GAUSSIAN PROCESS LANDMARKING FOR
THREE-DIMENSIONAL GEOMETRIC MORPHOMETRICS∗

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Abstract. We demonstrate applications of the Gaussian process-based landmarking algorithm proposed in [T. Gao, S.Z. Kovalsky, I. Daubechies 2018] to geometric morphometrics, a branch of evolutionary biology centered at the analysis and comparisons of anatomical shapes, and compares the automatically sampled landmarks with the “ground truth” landmarks manually placed by evolutionary anthropologists; the results suggest that Gaussian process landmarks perform equally well or better, in terms of both spatial coverage and downstream statistical analysis. We provide a detailed exposition of numerical procedures and feature filtering algorithms for computing high-quality and semantically meaningful diffeomorphisms between disk-type anatomical surfaces.

Key words. Gaussian Process, Experimental Design, Active Learning, Manifold Learning, Geometric Morphometrics

AMS subject classifications. 60G15, 62K05, 65D18

1. Introduction. Computing one-to-one correspondences between objects, or registration, plays an important role in a wide range of scientific disciplines, such as functional Magnetic Resonance Imaging (fMRI) in medical studies [61], shape matching in computer graphics [80], remote sensing in geophysical sciences [54], to name just a few. When the number of objects in comparison is prohibitively large, or when the information encoded in each data object is noisy or redundant, a common strategy is to reduce the comparison to alignment or assignment between sets of extracted features, which are typically compact and informative representations of the original data constructed with domain knowledge. Examples of this sort include the scale-invariant feature transform (SIFT) in computer vision [48] and anatomical landmarks in statistical shape analysis [20] and computational anatomy [37].

Geometric morphometrics is a subfield of evolutionary biology that focuses on quantifying the (dis-)similarities between pairs of two-dimensional anatomical surfaces based on their spatial configurations [83]. The main objects of study are anatomical surfaces, such as bones and teeth, of extinct and extant animals of particular interest to evolutionary biologists. These analyses typically start with manually identifying an equal number of geometrically or semantically meaningful feature points, or landmarks, on each specimen in a potentially large collection of anatomical surfaces; the landmarks are certified by domain experts to be in consistent one-to-one correspondences. Pairwise or group-wise Procrustes analysis [30, 20, 31] is then applied to quantitatively compare the specimens in the collection. These quantitative comparison results are subject to further statistical analysis to draw biologically relevant conclusions. With the advancement of digitization technologies, discrete triangular

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mesh representations of continuous biological surfaces are readily available and are used for high-throughput analysis of large collections of specimens. This paper focuses on two-dimensional surfaces represented by discrete triangular meshes.

In [25] a novel landmarking procedure was proposed, motivated by the analogy between Gaussian process experimental design and the landmark selection procedure in geometric morphometrics. In a nutshell, the method considers a Gaussian process on the anatomical surface, with a variance-covariance structure specified by the heat kernel of the underlying Riemannian manifold. The landmarks are then selected successively, each time picking a new landmark as the point on the surface with largest variance conditioned on all the previously selected landmarks. Surprisingly, with a simple reweighting modification for the construction of the kernel function, the landmarks generated from this procedure closely resembles the “ground truth” landmarks manually placed by evolutionary anthropologists. As we will demonstrate in this paper, these Gaussian process landmarks perform equally well or better compared with the ground truth landmarks, in terms of both spatial coverage and downstream statistical analysis.

The rest of this paper is organized as follows. Section 2 provides background materials on geometric morphometrics, Gaussian processes, and the Gaussian process landmarking algorithm proposed in [25]; Section 3 explains the importance of using a reweighted kernel in the Gaussian process landmarking algorithm for our application; Section 4 details all the numerical procedures in this application; Section 5 closes with comments and discussions.

2. Background.

2.1. Geometric Morphometrics: Old and New. Statistical shape analysis, often termed geometric morphometrics in the context of comparative biology, is the quantitative analysis of variations and correlations among biological forms through the Cartesian coordinates of “landmarks”—biologically informative, repeatable, and in some sense corresponding anatomical loci—on surfaces representing anatomy of biological organisms [1, 83]. In order for the comparisons across specimens to be meaningful, practitioners in this field often require that the landmarks be consistently annotated on each specimen in a manner reflecting the “operational homology” or “biological correspondence” (as discussed in [13]) across individuated traits inherited from a common ancestry [51]. For instance, in the generalized Procrustes analysis (GPA) framework [30, 20, 31, 2], the Procrustes distance between two surfaces $S_1, S_2$ is computed using the following procedure:

(i) