Large-scale Urban Reconstruction with Tensor Clustering and Global Boundary Refinement

Charalambos Poullis, Member, IEEE

Abstract—Accurate and efficient methods for large-scale urban reconstruction are of significant importance to the computer vision and computer graphics communities. Although rapid acquisition techniques such as airborne LiDAR have been around for many years, creating a useful and functional virtual environment from such data remains difficult and labor intensive. This is due largely to the necessity in present solutions for data dependent user defined parameters. In this paper we present a new solution for automatically converting large LiDAR data pointcloud into simplified polygonal 3D models. The data is first divided into smaller components which are processed independently and concurrently to extract various metrics about the points. Next, the extracted information is converted into tensors. A robust agglomerate clustering algorithm is proposed to segment the tensors into clusters representing geospatial objects e.g. roads, buildings, etc. Unlike previous methods, the proposed tensor clustering process has no data dependencies and does not require any user-defined parameter. The required parameters are adaptively computed assuming a Weibull distribution for similarity distances. Lastly, to extract boundaries from the clusters a new multi-stage boundary refinement process is developed by reformulating this extraction as a global optimization problem. We have extensively tested our methods on several pointcloud datasets of different resolutions which exhibit significant variability in geospatial characteristics e.g. ground surface inclination, building density, etc and the results are reported. The source code for both tensor clustering and global boundary refinement will be made publicly available with the publication.

Index Terms—pointcloud tensor field, parameter-free clustering, LiDAR reconstruction, boundary refinement

1 INTRODUCTION

ARGE-scale urban reconstruction has long been of great 2 interest and significant importance to the computer 3 vision and computer graphics communities. Specifically, following the recent advances in virtual and augmented 5 reality technologies there has been an increasing demand for robust and efficient methods for generating these virtual environments. Rapid acquisition techniques such as 8 airborne LiDAR have been around for many years and are indeed capable of capturing very large areas in a single 10 deployment however the difficulties arising with processing 11 the captured data considerably limit their uses. For one, 12 as with every scanning, noise is introduced in the mea-13 surements due to possible system error or sensor miscal-14 ibration. In addition, the zig-zag nature of the scanning 15 almost always produces spurious measurements at object 16 boundaries which manifest themselves as jagged edges in 17 the data. Another significant limiting factor is the resolution 18 or sampling density of the captured data which depends on 19 the sampling rate of the LiDAR sensor as well as the flying 20 21 altitude of the aircraft as explained in [14].

Even with the tremendous advances in remote sensing technologies given the above mentioned capture characteristics, processes in current practice still require trained personnel with extensive experience in order to produce models useful in the end application, i.e. lightweight, polygonal 3D models. The process is expensive since it requires manual or at best semi-automatic work, and is human effort

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intensive and slow. A primary cause for this is the fact that existing state-of-the-art systems for large-scale urban 2 reconstruction require a plethora of parameters [e.g. number 3 of user-defined non-adaptive parameters in ([10] \simeq 15, [22] 4 $\simeq 12$) which have to be carefully tweaked by the user since 5 these often depend on the input data characteristics. This 6 sensitivity to the input data and the large choice of multiple 7 parameters are always major concerns in the application 8 of these methods in practice. Their optimal values are not 9 known and cannot be easily computed either. This makes it 10 necessary for the user/operator to experimentally search in 11 this large parameter space for values that yield the desired 12 quality of 3D models, resulting in making this process so 13 difficult and time consuming. Another serious limitation 14 with existing solutions is scalability; primarily in terms of 15 the size of area which can be successfully processed and 16 secondly, in terms of performance i.e. how long it takes to 17 generate the results. 18

To summarize, there still exists a wide gap between the current state-of-the-art and the desired goal of automated large-scale urban reconstruction of real-world areas, for applications requiring digital 3D environments.

In this paper we address the difficult problem of large-23 scale urban reconstruction and propose a novel and au-24 tomatic solution for generating simplified, polygonal 3D 25 models. The proposed technique takes as input the raw, 26 unstructured, incomplete, and noisy pointcloud captured by 27 an airborne LiDAR scanner during multiple sweeps, and 28 first separates it into a set of smaller components. Each 29 component is a resampled XYZ map containing a different 30 part of the data as in [14] which is then processed inde-31 pendently and concurrently to extract various metrics about 32 the points in the maps. This technique adaptively calculates 33

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C. Poullis is the Director of the Immersive and Creative Technologies Lab at the Department of Computer Science and Software Engineering, Concodia University, Montreal, Quebec, Canada.
 E-mail: charalambos@poullis.org - see http://www.poullis.org

the number of XYZ maps according to the user-suggested map resolution. Next, information extracted directly from 2 this data is encoded using a tensorial representation. This 3 representation allows multiple types of information about each point to be encoded at the same time. A robust ten-5 sor clustering algorithm is proposed for segmenting the 6 tensors into clusters representing geospatial objects such as buildings, cars, trees, etc. Most significantly, our clustering 8 method has no input data dependencies and does not re-9 quire any user defined parameter making it distinct from 10 earlier solutions for this problem. It is based on adaptive 11 computation of the statistical parameters of the underlying 12 distribution for points belonging to a cluster. The result of 13 the clustering is a set of contiguous clusters of points which 14 are further processed in order to extract the boundaries. 15 This is achieved with a new multi-stage boundary extraction 16 and refinement process which reformulates the boundary 17 extraction as a global optimization problem. Unlike existing 18 work, our proposed technique requires no user interaction, 19 makes no assumptions on the input data, such as requir-20 ing Manhattan-style building orientations, and is able to 21 process the entire data without any user inputs, i.e. does 22 not require like in earlier work that the user accurately 23 marks the boundaries of every single building before re-24 constructing it. Our technique has been extensively tested 25 on five different datasets of different resolutions/densities 26 which have significant variability in the ground surface 27 elevation/inclination, building density, type, etc. 28

The rest of the paper is organized as follows: Section 2 29 provides a brief overview of the state-of-the-art in the area 30 of large-scale reconstruction. In Section 3 we provide a tech-31 nical overview of the proposed solution. The extraction of 32 the information from the pointcloud data and its encoding 33 to tensors is presented in Section 4. In Section 5 we present 34 Tensor Clustering and in Section 6 the global boundary 35 refinement. Section 7 reports on experimental results and 36 evaluation, and lastly, Section 8 has the conclusion and 37 future work. 38

2 **RELATED WORK**

Many algorithms and systems have already been proposed 40 for the problem of urban reconstruction by researchers in 41 computer vision and graphics communities. A comprehn-42 sive summary can be found in [13]. The most relevant to 43 the proposed work are categorized according to the line of 44 45 approach they followed and summarized below.

First, there are techniques which use geometric primi-46 tives. In [17] the authors proposed a system which included 47 a minimal set of three parameterized-primitives which the 48 operator could use to model any type of linear and non-49 linear surfaces. Fast forward to the current state-of-the-art, 50 there are techniques [8] and [7] which given a set of points 51 automatically produce a set of primitives describing the site. 52 However, these approaches require that the input points 53 correspond only to a single building. In fact, one of the 54 55 most difficult tasks of reconstructing large urban areas is the automatic detection of buildings and other components. 56 Which is why, these approaches combine manual detection 57 with automatic extraction. 58

A different line of approach uses symmetries and regularities. Extensive work using this approach has already been done and has shown to yield impressive results, but mostly for small scale objects [12]. In [21] this same approach is used to propose a system for urban reconstruction. However, again, solutions based on this approach require that the detection is performed manually.

