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Geometry Preserving Sampling Method Based on Spectral Decomposition for Large-Scale Environments

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In the context of 3D mapping, larger and larger point clouds are acquired with lidar sensors. Although pleasing to the eye, dense maps are not necessarily tailored for practical applications. For instance, in a surface inspection scenario, keeping geometric information such as the edges of objects is essential to detect cracks, whereas very dense areas of very little information such as the ground could hinder the main goal of the application. Several strategies exist to address this problem by reducing the number of points. However, they tend to underperform with non-uniform density, large sensor noise, spurious measurements, and large-scale point clouds, which is the case in mobile robotics. This paper presents a novel sampling algorithm based on spectral decomposition analysis to derive local density measures for each geometric primitive. The proposed method, called Spectral Decomposition Filter (SpDF), identifies and preserves geometric information along the topology of point clouds and is able to scale to large environments with a non-uniform density. Finally, qualitative and quantitative experiments verify the feasibility of our method and present a large-scale evaluation of SpDF with other seven point cloud sampling algorithms, in the context of the 3D registration problem using the Iterative Closest Point (ICP) algorithm on real-world datasets. Results show that a compression ratio up to 97 % can be achieved when accepting a registration error within the range accuracy of the sensor, here for large scale environments of less than 2 cm.

Keywords: sampling, spectral decomposition, large-scale environments, tensor voting, iterative closest point (ICP), registration, 3D mapping, lidar

1. INTRODUCTION

Light Detection And Ranging (Lidar) sensors has recently been widely democratized in robotics applications. Indeed, these sensors are able to acquire an efficient representation of the environment (i.e., a point cloud), which can be used in localization algorithms, 3D mapping or environments inspection (Stumm et al., 2012). Prior work on lidar-based registration algorithms have been recently used to create larger and larger 3D maps (Pomerleau et al., 2014). As an example, **Figure 1** shows the map of the *Grand Axe* of Laval University campus, where only a few minutes of data collection lead to a number of points at the limit of the real-time computation capability. The sensor used for this map, the Velodyne HDL-32E, yields up to 1.39 million points per second.

Although pleasing to the eye, dense maps are not necessarily tailored for practical applications. Indeed, such point clouds are heavy to process and transmit: in scenarios where the processing power and bandwidth are critical resources, point clouds need to be compressed or sampled before any other manipulation. For instance, search & rescue missions often lead the robot in areas where the bandwidth is very narrow. As the tele-operator needs quick feedbacks on where the robot is heading, having access to point clouds with low memory footprint to send through the network without losing information is a necessity.

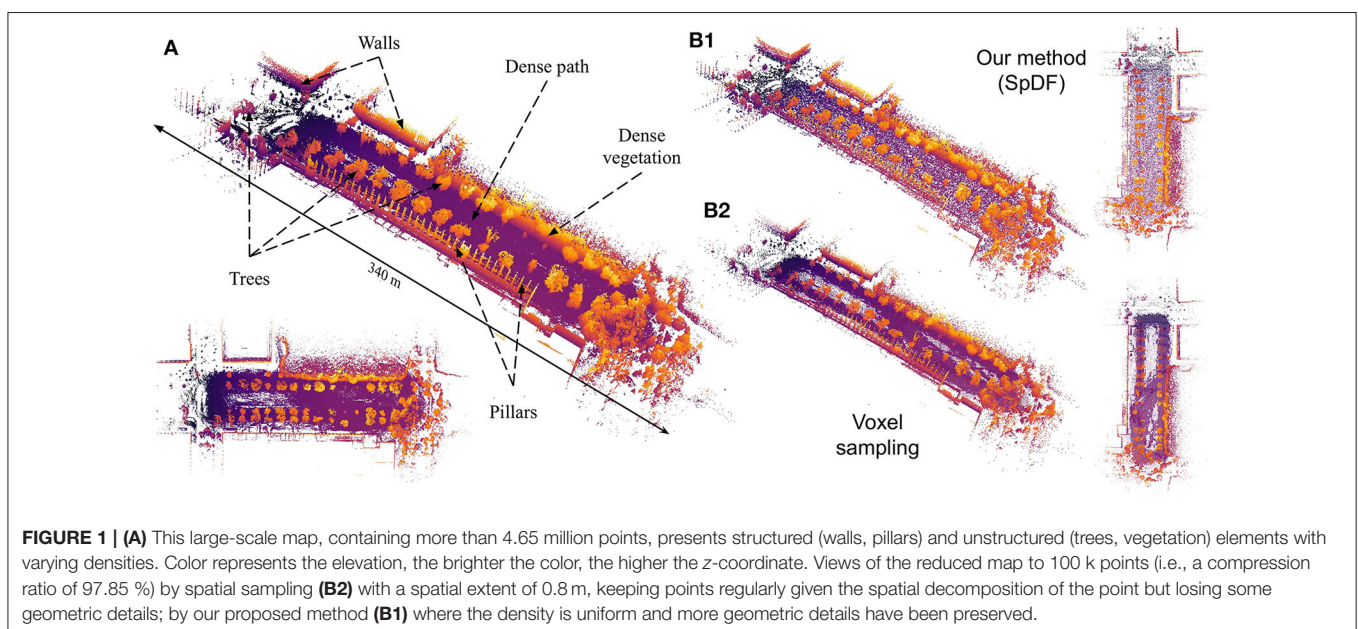
For any of these tasks, the robot needs to merge the different acquired point clouds to create a dense 3D map as it is a necessary information for most of robotics applications. The Iterative Closest Point (ICP) is one of the main algorithms for creating such maps, introduced by Besl and McKay (1992) and Chen and Medioni (1992). It is still considered a strong solution for registration in mobile robotics as shown by Pomerleau et al. (2015). As we want to sample the point cloud, we need to make sure that such action does not hinder the ICP process, since the contrary would lead to noisy, impractical maps. On the one hand, limiting the growth by reducing the number of points will enlarge the spectrum of real-time applications. On the other hand, reducing the number of points too aggressively can lead to unworkable localization and mapping and might discard potential critical information. For example, keeping the edges of objects while doing surface inspection is essential to detect cracks, whereas sampling very dense areas of very little information such as the ground would lead to a substantial compression. Under these considerations, geometric primitives seem a good approach to capture the details along the topology (Stumm et al., 2012). This information can be retrieved by the methodology of Tensor Voting introduced by Guy and Medioni (1997) and Medioni et al. (2000).

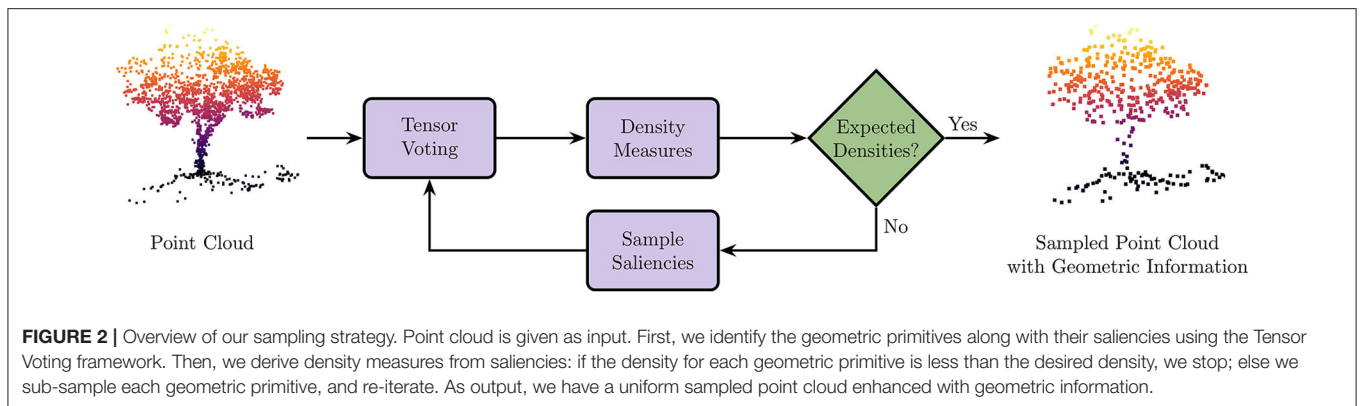
As opposed to solutions for registration-based object reconstruction, we will consider large-scale 3D environments, which are still challenging even for the state-of-the-art sampling methods, because of the uneven density coming from the radial distribution of lidars. Given these working hypotheses, contributions of this paper are 2-fold:

1. A novel sampling method, called Spectral Decomposition Filter (SpDF), based on spectral decomposition analysis. This method identifies and preserves geometric information along the topology of point clouds and is scalable to large environments.
2. A large-scale evaluation of current sampling strategies relying on more than 2.45 million registrations in different types of environments (indoor/structured, outdoor/unstructured), including large-scale outdoors environments from real-world datasets.

A visual overview of our sampling strategy is given by the **Figure 2**. A point cloud is given as input. First, we identify the geometric primitives along with their saliencies using the tensor voting framework. Then, we derive density measures from saliencies: if the density for each geometric primitive is less than the desired density, we stop; else we sub-sample each over-represented geometric primitive, and re-iterate. As output, we have a uniform sampled point cloud enhanced with geometric information.

This paper is organized as follows. First, we review the existing sampling strategies of the literature in section 2. Secondly, we briefly summarize the theory of tensor voting for completeness in section 3. Then, we show how to derive new density measures from the output of the tensor voting process in section 4, and how these new measures are used to efficiently sample point clouds in section 5. Finally, we present our experimental setup in section 6 and analyze the results of our method along





with the evaluation of seven other sampling strategies from the state-of-the-art in section 7.

