Stochastic Poisson Surface Reconstruction

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We introduce a statistical extension of the classic Poisson Surface Reconstruction algorithm for recovering shapes from 3D point clouds. Instead of outputting an implicit function, we represent the reconstructed shape as a modified Gaussian Process, which allows us to conduct statistical queries (e.g., the likelihood of a point in space being on the surface or inside a solid). We show that this perspective: improves PSR’s integration into the online scanning process, broadens its application realm, and opens the door to other lines of research such as applying task-specific priors.

CCS Concepts: • Computing methodologies → Point-based models; Volumetric models; Reconstruction.

Additional Key Words and Phrases: surface reconstruction, uncertain geometry processing, Gaussian processes

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1 INTRODUCTION

Surface reconstruction refers to the process of converting a point cloud (the most common real-world raw 3D capture format) into another shape representation, such as a mesh or an implicit function, for use in downstream applications. This is an underdetermined process filled with uncertainty, not just due to a point cloud’s discrete nature and lack of topological information but also because of real-world challenges like scan occlusions or measurement error.

The de facto standard geometry processing algorithm for this task is Poisson Surface Reconstruction (PSR) [Kazhdan et al. 2006], which solves a partial differential equation to reconstruct a function \( f_{PSR} \) whose zero levelset \( f_{PSR} = 0 \) defines the desired surface. Due to its speed, quality and simplicity, PSR remains relevant and has seen uses in fields as varied as digital heritage preservation [Andrade et al. 2012], topography [Gupta and Shukla 2017], medicine [Palomar et al. 2016] and autonomous driving [Vizzo et al. 2021].

Unfortunately, PSR lacks the statistical formalism to quantify the uncertainties of the surface reconstruction process. The magnitudes of \( f_{PSR} \) outside of the zero levelset are arbitrary (see Fig. 2) and \( f_{PSR} \)

Fig. 2. PSR values outside of zero are arbitrary (e.g., they depend on grid size, sampling rate and scale) and contain no direct statistical information.
alone cannot provide an answer to statistical questions crucial to the reconstruction process like “how confident can one be of the values of \( f_{\text{PSR}} \)?” or “where should one aim the scanner next to optimize information gain?” Similarly, it cannot respond to queries like “what is the probability of a point \( p \) being contained in the shape?”, critical for collision detection or ray casting applications.

In this paper, we introduce Stochastic Poisson Surface Reconstruction (SPSR), a statistical derivation of PSR as conditional probability distributions in a Gaussian Process (GP). Instead of just an implicit function value, we endow every point in space with a Gaussian distribution of possible values. We propose an algorithm to compute the mean and variance that fully determine this distribution (see Fig. 1), allowing us to answer any statistical queries.

This extension vastly broadens the use cases of Poisson Surface Reconstruction, as we highlight with prototypical examples of surface point cloud repair, ray casting, next-view planning and collision detection. Furthermore, we show that by understanding PSR from this new perspective, we can borrow from the Gaussian Process literature to modify it by incorporating task-specific priors, opening several promising lines of future research.

2 RELATED WORK

A complete survey of vast research areas like uncertainty quantification or surface reconstruction is beyond the scope of this work. Instead, we focus this section on setting our Stochastic PSR in its context of bridging the gap between PSR and Gaussian Processes.

2.1 Surface Reconstruction from Point Clouds

Point clouds are a common raw format for 3D geometry acquired form the real world. However, most applications in fields like rendering, simulation and geometry processing require more structured representations like triangle meshes. Thus, reconstructing surfaces from point clouds is a well-studied, fundamental problem in Computer Graphics (see [Berger et al. 2017] for an exhaustive survey). While some methods convert point clouds directly to meshes (e.g., by dictionary learning [Xiong et al. 2014]) or simple primitives (e.g., [Monszpart et al. 2015; Nan et al. 2010]), we focus on those that extract the shape as the zero levelset of a reconstructed function \( f \).

Within these, a common separation is made between local and global algorithms. Local algorithms prioritize performance in speed and memory; for example, by fitting linear [Hoppe et al. 1992], polynomial [Alexa et al. 2001] or higher-order [Fuhrmann and Goesele 2014] functions to restricted point subsets. By their nature, these are more susceptible to oscillations far from the sample points. To cope, global algorithms (e.g., [Jacobson et al. 2013]) allow \( f \) to be influenced by every point in the cloud (hierarchical fast summation structures can help reduce computation [Barill et al. 2018]).

Poisson Surface Reconstruction (PSR) [Kazhdan et al. 2006] captures the best features of the global (robustness) and local (performance) methods by computing \( f \) in two steps. First, a vector field \( \tilde{V} \) is interpolated from the point cloud using only local information. Then, \( f \) is obtained from \( \tilde{V} \) via a global sparse PDE solve, which is discretized as a linear system and can be solved very efficiently using an adaptative grid structure. PSR has been improved since its publication; for example, Kazhdan and Hoppe [2013] combine both steps to improve noise robustness, Kazhdan et al. [2020] include envelope constraints and Peng et al. [2021] formulate the Poisson solve in a differentiable way. Nonetheless, the original 2006 publication is still one of the few showing robustness in every metric considered by Berger et al. [2017] more than a decade later.

Our Stochastic PSR inherits all the benefits of the original PSR, supplying it with a complete statistical formalism that extends it and its application realm (see Fig. 3). Our contributions are orthogonal to the specific grid structure used (see [Kazhdan and Hoppe 2019]).

Our algorithm can output the probability of any point in space being inside the sampled domain (see Fig. 1). While this resembles the shape representations proposed by occupancy networks [Mescheder et al. 2019] and neural radiance fields [Mildenhall et al. 2020], we note that our algorithm produces this quantity as a direct byproduct of a fully determined statistical distribution which can answer many other statistical queries, like boundary probabilities (see Fig. 5) or regional probabilities (see Fig. 12).