Finally, a rather different approach by [20] used inverse constructive solid geometry techniques. Rather than using boolean operations on simple primitives to generate a complex structure, they start off with a point cloud representing the indoor area of a structure and decompose that into layers which are then grouped into higher-order elements.

Perhaps the closest work related to the proposed tensor clustering is the work of [15] on multi-type feature extractor and classifier. Information extracted from color satellite images is represented using tensors. Next, the tensors are decomposed and pixels are classified as junctions, curves or surfaces via graph-cut optimization. In our work, we employ 3D data and propose a different way to encode extracted information into tensors. Furthermore, there is no classification but instead all comparisons are performed in terms of the tensorial representation of each point.

In summary, previous solutions require extensive user input, in the form of manual identification of components in the urban area and/or in the form many threshold 26 values to be tweaked on a case by case basis. As a result, these solutions do not scale well to large urban areas. In contrast to the above work, in this paper, we present an automatic urban reconstruction system which requires no user interaction, and yet efficiently generates accurate urban reconstructions. Our technical contributions are:

- An elegant representation of all the multiple informa-33 tion at each point in a LiDAR depth map encoded in 34 the form of a single-second order symmetric tensor. 35 The tensor encapsulates into it the property of a 36 point belonging to the surface, curve and junction 37 categories without the need for various user specified 38 thresholds. This formulation which fuses per point 39 information into a single entity leads to considerable 40 simplification of the similarity comparison between 41 points. 42
- A robust clustering technique for depth maps captured by LiDAR scanners which retains important details without the need for user defined thresholds and yields better results than earlier solutions.
- An innovative method of computing the parameters needed for region growing to group points into clusters based on adaptive computation of per-point and per-cluster statistical parameters. The Weibull probability distribution function (pdf) is assumed for points within a cluster. The Weibull pdf parameters are dynamically updated as new points get added to the cluster. The clustering results achieved are superior to previous results.
- A new multi-stage method of boundary extraction and refinement which reformulates this as a solution to a global optimization problem.

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3 TECHNICAL OVERVIEW

Our solution involves the clustering first, followed by the 2 boundary extraction. An overview of the proposed clusз tering technique is shown in Figure 1. The LiDAR data is 4 essentially an XYZ map consisting of 3D points. A crucial 5 step in the 3D reconstruction using this data is to seg-6 ment these points into surface patches and boundary curve 7 segments. For this, firstly, a set of per-point metrics are 8 computed from the input XYZ map and these are encoded into tensors. The per-point metrics include, the normal, 10 height, local height variation and local normal variation. In 11 addition, edge saliency is computed for multiple directions 12 using multi-scale, multi-frequency filtering with Gabor jets. 13 The encoding into the tensorial representation is based on 14 the following reasoning. Surfaces should have low/constant 15 height variation and low/constant normal variation and 16 low curve response, while curves should have high height 17 variation and high normal variation and high curve re-18 sponse. This formulation is presented in Section 4. The ten-19 sors are then clustered together based on their similarity. 20 This clustering is presented in Section 5. The method for 21 adaptive computation of per-point and per-cluster statistics 22 used in region growing for cluster formation is discussed 23 in 5.3 using a Weibull distribution of the points. Lastly, 24 given a set of observations/samples, the estimation of the 25 shape α and scale β parameters of the Weibull distribution 26 is performed using Maximum-Likelihood Estimation and 27 described in detail in the Appendix. 28



Fig. 1: An overview of the proposed clustering technique.

The clusters are then further processed to extract region boundaries. The pipeline for the boundary extraction and refinement process is shown in Figure 2. Firstly, the clusters' boundaries are extracted and grouped into neighbourhoods as described in Section 6.2. Next, the dominant orientations of each cluster are computed using Principal Component Analysis (PCA) and are used to compute the globally dominant orientations for the entire scene. To avoid assumptions on the number and type of orientations we employ Gaussian Mixture Models (GMMs) and use a Minimum Description Length (MDL) criterion to specify the number of Gaussian components. The boundaries are then refined based on the globally dominant orientations and a global optimization step which ensures that the boundary positions are optimal as described in Section 6.3.

Finally, simplified, polygonal 3D models are created from the boundaries.

4 ENCODING USING THE TENSORIAL REPRESEN- 12 TATION 13

The XYZ map contains points in 3D Euclidean space. The objective of this phase is the clustering of similar neighbouring points. As has been current practice, this has involved computing of various metrics at each point in order to determine the similarity between them. Typically, these metrics can be derived directly from the XYZ map containing the points and many have already been reported: height h, normal \vec{N} , surface fitness error, height variation, normal variation, etc. Once the metrics have been computed the decision whether two points P_1 and P_2 are similar or not is expressed as either a combination $D_{component-wise}$ of the results of the per-metric comparisons:

$$D_{component-wise} = d_h(H_{P^1}, H_{P^2}) < \tau_H \bigwedge$$
$$d_N(N_{P^1}, N_{P^2}) < \tau_N \bigwedge \dots \quad (1)$$

or as $D_{combined}$ the result of a single comparison between two N-dimensional feature descriptors $f_{P_1} = \langle H_{P^1}, N_{P^1}, ... \rangle$ and $f_{P_2} = \langle H_{P^2}, N_{P^2}, ... \rangle$ containing the N metrics at each point:

$$D_{combined} = d_f(f_{P_1}, f_{P_2}) < (\tau_H, \tau_N, ...)$$
(2)

where d_h , d_N , d_f are distance functions for the height, normal, and feature descriptors respectively.

In either case, there is an explicit requirement that a set of thresholds τ_H, τ_N, \dots is specified which renders most proposed techniques dependent both on the dataset and also on the user to provide the right values for each of the thresholds. Further, this requirement implies the inherent assumption that the linear hyper-plane defined by the specified thresholds divides the N-dimensional feature space into two parts where all points lying above the hyper-plane are similar and all below are not similar. Although for low dimensional feature spaces this may often be true, when dealing with higher dimensions this does not hold and can negatively impact the accuracy of the results.

In this work, we eliminate the above requirement and in-32 stead propose a third option: combine the metrics computed 33 at each point into a single entity: a second-order symmetric 34 tensor. This choice was based on the fact that a second-order 35 symmetric tensor can encode information about multiple 36 geometric types passing through each point and therefore 37 can encode all the information extracted by the metrics. 38 More importantly, it enables us to present a solution which 39 does not require the user to guess even a single threshold 40 value for it to function. 41

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Fig. 2: An overview of the proposed boundary refinement technique.

