Title
Cluster-Based Generation of Hierarchiacal Surface Models

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Cluster-Based Generation of Hierarchical Surface Models

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Abstract

We present a highly efficient, automatic method for the
generation of hierarchical surface triangulations. Given a
set of scattered points in three-dimensional space, without
connectivity information, our method reconstructs a valid
triangulated surface model in a two-step procedure. First,
we apply clustering to the set of given points and iden-
tify point subsets in locally nearly planar regions. Second,
we construct a surface triangulation from the output of the
clustering step. The output of the clustering step is a set
of 2-manifold tiles, which locally approximate the under-
lying, unknown surface. We construct the triangulation of
the entire surface by triangulating the individual tiles and
triangulating the "gaps" between the tiles. Since we apply
point clustering in a hierarchical fashion we can generate
model hierarchies by triangulating various levels resulting
from the hierarchical clustering step.

1. Introduction

Surface reconstruction is concerned with the extraction
of shape information from point sets. Often, these point
sets describe complex objects and are generated by scan-
ning physical objects, by sampling other digital represen-
tations, or by merging data from different sources. Conse-
quently, they might embody incompleteness, noise and re-
dundancy, which makes a general approach for reconstruc-
ting surfaces a challenging problem. In many instances, the
described objects are characterized by high complexity and
high level of detail. Different levels of representation are
needed to allow rapid rendering and interactive exploration
and manipulation. Surface reconstruction problems arise in
a wide range of scientific and engineering applications. The
most important application is reverse engineering, i.e., the
reconstruction of surfaces from digitized data.

We introduce an extremely fast surface reconstruction
technique that is based on cluster analysis. Our approach
generates a multiresolution representation of reconstructed
surfaces from arbitrary point sets. Furthermore, our method
allows us to control the level of detail locally.

Our paper is structured as follows: In Section 2, we
briefly summarize related work in the fields of surface re-
construction, multiresolution modeling and clustering. In
Section 3, we provide an overview of our approach, which
is presented in greater detail in Section 4. The applica-
tion of our surface reconstruction approach to a variety of data
sets is described in Section 5. We conclude with summariz-
ing the results in Section 6 and give an outlook on future
work.
2. Related Work

2.1. Surface Reconstruction

The goal of surface reconstruction methods can be described like this:

*Surface Reconstruction: Given a set of sample points X assumed to lie on or near an unknown surface U, construct a surface model S approximating U.*

Most surface reconstruction methods require additional topological knowledge, such as connectivity between data, surface topology, or orientation.

*Parametric reconstruction* represents the reconstructed surface by mapping a two-dimensional parameter domain to a surface in three-dimensional space. This method usually requires knowledge of the topological type of the surface. Moreover, in order to converge to a valid model, this method also requires an initial embedding that is sufficiently close to the original surface and assumes a "good" parameterization that may be difficult to construct.

*Function reconstruction* methods deal with surfaces that are graphs of bivariate functions \( f(x, y) \). Various applications are concerned only with this surface type, including digital elevation maps, digital satellite imaging, and medical imaging. It is also possible to apply these non-parametric methods in a local manner to general two-manifold data, which we can locally represent by a function \( f(x, y) \).

*Constriction methods* attempt to find a surface mesh interpolating a set of data points in three-dimensional space without any given topological information. A three-manifold (tetrahedral) triangulation of the points (often Delaunay triangulation) is constructed first. The boundary of the triangulation is a mesh that defines the convex hull of the points. Since many surfaces are not convex, "undesired" elements must be eliminated. Therefore, one has to use iterative techniques creating a new triangulation by removing "undesired" tetrahedra. The result of these methods is a closed surface mesh.

Only a few methods developed fairly recently [15] [14] can create valid topological and geometrical models from only three-dimensional coordinates of sample points. Work in this field includes Edelsbrunner and Muecke [4], who generalize the notion of convex hull to that of an alpha hull (\( \alpha \) hull). An interesting, directly related concept is the concept of alpha shapes (\( \alpha \) shapes) used for the approximation of shapes in three-dimensional space (or even higher dimensions). These shapes are derived purely from finite sets of scattered, unorganized points. Applications for alpha shapes are automatic mesh generation, cluster identification for point sets in three-dimensional space, and modeling complex molecular structures.

Alpha shapes can be viewed as a generalization of the Delaunay triangulation. Ignoring certain degenerate cases, the Delaunay triangulation of a point set in three dimensions is characterized by the fact that the sphere passing through the four vertices of any tetrahedron does not contain any other point but the four vertices. The Delaunay triangulation defines a complex of edges, triangles, and tetrahedra. Given a specific alpha value, an edge in this complex belongs to the alpha shape if the radius of the smallest circle passing through the edge’s end points is less than alpha. Similarly, a triangle (tetrahedron) in the complex belongs to the alpha shape if the radius of the smallest sphere passing through the triangle’s (tetrahedron’s) vertices is less than alpha. The Delaunay triangulation itself has an associated alpha value of infinity. Gradually decreasing the alpha value towards zero leads to structures consisting of increasingly “isolated sub-complexes,” e.g., strings of edges, chains of triangles, groups of connected tetrahedra, and isolated points.

The alphashape approach has great potential for a general surface reconstruction paradigm. The alpha-shape approach will, in general, lead to geometric model with relative “thickness”, i.e., it may locally describe a three-manifold region. This usually happen when the samples are noisy or when the underlying surface is not sufficiently smooth.

2.2. Multiresolution Modeling

A multiresolution model is a model defined by a set of detail levels of an object, which can be used to efficiently access any one of those levels on demand. Surface simplification is a particular form of multiresolution modeling where the goal is to take a polygonal model as input and generate a simplified model, i.e., an approximation of the original model. A variety of methods have been developed, including:

- **Image Pyramids.** They provide a successful and fairly simple multiresolution technique for raster images [18].

- **Volume Methods.** They allow multiresolution representations for models that are acquired as volumes and will be rendered as volumes. If the simplified volumes must be rendered using polygon-based rendering, then these volume methods may become less attractive [11].

- **Decimation Techniques.** There are vertex, edge, face and cell decimation techniques, which are all iterative simplification algorithms. In each step, an element is selected for removal according to one of several rules, and the algorithm must locally re-triangulate the area. Most algorithms were developed to reduce
the density while preserving topology. Such algorithms become computationally expensive for large data sets [20] [13] [17] [7] [8].

- **Vertex Clustering.** This is a method that subdivides an object's bounding box into an octree-like grid structure, and all the vertices included in a single grid cell are clustered together and replaced by a single vertex. This is a very fast and general method, but it is often affected by a severe loss of detail and by distortions in the model [19].

- **Simplification Envelopes.** Such methods provide a global error measure for approximation quality of the simplified model, and they preserve topology. However, these methods require the original model to be an oriented manifold and can have problems with sharp edges [2].

- **Wavelet Methods.** They typically require a tensor product structure for a mesh to be represented at multiple resolution levels. Recently, different approaches have been introduced to overcome topology restrictions [3] [23].

### 2.3. Clustering

Clustering is a classical data analysis technique that has thoroughly been studied over the last few decades and has also been adopted as a standard technique in the emerging field of data mining [1]. So far, very little research has been done on applying data mining and clustering techniques to visualization problems. Vertex clustering, briefly described in 2.2, is one of the very few applications of clustering to visualization. Conversely, visualization techniques can be applied to the output of cluster analysis to present extracted patterns effectively [12] [21].

### 3. Overview of Approach

Starting with scattered point data in three-dimensional space, our method is able to generate a multi-resolution model extremely quickly and automatically reconstruct a valid geometrical and topological model. Our method is unique in two aspects: (1) It does not require any connectivity information to be supplied with the sample points, and (2) it is significantly faster than most currently used methods. We achieve this by using a clustering methodology tailored to the specific needs of surface reconstruction from scattered data at multiple levels of detail.

Knowing that the data points originate from some unknown underlying two-dimensional manifold in three dimensions, we associate points with a certain cluster based on coplanarity and distance checks. By using more or less restrictive conditions regarding cluster membership, more or less points are associated with the individual clusters, and we thus have a means for controlling the levels of detail. Each cluster is essentially characterized by a closed polygon in three-dimensional space and an associated normal vector. This information suffices to locally characterize the underlying surface. These clusters can thus be thought of as "tiles"—unfortunately unconnected—approximating the surface from which all sample points originated.