2.2 Gaussian Processes

A Gaussian Process (GP) is an infinite collection of joint normal distributions [Doob 1944; Dudley 2018], usually parametrized by a continuous parameter like time or space [Kac and Siegert 1947]. Recently, Gaussian processes have been used as a tool in unsupervised
Our Input PSR vector field whose parameters minimize a chosen distribution loss. (see model for their individual uncertainty to weigh their contributions [Bosse and Zlot 2008; Landry et al. 2019] and, most recently, for cloud registration. Specifically, a lot of work has been dedicated to the uncertainty of a regression model, a field which has seen significant growth since recent advances in deep learning (see [Abding et al. 2005; Raissi et al. 2017] for examples), even suggested it has been grasping [Dragiev et al. 2011], next view planning [Hollinger et al. 2007] similarly measure topological reconstruction uncertainty to identify where user-provided disambiguation is most needed.

Closer to our application, Curless and Levoy [1996] compute histograms in logarithmic scale. Closer to our application are Gaussian Process Implicit Surfaces (GPIS) [Williams and Fitzgibbon 2006]. These algorithms exchange the function that implicitly defines a surface for a Gaussian distribution \( f(x) \sim N(\mu(x), \sigma(x)) \), and compute \( \mu(x) \) and \( \sigma(x) \) by studying the posterior GP distribution given observed points. One can recover a surface by extracting the zero levelset of \( \mu \); however, the information contained in \( \sigma \) also finds use in tasks like robotic grasping [Dragiev et al. 2011], next view planning [Hollinger et al. 2012] and segmentation [Ramon Soria et al. 2017; Shin et al. 2017].

GPIS present the same global/local dilemma as other surface reconstruction algorithms. If the interpolation is done using all the point cloud information (this is encoded in the support of the GP covariance function), the method suffers in performance; if not, in robustness. By using the insights of Poisson Surface Reconstruction, our Stochastic PSR instead uses a local Gaussian Process to build the distribution of the gradient field \( \nabla f(x) \), and then recover the full distribution of \( f(x) \) with a global PDE solve. While using a GP to interpolate vector-valued data is unorthodox, we note it has been done before; e.g., for fluid velocity information [Lee et al. 2019].

Gaussian processes are a more traditional approach to quantifying the uncertainty of a regression model, a field which has seen significant growth since recent advances in deep learning (see [Abdar et al. 2021] for a survey). Often, the posterior distribution of a given neural network is approximated by another Bayesian network whose parameters minimize a chosen distribution loss. (see e.g., [Xue et al. 2019]). Alternatively, the posterior may be learned directly from the observed data (see e.g., [Shen et al. 2021]).

### 2.3 Stochastic Geometry Processing

Our algorithm stands among many similar works that add statistical formalism to standard geometry processing techniques, like point cloud registration. Specifically, a lot of work has been dedicated to computing the covariances in the pairwise iterative closest point [Bosse and Zlot 2008; Landry et al. 2019] and, most recently, for general multi-scan registration [Cao et al. 2018; Huang et al. 2020].

Closer to our application, Curless and Levoy [1996] compute truncated signed distance fields for each sensor position and use a model for their individual uncertainty to weigh their contributions to the final implicit reconstruction. Pöthkow et al. [2011] model grid samples of an implicit function as normal distributions and calculate the individual probabilities of each voxel marching cubes configuration to quantify the contouring uncertainty. Sharf et al. [2007] similarly measure topological reconstruction uncertainty to identify where user-provided disambiguation is most needed.

Pauyl et al. [2004] work is the most similar to ours. It takes a point cloud as input and uses a surface smoothness prior to compute the likelihood of any point in space lying on it (see Fig. 5, left). Instead, our proposed algorithm assumes samples to lay on the boundary of a solid, and impose solid smoothness prior (examined further in Section 5). Further, while our algorithm can also output a surface likelihood quantity (see Fig. 5, center), it is only part of a full statistical distribution of the solid (see Fig. 5, right).

### 3 BACKGROUND

By posing PSR as a Gaussian Process, our work combines these two concepts. We begin by reviewing them individually.

#### 3.1 Gaussian Processes

Intuitively, a Gaussian Process is an extension of the multivariate normal distribution to an infinite number of dimensions. Formally, let \( \mathcal{A} = \{ A(x) \}_{x \in \Omega} \) be a collection of random variables parametrized by some continuous parameter \( x \). \( \mathcal{A} \) is said to be a Gaussian Process if any finite subset of \( \mathcal{A} \) follows a multivariate Gaussian distribution. Equivalently, \( \mathcal{A} \) is a Gaussian Process if for any two \( x, x' \in \Omega \),

\[
A(x), A(x') \sim \mathcal{N}
\begin{pmatrix}
  m(x) \\
  m(x')
\end{pmatrix}
\begin{pmatrix}
  k(x, x) & k(x, x') \\
  k(x', x) & k(x', x')
\end{pmatrix}
\]

(1)

for some mean and covariance functions \( m : \Omega \rightarrow \mathbb{R}, k : \Omega \times \Omega \rightarrow \mathbb{R} \). These two functions uniquely determine the Gaussian Process \( \mathcal{A} \).

