against highly, non-unifomly distributed and topologically complex datasets allowing its usage in various application fields.

We showed the quality of our approach on a variety of examples in different domains, and the quantitative re-
sults confirmed our expectation of the linear complexity behaviour. We think that the simplicity of the described process combined with the practical implementation issues given in this paper makes our approach highly accessible.

Our new approach intrigues us in various areas for current and future resarch. For example, the hierarchy of regions has useful information only into its leafs. We are currently investigating how the recursive domain decomposition method can be exploited in order to define a multiresolution representation that can be used not only for progressive reconstruction, but also for level-of-detail evaluation and visualization.

Moreover, we are currently exploiting the locality of our reconstruction scheme to define a point-based modeling environment in order to improve previous work by Turk et al. [Turk02] and ourselves [Reute03]. Not only the locality of the reconstruction process, but also the constant evaluation time of the reconstruction function makes our new approach attractive for interactive modelling applications. Herein, we are also using our approach to create 3D procedural textures from the attributes of the nonuniformly distributed points.

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