This paradigm leads to an extremely efficient algorithm for tile generation at multiple levels of detail. Furthermore, the membership criterion used to associate samples with a cluster could easily be changed to accommodate hierarchical representations of vector fields in two or three dimensions or inherently three-manifold data (volumetric data).

We apply the Delaunay triangulation to the set of tile center points leading to a set of tetrahedra. At this point, we have established a topologically complete representation which, unfortunately, contains "too much connectivity" information: After all, we know that our samples belong to some underlying surface and thus the triangulation that we need for the tile centers must define a two-manifold surface. Using a set of rules, we produce a model that, in the end, will describe a true two-manifold surface triangulation in space.

![Figure 1. Illustration of the principal phases of the algorithm.](image)

The input data set consists of a list of points in space (without connectivity). The output is a valid geometrical and topological surface model for the given samples. Our clustering-based surface reconstruction algorithm consists of two stages:

- **Stage 1: Tile generation**

This stage uses the input data and applies a clustering algorithm that splits the original data set in quasi-planar tiles according to a user-defined error tolerance.
The goal of the clustering process is to construct a hierarchy of representations of the model based on the surface description by tiles. The output of the first stage is a set of tiles representing clusters of nearly coplanar points.

- Stage 2: Reconstruction

In this stage, we fill the gaps between tiles by using a triangulation algorithm, i.e., we construct the missing connectivity information. For the triangulation step, we consider only the boundary of the tiles. Since the triangulation algorithm will lead to a model whose boundary is the convex hull of the entire point set, a post-processing phase is necessary to delete "undesired" triangles/tetrahedra from our model.

4. Surface Reconstruction

4.1. Clustering

The input for our clustering process is a set of scattered points in three-dimensional space. Our method for creating the cluster hierarchy is based on an incremental and divisive paradigm. Initially, all points are placed in one cluster, which is recursively split. A cluster $C$ is a subset of the given data set. The center of a cluster $C_{\text{center}} = (c_x, c_y, c_z)$ is defined as the geometric mean of the points $P_i = (x_i, y_i, z_i)$, $i = 1, 2, \ldots, k$, associated with a cluster of size $k$. At each stage of the clustering process, every point is associated with exactly one cluster, which is the cluster with its center being the closest in terms of Euclidean distance. The internal error of a cluster is the sum of the distances from the cluster center to the associated points, i.e.,

$$Error_{\text{internal}}(C) = \sum_{p[i] \in C} \| p[i] - C_{\text{center}} \|.$$

The global error is defined as the sum of the internal error over all clusters. In each iteration of the clustering process, the cluster $C_i$ with the highest internal error is split into two clusters. Each iteration decreases the global error and the maximum internal error. The centers of the generated clusters are determined by adding/subtracting an offset vector to/from the center of the original cluster. This offset vector lies either in the direction of highest deviation of the cluster data or in the direction of maximum variance. The direction of maximum variance is computed by performing principal component analysis (PCA) for the 3-by-3 covariance-matrix $M$, given by

$$M = \Delta^T \cdot \Delta,$$

where $\Delta$ is the cluster's normalized $k$-by-$3$ data matrix $\Delta$, defined as

$$\Delta = \begin{bmatrix} x_1 - c_x & y_1 - c_y & z_1 - c_z \\ x_2 - c_x & y_2 - c_y & z_2 - c_z \\ \vdots & \vdots & \vdots \\ x_k - c_x & y_k - c_y & z_k - c_z \end{bmatrix}.$$

We compute the eigenvalues and "eigendirections" for $M$. The direction of maximum variance is equivalent to the eigendirection with the largest eigenvalue. Using the direction of maximum variance is generally more accurate, but this is only feasible when the number of points is relatively small e.g., less than 500. Therefore, a threshold variable $\tau_{\text{PCA}}$ is used to control which splitting mechanism is used. If the number of points associated with the cluster $C_i$ to be split exceeds $\tau_{\text{PCA}}$, the cluster $C_i$ is split in the direction of highest deviation. Otherwise, the eigenvector with the highest eigenvalue is used to determine the centers of the new clusters that result from splitting $C_i$. After splitting a cluster, a local recategorization scheme is used to improve the quality of the classification.