Gaussian Processes are a particularly useful tool for supervised learning tasks (see Fig. 4). Assume training observations \( T = \{ (x_i, a_i) \}_{i=1}^n \), where each \( a_i \) is an observation of the distribution \( A(x_i) \), and consider an unobserved (test) point \( x \). By definition of GP, the joint distribution of \( \{ A(x), A(x_1), \ldots, A(x_n) \} \) is

\[
\mathcal{N}
\begin{pmatrix}
  m(x) \\
  m(x_1) \\
  \vdots \\
  m(x_n)
\end{pmatrix}
\begin{pmatrix}
  k(x, x) & k(x, x_1) & \cdots & k(x, x_n) \\
  k(x_1, x) & k(x_1, x_1) & \cdots & k(x_1, x_n) \\
  \vdots & \vdots & \ddots & \vdots \\
  k(x_n, x) & k(x_n, x_1) & \cdots & k(x_n, x_n)
\end{pmatrix}
\]

(2)

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which we write as

$$A(x), A(x_1), \ldots, A(x_n) \sim \mathcal{N}
\begin{pmatrix}
\mu_1 \\
\mu_2 \\
\mu_3
\end{pmatrix},

\begin{pmatrix}
k_1 & k_2 & k_3 \\
k_2 & k_4 & k_5 \\
k_3 & k_5 & k_6
\end{pmatrix},
$$

(3)

where it is relevant to note that $K_3$ depends only on the training data and $k_2$ depends on both training and test sets. By Bayes’ theorem, this means the distribution of $A(x)$ conditioned on the observations $\{(x_1, a_1)\}_{n=1}^N$ is

$$A(x) | T \sim \mathcal{N}(m_1 + k_2^T k_3^{-1}(a - m_2), k_1 - k_2^T k_3^{-1}k_2)
$$

(4)

One usually assumes $m = 0$ (otherwise, the Gaussian process $\mathcal{A} - m$ is considered), and writes

$$A(x) | T \sim \mathcal{N}(k_2^T k_3^{-1}a, k_1 - k_2^T k_3^{-1}k_2).
$$

(5)

In the case of noisy observations, a term $\sigma^2_n I$ (where $\sigma^2_n$ is the noise variance) is added to $K_3$.

3.2 Poisson Surface Reconstruction

Any input oriented point cloud of the surface of a solid shape $\Omega$ can be written as a set of observations $s \in S$, each of them storing a position $p_s$ and a (normalized) orientation $\mathbf{N}_s$. Poisson Surface Reconstruction [Kazhdan et al. 2006] aims to build a function $f_{PSR} : \mathbb{R}^3 \rightarrow \mathbb{R}$ which takes positive values inside $\Omega$ and negative values outside of it, thus making the zero levelset $f_{PSR} = 0$ the reconstructed surface $\partial \Omega$.

PSR begins by building a grid $O$. Then, they define the following vector field for any $q \in \mathbb{R}^3$ by using the grid structure to interpolate the orientation observations:

$$\tilde{V}_{PSR}(q) = \sum_{s \in S} \frac{1}{W(p_s)} \sum_{o \in B(s)} a_{o,p_s} F_o(o) \mathbf{N}_s
$$

(6)

where $B(s)$ are the eight closest grid nodes to $s$, $a_{o,p_s}$ is the trilinear interpolation weight for $p_s$ at $o$, $W$ is a measure of volumetric sampling density and $F_o$ is a compactly-supported approximation of a Gaussian kernel centered at the $o$-th node.

The fundamental observation of PSR is that once $\tilde{V}$ has been constructed, the desired implicit function $f$ satisfies

$$\Delta f_{PSR}(x) = \nabla \cdot \tilde{V}_{PSR}(x), \quad \forall x \in O
$$

(7)

Eq. (7) is underdetermined, as wildly different functions can have the same Laplacian. This ambiguity is resolved in part during discretization. Kazhdan et al. [2006] suggest building $O$ as an adaptive grid of a bounding box of the point cloud. Then, they use the finite element method to build discrete Laplacian and divergence operators $L$ and $Z$ and solve

$$Lf_{PSR} = Z \tilde{V}_{PSR}.
$$

(8)

This discretization imposes zero Neumann $\nabla f \cdot \tilde{n} = 0$ boundary conditions on the boundary of $O$. Even so, Eq. (8) only determines $f$ up to translation. To account for this, one valid $f_{PSR}$ is computed and then shifted so its values near the observation points are zero on average. The solution $f_{PSR}$ contains the implicit function $f_{PSR}$ evaluated at each grid node, and its zero levelset can be extracted using contouring algorithms like Marching Cubes [Lorensen and Cline 1987] or Dual Contouring [Ju et al. 2002].

4 STOCHASTIC POISSON SURFACE RECONSTRUCTION

We introduce our Stochastic PSR, which extends the traditional PSR from Kazhdan et al. [2006] with the statistical formalism of a Gaussian Process. We begin by defining the PSR semicovariance as

$$k_{PSR}(x, y) = \sigma_g \sum_{o \in B(x)} a_{o,x} F_o(y),
$$

(9)

where $\sigma_g$ is a scalar parameter. Then, the vector field interpolation in Eq. (6) can be written as

$$\tilde{V}_{PSR}(q) = \sum_{s \in S} k_{PSR}(p_s, q) \frac{1}{\sigma_g W(p_s)} \mathbf{N}_s.
$$

(10)

By term identification, it may be tempting to interpret this interpolation directly as the posterior mean of a Gaussian Process $k_{PSR}(p_s, q)$ (see Eq. (5)), with normals as the observed values $\mathbf{N}_s$, $\sigma_g$ as the variance, $W(p_s)$ as the entries of $K_2$ and division by $\sigma_g W(p_s)$ as the role of multiplication by $K_2^{-1}$. However, a valid Gaussian distribution must have a symmetric covariance, and $k_{PSR}(p_s, q) \neq k_{PSR}(q, p_s)$ in general (hence the semi in “PSR semicovariance”). To circumvent this, we define the Stochastic PSR covariance to be its symmetrized version

$$k_{SPSR}(x, y) = k_{SPSR}(y, x) = \frac{1}{2} (k_{PSR}(x, y) + k_{PSR}(y, x)),
$$