2. RELATED WORKS

A point-sampled surface is a good representation for analyzing the properties of 3D shapes (Alexa et al., 2001). Unfortunately, most point clouds obtained in robotics context are noisy, sparse, large, and have an uneven density. An important step during the process of analyzing point clouds is to remove noise and outliers. This can be done using filtering algorithms. An extensive review of these algorithms has been realized by Han et al. (2017). Point cloud simplification is related to the problem addressed by the computer vision field but aims to accelerate graphic rendering. A lot of methods based on meshing are used to address this problem (Hoppe, 1996; Li and Zhu, 2008). Technically, these methods can be directly extended to point cloud representation, but most of the algorithms perform an expensive dataset meshing pre-step. A review and comparison of mesh simplification algorithms has been done by Cignoni et al. (1998). Mesh-free algorithms have also been developed to directly simplify point clouds. For instance, Pauly et al. (2002) introduced, analyzed and quantitatively compared a number of surface simplification methods for point-sampled geometry. More recently, Leal et al. (2017) presented a novel method for point cloud simplification using an estimated local density of the point cloud, requiring no additional mesh estimation procedure. Most of these algorithms have been designed in context of computer graphics applications. They compare themselves only with few others strategies, and rely most of the time on evaluations conducted on small, uniform, dense, and complete point clouds such as object models. These working hypotheses usually break in the case of most robotics applications, where point clouds are noisy, incomplete, and sparse. In addition, the sampling process can be addressed by signal processing strategies. Indeed, a point cloud can be considered as a manifold sample. Pauly and Gross (2001) introduced the concept of local frequencies on geometry in order to be able to use all existing signal processing algorithms. Oztireli et al. (2010) proposed a new method to find optimal sampling conditions based on spectral analysis of manifolds. However, these methods stand under the hypothesis of smooth manifolds, which is

usually not the case in maps acquired with lidar sensors in robotics, and rely on evaluations conducted on object models data only.

Sampling algorithms aim to decrease the complexity of point cloud processing (e.g., the computation time) by reducing the number of input points. For instance, ICP algorithm complexity depends on the number of point to process. There are different strategies for points selection that can be categorized as global methods (e.g., uniform and random sampling, spatial sampling), local methods (e.g., using geometric information or density information), and feature-based methods. Feature-based methods such as Fast Point Feature Histogram (FPFH) introduced by Rusu et al. (2009), or Feature-Preserved Point cloud Simplification (FPPS) presented by Zhang et al. (2019) use features which describe the local geometry around a point. It reduces then the number of points by grouping them to describe the neighborhood. These methods provide improvements only with point clouds where features are distinctive which is hard to obtain with noise or incomplete data (Mellado et al., 2014). Hence, this paper will only focus on global and local methods. The most widely used methods are based on octree or voxel representations of point clouds (Schnabel and Klein, 2006; Elseberg et al., 2013; Hornung et al., 2013; Fossel et al., 2017). They take into consideration the spatial distribution of the points by regularly partitioning the space into cells aligned with the xyz -basis. Then, they reduce the number of points by taking the most representative point in each cell, e.g., the centroid. Spatial segmentation methods however do not take into account distinctiveness between the points within cells. For instance, when several geometric features are present in the same cell, the information is lost, which is the case of details in dense areas. They lack of fine grained control over the number of output points and over the level of details.

Another category of methods analyzes geometric primitives to sample relevant points in point clouds. Adaptive sampling strategy based on local density estimation has been proposed by Al-Durgham (2014) in context of registration. Al-Rawabdeh et al. (2020) further extended the latter by sub-sampling on the Gaussian sphere, where normals have been estimated by Principal Component Analysis (PCA), but evaluated their method only on precise point clouds and only against random sampling. Rusinkiewicz and Levoy (2001) proposed a method based on

normals analysis named Normal-Space Sampling (NSS). Points are sampled uniformly according to their normal orientations. They state that it helps convergence for scenes with small, sparse features but by their nature, normals cannot help to handle rotational uncertainties. Rodolà et al. (2015) defined the concept of relevance based on curvature to sample points but such primitives are often noisy and must be processed carefully (Kalogerakis et al., 2009). They compared themselves with the latter strategy and against uniform sampling, in context of object model registration through ICP. Kwok (2018) extended the work on normal space to handle rotational error by introducing a dual normal space to constrain both translation and rotation. He evaluated his method Dual Normal-Space Sampling (DNSS) against several methods, including feature-based and normal-based method, but only on uniformly sampled mesh models. Both NSS and DNSS do not take into account the spatial distribution of the sampling points as they only analyze the normals distributions. Points are not guaranteed to be kept uniformly in space, leading to less accurate results in large-scale sparse point clouds. Gelfand et al. (2003) presented a method based on covariance analysis, Covariance Sampling (CovS), to perform stability analysis in order to select geometrically stable points that can bind the rotational component as well as the translation. An improvement of CovS has been proposed in the context of manufacturing by Kwok and Tang (2015). No evaluation have been conducted on real-world large-scale point cloud for these methods. The authors stated that their proposed approach may suffer from high levels of noise, as large noises make some originally smooth areas strongly constraining. Given our results, previous methods cannot handle noisy, large-scale and density-varying point clouds when they are used to reduce the number of points. Eventually, similar to our method but subsequent to our previous work (Labussière et al., 2018), Ervan and Temeltas (2019) also proposed a sampling algorithm based on a modified tensor voting framework to preserve geometric primitives while down sampling dense areas taking most salient points as representatives. Only qualitative results are given on only one scan, and no comparison have been made. The method presented in this paper is able to both reduce the number of point and retrieve the geometric information in large sparse noisy point clouds. Contrarily to other strategies, we presents a large-scale evaluation of these strategies along with our method in context of large-scale 3D environments.

Although several new strategies have been proposed in the recent years, the most used point cloud processing software such as CloudCompare, the Point Cloud Library (PCL), or libpointmatcher, a popular modular library implementing the Iterative Closest Point (ICP), mostly still rely on either random, uniform or spatial sampling to reduce the number of points.

3. TENSOR VOTING: THEORY

Medioni et al. (2000) introduced Tensor Voting (TV) as a methodology to infer geometric information (e.g., surface, curve,

and junction descriptions) from sparse 3D data¹. The algorithm is based on tensor calculus for data representation and tensor voting for data communication. Theory related to TV will be summarized in this section for completeness.

3.1. Tensor Representation

To capture the first order differential geometry information and its saliency, each datum can be represented as a second order symmetric tensor in the normal space. In 3D, such a tensor can be visualized as an ellipsoid with a shape that defines the nature of the information and a scale that defines the saliency of this information. A second order symmetric tensor \mathbf{K} is fully described by its associated spectral decomposition using three eigenvectors \mathbf{e}_1 , \mathbf{e}_2 , and \mathbf{e}_3 , and three corresponding ordered positive eigenvalues $\lambda_1 \geq \lambda_2 \geq \lambda_3$. This tensor can be decomposed in three basis tensors, resulting in

$$\mathbf{K} = (\lambda_1 - \lambda_2) \mathbf{S} + (\lambda_2 - \lambda_3) \mathbf{P} + \lambda_3 \mathbf{B}, \quad (1)$$

with

$$\begin{aligned} \mathbf{S} &= \mathbf{e}_1 \mathbf{e}_1^T, \\ \mathbf{P} &= \sum_{d=1}^2 \mathbf{e}_d \mathbf{e}_d^T, \\ \mathbf{B} &= \sum_{d=1}^3 \mathbf{e}_d \mathbf{e}_d^T, \end{aligned} \quad (2)$$

where \mathbf{S} describes the stick tensor, \mathbf{P} the plate tensor, and \mathbf{B} the ball tensor.

3.2. Voting Process

The main goal of Tensor Voting is to infer information represented by the tensor \mathbf{K}_i at each position \mathbf{x}_i by accumulating cast vote \mathbf{V} from its neighborhood \mathcal{N} , following

$$\mathbf{K}_i = \sum_{\mathbf{x}_j \in \mathcal{N}(\mathbf{x}_i)} \mathbf{V}(\mathbf{x}_i, \mathbf{x}_j). \quad (3)$$

This process can be interpreted as a convolution with a predefined aligned voting field. The voting fields encode the basis tensors and are derived from the 2D stick field by integration (see Medioni et al., 2000 for more details). Each input point is encoded into a tensor. First, if no direction is given, the tensor encodes a unit ball \mathbf{B} . Second, if tangents are provided, the tensor encodes a plate \mathbf{P} . Finally, if normals are available, the tensor encodes a stick \mathbf{S} . In a case where no direction is given, a first pass of refinement is done to derive the preferred orientation information. Each tensor then broadcasts each of its independent elements using an appropriate tensor field:

$$\mathbf{V}(\mathbf{x}_i, \mathbf{x}_j) = \mathbf{V}_S(\mathbf{x}_i, \mathbf{x}_j) + \mathbf{V}_P(\mathbf{x}_i, \mathbf{x}_j) + \mathbf{V}_B(\mathbf{x}_i, \mathbf{x}_j), \quad (4)$$

where \mathbf{V}_S (resp. \mathbf{V}_P and \mathbf{V}_B) is the vote generated by the tensor field associated to \mathbf{S} (resp. \mathbf{P} and \mathbf{B}).

¹The original formulation can be extended to n -D data (Tang et al., 2001).

3.3. Vote Interpretation

The resulting generic second order symmetric tensor \mathbf{K} is then decomposed into elementary components to extract the saliencies and the preferred direction. The interpretation of these values is given in **Table 1**. We can then infer geometric primitives, but the procedure to extract the salient features corresponding to local maxima of the three saliency maps will not be discussed here.