The points to be reclassified are given by all points that are assigned to the split cluster or to the Gabriel neighbors of that cluster. Hierarchical clustering is illustrated for curve reconstruction in Figure 2.

![Figure 2. Cluster splitting.](image)

Two clusters are Gabriel neighbors when they are connected in the Gabriel graph constructed from the cluster centers. Two points $p$ and $q$ are connected in the Gabriel graph when only $p$ and $q$—but no other point—are contained in the hypersphere with the midpoint of $p$ and $q$ as its center and the distance from $p$ to $q$ as its diameter. (The Gabriel graph of a point set is a subset of its Delaunay graph and a superset of its relative neighborhood graph [16].)

After each reclassification step we update the local neighborhood. The cluster centers are moved to reflect the changes in the point-cluster association due to local reclassification. The Gabriel graph is updated locally and another
cluster is split subsequently. The clustering process terminates when the smallest eigenvalues of all clusters do not exceed the threshold $\tau_{PCA}$.

Algorithm 1: Creating the cluster hierarchy

Input: set of points
Output: set of clusters
while $\tau_{PCA}$ is exceeded {
    determine cluster $C_s$ to be split;
    split cluster $C_s$ into two new clusters and define their centers;
    create list of clusters in the local neighborhood of split cluster;
    perform local reclassification;
    update neighborhood;
}

4.2. Tile Generation

At this point, we have generated a set of clusters that partition the original data set. Since we choose a relatively small value for $\tau_{PCA}$, the points associated with a certain cluster are nearly coplanar. Thus, the point clusters have an almost flat shape. For each cluster $C_i$, the cluster center and the two eigendirections with the two largest eigenvalues define a plane $P_i$ that locally minimizes the sum of the plane-to-point distances of the associated points. We project all points $p_k$ associated with cluster $C_i$ into the plane $P_i$ and compute the convex hull $H_i$ for the projected points $p'_k$ in the associated plane. We map the points $p'_k$ on $H_i$ back to their original locations in three-dimensional space. The result is a closed, nearly planar polygon in three-dimensional space, defining the so-called tile $T_i$ of cluster $C_i$ consisting of a set of points $p''_k$ lying in $H_i$. For each cluster we generate a pseudo tile $T'_i$ by replacing the points $p''_k$, defining the tile boundary, by their original points $p'_k$.

4.3. Reconstruction

Once tile generation is completed, each cluster $C_i$ has two corresponding representations by its tile $T_i$ and its pseudo tile $T'_i$. The normal directions of $T_i$ and $T'_i$ are defined by the cluster’s eigendirection with the smallest eigenvalue.

Figure 5. Triangulation of tiles and between tiles.

A "connected model" is obtained by $T_1$ with a set of triangles $T_2$ (gap triangulation) that fills the gaps between the tiles. To determine $T_2$ we apply a Delaunay triangulation algorithm to the boundary points of the pseudo tiles.\footnote{The result of the Delaunay triangulation step is a "true volumetric" triangulation of the input points, i.e., a set of tetrahedra, from which we have to eliminate "undesired" edges, triangles, tetrahedra according to a set of heuristics.}
The algorithm we use employ a variant of the randomized incremental-flip algorithm developed by Edelsbrunner and Shah [5]. Based on the characteristics of the data, we use two different variants of the Delaunay triangulation:

- For functional data (i.e., data sets where each point of the input lies on the graph of a bivariate function \( f(x, y) \)) we project the points in the x-y plane and apply the two-dimensional Delaunay triangulation algorithm. We use the resulting triangulation to establish the connectivity of the points in three-dimensional space (Figure 6).

- For an arbitrary data set, the three-dimensional Delaunay triangulation algorithm is directly applied to the boundary points of the pseudo tiles.