(11)

which analogously defines a Stochastic PSR vector field

$$\tilde{V}_{SPSR}(q) = \sum_{s \in S} k_{SPSR}(p_s, q) \frac{1}{\sigma_g W(p_s)} \mathbf{N}_s.
$$

(12)

This variance symmetrization will be our only deviation from the traditional Poisson Surface Reconstruction by Kazhdan et al. [2006] (see Fig. 8). Strictly speaking, all the observations that follow in this paper can only be said to apply to $V_{SPSR}$; however, note that $V_{PSR}$ and $V_{SPSR}$ are visually indistinguishable (see Fig. 7) and their difference vanishes under refinement (see proof in Appendix A).
assumptions of a Gaussian Process with \( m = 0 \) and covariance \( k_{\text{SPSR}} \). Indeed, one needs only to exchange \( A \) for the vector-valued \( \tilde{\mathbf{v}} \) and the observations \( \{(x_i, a_i)\} \) for \( \{(p_o, \bar{N}_i)\} \) in Eq. (5) to obtain the conditional probability distribution

\[
\tilde{\mathbf{v}}(q) | S \sim \mathcal{N}(k_1^T \mathbf{K}_3^{-1} \bar{\mathbf{N}}_s, k_1 - k_2^T \mathbf{K}_3^{-1} k_2)
\]  

where

\[
k_1 = k(q, q), \quad k_2 = (k(q, p_o))_{q \in S} \in \mathbb{R}^{|S|},
\]

\[
\mathbf{K}_3 = (k(p_o, p_{o'}))_{o,o' \in S} \in \mathbb{R}^{|S| \times |S|}.
\]  

The main computational cost in computing the mean and variance of the conditional distribution is the inversion of \( \mathbf{K}_3 \), a matrix whose size scales with the number of points in the cloud. One could avoid this by assuming that the samples \( S \) are independent, which would result in approximating \( \mathbf{K}_3 \) by \( \sigma^2 I \). However, completely discarding the sample interdependency could have adverse effects: in practice, it would make densely sampled regions have an outsized effect over more sparsely sampled ones (see Fig. 9, center right).

We introduce a better approximation: we assume each sample is independent, but with variance proportional to sampling density. Intuitively, this means we account for the interdependence by trusting each individual sample in a densely sampled region less than in a sparsely sampled one (see Fig. 9, right). We justify this choice mathematically in the general GP context in Appendix B.

We are not aware of any previous work that proposes this approximation, which addresses one of the main performance limitations of general Gaussian Processes. Numerically, this resembles the mass matrix lumping step often carried out in the Finite Element literature (see e.g., [Zienkiewicz et al. 2005] Chap. 16.2.4.), so we call it the (diagonal) lumped covariance matrix

\[
\mathbf{D} := \text{diag}(\sigma_g w) \approx \mathbf{K}_3
\]  

where \( w \) is the local sampling density at each sample point, which we compute as described by Kazhdan et al. [2006]. As is usual with Gaussian processes, we sum \( \sigma_g I \) if the sampled point cloud is assumed to contain noise with variance \( \sigma_g \) in the normal vector.

Then, the conditional distribution becomes

\[
\tilde{\mathbf{v}}(q) | S \sim \mathcal{N}(k_1^T \mathbf{D}^{-1} \bar{\mathbf{N}}_s, k_1 - k_2^T \mathbf{D}^{-1} k_2)
\]  

Importantly, note that the mean of this distribution is exactly the Stochastic PSR vector field we introduced in Eq. (12), i.e.,

\[
\tilde{\mathbf{v}}(q) | S \sim \mathcal{N}(\tilde{\mathbf{v}}_{\text{SPSR}}(q), k_1 - k_2^T \mathbf{D}^{-1} k_2).
\]  

In other words, this critical reinterpretation of PSR has allowed us to extend the interpolated PSR vector field into a complete statistical distribution whose mean \( \tilde{\mathbf{v}}_{\text{SPSR}} \) is nearly identical to \( \tilde{\mathbf{v}}_{\text{PSR}} \) (see Fig. 7, which we can now understand as comparing the traditional PSR vector field to the mean of our lumped Gaussian Process).

Ideally, however, we would want a similar statistical formalism not for \( \tilde{\mathbf{v}} \) but for the implicit function \( f \), which would allow us to formulate statistical queries meaningful to the reconstruction task, e.g., \( P(f(q) < 0) \).

Fortunately, transferring our statistical understanding of \( \tilde{\mathbf{v}} \) to \( f \) is just a matter of Gaussian arithmetic. First, note that a reasoning identical to the one above leads to the joint conditional probability of \( \tilde{\mathbf{v}} \) at all grid nodes being

\[
\tilde{\mathbf{v}}(o_1), \ldots, \tilde{\mathbf{v}}(o_3) | S \sim \mathcal{N}(\mathbf{V}_{\text{SPSR}}, \mathbf{K}_1 - \mathbf{K}_2^T \mathbf{D}^{-1} \mathbf{K}_2),
\]  

where

\[
\mathbf{K}_1 = (k(o, o')) \in \mathbb{R}^{|O|\times|O|}, \quad \mathbf{K}_2 = (k(o, p_o)) \in \mathbb{R}^{|O| \times |S|}
\]  

and

\[
\mathbf{V}_{\text{SPSR}} = (\mathbf{V}_{\text{SPSR}}(o) \in \mathbb{R}^{|O| \times 3}.
\]  