3.4. k-Nearest Neighbors Closed Form Tensor Voting

Although tensor voting is a robust technique for extracting perceptual information from point clouds, the complexity of its original formulation makes it difficult to use in robotics applications. We use the closed-form (CFTV) formulation proposed by Wu et al. (2012) for efficiency. The generic second order symmetric tensor is then computed given

$$\mathbf{K}_i = \sum_{\mathbf{x}_j \in \mathcal{N}(\mathbf{x}_i)} \mathbf{S}_{ij} \quad \text{with} \quad \mathbf{S}_{ij} = c_{ij} \mathbf{R}_{ij} \mathbf{K}_j \mathbf{R}'_{ij}, \quad (5)$$

and

$$\begin{aligned} \mathbf{R}_{ij} &= \left(\mathbf{I} - 2\mathbf{r}_{ij} \mathbf{r}'_{ij} \right), & \mathbf{R}'_{ij} &= \left(\mathbf{I} - \frac{1}{2} \mathbf{r}_{ij} \mathbf{r}'_{ij} \right) \mathbf{R}_{ij}^T, \\ \mathbf{r}_{ij} &= \frac{\mathbf{x}_i - \mathbf{x}_j}{\|\mathbf{x}_i - \mathbf{x}_j\|}, & c_{ij} &= \exp\left(-\frac{\|\mathbf{x}_i - \mathbf{x}_j\|^2}{\sigma}\right), \end{aligned} \quad (6)$$

where c_{ij} is a decay function and controls the strength of the vote given the distance between the two positions and the scale parameter σ ; \mathbf{r}_{ij} is the normalized vector from \mathbf{x}_j in the direction of \mathbf{x}_i ; and \mathcal{N} is the neighborhood retrieved using an efficient k -Nearest Neighbors (k -NN) search (e.g., with a kD -tree). As the input is generally not oriented, we still have to do a first pass by encoding \mathbf{K}_j as a unit ball to derive a preferred direction. Then, we do a second pass by encoding points with the tensors previously obtained, but with the ball component disabled as suggested by Wu et al. (2012), such as $\mathbf{K}_j = (\lambda_1 - \lambda_2) \mathbf{S}_j + (\lambda_2 - \lambda_3) \mathbf{P}_j$. Once the generic tensor is computed, we decompose and interpret it as shown above.

4. DERIVATION OF DENSITY MEASURES

Based on tensor voting theory, this paper presents a novel density measure for each geometric primitive. By doing a first pass of TV

TABLE 1 | Interpretation of saliencies and preferred directions obtained by the tensor voting framework, where the λ_d are the eigenvalues associated to the eigenvectors \mathbf{e}_d obtained from spectral decomposition of the resulting tensors. Predominating saliency determines the affected geometric primitive.

	Geom. Primitive	Tensor	Saliency	Normals
Surface-ness	Surface	Stick \mathbf{S}	$\lambda_1 - \lambda_2$	\mathbf{e}_1
Curve-ness	Curve	Plate \mathbf{P}	$\lambda_2 - \lambda_3$	$\mathbf{e}_1, \mathbf{e}_2$
Point-ness	Junction	Ball \mathbf{B}	λ_3	$\mathbf{e}_1, \mathbf{e}_2, \mathbf{e}_3$

using the closed-form with an k -NN search (Equation 5), we are able to derive more information from the tensors. In fact we can show that $0 \leq \lambda_d \leq k, \forall d \in \{1, 2, 3\}$, where k is the number of neighbors. As the strength of the vote through the decay function is directly dependent on the distance, we have $\lambda_d = k$ when all neighbors are at a distance $\delta = 0$. Given this observation, the lambdas can be considered as an indicator of local density.

In the following, the λ_d are normalized by k . We can compute the expected normalized vote strengths ξ_D at a position where the density would be uniform in a D -hyperball of radius ρ to derive the density measures. The strength of the vote c_{ij} is only dependent on the distance δ between \mathbf{x}_i and \mathbf{x}_j (Equation 6) such as $\delta = \|\mathbf{x}_i - \mathbf{x}_j\|^2$. Therefore only the decay function $c(\delta) = \exp(-\delta^2/\sigma)$ is taken into account. We compute the expectation of the decay function given points following a uniform spatial distribution in a D -hyperball of radius ρ . In order to achieve this distribution, we can generate samples of the distance δ through the random variables $X \sim \mathcal{U}_{[0,1]}$, where $\mathcal{U}_{[0,1]}$ is the uniform distribution between 0 and 1, by using the inverse of the cumulative distribution function (CDF) corresponding to the surface area of this hyperball. A random sample is generated by mapping random numbers in the range $[0, 1]$ through the application $\delta(x) = \rho \cdot x^{\frac{1}{D}}$. We then compute the expected value of this distribution such as

$$\begin{aligned} \mathbb{E}[c(\delta(X))] &= \int_{-\infty}^{\infty} \text{pdf}_X(x) \cdot c(\delta(x)) \, dx \\ &= \frac{D}{2} \left(\frac{\rho^2}{\sigma}\right)^{-\frac{D}{2}} \left(\Gamma\left(\frac{D}{2}\right) - \Gamma\left(\frac{D}{2}, \frac{\rho^2}{\sigma}\right)\right), \end{aligned} \quad (7)$$

where σ is the scale of the vote, $\Gamma(\cdot)$ is the gamma function and $\Gamma(\cdot, \cdot)$ is the incomplete gamma function. For $D \in \{1, 2, 3\}$, the expected kernel strengths are given by

$$\begin{aligned} \xi_1 &= \frac{1}{4\rho} \sqrt{\pi\sigma} \operatorname{erf}\left(\frac{\rho}{\sqrt{\sigma}}\right) \\ \xi_2 &= \frac{\sigma}{\rho^2} \left(1 - \exp\left(-\frac{\rho^2}{\sigma}\right)\right) \\ \xi_3 &= \frac{3\sigma}{4\rho^3} \left(\sqrt{\pi\sigma} \operatorname{erf}\left(\frac{\rho}{\sqrt{\sigma}}\right) - 2\rho \exp\left(-\frac{\rho^2}{\sigma}\right)\right), \end{aligned} \quad (8)$$

where $\operatorname{erf}(\cdot)$ is the Gauss error function, ξ_1 is the expected strength in the plate case ($D = 1$), ξ_2 is the expected strength in the stick case ($D = 2$) and ξ_3 is the expected strength in the ball case ($D = 3$).

The associated eigenvalues $\hat{\lambda}_d$ can be derived if we consider, for each component, the ideal cases illustrated by **Figure 3** where each voter strength is ξ_D . By developing (Equation 5) with $c_{ij} = \xi_D$, considering an infinite number of neighbors, thus taking \mathbf{r}_{ij} as the integral variable on the considered domain (i.e., all possible orientations in D dimensions), we compute for each case the expected eigenvalues $\hat{\lambda}_d$. For instance, in the case where points are uniformly distributed on a sphere (i.e., $D = 3$ as illustrated by **Figure 3A**) of radius such that $c_{ij} = \xi_3$, the integral variable is given by the normalized

vector $\mathbf{r} = [\sqrt{1-u^2} \cos(\theta) \sqrt{1-u^2} \sin(\theta) u]^T$ with $(\theta, u) \in [0, 2\pi] \times [-1, 1]$, and the resulting normalized tensor by

$$\begin{aligned} \bar{\mathbf{K}} &= \eta \int_{\Gamma} \xi_D (\mathbf{I} - 2\mathbf{r}\mathbf{r}^T) \left(\mathbf{I} - \frac{1}{2}\mathbf{r}\mathbf{r}^T \right) (\mathbf{I} - 2\mathbf{r}\mathbf{r}^T)^T d\mathbf{r} \\ &= \frac{1}{4\pi} \int_0^{2\pi} \int_{-1}^1 \xi_3 (\mathbf{I} - 2\mathbf{r}\mathbf{r}^T) \left(\mathbf{I} - \frac{1}{2}\mathbf{r}\mathbf{r}^T \right) (\mathbf{I} - 2\mathbf{r}\mathbf{r}^T)^T du d\theta \\ &= \frac{\xi_3}{4\pi} \begin{bmatrix} \frac{20}{6}\pi & 0 & 0 \\ 0 & \frac{20}{6}\pi & 0 \\ 0 & 0 & \frac{20}{6}\pi \end{bmatrix} = \begin{bmatrix} \frac{5}{6}\xi_3 & 0 & 0 \\ 0 & \frac{5}{6}\xi_3 & 0 \\ 0 & 0 & \frac{5}{6}\xi_3 \end{bmatrix} = \begin{bmatrix} \hat{\lambda}_3 & 0 & 0 \\ 0 & \hat{\lambda}_2 & 0 \\ 0 & 0 & \hat{\lambda}_1 \end{bmatrix} \end{aligned} \tag{9}$$

where Γ is the integration domain of \mathbf{r} , and $\eta = (\int_{\Gamma} d\mathbf{r})^{-1}$ is the normalization constant since the eigenvalues are normalized by the number of neighbors. Finally, we have $\hat{\lambda}_1 = \hat{\lambda}_2 = \hat{\lambda}_3 = \frac{5}{6} \xi_3$ and therefore the expected point-ness saliency is $\frac{5}{6} \xi_3$. In a similar fashion for the case $D = 2$ illustrated by **Figure 3B**, points are uniformly distributed on a circle of radius such that $c_{ij} = \xi_2$, the integral variable is given by $\mathbf{r} = [\cos(\theta) \sin(\theta) 0]^T$ with $\theta \in [-\pi, \pi]$, and the expected surface-ness saliency is given by $\frac{1}{4} \xi_2$. For the case $D = 1$ illustrated by **Figure 3C**, points are uniformly distributed on line segment endpoints along the x -axis, of length such that $c_{ij} = \xi_1$, we only have to consider the integral variable $\mathbf{r} \in \left\{ [1 \ 0 \ 0]^T, [-1 \ 0 \ 0]^T \right\}$, and the expected curve-ness saliency is given by $\frac{1}{2} \xi_1$.