![Figure 6. Triangulation of functional data.](image)

Since the boundary of the Delaunay triangulation describes the convex hull of the point set, we have to remove certain elements of the mesh (triangles or tetrahedra) that “do not belong” to the desired surface. Our approach to reduce the Delaunay triangulation to \( T_2 \) is related to the alpha shape definition by Edelsbrunner and Mücke [4]. We remove triangles from the Delaunay triangulation when

- all points lie in one tile or
- the points do not fit in a sphere of radius \( b \),

where the value of \( b \) is chosen by locally adjusting the global alpha threshold depending on the tile area.

In the original alpha shape approach, the quality of the reconstructed model is very sensitive to the alpha value, which is used globally to identify triangles to be removed. When using an alpha value that is too large, “undesired” triangles do not get removed, while an \( \alpha \)-value that is too small might remove features that do belong to the model. A valid surface model cannot be reconstructed in many cases, due to an inadequately chosen alpha value. Our clustering-based approach yields additional information (e.g., tile area, connectivity, local density of point distribution) that can be used to locally refine the alpha value.

Since we have “almost defined a topology” after the tile generation step, completing the reconstruction is a far simpler problem than applying the alpha shape approach directly to the original set of points. Because the gaps between the tiles are relatively small, one can choose a very small alpha value without removing important features of the model. However, long, skinny triangles along the tile boundaries can result due to small alpha values. We deal with this problem by inserting vertices along boundary edges when they exceed a certain length.

![Figure 8. Removing “undesired” triangles.](image)

With this enhancement, our approach is relatively insensitive to changes of the alpha value. Since a wide range of alpha values yields a consistent model, the global alpha value can be determined automatically as a linear function of the average edge length. However, when a data set contains a large number of discontinuities (the underlying surface is actually discontinuous) we propose to select the alpha value manually. Removing the undesired triangles from the Delaunay triangulation of the tile boundary points yields the set of triangles \( T_2 \) that fill the gap between tiles. By merging \( T_1 \) and \( T_2 \) we obtain a consistent model.

Algorithm 2 Reconstruct model

Input: set of pseudo tiles
Output: consistent model
reconstruct model { 
  generate \( T_1 \) by triangulating pseudo tiles;
  generate \( T_2 \):
  if boundary edge length > \( \epsilon \)
    then insert additional vertices;
    triangulate vertices using Delaunay triangulation;
    remove undesired triangles;
  merge \( T_1 \) and \( T_2 \);
}

5. Results

We have applied our method to a variety of data sets. The number of tiles produced by our algorithm is—for a given data set—a function of the threshold for the smallest eigenvalue of a cluster (see 4.1). The error \( E_k \) of a point \( p_k \) of the original data set associated with cluster \( C_i \) is defined as the distance of \( p_k \) to the plane \( H_i \) containing tile \( T_1 \), i.e.,

\[
E_k = \text{Dist} (p[k], H_i).
\]  

The global error \( E_{global} \) is defined as the sum of the errors of all points in the original data set, i.e.,

\[
E_{global} = \sum_{p[k]} \text{Dist} (p[k], H_i).
\]

To measure the root-mean-square (RMS) error, we compute the object diameter, which is the diameter of the smallest sphere that includes the object. We have tested our approach for five different data sets, which are listed in Table 1.

### Table 1. Number of points and object diameters

<table>
<thead>
<tr>
<th>Data set</th>
<th>Original # of pts.</th>
<th>Object diameter(^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rabbit</td>
<td>35929</td>
<td>0.1557</td>
</tr>
<tr>
<td>3Holes</td>
<td>4000</td>
<td>208</td>
</tr>
<tr>
<td>Car</td>
<td>20621</td>
<td>0.8</td>
</tr>
<tr>
<td>Step function(^3)</td>
<td>8000</td>
<td>1</td>
</tr>
<tr>
<td>Mt. St. Helens</td>
<td>151728</td>
<td>567,602</td>
</tr>
</tbody>
</table>

We obtain an RMS error measure \( E_{RMS} \) by dividing the global error by the product of object diameter and number of points in the original data set, i.e.,

\[
E_{RMS} = \frac{E_{global}}{\text{number of points} \times \text{object diameter}}.
\]

The listed execution times were obtained for an SGI O2 workstation with a 180 MHz R5000 processor. Table 2 summarizes the performance and quality information for our five data sets.