For simplicity, denote \( \mathbf{K}_V = \mathbf{K}_1 - \mathbf{K}_2^T \mathbf{D}^{-1} \mathbf{K}_2 \), and let \( \mathbf{v} \) be the concatenated vector of \( \tilde{\mathbf{v}} \) values as in the previous section. Then, Eq. (19) becomes

\[
\mathbf{v} | S \sim \mathcal{N}(\mathbf{V}_{\text{SPSR}}, \mathbf{K}_V).
\]  

By linearity and bilinearity of the mean and covariance, respectively, we know

\[
\mathbf{Z} \mathbf{v} | S \sim \mathcal{N}(\mathbf{Z} \mathbf{V}_{\text{SPSR}}, \mathbf{Z} \mathbf{K}_V \mathbf{Z}^T)
\]  

and thus, since the vector \( f \) of evaluations of \( f \) satisfies \( \mathbf{L} f = \mathbf{Z} \mathbf{v} \),

\[
f | S \sim \mathcal{N}(\mathbf{L}^{-1} \mathbf{Z} \mathbf{V}_{\text{SPSR}}, \mathbf{L}^{-1} \mathbf{Z} \mathbf{K}_V \mathbf{Z}^T (\mathbf{L}^T)^{-1}).
\]  

Similarly to the traditional PSR, we shift the mean values to be zero near the sample points and shift the variance values to have a minimum diagonal entry of zero. We write the above distribution as

\[
f | S \sim \mathcal{N}(\mathbf{f}_{\text{SPSR}}, \mathbf{K}_f).
\]  

This statement is our principal contribution, as it extends the traditional PSR implicit function \( f_{\text{PSR}} \) into a full statistical distribution whose mean is almost identical to \( f_{\text{PSR}} \). Of all the information contained in \( \mathbf{K}_f \), a particularly useful quantity is its diagonal \( \sigma_f^2 \), which contains the variances of each entry of \( f \) (see Fig. 3).
4.1 Statistical queries

Eq. (25) contains all the information we need to respond to statistical queries fundamental to the reconstruction process; e.g., the probability of a given point being inside the shape (see Fig. 10)

\[
p(\alpha_i \in \Omega) = p(f_i \leq 0) = \text{CDF}_{\text{SPSR}, \alpha_i}(0) \tag{26}
\]

or the probability density of being on the surface (see Fig. 13)

\[
p(\alpha_i \in \Omega) = p(f_i = 0) = \text{PDF}_{\text{SPSR}, \alpha_i}(0) \tag{27}
\]

where \(CDF_{\mu,\sigma^2}\) and \(PDF_{\mu,\sigma^2}\) are the cumulative distribution and probability density functions, respectively, of a Gaussian distribution with mean \(\mu\) and variance \(\sigma^2\). Similar Gaussian arithmetic lets us compute other quantities like confidence intervals (see Fig. 14).

All these queries would be impossible to answer from PSR’s \(f_{\text{PSR}}\) alone. We show this in the didactic example in Fig. 6, where vectors pointing upward are sampled from a normal distribution centered at the middle horizontal line. As more samples get added, \(f_{\text{PSR}}\) converges to a smooth transition between positive and negative values; however, our variances are progressively decreasing, making \(P(x \in \Omega)\) converge in certainty as expected.

We formalize this observation further by defining an integrated statistical quantity, the total uncertainty \(U_{\text{SPSR}}\), which we define as

\[
U_{\text{SPSR}} = \int_B (0.5 - |P(x \in \Omega) - 0.5|) \, dx \tag{28}
\]

where \(B\) is a bounding box around the point cloud. \(U_{\text{SPSR}}\) can be intuitively interpreted as a global measure of reconstruction quality which converges to zero when the surface has been reconstructed with absolute certainty (see Fig. 15). We pose that this quantity can be used as a stopping criteria to guide the scanning process (see Fig. 14).

The off-diagonal entries of \(K_f\) quantify the correlation between the values of \(f\) at different points in \(O\). This allows for joint probability queries; for example, computing the likelihood of a whole region in space \(R\) intersecting \(\Omega\) (see Fig. 12), a quantity of immediate relevance to collision detection applications.

To compute it, we can randomly sample \(R_k = \{r_1, \ldots, r_s\} \subset R\). If \(W\) is the linear interpolation matrix from \(O\) to \(R_k\), then distribution arithmetic says

\[
f(r_1, \ldots, r_s) | S \sim N(Wf_{\text{SPSR}}, W^T K_f W). \tag{29}
\]

4.2 Space reduction with eigenspace analysis

All this additional information comes at a computational cost. Specifically, the main performance bottleneck is the computation of the covariance matrix

\[
K_f = L^{-1} Z K_y Z^T (L^T)^{-1}, \tag{30}
\]

a matrix equation that amounts to solving \(2 \times |O|\) linear systems. Even making use of efficient prefactorizations of \(L\), this step is impracticable for large 3D grids with sizes in the millions of cells.

To circumvent this, we will work in the reduced space of Laplacian eigenvectors. For a cuboid with lengths \(\ell_1, \ell_2, \ell_3\) and zero Neumann boundary conditions, these are known [Gottlieb 1985] to respond to the analytic function

\[
\psi_{M,N,N}(x, y, z) = \cos \left( \frac{M \pi x}{\ell_1} \right) \cos \left( \frac{N \pi y}{\ell_2} \right) \cos \left( \frac{N \pi z}{\ell_3} \right), \tag{31}
\]

with associated eigenvalues

\[
\lambda_{M,N,N} = \pi^2 \left( \frac{M^2}{\ell_1} + \frac{N^2}{\ell_2} + \frac{N^2}{\ell_3} \right). \tag{32}
\]
Specifically, we will evaluate these analytical expressions into $E \in \mathbb{R}^{|O| \times k}$, the matrix of the $k$ lowest-magnitude eigenvectors of $L$, and $D_e$, the diagonal eigenvalue matrix. Then, we approximate $K_f$ by projecting $ZK_fZ^\top$ to the eigenspace, solving the matrix equation in this reduced space, and reprojecting; i.e.,

$$K_f \approx ED_e^{-1}E^\top (ZK_fZ^\top) ED_e^{-1}E^\top. \tag{33}$$

In all our examples, we fix $k = 3000$.