We are now able to interpret the saliencies obtained by a first pass of the closed-form TV where every point has been encoded as ball tensor (i.e., $\mathbf{K}_j = \mathbf{I}, \forall j$) as a measure of local density. We can therefore compare the values with the expected saliencies, summarized in **Table 2**, to control the density of each geometric primitive.

5. SPECTRAL DECOMPOSITION FILTER (SPDF): OVERVIEW

The method presented in this article, namely SpDF, aims to reduce the number of points while preserving as much as possible the topology of the point cloud using geometric primitives (i.e., curve, surface, and junction). Note that it is not limited to plane, line and point as the tensor voting framework allows to detect more generic geometric primitives. A major challenge in robotics

applications is the non-uniformity of scans acquired with lidar sensors. In fact most of sampling algorithms are designed for uniform point clouds. This problem is addressed by proposing a new efficient strategy to make the density uniform for each of the three geometric primitives we consider. SpDF can be divided into two main steps: (1) making the density uniform for each geometric primitive; and (2) labeling and rejecting outliers according to the confidence in the geometric information. A visual overview of the algorithm is given in **Figure 2**.

5.1. Making the Density Uniform

Using the new local density measure on each geometric primitive, the point cloud can be made uniform as follows. An iterative procedure allows to progressively decimate primitives where the saliencies are higher than the expected values reported in **Table 2**. The saliencies are recomputed using TV with tensors encoded as unit balls. The algorithm stops when the number of points is stable, which means that the saliencies distributions have converged below the expected values, as shown by **Figure 4A**. This figure shows the convergence of the saliencies below their expected values (represented by the vertical dashed lines), where the top-histogram represents the initial saliencies distribution and the bottom-histogram shows the resulting distribution after making the density uniform. Therefore, the densities are uniform around each primitive allowing us to detect them more clearly. Otherwise, most dense areas will be detected as junction because noise will predominate. An example of the result of making the density uniform (for $k = 50$ and $\sigma = \rho = 0.2$) is given by **Figure 4B** for a point cloud from 370 to 40k points (i.e., a compression ratio of 89%). The resulting point cloud have uniform density, and geometric primitives are preserved. All

TABLE 2 | Expected eigenvalues and saliencies in the case of a uniform density in a D -hyperball, which can be interpreted as density measures and allow to control the density of each geometric primitive.

	D	Eigenvalues	Saliency
Curve-ness (P)	1	$\hat{\lambda}_1 = \hat{\lambda}_2 = \xi_1$ and $\hat{\lambda}_3 = \frac{1}{2} \xi_1$	$\frac{1}{2} \xi_1$
Surface-ness (S)	2	$\hat{\lambda}_1 = \xi_2$ and $\hat{\lambda}_2 = \hat{\lambda}_3 = \frac{3}{4} \xi_2$	$\frac{1}{4} \xi_2$
Point-ness (B)	3	$\hat{\lambda}_1 = \hat{\lambda}_2 = \hat{\lambda}_3 = \frac{5}{6} \xi_3$	$\frac{5}{6} \xi_3$

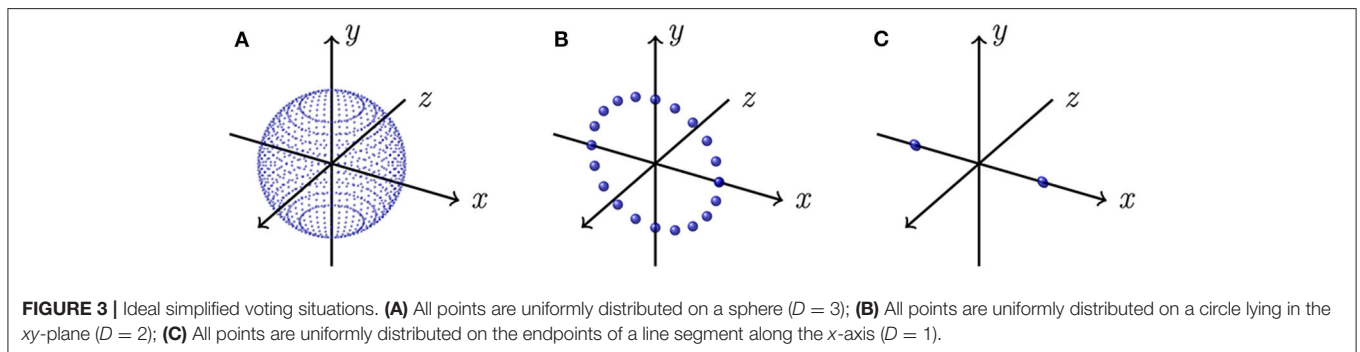
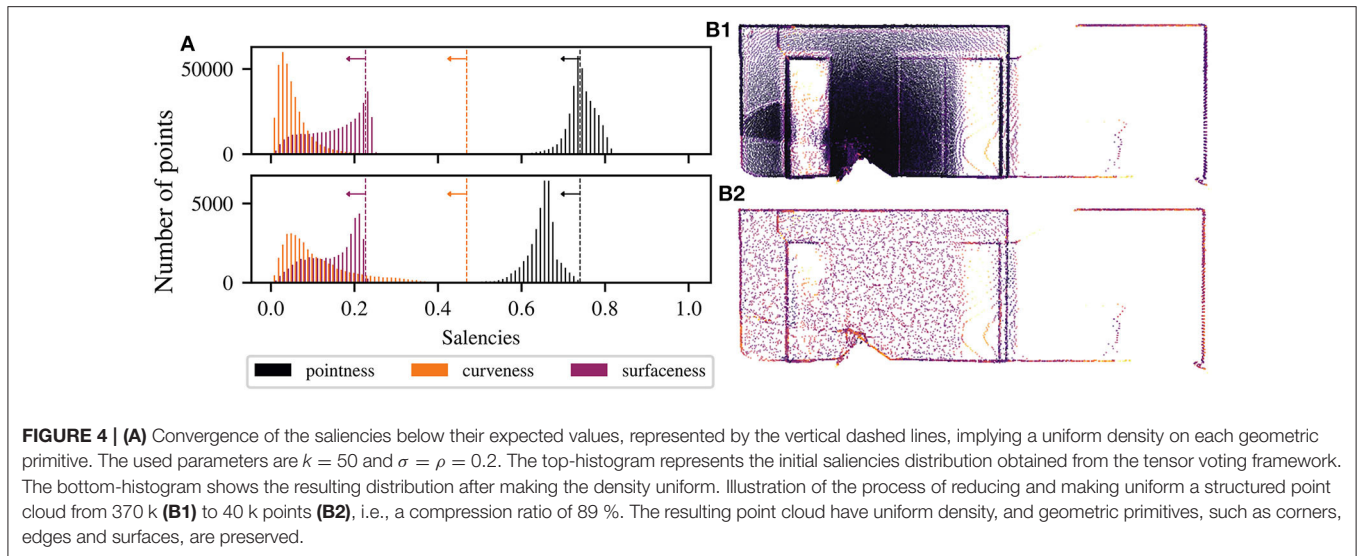


FIGURE 3 | Ideal simplified voting situations. **(A)** All points are uniformly distributed on a sphere ($D = 3$); **(B)** All points are uniformly distributed on a circle lying in the xy -plane ($D = 2$); **(C)** All points are uniformly distributed on the endpoints of a line segment along the x -axis ($D = 1$).



planes have then the same density, edges have been kept at the same density, and less points are wrongly identified as junction.

Finally, to control the density, the user only has to tune the parameter ρ , i.e., the radius of uniformity within the D -hyperball (where D is the dimension of the considered geometric primitive). Indeed, the expected vote strengths ξ_D is normalized, therefore σ (the spatial extent of the neighborhood) and k (the number of points to consider in the neighborhood) do not influence the density once set. In our experiment we set $\sigma = 20$ cm, $k = 50$, which is a good compromise between precision and time complexity, and ρ varies from 10 to 1.35 m. It means that once sampled, we expect a density of k points uniformly distributed within a ball of radius ρ for junctions, within a disk of radius ρ for surfaces, and on a segment of half-length ρ for curves.

5.2. Rejecting Outliers and Identifying Geometric Primitives

Given the saliencies computed with a last iteration of TV with the ball component disabled, each point is then labeled into *junction*, *curve*, or *surface* following the TV methodology, and the saliency associated (respectively point-ness, curve-ness, or surface-ness) encodes the confidence in this labeling. It provides a high level description in terms of geometry. Points with a confidence higher than t % of the maximum confidence of the considered geometric primitive are kept (we used in our experiments $t = 10$ %). This heuristic allows to reject outliers having a low confidence in their measure. At the end of the process, the point cloud is uniform, outliers have been rejected, and each point is labeled as surface, curve or point. SpPDF is then able to reduce the point cloud while enhancing it with geometric information.

6. EXPERIMENTAL SETUP

To validate our sampling method, we investigate its impact on a 3D registration process, and evaluate it along with several

sampling methods. Indeed, registrations of point clouds is a mandatory step in most of robotics applications.

6.1. Registration Based on the Iterative Closest Point (ICP) Algorithm

Registration is the process of aligning the frames of two point clouds, the reference P and the reading Q , by finding the rigid transformation $T \in SE(3)$ between them by a minimization process. The transformation can be determined through the ICP algorithm introduced by Besl and McKay (1992) and Chen and Medioni (1992). The original algorithm only works well in ideal cases. To improve the robustness of the original formulation, several variants have been proposed. In order to leverage geometric information, such as normals, Rusinkiewicz and Levoy (2001) introduced an efficient variant of the ICP algorithm based on the *point-to-plane* formulation of the error minimization. Segal et al. (2009) proposed a generalization of the ICP algorithm, called Generalized-ICP, which takes into account the locally planar structure of both scans in a probabilistic model. Stoyanov et al. (2012) presented a novel approach to point cloud registration, based on minimizing the distance between Gaussian distributions. Point set are locally represented by their Normal Distribution Transform (NDT), which are a special case of Gaussian Mixture Model for representing the probability to find a surface point at a specific position in space. Pomerleau et al. (2015) highlight that surface reconstruction is expensive to compute, but at the same time highlight the fact that leveraging geometric information when using *point-to-plane* error leads to a faster convergence rate. Most of the current strategies rely on geometric information to minimize the error of the registration process, requiring then to process the point cloud to extract these information.