In the following section we describe the steps required to extract and encode the information into tensors. First, 2 we begin by computing several per-point metrics from the 3 input XYZ map required for subsequent processing. This 4 is described in Section 4.1. Next, the metrics are encoded 5 into tensors and a tensor field representing the 3D scene 6 is computed. This is explained in Section 4.2. Finally, the information encoded in the tensors is used for clustering the points into surface patches and boundary lines. 9

4.1 Per-point Metrics 10

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In order to proceed with the classification, the following 11 per-point metrics are first computed using the XYZ map 12 containing the points in raster form: 13

Normal computation. By taking into account the 14 local 8-neighbourhood at each point P, a normal N_P 15 is defined as the mean of the eight neighbouring 16 normals. In our experiments the neighbourhood is 17 defined as the 3×3 neighbouring points around 18 point *P* since it provides the fastest and best results. 19 The neighbouring normals are computed as the cross 20 product of two vectors formed by connecting con-21 secutive neighbouring points $P_i, P_{(i+1) \mod 8}$, where 22 $0 \le i \le 8$, to point *P* and is given by, 23

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$$N_{P_{i,(i+1) \mod 8}} = (P_i - P) \times (P_{(i+1) \mod 8} - P).$$

25 Hence, the normal at point *P* is defined as
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$$\vec{N}_P = \frac{1}{2} \sum_{i=0}^{8} (N_{P_{i,(i+1)} \longrightarrow 8})$$

 $N_P = \frac{1}{8} \sum_{i=0}^{\infty} (N_{P_{(i,(i+1) \mod 8)}})$ Height variation. The height variation is defined as 27 the local neighbourhood height variation of point P28 and is given by, 29

$$H_{var}^{P} = \frac{||h_{P} - h_{min}||}{(h_{max} - h_{min})}$$
(3)

where h_P is the height at point P, and h_{min}, h_{max} 30 are the minimum and maximum heights in the 31 neighbourhood, respectively. In our experiments the 32 neighbourhood is defined as the 7×7 neighbouring 33 points around point P. 34

Normal variation. Similar to the height variation, the 35 normal variation is defined as the local neighbour-36 hood normal variation of point P and is given by, 37

$$N_{var}^{P} = ||nd_{max} - nd_{min}|| \tag{4}$$

where $nd_{max} = 1 - d_{max}$ and $nd_{min} = 1 - d_{min}$. 39 d_{max} and d_{min} are the maximum and minimum 40 dot products respectively, between vectors formed 41 by connecting consecutive neighbouring points to 42 point P. For example, the N_{var}^P at point P located 43 in a neighbourhood where all points have similar(or 44 equal) normal orientation will have a value closer (or 45 equal) to zero. 46

Gabor response. A bank of Gabor jets is applied on the input XYZ map at different frequencies (i.e. $\aleph_f = 5$) and orientations (i.e. $\aleph_{\Theta} = 16$). Since the XYZ maps are essentially depth maps (with X,Y coordinates) the Gabor jets respond to oriented depth discontinuities. The resulting response r_P^{θ} at each point P corresponding to the same orientation θ but different frequencies are added together to form a per-orientation response image. The combination of the multiple-scales per orientation accounts for features appearing at different scales.

The metrics H_{var}^P, N_{var}^P and r_P^{θ} are normalized and range between [0, 1].

4.2 Tensor field computation

The per-point metrics described in Section 4.1 are encoded 15 into a second-order symmetric tensor T_P for each point P. A second-order symmetric tensor T is defined as T =17 $\lambda_1 \vec{e_1} \vec{e_1}^T + \lambda_2 \vec{e_2} \vec{e_2}^T + \lambda_3 \vec{e_3} \vec{e_3}^T$ where $\lambda_1 \geq \lambda_2 \geq \lambda_3 \geq 0$ 18 are eigenvalues, and $\vec{e}_1, \vec{e}_2, \vec{e}_3$ are the eigenvectors 19 corresponding to $\lambda_1, \lambda_2, \lambda_3$ respectively. Using the Spectral 20 theorem, the tensor T can be decomposed into a linear 21 combination of three basis tensors(ball, plate and stick) as 22 in Equation 5. 23

$$T = (\lambda_1 - \lambda_2)\vec{e}_1\vec{e}_1^T + (\lambda_2 - \lambda_3)(\vec{e}_1\vec{e}_1^T + \vec{e}_2\vec{e}_2^T) + \lambda_3(\vec{e}_1\vec{e}_1^T + \vec{e}_2\vec{e}_2^T + \vec{e}_3\vec{e}_3^T)$$
(5)

In Equation 5, $(\vec{e}_1 \vec{e}_1^T)$ describes a stick(surface) with associated saliency $(\lambda_1 - \lambda_2)$ and normal orientation $\vec{e}_1, (\vec{e}_1 \vec{e}_1^T + \vec{e}_1)$ $\vec{e}_2 \vec{e}_2^T$) describes a plate(curve) with associated saliency (λ_2 – λ_3) and tangent orientation \vec{e}_3 , and $(\vec{e}_1\vec{e}_1^T + \vec{e}_2\vec{e}_2^T + \vec{e}_3\vec{e}_3^T)$ describes a ball(junction) with associated saliency λ_3 and no orientation preference.

A tensor T_P is computed for each point P as the weighted sum of \aleph_{Θ} tensors corresponding to the orientations of the Gabor jets and is defined as,

$$T_P = \frac{1}{\aleph_{\Theta}} \sum_{\theta=1}^{\aleph_{\Theta}} T_{\theta}$$
(6)

where T_{θ} is the tensor corresponding to the Gabor filter orientation θ and is calculated as described in the next section.

4.2.1 Eigenvectors

The eigenvectors $\vec{e}_1, \vec{e}_2, \vec{e}_3$ of the tensor T_{θ} form an or-37 thonormal basis system in which the normal orientation \vec{e}_1 38 is aligned with the normal N_P at point P, and the tangent 39 orientation \vec{e}_3 is aligned with the orientation $\frac{2\pi}{\theta}$ of the Gabor 40 jet. \vec{e}_2 is computed as the cross product of \vec{e}_1 and \vec{e}_3 and 41 finally, \vec{e}_3 is recalculated as the cross product of \vec{e}_1 and 42

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C. POULLIS: LARGE-SCALE URBAN RECONSTRUCTION WITH TENSOR CLUSTERING AND GLOBAL BOUNDARY REFINEMENT

 \vec{e}_2 . The recalculation of \vec{e}_3 is essential to ensure a proper orthonormal basis system, since the initial orientation θ is 2

possible to be a projection of the actual vector. Hence, the 3

eigenvectors are given by, 4

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$$\vec{e}_1 = \vec{N}_P \tag{7}$$

$$\vec{e}_3 = \langle \cos(\theta), \sin(\theta), 0 \rangle \tag{8}$$

$$\vec{e}_2 = \vec{e}_1 \times \vec{e}_3 \tag{9}$$

$$\vec{e}_3 = \vec{e}_1 \times \vec{e}_2 \tag{10}$$

4.2.2 Eigenvalues

In what follows, let M_c be defined as the magnitude of 9 the vector $\vec{c} = \langle r_P^i, H_{var}^P, N_{var}^P \rangle$. In order to determine the 10 eigenvalues we define three equations as follows. 11