Due to the dimensionality of the above matrices, the mostly computationally expensive step in Eq. (33) is the product $ECE^\top$, where $C \in \mathbb{R}^{k \times k}$ results from multiplying all middle matrices in Eq. (33). We avoid this step by noting that it is rare that one is interested in all entries of $K_f$. As seen in Section 4.1, most statistical queries require only the diagonal $\sigma^2_f$, which we can compute directly and efficiently as $(EC) \cdot E$, where $\cdot$ denotes row-wise dot product. In the case of joint probability queries that require off-diagonal knowledge of $K_f$, we first construct the selection matrices $S, S^\top$ that extract the covariance rows and columns relevant to the specific application, and directly compute $E'C'E^\top$, where $E' = SE$. Thus, we eventually avoid ever computing (or storing) the full $K_f \in \mathbb{R}^{|O| \times |O|}$.

Dimension reduction has been used for implicit reconstruction before; e.g., the Fourier coefficients computed by Kazhdan [2005]. We limit our approximation to the covariance computation only.

### 4.3 Adaptive discretization

The statistical formalism we contribute is agnostic to where the values of $f$ and $\tilde{V}$ are stored and to the discretization scheme used to solve the Poisson equation. Kazhdan et al. [2006] suggest building an adaptive grid which is finer near the sampled point cloud. This choice is justified through their sole goal of accurately recovering the zero levelset of $f$.

Our Stochastic PSR extends the traditional PSR by providing volumetric statistical information, which is also relevant and non-trivial in regions away from the zero levelset of our mean function $f_{SPSR}$ or the point cloud (see, e.g., the probability in Fig. 10). Further, computing statistical queries accurately away from the mean-zero levelset can be critical for applications that our formalism newly allows like collision detection or ray casting (see Fig. 24). Thus, the choice of an adaptive grid structure is less obvious for our algorithm. Nonetheless, we show in Fig. 16 our algorithm working as expected on a graded quadtree grid structure, following the finite difference discretization proposed by Bickel et al. [2006]. While orthogonal to our contribution, we believe exploring different adaptive discretization strategies (e.g., narrowing in on regions of higher uncertainty) to be a promising avenue for future work.

## 5 BEYOND POISSON SURFACE RECONSTRUCTION

A significant benefit of our interpretation of Poisson Surface Reconstruction as a Gaussian Process is that we may borrow from the vast literature on the latter to study variations of the traditional PSR. A prototypical example of this is incorporating task-specific priors.

In Section 3.1, we mentioned that most Gaussian Process consider $m = 0$ and are thus fully determined by the covariance $k$ (this is often known as kriging in the GP literature). Indeed, it was by assuming $m = 0$ that we recovered the PSR vector field interpolation as the mean of the GP posterior distribution. In fact, the choice of $m$ is a prior imposed on the joint distribution before any data is observed. Thus, PSR can be understood to have a zero-gradient prior.

Incidentally, this observation also provides an answer to an often asked question about PSR; namely, why PSR, which is posed as recovering a function which is zero on the surface and satisfies certain...
norm-one gradients, produces an output so different from a Signed Distance Field (SDF). From this novel Gaussian Process perspective, we can understand that PSR is, via its \( m = 0 \) prior, encouraging zero-norm gradients away from the sample positions, while an SDF requires Eikonal norm one gradients almost everywhere.

**Primitive geometric priors.** Martens et al. [2016] suggest using simple geometric primitives as priors in the context of GP implicit surfaces. Since the authors are directly learning the surface’s implicit representation, their suggested priors \( m \) are scalar, SDF-like functions. In our case, where learning is carried out in the gradient space and then transferred to an implicit function via a PDE solve, we can use gradient vector-valued priors.

In Fig. 19, we exemplify the effect of a simple spherical prior

\[
m(x) = \alpha \frac{x - c}{\|x - c\|},
\]

which we pose can be a useful tool for favouring closed reconstructions over open ones. \( \alpha \) is a parameter tuning the strength of the prior and \( c \) is the center of the sphere, which we set to be the average of all point cloud positions. In Fig. 18, we show less didactical uses of this same prior, along with a similar ellipsoidal one.

6 IMPLEMENTATION DETAILS

We implemented our algorithm in Python, using libigl [Jacobson et al. 2018] for common geometry processing subroutines. We rendered all our figures in Blender, using BlenderToolbox [Liu 2022]. All our results were produced on our machine with Intel Xeon CPU E5-2637 v3 3.50Hz (16 cores) with 64GB of RAM.

For parameter standardization, we normalized all our examples to fit a length-one cube as a preprocessing step. Unless specified otherwise, we use a 100\(^3\) uniform grid (100\(^2\) in the two-dimensional examples) such that \( L \) and \( Z \) become the usual finite difference Laplacian and divergence matrices. We fix \( \sigma = 0.02, \alpha = 0.05 \) and \( k = 3000 \). We use the same function \( F_\alpha \) as Kazhdan et al. [2006], obtained by convolving a box filter with itself three times.

For memory efficiency, we use SciPy’s biconjugate gradient method to solve for the distribution mean \( f_{SPSR} \), which accounts for approximately 5\% of our runtime (around 10 seconds in all 3D examples). Our main computational bottleneck is Eq. (33) (specifically, the projection of \( K_f \) into the reduced space), covering 90\% of our runtime (around two minutes in all 3D examples).