To evaluate the accuracy of the registration, we calculate separately the translation error part, ε_t , and the rotational error part, ε_r , the same way it is done by Pomerleau et al. (2013). Given the ground-truth transformation T_{gt} and its

corresponding transformation found by the registration solution T , the remaining error ΔT is defined as

$$\Delta T = \begin{bmatrix} \Delta R & \Delta t \\ \mathbf{0} & 1 \end{bmatrix} = T T_{gt}^{-1}, \quad (10)$$

and ε_t is then given as the Euclidean norm of the translation Δt , and ε_r is defined as the Geodesic distance directly from ΔR , such as:

$$\varepsilon_t = \|\Delta t\| \quad \text{and} \quad \varepsilon_r = \arccos\left(\frac{\text{trace}(\Delta R) - 1}{2}\right). \quad (11)$$

As ICP needs a prior for fine registration (i.e., the initial transformation \tilde{T}) to compute the transformation between two point clouds, we applied a uniform perturbation on the ground-truth transformation T_{gt} using Lie algebra, such that $\tilde{T} = \exp(\zeta) T_{gt}$, with $\zeta \in \mathfrak{se}(3)$ and $\exp(\cdot)$ being the standard matrix exponential. For our experiments, a perturbation sampled from a uniform distribution of 50 cm was applied on the translation, and from a uniform distribution of 20° on the rotation. During the data filtering step, we applied the evaluated filter on both the reading and the reference. The data association is conducted by matching the two closest neighbors. We rejected the outliers in the matching process according to a trimmed distance. We limited the scope our experiments to a *point-to-plane* version of ICP, as it tends to perform better in those datasets as highlighted by Pomerleau et al. (2013). The minimization process stops when the number of iterations reached 150, or when the differential translation error is less than 1 cm and the differential rotation error is less than 0.001 rad.

6.2. Details on the Real-World Datasets

Working under the hypothesis of robotics applications, this paper presents an in-depth evaluation of sampling algorithms on (1) structured with 45 pairs of scans, (2) semi-structured with 32 pairs of scans, and (3) unstructured point clouds with 32 pairs of scans, using the datasets ‘‘Challenging data sets for point cloud registration algorithms’’ (Pomerleau et al., 2012)². The structured environment is a map of an apartment with approximate dimension of $17 \times 10 \times 3$ m, with an average 365 k points per scan. The semi-structured environment is a map of a park where there is grass, paved small roads, sparse trees, and containing a gazebo made of rock walls and a wood ceiling covered with vines trees. The dimension are $35 \times 45 \times 16$ m, with an average of 170 k points per scan. Finally, the unstructured environment is a map of a wood, mainly constituted of vegetation (tree, bushes, etc.). The only structured element is a small paved road that crosses the wood. The dimension are $36 \times 60 \times 22$ m, with an average of 178 k points per scan. The used sensor is an Hokuyo UTM-30LX, a time-of-flight sensor with a minimum range of 0.1 m and a maximal range of 30 m. As indicated by Pomerleau et al. (2012), the specifications of the sensor give a range accuracy varying from ± 1 cm for distances within [0.1, 10] m to ± 3 cm for distances within [10, 30] m.

²Details on dataset can be found online at <https://projects.asl.ethz.ch/datasets/doku.php?id=laserregistration:laserregistration> (last accessed July 27, 2020).

TABLE 3 | Ranges [a; b] of parameters influencing the number of points, where a gives the smallest number of points (here, 1,000 points) and b preserves all the points (with n being the total number of points), for each evaluated method with their description.

Method	Parameter description	Range
Random (baseline)	prob. to keep point	[0.004 ; 1.]
NSS	nb. of points to keep	[1000 ; n]
CovS	nb. of points to keep	[1000 ; n]
SSNormal	nb. of neighbors to merge	[253 ; 3.]
Octree	nb. max of points by cell	[1000 ; 1.]
Voxel	size max of the cell in m	[2.49 ; 0.001]
Max Density	nb. max of points by m^3	[16.8 ; 506 k]
SpDF (ours)	radius of uniformity in m	[1.35 ; 0.1]

6.3. Overview of the Evaluated Methods

To evaluate the impact of the number of points on the registration process, several methods from the state-of-the-art had been implemented in the open-source modular library for ICP named `libpointmatcher`, introduced by Pomerleau et al. (2013) and available online³. The eight evaluated filters, summarized in **Table 3**, are the following:

- Random sampling, chose as our baseline for its simplicity and because it is still one of the most used solution; point cloud is reduced by dropping points given a fixed probability.
- One neighbor-based method from the `libpointmatcher`, the Sampling Surface Normal filter, `SSNormal`; it recursively decomposes the point cloud into boxes until each box contains at most a given number of points, and select one point if normal estimation can be conducted within the box.
- Two variants of spatial segmentation, `Voxel` and `Octree` (centroid); `Voxel` sub-divides the space with a fixed spatial extent whereas `Octree` uses the number of points per cell as a criterion to stop the partition. Both take the centroid in each cell to sub-sample the point cloud.
- Two normal-based methods, `CovS`, the Covariance Sampling method from Gelfand et al. (2003) and `NSS`, the Normal Space Sampling method from Rusinkiewicz and Levoy (2001).
- Two density-based method, the `MaxDensity` method from the `libpointmatcher` and eventually, the proposed method in this paper, `SpDF`; `MaxDensity` aims to homogenize the density of a point cloud by rejecting a sub-sample of points in high-density regions. Points are only considered for rejection if they exceed a density threshold, otherwise they are preserved. It relies on a spherical approximation to compute density. `SpDF` also aims to homogenize the density but on each different geometric primitive. It leverages the tensor voting framework to identify these primitives and to derive density measures for each. It therefore reduces the number of points and enhances the point cloud with geometric information, such as the

³The library is available at <https://github.com/ethz-asl/libpointmatcher>. The filters documentation is available at <https://libpointmatcher.readthedocs.io/en/latest/DataFilters/> (last accessed August 06, 2020).

normals. Hence, SpDF can be seen as an improvement of the MaxDensity method.

The evaluated methods are similar to the most used strategies available in point cloud processing software such as CloudCompare and PCL. For each method, we resumed in Table 3 the range $[a; b]$ of parameters influencing the number of points, where a gives the smallest number of points, here, 1.000 points, and b preserves all the points, with n being the total number of points. We performed 2.500 registrations using the ICP algorithm (5.000 for our baseline Random) across the range for each method on each pair of scans, for each dataset, accumulating more than 2.45 million registrations in different types of environments.

7. RESULTS AND DISCUSSION

7.1. Evaluation on Registration Accuracy

First, we compare the different sampling methods on the registration accuracy. Figure 5 presents the translation and the rotational errors as functions of the compression ratio of the point clouds for all environments concatenated. At 0%, all points have been kept and at 100%, all points have been removed. The gray area represents the errors lower than our baseline (Random), the red solid-line corresponds to SpDF and the dashed-lines to the other methods.

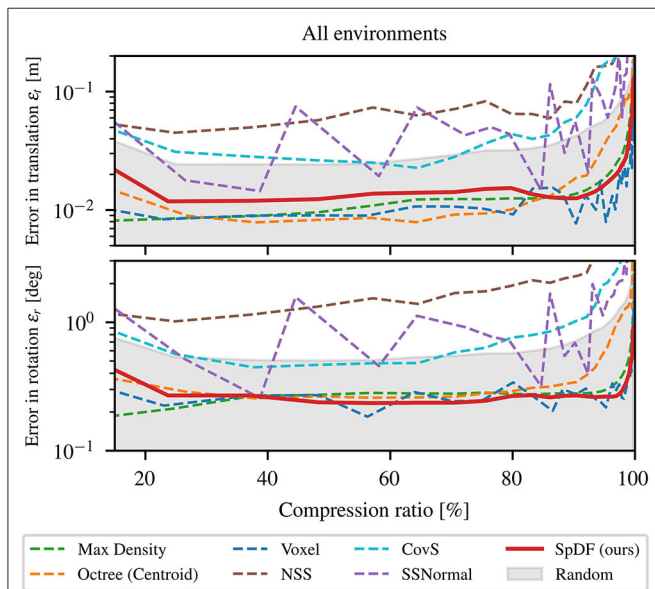
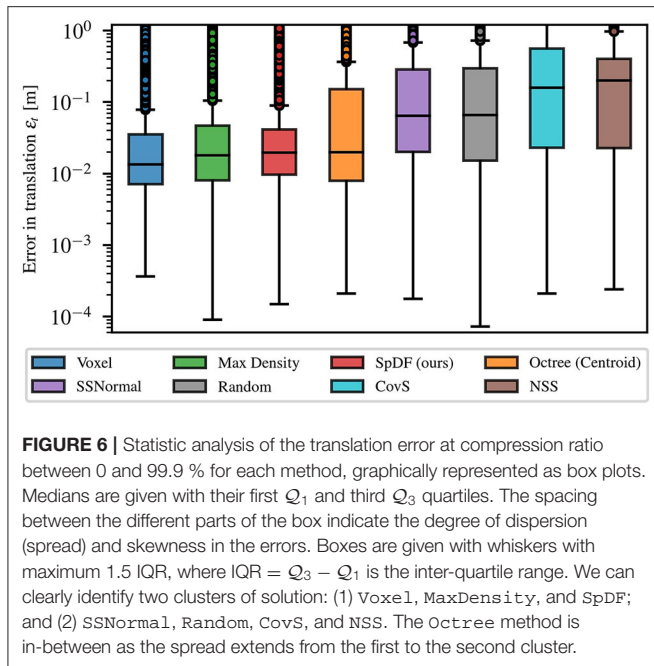


FIGURE 5 | Influence of the number of points (i.e., compression ratio) on the registration process for all environments concatenated. The gray area represents the errors less than the baseline. SpDF (ours) is displayed in solid-line. The top-graph represents the error in translation ϵ_t in m; The bottom-graph represents the error in rotation ϵ_r in deg. Both translation and rotational errors show the same patterns. Only spatial methods (Voxel and Octree) and density-based methods (MaxDensity and SpDF (ours)) outperform the baseline, providing a translation error less than 2 cm, and a rotation error of less than 0.3° , which tend to the minimal reachable errors for the ICP algorithm.