The tensor as defined in Equation 6 is modelled after 12 Gabor responses (the response is high only for a tensor 13 belonging to an edge in a specific orientation), its "junction-14 ness" or junction-saliency should be 0 and hence we set 15 $\lambda_3 = 0$. This provides us our first equation. The other two 16 equations are derived based on the following observations: 17

- Points lying on a curve produced by a depth dis-18 continuity - which is the always the case for XYZ 19 maps - have high response to the Gabor filters and 20 high normal and height variation. Thus, a point on a 21 curve will have a maximal M_c . The vector \vec{c} measures 22 the "curve-ness" or curve-saliency of a point and the 23 range of its magnitude is $0 \le M_c \le \sqrt{3}$]. 24
- On the other hand, points lying on a surface have no 25 (or low) response to the Gabor filters and low normal 26 and height variation. Thus, a point on a surface will 27 have a minimal M_c . Specifically, the "surface-ness" 28 or surface-saliency of a point on a surface is defined 29 as $M_s = \frac{\sqrt{3} - M_c}{\aleph_0}$ where \aleph_{Θ} is the number of Gabor 30 filter orientations. Note that the division by the num-31 ber of Gabor filter orientations is imperative because 32 points on curves have a high response to only one 33 filter orientation; whichever is aligned to the curve. 34 Although the resulting tensor at each point is the 35 sum of \aleph_{Θ} tensors, $\aleph_{\Theta} - 1$ of those will not have high 36 curve-saliency but will instead have high surface-37 saliency. Hence, the normalization ensures that there 38 is no unjustified increase in surface-saliency. 39

From Equation 5, we note that the surface-saliency of a 40 41 point is given by $\lambda_1 - \lambda_2$ and the curve-saliency of a point is 42 given by $\lambda_2 - \lambda_3$. Hence, following the above observations we get the other two equations with the unknown eigenval-43 ues: $\lambda_1 - \lambda_2 = M_s$ and $\lambda_2 - \lambda_3 = M_c$. 44

Solving the three equations for the three unknown eigen-45 values we get 46

$$\langle \lambda_1, \lambda_2, \lambda_3 \rangle = \langle \frac{\sqrt{3 - M_c + M_c * \aleph_\Theta}}{\aleph_\Theta}, M_c, 0 \rangle$$
 (11)

where \aleph_{Θ} is the number of Gabor filter orientations. Figure 3 47

shows the relation between the eigenvalue differences λ_1 – 48 λ_2 and $\lambda_2 - \lambda_3$ with respect to the magnitude M_c , used to



Fig. 3: Relation between the eigenvalue differences $\lambda_1 - \lambda_2$ and $\lambda_2 - \lambda_3$ with respect to the magnitude M_c which is used to calculate the eigenvalues.

compute the eigenvalues. The λ values have been scaled by a factor of $(1/\sqrt{3})$ to get them in the 0-1 range. As it can be seen, a point lying on a curve will have a high M_c . Therefore the eigenvalue corresponding to the curve-saliency will be higher i.e. $\lambda_2 - \lambda_3$.

Figure 4 shows the results of the above tensor encoding for a synthetic image. The geometric interpretation of the tensor is an ellipsoid in 3D space. There are three basis cases which define the possible variations of the ellipsoid:

- 1) Tensors corresponding to points on a curve appear as ellipsoids with a plate-like shape where the normal to the plate is the tangent to the curve. This tangent corresponds to the eigenvector corresponding to the smallest eigenvalue i.e. $\vec{e_3}$ of the tensor.
- Tensors corresponding to points lying on a surface 2) appear as ellipsoids with a stick-like shape where the orientation of the stick is the normal to the surface. This normal corresponds to the eigenvector with the largest eigenvalue i.e. $\vec{e_1}$ of the tensor.
- 3) Tensors corresponding to points where there is no curve nor surface appear as perfect spheres/balls since there is no preference towards a particular orientation at those locations.

Examples are shown in Figure 4 for the first two cases; the red straight line drawn shows a sequence of neighbouring tensors corresponding to points on the same straight line and the red planar surface drawn shows neighbouring tensors corresponding to points lying on the same surface.

An alternative validation procedure would be to apply the Spectral theorem to decompose the tensors into the three basis tensors i.e. stick - Figure 5a, plate - Figure 5b, and ball - Figure 5c. The eigenvalue differences can then be used to classify each point into a surface, curve or junction as described in [11] and is shown in Figure 5.

5 CLUSTERING

Clustering is performed on the tensor field using a region 36 growing approach. Tensors in the same region are grouped 37 together based on their similarity. To achieve this, we first 38 define a similarity measure between two tensors, then in-39 troduce a comparison condition for deciding when a new 40 tensor should be added to an existing cluster, and finally 41

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Fig. 4: Visual interpretation of the encoded tensors for the synthetic depth map shown. Tensors corresponding to points on a curve appear as ellipsoids with a plate-like shape where the normal to the plate is the tangent of the curve; one such case is shown with a straight red line. Tensors corresponding to points on a surface appear as ellipsoids having a stick-like shape where the direction of the stick is the normal to the surface; one such case is shown with a red plane.

use this condition to grow the cluster by applying it to all
neighbour tensors of that cluster. A unique feature of this
method is that even though we have all the different per
point metrics as input, unlike all other methods, we do not
need to ask the user to define a large number of threshold
values. All the terms in the comparison are adaptively
computed for each new tensor.

8 5.1 Similarity measure

9 We define the similarity measure between two tensors T_i 10 and T_j as,

$$d_{(T_i,T_j)} = 1 - \frac{trace(T_i,T_j)}{||T_i|| \cdot ||T_j||}$$
(12)

where ||.|| is the Frobenius norm. The range of this similarity 11 measure is $0 \leq d_{(T_i,T_i)} \leq 1$ and $d_{(T_i,T_i)} = 0$ iff the 12 two tensors being compared are identical. Although several 13 other similarity measures have been reported (e.g. [5], [2], 14 15 [9]), this measure, which was first introduced by [6], is preferred because it does not require the eigen-value/-vector 16 decomposition of the two tensors, and hence is faster to 17 calculate. Figure 6b shows the similarity variation measured 18 as the maximum minus the minimum similarity between the 19 tensors in the 8-neighbourhood of each point in the depth 20 map shown in Figure 6a. 21

22 5.2 Deciding whether two Tensors are Similar

In the region growing method for clustering, typically at 23 this point various thresholds need to be defined to deter-24 mine whether two tensors are similar or not. As mentioned 25 before, in contrast to existing techniques, we do not require 26 user-defined thresholds at all. Instead we propose a new 27 technique for adaptively computing a comparison condition 28 using the per-candidate and per-cluster statistics as its basis. 29 This is described next. 30

5.2.1 Per-cluster statistics

Assume that a cluster C contains N tensors T_i^C , $1 \le i \le N$ at iteration/time t. First, we compute the cluster's average tensor $\overline{T^t} = \frac{1}{N} \sum_{i=1}^{N} T_i^C$ at iteration/time t. Second, we calculate the probability distribution function (pdf) ϕ_W^C of the similarity distances between the tensors T_i^C contained in the cluster and the average tensor $\overline{T^{t_i}}$ at the time t_i that the tensor T_i^C was added to the cluster C. Thus the pdf gets continuously updated as new tensors get added to the cluster and in turn influences which new tensors can be added to the cluster in an adaptive manner.