### Table 2. Data sets: 1-Rabbit, 2-3Holes, 3-Car, 4-step function\(^3\), 5-Mt. St. Helens

<table>
<thead>
<tr>
<th>Set</th>
<th>( \tau_{PCA} ) sim</th>
<th>( t_{cluster} )</th>
<th>( N_{tiles} )</th>
<th>( N_{points} )</th>
<th>( E_{RMS} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.006</td>
<td>7.83</td>
<td>50</td>
<td>1110</td>
<td>1.68</td>
</tr>
<tr>
<td>1</td>
<td>0.002</td>
<td>15.05</td>
<td>354</td>
<td>4764</td>
<td>0.049</td>
</tr>
<tr>
<td>1</td>
<td>0.001</td>
<td>18.94</td>
<td>727</td>
<td>8016</td>
<td>0.026</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>0.97</td>
<td>59</td>
<td>762</td>
<td>1.7</td>
</tr>
<tr>
<td>2</td>
<td>5</td>
<td>1.24</td>
<td>91</td>
<td>1002</td>
<td>0.9</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>1.77</td>
<td>202</td>
<td>1815</td>
<td>0.39</td>
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<tr>
<td>3</td>
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<td>8.37</td>
<td>517</td>
<td>5174</td>
<td>0.064</td>
</tr>
<tr>
<td>4</td>
<td>0.001</td>
<td>1.63</td>
<td>60</td>
<td>715</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>80</td>
<td>43.08</td>
<td>154</td>
<td>3100</td>
<td>0.0034</td>
</tr>
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<td>5</td>
<td>40</td>
<td>51.06</td>
<td>382</td>
<td>6307</td>
<td>0.0025</td>
</tr>
<tr>
<td>5</td>
<td>20</td>
<td>74.70</td>
<td>1391</td>
<td>17513</td>
<td>0.0014</td>
</tr>
</tbody>
</table>

Figures 9–13 show reconstructed models for our five test data sets. Figures 9 and 10 show tiles and reconstructed, triangulated surfaces. Figure 9 demonstrates how our method can be used to preserve or remove a discontinuity.

Figures 11, 12, and 13 illustrate the power of our technique to generate multiresolution models. Three examples show that our algorithm can handle extremely large data sets (Mt. St. Helens) and surfaces that are topologically complicated (3Holes, Rabbit).

6. Conclusions

The algorithm we have presented allows the generation of a hierarchy of surface models from discrete point sets without known connectivity information. While we have demonstrated the power of our approach only for surface models, we are quite confident that the same clustering paradigm, when applied to more general two- or three-manifold, or even time-varying data, would significantly speed up the process of computing level-of-detail representations.

We plan to extend our approach to the clustering of more general scattered data sets describing scalar and vector fields, defined over either two-dimensional or three-dimensional manifolds.
dimensional domains. Faster algorithms for the generation of data hierarchies for scientific visualization will become more important as our ability to generate ever larger data sets increases: Computing a data hierarchy prior to the application of a visualization algorithm should not require minutes or hours but seconds instead. We believe that our clustering methodology provides one viable answer to this problem.

Currently, we are working on a scalable parallelization of our approach. We believe that this parallelization will allow us to analyze data sets that consist of several millions of points in real time.

Furthermore, we plan to extend our algorithm to ensure that all tetrahedra are removed, such that the reconstructed model is a true two-manifold representation. Regarding the triangle elimination phase, we will develop means to guarantee that no "holes" are inserted into the model.

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References


Figure 9. Discontinuous example (step function); (a) tiles, (b) discontinuous triangulation, (c) continuous triangulation.

Figure 10. Car data set; (a) tiles, (b) reconstructed model.

Figure 11. Three resolution levels of Mt. St. Helens data set (154, 382 and 1391 tiles); (a)–(c) tiles, (d)–(f) reconstructed surface.
Figure 12. Four resolution levels of three-hole data set 59, 91, 202 and 402 tiles); (a)–(d) tiles, (e)–(h) reconstructed model.

Figure 13. Three resolution levels of rabbit data set (50, 354 and 727 tiles); (a)–c) tiles, (d)–(f) reconstructed model.