Both translation (ϵ_t) and rotational (ϵ_r) errors show the same patterns. From 0 to 25%, corresponding to more or less 100 k points, the errors decrease as the compression ratio increases. From 25 to 80%, the errors are almost constant or grow at a really slow rate. Beyond 80%, the errors start to increase exponentially. In the first situation, with more points, the dense areas predominate in the minimization process, leading to less accurate results. In the last situation, with only a few points, the minimization process cannot converge efficiently as not enough information is available to constrain the process. Both situations lead to less accurate results. The evaluated methods are significantly more accurate than the baseline except NSS, CovS, and SSNormal, which perform worse than Random, with a median error of 20 cm against 7 cm for the translation. This confirms their inability to manage uneven density and large-scale point cloud performing poorly for all types of environment. These algorithms need to be adapted for an application in the context of robotics, with large-scale, uneven environments.

Spatial methods perform well on all datasets, providing a median translation and rotation errors which tend to the minimal reachable errors for the ICP algorithm on these kind of maps. However, Octree starts diverging sooner than the others, around 90%, corresponding to approximately 30 k points, when the number of points decreases. With Octree, we obtained a translation error of 1.9 cm and a rotational error of 0.4° . It performs well to a certain extent as it is able to preserve the spatial distribution for a large number of points, but suffers from the uneven density distribution for a small number of points. The Voxel method performs well as it preserves the spatial distribution whatever the density, with a translation error of 1.2 cm and a rotational error less than 0.27° . Using only the density, MaxDensity leads to more accurate alignments than the baseline showing a translation error of 1.6 cm and a rotational error less than 0.3° . Finally, SpDF with an error in translation of 1.8 cm and an error in rotation of 0.278° , shows the best results from all the evaluated methods along with MaxDensity and Voxel, which are equivalent in terms of errors. The proposed method inherits the qualities of the density-methods and manage efficiently large point clouds with uneven densities by maintaining a given density for each geometric primitive.

Figure 6 gives a statistic analysis of the translation error at compression ratio between 0 and 99.9% for each method, graphically represented as box plots. Medians are given with their first Q_1 and third Q_3 quartiles. The spacing between the different parts of the box indicate the degree of dispersion (spread) and skewness in the errors. Boxes are given with whiskers with maximum 1.5 IQR, where $IQR = Q_3 - Q_1$ is the inter-quartile range. Points outside the boxes represent the outliers. We can clearly identify two clusters of solutions: (1) Voxel, MaxDensity, and SpDF; and (2) SSNormal, Random, CovS, and NSS. The Octree method is in-between as the spread extends from the first to the second cluster. The second cluster presents a median error between 6.5 and 20 cm with a large spread of more than 50 cm. The first cluster presents a median error between 1.2 and 1.8 cm but with an equivalent degree a dispersion of 2.8 cm. The three methods are therefore equivalent

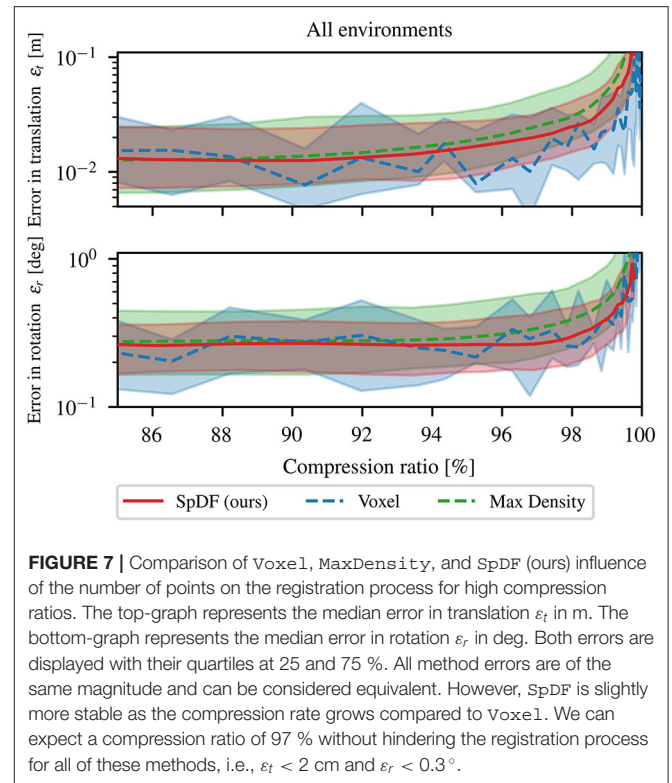


in terms of error. In the following, we will conduct the analysis only on the first cluster of solutions, i.e., on *SpDF* along with *MaxDensity* and *Voxel*.

7.2. Comparison for High Compression Ratio and for Each Type of Environment

Secondly, we analyze the behavior for high compression ratio. The three best methods have been compared for compression ratio between 85 and 99.9 %. Errors in translation and rotation are illustrated by **Figure 7**. Both errors are displayed with their quartiles at 25 and 75 %. All method errors are of the same magnitude, even if *MaxDensity* performs slightly worse as the compression ratio grows. Rotational error is constant and less than 0.3° until 97 % and start growing exponentially afterward. The same behavior is observed for the translation errors which are less than 2 cm until a compression of 92 %. We believe that the error tends to the minimal reachable error that can be obtained with the ICP algorithm, making it hard to state that one method outperforms the others. However, *SpDF* and *MaxDensity* are slightly more stable as the compression rate grows compared to *Voxel*. One explanation is that spatial sampling depends on the origin of the sampling and on the frequency: the larger the spatial extent of the voxel, the greater the shift in the point cloud. Sampled point in the reading and reference point cloud might not represent the same part of the point cloud, leading to less stable results while matching. Finally, density-based methods as ours are able to perform well in context of 3D registration because they are able to deal with noisy and non-uniform point clouds.

These three methods are equivalent in terms of translation and rotation errors. Statistical analysis reported in **Figure 8** confirms this hypothesis. On this range of compression ratio, we can expect a translation errors of: 2.37 cm with a lower quartile



$Q_1 = 1.05$ cm, and upper quartile $Q_3 = 5.11$ cm, i.e., an IQR of 4.06 cm for *Voxel*; for *SpDF*, an error of 2.40 cm with a lower quartile $Q_1 = 1.21$ cm, and upper quartile $Q_3 = 4.59$ cm, i.e., an IQR of 3.37 cm; and of 3.59 cm with a lower quartile $Q_1 = 1.63$ cm, and upper quartile $Q_3 = 9.62$ cm, i.e., an IQR of 7.99 cm for *MaxDensity*. *MaxDensity* provides a higher error and spread than the two others methods. *Voxel* and *SpDF* are equivalent in terms of median errors, but *SpDF* is slightly more stable as the spread is lower. On this range of compression ratio, we can expect a rotation errors of: 0.40° with a lower quartile $Q_1 = 0.23^\circ$, and upper quartile $Q_3 = 0.82^\circ$, i.e., an IQR of 0.59° for *Voxel*; for *SpDF*, an error of 0.31° with a lower quartile $Q_1 = 0.19^\circ$, and upper quartile $Q_3 = 0.50^\circ$, i.e., an IQR of 0.31° ; and of 0.48° with a lower quartile $Q_1 = 0.26^\circ$, and upper quartile $Q_3 = 1.15^\circ$, i.e., an IQR of 0.89° for *MaxDensity*. Compared to *MaxDensity* and *Voxel*, *SpDF* gives the lowest median error with the smallest spread, but the difference is not significant enough and all three methods can be considered equivalent.