7 EXPERIMENTS

7.1 Deviation from Kazhdan et al. [2006]

Our sole deviation from traditional Poisson Surface Reconstruction as introduced by Kazhdan et al. [2006] is the symmetrization of the covariance in Eq. (37). Theoretically, it is clear that both \( k_{PSR} \) and \( k_{SPSR} \) converge to the same symmetric \( F_\alpha(y) \) in the limit of grid refinement. However, we also justify the step by showing that the difference between these two is small even at a very coarse resolution in Fig. 8. In PSR, \( k_{PSR} \) is used to reconstruct a vector field \( \tilde{V}_{PSR} \). In Fig. 7, we show that the interpolated vector field \( \tilde{V}_{SPSR} \) obtained with the symmetrized covariance is visually identical. In Fig. 3, we show that this applies also to the implicit function \( f \) obtained through the Poisson solve.

7.2 Statistical quantities

Our Stochastic PSR exchanges the functional PSR output for a fully defined statistical distribution. To explore just how much more information is contained in this distribution, we show different statistical quantities meaningful to the shape reconstruction process.

In Fig. 10, we use our computed mean and variance to evaluate each point’s cumulative distribution functions at zero, which returns the probability of any point in space being contained in \( \Omega \). Similarly, evaluating the probabilistic density function at zero returns a measure of surface probability, as we show in Fig. 13. We...
8 APPLICATIONS

Our main contribution is a novel theoretical understanding of Poisson Surface Reconstruction that expands it beyond its traditional application realm. We explore this with prototypical results.

8.1 Collision detection

Statistical information is critical for robots or autonomous vehicles, which may capture their surroundings as point clouds with partial occlusions. When a car is deciding whether to swerve, it does not want to know only if the manoeuvre is secure for the most likely scenario of its surroundings (the traditional PSR output or the SPSR mean); rather, it must account for uncertainty and conclude whether the manoeuvre is safe in the vast majority of possible scenarios.

We exemplify this in Fig. 20, where we simulate a 3D scan of a street and use joint probability queries to compute the collision chance of an automated vehicle along a given trajectory. We compare this chance against a baseline (in grey), which only checks for intersections between the car mesh and the traditional PSR zero isosurface extracted with Marching Cubes [Lorensen and Cline 1987]. We include synthetic Gaussian noise both on the input point positions and normal vectors. In Fig. 22, we show how one can threshold our computed probabilities \( p(x \in \Omega) \) to obtain isosurfaces for use in collision detection applications.

8.2 Volume rendering

The arbitrary units in the traditional PSR output mean that it cannot be interpreted as a volume rendering occupancy. Fortunately, our Stochastic Poisson Surface Reconstruction allows us to compute \( p(x \in \Omega) \), values between 0 and 1 which can be interpreted as occupancies (this reinterpretation is common in the Neural Radiance Field literature, e.g., [Mildenhall et al. 2020]). In Fig. 24, we use standard volume rendering techniques (see [Pharr et al. 2016], Chap. 11) to sample the free path of a ray aimed at a partial reconstruction of a Space Wizard vehicle from two different directions. As expected, rays intersecting regions with higher uncertainty produce a wider spread of paths, as evidenced by the orange and blue histograms.

8.3 Reconstruction

Scanning integration. Surface reconstruction is the flagship application of PSR. Despite this, PSR cannot be fully integrated into the scanning pipeline, as it fails to provide basic feedback like when a point cloud is dense enough or where to scan next. In Fig. 15, we show that our introduced total uncertainty \( u \) converges to zero as points are added, giving a clear measure of scan quality. We show how one can use thresholds on \( u \) as a scanning stopping criteria in Fig. 17, obtaining reconstructions of similar quality for the same threshold value. In Fig. 23, we show our algorithm’s output on a real-world depth scan obtained with a smartphone app.

The ability to simulate ray casting on our reconstructed shapes also allows us to score different potential scan positions, as we show in Figs. 25 and 26. We begin by simulating a scan on a given groundtruth object using traditional ray tracing in a cone around some predetermined camera positions (see Fig. 25) placed on the left side of a race car. We include synthetic Gaussian noise both on the input point positions and normal vectors. This lets us obtain a point cloud \( P \) which we can input to our algorithm and compute its total uncertainty \( U_{SPSR}(P) \). For each potential new scan position, we cast a ray and add the resulting intersection \( p \) as a new point to the input point cloud, defining the camera score as the change in total uncertainty from adding the new point (see Fig. 26),

\[
s = |U_{SPSR}(P) - U_{SPSR}(P \cup \{p\})|
\]

We repeat this process ten times for each potential camera position, to account for the statistical variability in the ray casting. A fundamental part of any scan feedback pipeline, this process helps us decide on a next best camera view. As expected, scores are highest for hypothetical cameras on the right side of the car.
Our probability allows for ray casting queries.

Point cloud repair. Converting to an implicit function or a triangular mesh is not needed for some downstream applications that can work directly on point clouds. However, raw scan clouds often contain holes due to occlusions or missing camera angles. In Fig. 21, we show how our algorithm can be used to produce possible surface points in the occluded regions by using a Metropolis-Hastings scheme to sample from our surface probability \( p(x \in \Omega) \).

Incorporating priors. Our novel statistical interpretation of PSR allows us to build on it even as it applies to its most direct purpose of recovering surfaces from point clouds; for example, by incorporating geometric priors as we exemplify in Fig. 19. In Fig. 18, we show how even very simple primitive geometric priors can be useful for recovering complex three-dimensional geometry.

9 LIMITATIONS & CONCLUSION

Our algorithm extends the traditional PSR by Kazhdan et al. [2006] by separating it into a GP vector field reconstruction and a PDE solve. It is unclear how our approach could be applied to the Screened PSR by Kazhdan and Hoppe [2013], which combines both steps to improve robustness in unsampled regions. We conjecture that the answer may lie in works on incorporating gradient observations to Gaussian processes (see 9.4 in [Williams and Rasmussen 2006]).