The pdf ϕ_W^C for a cluster *C* is modeled as a Weibull distribution. It has already been shown [1] that Extreme Value distributions and in particular the Weibull distribution significantly outperforms other distributions such as Gaussian, Student-t, etc, in cases where the observations represent *similarities* and therefore are closer (or equal) to zero, leading to zero-mean and/or zero-variance. This is clearly brought out in Figure 7, which shows three distributions being applied on the same set of tensors corresponding to the surface points of the area shown in Figure 8a; two extreme value distributions.

Thus, the pdf ϕ_W^C for a cluster *C* is given by,

$$\phi_W^C = \left(\frac{\alpha}{\beta}\right) \left(\frac{x}{\beta}\right)^{a-1} e^{-\left(\frac{x}{\beta}\right)^{\alpha}} \tag{13}$$

where $\alpha > 0$ is the shape parameter, $\beta > 0$ is the scale parameter, and x is the observation i.e. the similarity measure defined in Equation 12. In practice, because of the many logarithm calculations involved when fitting the Weibull distribution, and the fact that some similarity measures are close to zero [and the logarithm of zero is undefined], x represents the shifted similarity measure given by $1 + d_{(T_i,T_j)}$ such that the minimum value of the range of the observations is 1, rather than 0.

The mean of ϕ_W^C is defined as $\mu_{\phi} = \beta \Gamma(1 + \frac{1}{a})$ and the variance as $\sigma_{\phi}^2 = \beta^2 [\Gamma(1 + \frac{2}{\alpha}) - \Gamma^2(1 + \frac{1}{\alpha})]$, where $\Gamma(.)$ is the gamma function given by $\Gamma(n) = \int_0^\infty e^{-x} x^{n-1} dx$.

Given a set of observations/samples, the estimation of the shape α and scale β parameters of the Weibull distribution is done using Maximum-Likelihood Estimation as described in detail in the Appendix.

5.3 Cluster Formation

Region growing proceeds as follows. A candidate tensor T_{new} considered for inclusion in cluster C is first compared with the average tensor T_C of the cluster and added to the cluster iff the probability of the similarity measure $\phi_C^W(d_{(T_{new},T_C)})$ is higher than the probability of the mean μ_C perturbed by the standard deviation σ . Thus if the following comparison condition is true,

$$\phi_C^W(\mu_C + \sigma_C) \le \phi_C^W(d_{(T_{new}, T_C)}) \le \phi_C^W(\mu_C) \quad (14)$$

the candidate tensor T_{new} is added to the cluster. The choice for this is based on the *empirical rule* for probability models which states that about 67% of the values are contained within one standard deviation of the mean. This has also been verified in practice and has proven to be stable over different datasets.

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Fig. 5: Application of the Spectral theorem: each tensor is decomposed into three basis tensors. Figures (a), (b) and (c) depict the points classified as surfaces, curves or junctions respectively, according to their eigenvalue differences as explained in Equation 5. Figure (d) shows the orientation corresponding to each type i.e. for surfaces it represents the normal to the surface, for curves it represents the tangent to the curve, and for junctions it appears black.



Fig. 6: (b) Similarity variation measured as the maximum minus the minimum similarity between the tensors in the 8-neighbourhood of each point in the depth map shown in (a). Note that the color curve for (b) has been adjusted for easier readability.

A cluster C_0 is initialized with the first tensor T_0 in the tensor field. All tensors in the 8-neighbourhood of T_0 are 2 considered for inclusion. A candidate tensor for which the з comparison in Equation 14 is true is added to the cluster C_0 and the statistics are updated. Moreover, the neighbours of 5 the newly added tensor are also considered for inclusion in C_0 . This process is repeated until all neighbouring tensors 7 [neighbours of all tensors contained in the cluster] are 8 considered for inclusion into the cluster C_0 . A new cluster C_1 is then initialized with a neighbouring tensor for which 10 the comparison was false, or had not been yet considered. 11 This process is repeated until all tensors are considered and 12 belong to some cluster ¹. 13

Typically region growing algorithms are very sensitive 14 to the initialization however in this case changing the initial 15 starting point will primarily affect the sequence in which 16 the clusters are being formed and not so much the final 17 outcome. It is exactly for this reason (i.e. to address the 18 problem of sensitivity/robustness) that a distribution of the 19 tensor similarities is calculated for each cluster rather than 20 a distribution of the per-cluster metrics e.g. height, normal, 21 etc. The samples used to calculate the distribution have very 22 small values and are close to zero e.g. < 0.03, hence the 23 decision for using an Extreme Value Distribution to model 24 these; and in particular the Weibull distribution which can 25 be calculated with a much smaller sample size than other 26

distributions. These (Weibull distribution and small sample size) are what make the region growing algorithm more robust to changes in the initialization, since a good fit for the distribution will be available even after the first few points.

Equation 14 formulates the comparison in such a way that after each addition to a cluster the comparison also is updated according to the latest cluster's statistics. Thus every comparison test is potentially different from the previous. In practice, during the initial stages of the clustering there are moderate changes in the mean μ_C and variance σ_C which leads to moderately different comparison tests; once a cluster contains enough tensors these deviations are diminished since fitting the distribution converges to the same/similar set of parameters. Figure 9 shows an example of a cluster's statistics during the initial iterations i.e. 5^{th} containing 5 samples, and the final i.e. 1925^{th} containing 1925 samples ².

As previously mentioned, fitting a Weibull distribution involves many logarithm calculations which as a result significantly increase the computation time. To address this, we re-fit the Weibull distribution at every new insertion until it reaches a stable level. Our experiments have shown that the changes occurring to the mean and variance of a cluster's distribution dramatically reduce after the first 50 samples. Thus, after the first 50 iterations, we opt out of recalculating the mean and variance unless (i) the ratio between the last two mean estimations and the ratios between the last two variance estimations is less than 95% or, (ii) an additional 50 samples have been added to the cluster *without* recalculating the mean and variance. This results in significant improvement in computation speed.

5.4 Cluster Refinement

Clusters resulting from actual geospatial features with less or no significance such as bumps on the ground, shingles, grass, etc., or from noise during the acquisition process are removed by merging as described next.

First, an adjacency graph is built based on the result of the tensor clustering. In addition to keeping track of neighbouring clusters, the graph also stores point-level information i.e. how many and which points are neighbouring.

The cluster refinement is an iterative process which merges every cluster C^- containing a small number of

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^{1.} An animation of the clustering algorithm is shown here

^{2.} An animation of the Weibull distribution parameter estimation with Maximum Likelihood is shown here



Fig. 7: Comparison between two extreme value distributions (Gumbel, Weibull) and the Gaussian distribution for the set of tensors corresponding to the surface points of the marked area shown in Figure 8a. As it can also be visually confirmed, the Weibull distribution can provide a more accurate representation.



Fig. 8: (a) The normalized XYZ depth map of a building. The three distributions were tested on the tensors corresponding to the surface points in the marked area. (b) The mesh corresponding to the depth map in (a). (c) Color-coded clusters. (d) The complete sparse boundary map corresponding to the clustered regions in (c). (e) The boundary positions after snapping and adjustment are shown in red. The optimized boundary positions are shown in green. A closeup is shown in (f).