Thirdly, we analyze the impact of the type of environment on the registration error for high compression ratio between 85 and 99.9 %. **Figure 8** gives a statistic analysis of the translation and rotation error for the top three methods, graphically represented as box plots. Medians are given with their first Q_1 and third Q_3 quartiles. Boxes are given with whiskers with maximum 1.5 IQR. Even if each environment shows its own kind of errors variations, the methods behave similarly independently of the types of environment. For the rotation errors, the semi-structured and unstructured present similar errors of 0.3° , with

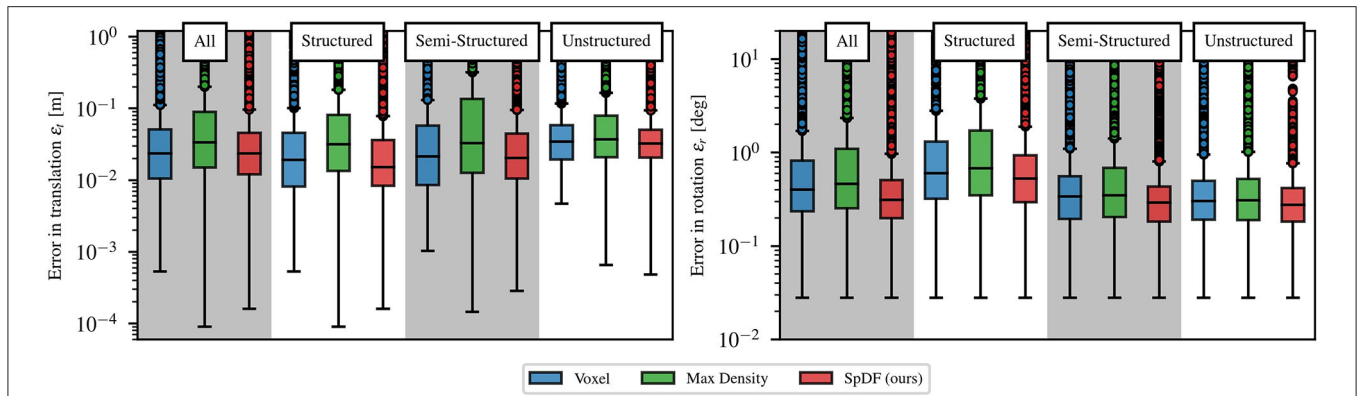


FIGURE 8 | Statistic analysis of the translation and rotation errors at high compression ratio between 85 and 99.9 % for each environment (structured, semi-structured, unstructured and all environments concatenated), for the top three methods, graphically represented as box plots. Medians are given with their first Q_1 and third Q_3 quartiles. The spacing between the different parts of the box indicate the degree of dispersion (spread) and skewness in the errors. Boxes are given with whiskers with maximum 1.5 IQR, where $IQR = Q_3 - Q_1$ is the inter-quartile range. All environments present similar errors, except in rotation for the structured environment which presents a slightly higher error. The spreads and medians are of the same order for the three methods.

similar dispersion of 0.3° . On the structured environment, the evaluated methods perform slightly worse, with an error of 0.6° with a larger spread of 0.8° . One explanation is that the latter environment usually presents a non-homogeneous distribution of the surfaces orientations (some orientations are preferred, e.g., the walls orientations) leading to less precise rotation estimation. One another explanation is that the structured environment is at a smaller scale than the others, which leads to a noisier estimation of the rotational component. Translation errors are of the same order for all the environments. We can expect an error of less than 3 cm with a 5 cm dispersion. The spread is smaller in the case of the unstructured environment than the other two. In particular, the spread of the MaxDensity is slightly higher than the others. Unsurprisingly, translation median errors are greater for the unstructured than the structured environment, as the *point-to-plane* cost function performs better in latter environments.

7.3. Computation Time Analysis

Computation time analysis has been conducted on an Intel[®] Core[™] i7-7820HQ CPU @ 2.90GHz, with 8 cores, on Ubuntu 16.04.3. Parallelization has been enabled when possible using then 8 threads. **Figure 9** provides a comparison of the computation time as function of the compression ratio, from 50 to 99.9%, for the following methods: Random, MaxDensity, Voxel, and SpDF. Voxel construction and the voting process have been parallelized. For each method, 2,500 samples have been used to sub-sample one scan per type of environment. The three types of environments are: structured, with 198 k points; semi-structured, with 134 k points; and unstructured, with 166 k points. The results are presented for all environments concatenated.

First, only the computation time of applying the filter has been measured, therefore not including the time required by the ICP process. As shown in **Figure 9**, the computation time of MaxDensity and Random is constant with respect to the

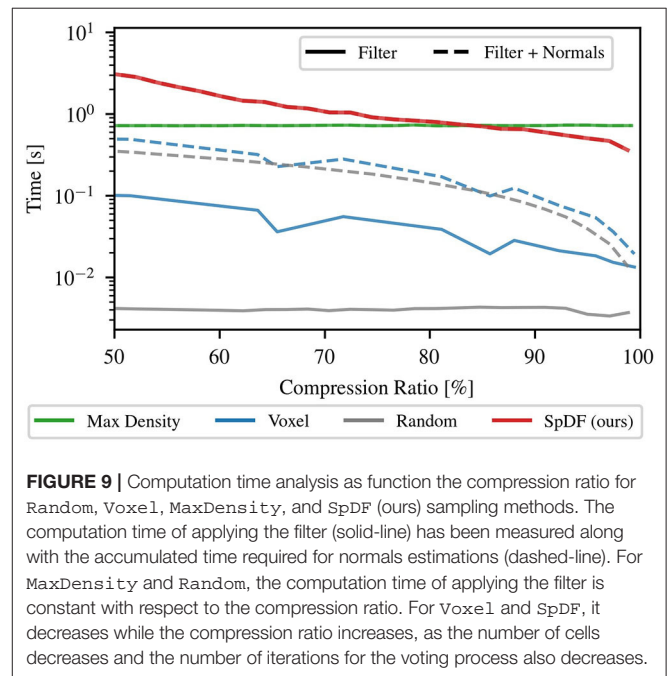


FIGURE 9 | Computation time analysis as function the compression ratio for Random, Voxel, MaxDensity, and SpDF (ours) sampling methods. The computation time of applying the filter (solid-line) has been measured along with the accumulated time required for normals estimations (dashed-line). For MaxDensity and Random, the computation time of applying the filter is constant with respect to the compression ratio. For Voxel and SpDF, it decreases while the compression ratio increases, as the number of cells decreases and the number of iterations for the voting process also decreases.

compression ratio. Random iterates only once on all the point cloud to decimate randomly the points, and MaxDensity computes the densities using the full point cloud, then randomly removes points with high density. For Voxel and SpDF, the computation time decreases while the compression ratio increases, as the number of cells decreases and the number of iterations for the voting process also decreases. Due to its iterative nature, SpDF is more costly and slower: for low compression ratios, one order of magnitude against MaxDensity, two order against Voxel, and three order against Random; for high compression ratios, MaxDensity is of the same order,

one order against `Voxel` and two order against `Random`. For instance, at a compression ratio of 95 %, for the semi-structured environment, we have the following computation times: $t_{\text{Random}} = 4.5$ ms, $t_{\text{Voxel}} = 20$ ms, $t_{\text{MaxDensity}} = 750$ ms, and $t_{\text{SpDF}} = 500$ ms. The evaluated strategies do only reduce the input point cloud, while `SpDF` is designed to reduce and enhance the point cloud with geometric information at the same time. Therefore, it is clear that the method has a higher computational time, but what we lose in computation time is gained in geometric information. As the computation time tends to the limit of real-time capabilities, efforts might be done to optimize the code and could be ported to GPU. Especially, porting the tensor voting and tensor decomposition parts on GPU, as done in Liu et al. (2012), provides significant improvement in computation time.

Secondly, considering the computational time of the whole pipeline of the ICP algorithm, we need to take into account the pre-processing time. In the case of the *point-to-plane* version it is required to compute the normals. **Figure 9** presents for each method the accumulated time needed for applying the filter and computing the normals. Normals estimation is conducted on the sampled point cloud. For `MaxDensity`, normals estimation is performed at the same time as the densities calculations, therefore there is no additional time. `SpDF` intrinsically computes the normals when applied, hence requiring no additional time either. For the other methods, the computation time is significantly increased for low compression ratio. At a compression ratio of 50 %, `Random` required a total computation time of 350 ms against 4.5 ms without normals estimation. The computation time of `Voxel` increases from 100 to 500 ms. The time required is now of the same order of magnitude as `MaxDensity`. The gap between `Voxel` and `SpDF` is now only of a factor 6. For high compression ratio, the difference is less significant. At a high compression ratio of 90 %, `Random` required a total computation time of 100 ms. The computation time of `Voxel` increases up to 200 ms, which is however still one order of magnitude quicker than `SpDF`. Note that when not enough points are available, the normals estimation is not robust and could hinder the registration process. `SpDF` leverages the tensor voting framework to provide a robust estimate of the normals.

7.4. Preserving Details and Making the Density Uniform on Real-World Large-Scale Data

Finally, we evaluate qualitatively `SpDF` on real-world large-scale datasets. **Figure 4** illustrates that `SpDF` is able to make the density uniform while preserving each geometric primitive, from an original point cloud of 370 k points where most points are concentrated in a small area, to a uniform point cloud of 40 k points, i.e., a compression ratio of 89 %. Edges and corners have been preserved, while dense surfaces have been made uniform.