Our work not only extends our knowledge of Poisson Surface Reconstruction from the perspective of a Gaussian Process. Indeed, we have also used PSR to expand our knowledge of Gaussian Processes, by introducing a covariance lumping technique which avoids costly matrix inversions at test time. We believe applying this in the broader GP context to be a promising avenue for future work.

As 3D acquisition becomes more accessible and attractive to consumers and the lines between real and virtual geometry get blurred, we believe quantifying and transmitting the uncertainties of the capture process to be one of the fundamental problems of this era. We hope our work inspires our geometry processing colleagues, whose workflow often involves PSR, to study how this statistical formalism carries over further down the shape processing pipeline.
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We will prove that the difference between the PSR semicovariance which, by definition of $k$,

$$\text{PSR}_k(x, y) = \sigma_g \sum_{o \in \mathcal{N}(x)} a_o \cdot F_o(y) \quad (36)$$

and our Stochastic PSR covaraiance

$$k_{\text{SPSR}}(x, y) = k_{\text{SPSR}}(x, y) - \frac{1}{2} (k_{\text{PSR}}(x, y) + k_{\text{PSR}}(y, x)) \quad (37)$$

vanishes at the limit of grid refinement. We can do so by showing that $k_{\text{PSR}}(x, y)$ converges to the gridless covariance function

$$k^*(x, y) = \sigma_g F_g(x) \quad (38)$$

Let $\lambda$ be the Lipschitz constant of $F$. Then, since $F$ is symmetric,

$$|F_o(y) - F_g(y)| = |F_o(o) - F_g(y)| \leq \lambda \|o - x\| \quad (39)$$

Since trilinear interpolation weights sum up to one, this means

$$|k_{\text{PSR}}(x, y) - k^*(x, y)| \leq \sigma_g \lambda \|o - x\| \quad (40)$$

Symmetrically, we get

$$|k_{\text{PSR}}(y, x) - k^*(x, y)| \leq \sigma_g \lambda \|o - x\| \quad (41)$$

which, by definition of $k_{\text{SPSR}}$, means

$$|k_{\text{SPSR}}(x, y) - k_{\text{PSR}}(x, y)| \leq \sigma_g \lambda \|o - x\| \quad (42)$$

**B JUSTIFICATION FOR COVARIANCE LUMPING**

Consider a simple Gaussian Process $\mathcal{A} = \{A(x) \}_{x \in \mathcal{D}}$ with zero mean and covariance function $k : D \times D \rightarrow \mathbb{R}$, with two different training observations $\{(x_1, a_1), (x_2, a_2)\}$ and one test point $x_3$. For clarity, let $k_{ij} = k(x_i, x_j)$. Then, since $k_{11} = k_{22}, k_{12} = k_{21}$, we have

$$K_3 = \begin{bmatrix} k_{11} & k_{12} & k_{12} \\ k_{12} & k_{22} & k_{21} \end{bmatrix} \quad , \quad K_2 = \begin{bmatrix} k_{11} \\ k_{12} \\ k_{21} \end{bmatrix} \quad (43)$$

Fig. 27. Cases where our lumping has correct asymptotics: (1) when training data are far from one another, (2) when test points are far from training data and (3) when training data are so close that their features are similar.

Traditionally, one uses these matrices to compute the GP posterior mean $k_2^T K_3^{-1} a$ and covariance $k_2^T K_3^{-1} k_2$. Let us write both of these as $k_2^T K_3^{-1} c$ for some generic feature vector $c$. Then,

$$k_2^T K_3^{-1} c = \frac{1}{k_{11} - k_{12}} \underbrace{((k_{11} k_{13} - k_{12} k_{23}) c_1 + (k_{11} k_{23} - k_{12} k_{13}) c_2)}_{= 0} + \underbrace{((k_{11} - k_{21}) k_{13} c_1 + (k_{11} - k_{21}) k_{23} c_2 + k_{12} (k_{23} - k_{13}) (c_1 - c_2))}_{\text{2nd term}}$$

or, equivalently,

$$k_2^T K_3^{-1} c = k_{13} c_1 + k_{23} c_2 + \frac{k_{12} (k_{23} - k_{13}) (c_1 - c_2)}{k_{11} - k_{12}} \quad (44)$$

Note that the denominator $k_{11} + k_{12}$ is (assuming $k$ is monotonically decreasing with distance) nothing but a measure of sampling density, meaning that we write the left term in the sum as

$$k_2^T K_3^{-1} c = k_2^T D^{-1} c + \frac{k_{12} (k_{23} - k_{13}) (c_1 - c_2)}{k_{11} - k_{12}} \quad (45)$$

where $D$ is our lumped covariance matrix. Therefore,

$$|k_2^T K_3^{-1} c - k_2^T D^{-1} c| \leq \frac{k_{12}}{k_{11} - k_{12}} |k_{13} - k_{12}| |c_1 - c_2| \quad (46)$$

Thus, while our lumping introduces error, it is bounded and has the correct asymptotic behaviour (see Fig. 27 when the training samples are far from one another ($k_{12} \rightarrow 0$), when the test samples are far enough from the training samples ($k_{23} - k_{13} \rightarrow 0$), and when the training samples are close enough that the training features are close ($c_1 - c_2 \rightarrow 0$). Assuming independent samples without accounting for sampling density would have meant approximating $k_{11} + k_{12}$ in Eq. (44) by a constant factor, instead of recognizing it as a measure of the relative positions of $x_1$ and $x_2$. This would have introduced further error and abandoned the asymptotic convergence.