Fig. 9: The shape and scale parameters of the distribution [and therefore the mean and variance] converge to the same values as the number of iterations [and therefore samples] increases. The results shown correspond to the surface shown in Figure 8a,8b.

points i.e. ≤ 5 to a cluster C' if and only if the following conditions hold true,

the cluster C' is a neighbour of C⁻ in the adjacency
 graph

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- the cluster C' has the highest number of neighbouring points from all neighbours of C⁻
- the cluster C' contains at least twice the points of C^-

⁸ If the above conditions are true, then the cluster C^- is ⁹ absorbed by C' and all the information in the clusters and ¹⁰ the adjacency map are updated to reflect the change.

As mentioned above, this is an iterative process which 11 is repeated until no merge has occurred. During our experi-12 ments the maximum number of iterations has been six with 13 an average number of four iterations, and the maximum 14 reduction in the number of clusters of around 90% with 15 an average reduction in the number of clusters of around 16 75%. Further to this, an additional average reduction of 17 35% occurs after the removal of vertical surfaces i.e. surfaces 18 for which the angle between their orientation and the nadir 19 direction is greater than 45° . 20

We tested our clustering method on several pointcloud datasets of different resolutions which exhibit significant variability in geospatial characteristics e.g. ground surface inclination, building density, etc.

25 6 BOUNDARY EXTRACTION AND REFINEMENT

The result of the clustering is the formation of a set of 26 contiguous regions. Perhaps one of the most difficult tasks 27 in urban reconstruction from LiDAR data is extracting 28 boundaries corresponding to depth discontinuities. Due to 29 the zig-zag scanning fashion of the LiDAR scanner, depth 30 discontinuities appear jagged in the captured data. Several 31 approaches have already been proposed [22], [14], [21] for 32 refining the boundaries however they all treat each cluster 33 of points individually. In this work we propose a different 34 technique for refining boundaries which unlike the existing 35 work reformulates this boundary refinement as a global 36 optimization problem based on the following two observa-37 tions; (i) all cluster boundaries [not on the captured image 38 boundary] between adjacent clusters are complementary to 39 each other and (ii) object boundaries must align to dom-40 inant directions. The boundary points of each cluster are 41 extracted, adjusted, refined and finally extruded to produce 42 3D lightweight polygonal models. The following subsec-43 tions further describe the boundary extraction process, the 44 subsequent refinement and optimization steps, and the final 45 extrusion to 3D models. 46

47 6.1 Boundary Extraction

The boundaries of each cluster $C_i, 0 \le i \le M$ where M is the number of clusters produced by the tensor clustering, are extracted as follows.

⁵¹ A 2D map is created for C_i marking all the points ⁵² contained in the cluster. The cluster's boundary points B^{C_i} ⁵³ are then retrieved from the map using the algorithm of ⁵⁴ Suzuki et al in [19]. The result is the dense set of the 2D ⁵⁵ image locations corresponding to the 3D exterior boundary ⁵⁶ points surrounding all points within the cluster. These are further reduced to a minimal set of the 2D image locations of the 3D boundary points after applying the iterative-endpoint fit algorithm of Douglas-Peucker in [4] as shown in Figure 8d. During the simplification only *colinear* points are removed from the set i.e. the threshold is very small and set to $\tau = 0.01$. This process is repeated until boundaries for all clusters have been extracted and simplified.

An example of the boundary extraction process is shown in Figure 10a. The points contained in a cluster representing part of a roof are shown with red in the normalized depth map. The dense boundary points are shown with a bright red color, which upon further reduction result in the simplified boundary points shown as blue pixels in the close-up in Figure 10b.



Fig. 10: An example of the boundary extraction process being applied to a cluster containing points representing part of a roof. The points are shown in red and the dense boundary points extracted are shown in bright red in (a). (b) shows a close up of the simplified boundary points in blue.

6.2 Snapping and Adjustment

Two adjacent clusters have complementary boundary components i.e. some of the boundary points in one cluster will correspond and complement some of the neighbouring cluster's boundary points. Refining the boundary points separately almost always leads to undesirable effects such as misalignment between the initially complementary boundaries; this problem often appears as holes in the resulting 3D models. In order to ensure that there is no misalignment between the final model's neighbouring boundaries, we group neighbouring boundary points together prior to the refinement. This helps in avoiding holes and making the generated geometry water-tight.

Boundary points B_i, B_j from any cluster which are within a user-defined radius i.e. $\tau_r = 2px$ from each other are grouped into a neighbourhood \aleph such that $\forall B_i, B_j \in$ $\aleph \Rightarrow ||B_i - B_j|| \leq \tau_r$. For all subsequent processing each boundary point is represented in terms of its *handle*:

- the neighbourhood's location in the image (u_{Rx}, v_{Ry}) computed as the average of the image locations[2D] of all points contained within the neighbourhood.
- the neighbourhood's X, Y components of its 3D position computed as the average X, Y components of all points within the neighbourhood.

The 3D position for each point B_i contained in a neighbourhood group is then snapped to the group's handle's X,Y 40 components but retaining the Z component of the original 41 3D point $(\bar{X}_{\aleph}, \bar{Y}_{\aleph_y}, Z_{B_i})$. This allows the representation of 42

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boundary points with the same X,Y but different vertical values in the same neighbourhood, e.g., two points on a 2 vertical wall may have the same X, Y components but 3 different Z components between the ground and roof points. After the snapping process, all boundary points are 5

expressed in terms of a set of handles. Figure 11 shows the 6 neighbourhood handles resulting from the sparse boundary points in Figure 8d. Each large color-coded sphere repre-8 sents the handle of a neighbourhood and the small spheres 9 with the same color [visible in the closeup] represent the ac-10

tual boundary points contained within the neighbourhood. 11



Fig. 11: Each large color-coded sphere represents the handle of a neighbourhood and the small spheres with the same color [visible in the closeup] represent the actual boundary points contained within it.

Refinement and Global Optimization 6.3 12

The position of each neighbourhood's handle is further 13 adjusted through a process which iterates between a re-14 finement step and a global optimization step. The following 15 sections describe these steps in detail. 16

6.3.1 Detection of Dominant Orientations and Refinement 17

Refinement involves (a) the detection of per-scene dominant 18 orientations and (b) the handle position refinement for each 19 neighborhood group based on the scene's detected domi-20 nant orientations 21

The dominant orientations present in the entire scene 22 are extracted. First, for each group, a set of vectors repre-23 senting boundary orientations is computed by subtracting 24 each pair of consecutive boundary points B_{prev}^{XY} and B_{next}^{XY} . 25 The vectors are kept unnormalized in order to account for 26 different weights i.e. vectors resulting from the subtraction 27 of pairs of boundary points whose distance is high will have 28 a higher weight. The vectors are 2D and only contain the X,Y 29 components of each 3D boundary point. Boundary points 30 close to the image boundaries are excluded so as not to 31 introduce spurious orientations corresponding to the image 32 boundaries. 33