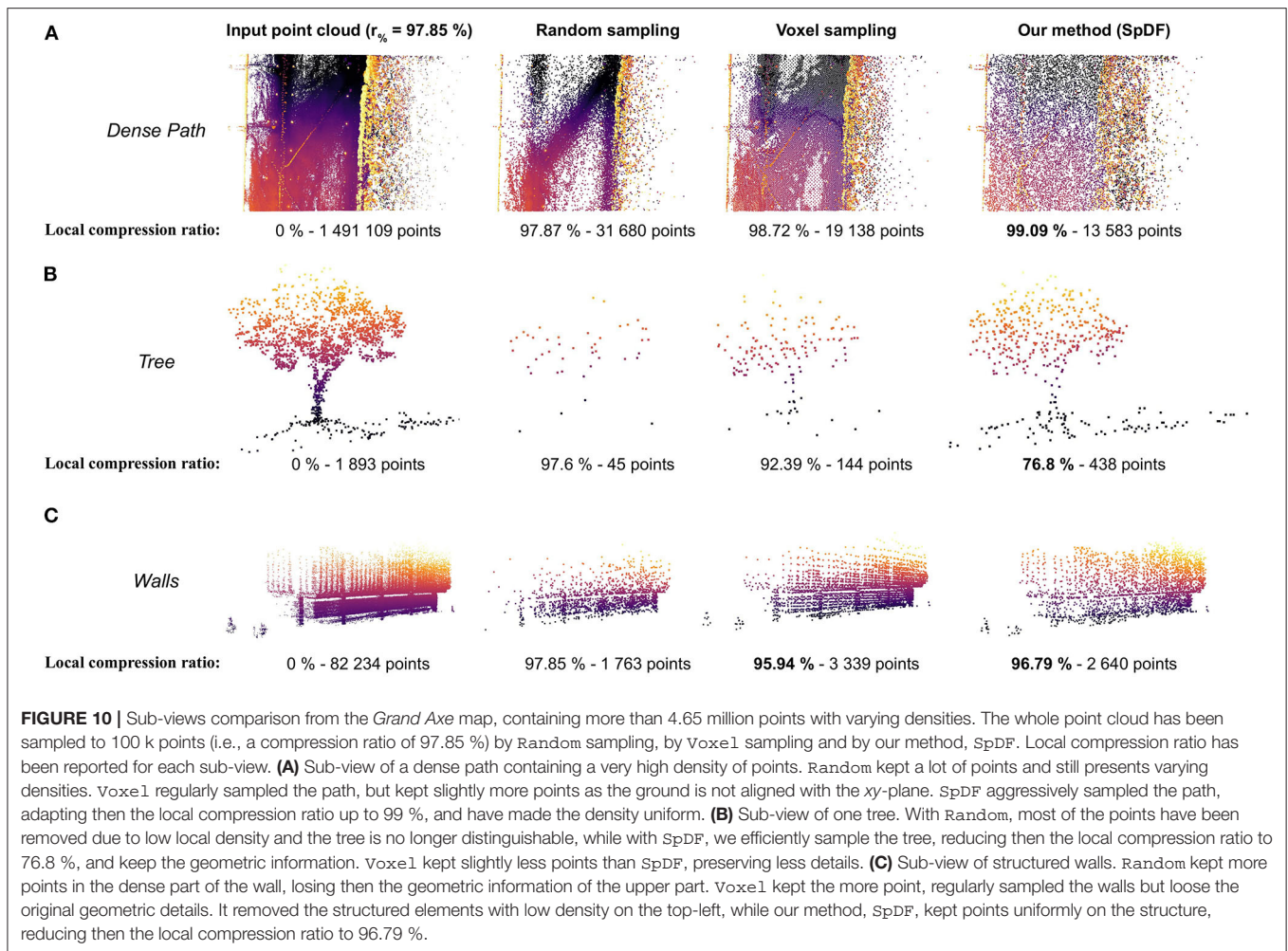
We evaluated the proposed method along with one of the top three methods, `Voxel` sampling with a spatial extent of 0.8 m on a real-world large-scale outdoor environment. **Figure 1** shows the qualitative result of a sampled point cloud from 4.65

million points to 100 k points, i.e., a compression ratio of 97.85 %. The map dimensions are approximately of $340 \times 100 \times 6$ m. **Figure 1B2** gives the full and top views of the map sampled by `Voxel`. The spatial sampling stops the voxel decomposition when the cell size is less than 0.8 m or when the cell contains less than 45 points. The decomposition is aligned with the *xyz*-basis. In this case, `Voxel` suffers from the discrepancy between the *x*, *y*, and *z* scales, as it creates the bounding box with a radius of the size of the bigger dimension. Furthermore, as the ground is not aligned with the *xy*-plane, it keeps points which are at different altitudes within the dense path. Finally, when geometric details are not dense enough such as the pillars of the top-left of the map, `Voxel` does not keep them. Contrarily, `SpDF` identifies the geometric primitives, and is able to sub-sample efficiently each primitive locally: the ground is uniformly sampled, and the pillars of the top-left of the map are preserved even if the density is low. From **Figure 1B1**, the full and top views of the map sampled by `SpDF` show that most of the details have been preserved and the density is uniform.

Figure 10 illustrates the effects of the proposed method on large-scale point cloud, presenting a comparison of sub-views from the *Grand Axe* map. Sub views have been extracted from the whole sampled point cloud by `Random` sampling, by `Voxel` sampling and by `SpDF`. Local compression ratio has been reported for each sub-view. **Figure 10A** presents a sub-view of a dense path containing a very high density of points. `Random` kept a lot of points and still presents varying densities. `Voxel` regularly sampled the path, but still kept slightly more points as the ground is not aligned with the *xy*-plane. `SpDF` aggressively sampled the path, adapting then the local compression ratio up to 99 %, and have made the density uniform. **Figure 10B** is a sub-view of one tree representing less than 0.04 % of the total point cloud. With `Random`, most of the points have been removed due to the low local density and the tree is no longer distinguishable. `Voxel` kept more points than `Random`, allowing to identify the tree within the point cloud, reducing the compression ratio to 92.39 %. However, less details have been preserved, and most of the ground have been removed. With `SpDF`, we efficiently sample the tree, reducing then the local compression ratio to 76.8 %, and keep most of the details. Finally, **Figure 10C** gives a sub-view of structured walls. `Random` kept more points in the dense part of the wall, then losing the geometric information of the upper part. It also removed the two small bushes next to the building, which are kept by the other methods. Even if `Voxel` sampled regularly the walls and kept more points, it does not differentiate the geometric primitives. For instance, structured elements have not been kept but replaced by regular samples, and the ones with a low density (on the left) have been removed. `SpDF` kept points uniformly on the structure, reducing then the local compression ratio to 96.79 %.

7.5. Discussion on the Choice of the Sampling Method

Our results show that sampling strategies designed for computer graphics applications cannot be extended to real-world large dataset. `CovS` and `NSS` cannot handle large scale map with noise,



while *SSNormal* and *Random* suffer from their inability to deal with varying densities.

Considering real-world large scale environments with spurious and noisy measurements, *Voxel*, *MaxDensity*, and *SpDF* can be used to sample the point cloud. In applications where computation time is critical, *Random* can still be considered as a strong solution. However, if the precision is critical, it is better to use *Voxel*, *MaxDensity*, or *SpDF* as they provide errors within the range accuracy of the used sensor. Here, in the context of large scale environments, it represents less than 2 cm in translation and less than 0.3 deg in rotation up to a compression ratio of 97 %. The *Voxel* method provides an efficient representation of the map by sampling regularly the point cloud space, with a computational time of less than 100 ms. However, one drawback of the spatial methods is the lack of fine grained control over the number of output points and over the level of details. In applications where geometric information is required, such as surface inspection scenario, *Voxel* does not guarantee to preserve the geometric details as it averages the points within each voxel. Density-based methods, such as *MaxDensity* are able to preserve further geometric information as they work locally. However,

MaxDensity does not differentiate geometric primitive during the density calculation and relies on spherical approximation. Our qualitative experiments show that *SpDF* by sampling locally on each geometric primitive further preserves the details and the topology of the point cloud. The major drawback of the proposed method is its high computational time at the limit of real-time capabilities, but does not required a pre-processing step. It provides a higher level description in terms of geometric primitives that could be used by another pipeline for scene interpretation, or directly in the minimization process as normals are provided in the *point-to-plane* version.

The general registration strategy selected for an application may require to balance advantages and drawbacks of the chosen solutions. Readers are invited to refer to Pomerleau et al. (2015) as they provide a clear overview of geometric registration in case of robotics applications, as well as a method to solve it in specific instances. Typical use-cases for ICP do not need to apply the filter at a high rate on all input scans. Most of the strategies rely on scan-to-map registration, where the maintained map is sup-sampled by an efficient sampling strategy according to the desired application. For instance, in case of surface inspection, we want to preserve geometric information, we can

then use the proposed method in this paper. At 10 Hz, input scans should use Random to maintain high rates. Reducing the map size increases all registration done to it and decreases data transmission bandwidth when it comes to sharing the map.

8. CONCLUSION

This paper presents a novel sampling algorithm, named Spectral Decomposition Filter (S_{PDF}), aiming at reducing the number of points while preserving the geometric information along the topology of large-scale point cloud with non-uniform density, large sensor noise, and spurious measurements. This method builds on spectral decomposition applied to point clouds in order to obtain a density better suited for robotics applications where geometric information are essentials. First, we identify the geometric primitives along with their saliencies using the tensor voting framework from the input point cloud. Then, we derive density measures from saliencies: if the density for each geometric primitive is less than the desired density, we stop; else we sub-sample each geometric primitive, and re-iterate. As output, we have a uniform sampled point cloud enhanced with geometric information.

We verified the feasibility of our method through quantitative and qualitative results. We presented a large-scale evaluation of S_{PDF} along with other seven point cloud sampling strategies from the state-of-the-art, in the context of the 3D registration problem using the ICP algorithm. Our results show that only spatial sampling strategies and density-based methods are able to manage such large 3D environments without hindering the registration process. In particular, S_{PDF} performs successfully on large-scale maps acquired with lidar sensor, where the density is non-uniform. We manage non-uniform densities by leveraging the new derived measures of density from saliencies for each geometric primitive which allows us to preserve the topology of the point cloud. Thus, by making the density uniform and leveraging the geometric information, the proposed method efficiently sub-samples large scale point cloud while simultaneously enhancing it. Even for high compression ratio (i.e., > 90 %), the process of registration is not hindered which enable a large spectrum of robotics applications. Indeed, for such applications, this sampling is usually critical to reduce bandwidth or computation complexity without losing accuracy during the registration process, whilst geometric information provides higher level of description.

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Computational time is however still high and at the limit of real-time capabilities. Future works will include efforts to optimize the code and the tensor voting part will be ported on GPU which should significantly improve the computation time (Liu et al., 2012). Furthermore, as we provide a second order symmetric tensor representation for each point (i.e., a Gaussian representation), future works will aim to leverage this information directly in the minimization process of the ICP, inspired by the *Point-to-Gaussian* cost function derived by Babin et al. (2019).

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

ML: conceptualization, methodology, software development, validation, formal analysis, visualization, and writing. JL: methodology, formal analysis, visualization, and writing. FP: supervision, review, and project administration. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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