Next, we perform Principal Component Analysis (PCA) 34 on the entire set of vectors derived from all the neighbor-35 hood boundary groups in the scene and reduce it into a 36 smaller representative set which accounts for most of the 37 variance in the original variables. In our experiments, the 38 maximum number of principal components (PCs) is set 39 40 to 5 although it has been observed that metropolitan areas following the Manhattan-style design contain primarily 41 clusters corresponding to axes-aligned building structures 42 and usually result in only 2-3 PCs. 43

Finally, a Gaussian Mixture Model $G^{2D}(\vec{v})$ is applied on the set of PCs. In order to avoid having to fix the number N_q of Gaussian distributions $g_i(\vec{v}), 1 \leq i \leq N_g$ contained in the $G^{2D}(\vec{v})$ we opt for using minimum-description length (MDL) [18] to adaptively determine this number. The result is the minimal set of the dominant orientations of all the neighbourhood boundary groups in the scene represented by the normalized means of each $q_i(\vec{v})$ contained in the $G^{2D}(\vec{v}).$

6.3.2 Refinement based on Dominant Orientations

The dominant orientations are used to refine the position of each neighbourhood's handle. First, for each vector $\vec{v}^B =$ $B_{next}^{XY} - B_{current}^{XY}$ resulting from subtracting two consecutive boundary points $B_{current}^{XY}$ and B_{next}^{XY} , we determine the g_{max} contained in the GMM $G^{2D}(\vec{v})$ in which \vec{v}^B is maximal,

$$g^{max}(\vec{v}^B) = argmax(g_i(\vec{v}^B))) \tag{15}$$

where $1 \leq i \leq N_q$ and $g_i \in G^{2D}$.

Once the g_{max} is determined, the vector \vec{v}^B is projected onto the vector representing the means $\mu_{g_{max}}$. This results in a refined position for one of the boundary points, i.e. B_{next}^{XY} . This process is repeated until the X and Y components of all the boundary points have been refined.

6.4 **Global Optimization Formulation**

The refinement based on the dominant orientations con-23 siderably improves i.e. linearizes, the appearance of the 24 boundaries. A final refinement using global optimization is 25 performed to ensure that the displacement introduced by 26 the previous steps also matches the observed data. For this 27 global optimization we define an error function E_f which 28 when maximized results in a mapping f for which the 29 boundary points are in their optimal positions. A boundary 30 location B is optimal when all the points along the bound-31 ary lines connecting the previous point B_{prev} , B, and the 32 next point B_{next} have maximal Gabor response. 33

The error function E_f is defined as a Gibbs potential,

$$E(f) = D(\bar{B}_{current}^{XY}) \times e^{-\frac{d}{\rho}}$$
(16)

where $d = ||B_{current}^{XY} - \bar{B}_{current}^{XY}||$ is the distance be-tween the optimized position $B_{current}^{XY}$ and the initial po-36 sition $B_{current}^{XY}$, ρ is the optimization search radius, and 37 $D(\bar{B}_{current}^{XY})$ is given by,

$$D(\bar{B}_{current}^{XY}) = \sum_{P=B_{prev}^{XY}}^{R(B_{prev}^{XY}, \bar{B}_{current}^{XY})} r_P + \sum_{P=\bar{B}_{current}^{XY}}^{R(\bar{B}_{current}^{XY}, B_{next}^{XY})} r_P$$
(17)

where R(.,.) is a function that rasterizes the line between 39 the two input points using Bresenham's algorithm and, r_P 40 is the Gabor response at location P. Intuitively, equation 17 41 gives a measure of how appropriate the optimized bound-42 ary position is by evaluating the Gabor responses along the 43 lines beginning and ending at that position. 44

Figure 8e shows the boundary points after snapping and 45 adjustment as red. The optimized points are shown as green 46 overlaid on the Gabor response image. 47

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6.5 Reconstruction

After the adjustment, refinement, and optimization, the 2 boundaries are extruded to the ground [ground elevation is 3 set to the minimum height in the scene], to form 3D models. The refined boundaries of each cluster can have arbitrary 5 [but not self-intersecting] shapes. In order to handle these 6 complex shapes we employ a variant of the line-sweep trian-7 gulation algorithm which can handle complex and concave 8 geometry robustly. Normal information about the boundary points is taken into account during the triangulation in order 10 to ensure smooth transitions between the planar surfaces of 11 the reconstructed model. Furthermore, texture coordinates 12 are computed for easier assignment of texture maps. 13

EXPERIMENTAL RESULTS & EVALUATION 7 14

We have extensively tested our methods on real data of 15 city-scale size. Several point clouds captured with airborne-16 LiDAR scanners have been used, those of which exhibiting 17 different characteristics are shown in this paper. Namely, 18 Baltimore, MD which covers an area of $16km^2$, Denver, 19 CO which covers an area of $14km^2$ and Oakland, ON 20 which covers an area of $17km^2$. Within each dataset there 21 is significant variability in terms of the geospatial object 22 density, sampling density, and area type i.e. rural, suburban, 23 24 urban.

Figure 12 and 13 clearly show that our techniques scale 25 up to large datasets.³. The city of Baltimore, MD consists of 26 36 components (each represented as an XYZ map of size 27 1024x1024) generated using the structuring algorithm in 28 [14]. Each component is processed independently and in-29 parallel using a Microsoft Azure Virtual Machine (Standard 30 DS15 v2) with 20 cores and 140GB memory. The processing 31 time for the clustering is determined by the time required 32 to process the slowest component, and for Baltimore that 33 was 22.75 hours . The city of Denver, CO consists of 20 com-34 ponents (each represented as an XYZ map of size 991x991) 35 and the processing time was 16.2 hours. Table 1 summarizes 36 these information and provides a comparison with the state-37 of-the-art large-scale modeling techniques in [16] and [14]. 38 As can be seen, our method does require more processing 39 time. This is primarily due to the computations involved in 40 recalculating the Weibull distribution (as explained in detail 41 in the Appendix). However, the clustering results obtained 42 using our approach outperform the other techniques in 43 terms of geometric accuracy, while not requirng any pa-44 rameter tweaking whatsoever. The geometric accuracy is the 45 equivalent to the surface fitting error and is defined as the 46 RMS distance of the fitted surface points from their original 47 48 position.

Figure 15 shows all intermediate results of our tech-49 nique for four test sites exhibiting variability in terms 50 of building/trees/cars/roads size and density. The depth 51 map is shown in the first column and the sum of the 52 gabor responses for all ie. 16 orientations is shown in the 53 second column. The third column shows the height and 54 normal variation map. The last two columns show the 55 resulting clusters after the application of tensor clustering 56 (fifth column) and the refined clusters generated by the 57

global boundary refinement (sixth column) after merging insignificant clusters i.e. < 4 points and, removing clusters corresponding to vertical surfaces $< 45^{\circ}$.

An additional comparison was performed with the approach presented in [21] and later extended by [22] which employs dual contouring on 2.5D pointcloud data. It should be noted that their approach requires a total of 12 userdefined thresholds. After extensive experimentation we found the optimal values for these thresholds and used them to generate the result in Figure 14. The original 10 captured data is assumed to be the ground truth. Each 11 reconstruction is compared against the ground truth by 12 computing the Hausdorff distance [3] for a fixed number 13 of sample points for both (i.e. 300K). A visualization is 14 shown in Figure 14a where the distances are shown with 15 RGB colors. The RMS error, the Hausdorff distances' range, 16 and the mean error for our reconstruction are 0.174456, 17 [0, 0.559475], 0.103138, respectively. For dual contouring 18 these values are 0.212459, [0, 0.562199], and 0.141765. As it 19 is evident, our method results in improved reconstructions. 20 Much of the errors occurring [note that the maximum error 21 is 0.559475] are due to noise in original data which when 22 compared with the planar surfaces may result in a high 23 error. 24

Figure 14 shows a visual comparison between our reconstruction (left), the original captured data (middle), and the dual contouring reconstruction (right). While, from such visual qualitative assessment, it may seem as though the results are quite similar, more careful visual inspection shows that for certain thin structures dual contouring seems to fail by creating vertical "ridges" in the reconstruction. Scaling to larger areas seems to be another problem with the dual contouring implementation. Despite the fact that we used the provided source code (without modifications other than experimenting to determine the optimal values of the 12 thresholds required), when running it on the same machine as the one we used for our method, the dual contouring program was unable to generate a 3D model for larger areas than this.

Our experiments have shown that the approach in 40 [22] performs best for small building-scale reconstructions 41 i.e. the input pointcloud represents a single building (or 42 sequence of attached buildings) and is cropped at the 43 buildings boundaries in a preprocessing step before being 44 processed. In cases of multiple disjoint buildings (such as 45 the own in Figure 14) dual contouring produces building 46 boundaries which may be visually pleasing but significantly 47 deviate from the original data; hence the high error at the 48 roofs/vertical walls. This can be attributed primarily to the 49 quadtree simplification. The proposed approach makes no 50 assumptions about the data and therefore all clusters are 51 processed the same regardless of the type of object they 52 represent e.g. building, tree, car, boats, etc. Furthermore, 53 [22] use triangulation based on the curvature to model 54 the clusters which, at the cost of increasing the geometry, 55 produce models which are closer to the original data even 56 in the presence of noise i.e. the noise is carried over the 57 the reconstructed models. In the proposed approach, each 58 cluster is modeled as a linear surface which drastically 59 reduces the geometry but may lead to higher deviations 60 from the original data; hence the higher error at the non-61

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^{3.} An animation of the reconstructed models is shown here



Fig. 12: The final clusters after tensor clustering. Baltimore, MD has a size of $16km^2$, the most complex part of which is shown here.

Dataset	Area (km^2)	# Comp.	Comp. Resolution	Processing time (hr)			Geometric Accuracy m^2		
				Our approach	[16]	[14]	Our approach	[16]	[14]
Baltimore	16	36	1024×1024	22.75	9	1.3	6.96×10^{-03}	9.72×10^{-01}	0.48×10^{-01}
Denver	14	19	991×991	16.2	3.7	1	4.826×10^{-03}	4.915×10^{-01}	0.311×10^{-01}

TABLE 1: Comparison table between our approach and state-of-the-art in [16] and [14]. Our approach requires more processing time for the same number of components primarily due to the recalculation of the Weibull distribution. However it is completely automatic, does not require any parameter value to be specified by the user and it produces superior results in terms of geometric accuracy.

¹ flat ground area in Figure 14.

8 CONCLUSION

We have presented a new method for automatic 3D reconstruction of large scale urban areas from raw lidar (point 4 cloud) data. Our most significant contribution, the elegant 5 second-order symmetric tensor representation for encoding all metric information about points, does not require any 7 user defined parameter for the entire region construction process, making our method automatic, effective and dis-9 tinctly different from previous methods. All earlier methods 10 are at best semi-automatic typically requiring the user to 11 carefully specify a plethora of parameter values which are 12 input data dependent and needed to produce usable recon-13 structions, making the process human effort intensive, dif-14 ficult and inefficient. In contrast, we show that our solution 15 works for any of the highly varied data sets we used to 16 test and evaluate our reconstruction process. Our process 17 includes a number of innovative techniques, a robust ag-18 glomerative tensor clustering technique for region finding, 19 adaptive computation of per-point and per-cluster statistical 20 parameters for the Weibull probability distribution function 21 (pdf), whose parameters are dynamically updated as new 22 points get added to the cluster, and a more accurate multi-23 stage region boundary extraction technique which reformu-24 lates it as a global optimization problem. We have tested 25 and evaluated our method extensively on large scale urban 26 areas from the United States with varying characteristics. To 27 the best of our knowledge no existing method can generate 28

this quality of reconstructions automatically for such large scale data. We plan to extend this work by using colour image data along with 3D lidar data, possibly using deep learning techniques to do the classification prior to 3D reconstruction. We will also investigate new methods for automating texture mapping to produce realistic 3D digital worlds.

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Fig. 13: (a), (b), (c), (e): Renderings of the reconstructed models for Baltimore, MD. (d), (f): Rendering of the reconstructed models in (c) and (e) respectively textured with the clustering results. Baltimore, MD has a size of $16km^2$.

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Fig. 14: (a) Visualization of the Hausdorff Distance between the original captured data and the result of our approach (left), and between the original captured data and the result of the approach in [22]. (b) 3D visualization of the reconstructions. For dual contouring, optimal values were determined via experimentation for the 12 required userdefined thresholds. (left) The result of our approach. (middle) Triangulation of the original captured data. (right) The result of dual contouring.

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Charalambos Poullis was born in Nicosia, Cyprus, in 1978. He received the B.Sc. degree in Computing and Information Systems with First Class Honors from the University of Manchester, UK, in 2001, and the M.Sc. in Computing Science with specialisation in Multimedia and Creative Technologies, and Ph.D. in Computer Science from the University of Southern California (USC), Los Angeles, USA, in 2003 and 2008, respectively. In 2010, after spending a year at the Department of Computer Science. University of

Cyprus as a Visiting Lecturer, he joined the Department of Multimedia and Graphic Arts, Cyprus University of Technology as a Lecturer, and in 2014 became an Assistant Professor. Since August 2015, he has been an Associate Professor with the Department of Computer Science and Software Engineering at the Faculty of Engineering and Computer Science at Concordia University where he also serves as the Director of the Immersive and Creative Technologies (ICT) lab.

His current research interests lie at the intersection of computer vision and computer graphics. More specifically, he is involved in fundamental and applied research covering the following areas: acquisition technologies & 3D reconstruction, photo-realistic rendering, feature extraction & classification, virtual & augmented reality.

More specifically, he is involved in fundamental and applied research covering the following areas: acquisition technologies & 3D reconstruction, photo-realistic rendering, feature extraction & classification, virtual & augmented reality. Charalambos is a member of the Association for Computing Machinery(ACM); Institute of Electrical and Electronics Engineers (IEEE) Computer Society; Marie Curie Alumni Association (MCAA); ACM Cyprus Chapter, where he also served in the management committee between 2010-2015; and British Machine Vision Association (BMVA). Charalambos has been serving as a regular reviewer in numerous premier conferences and journals since 2003.

37

38

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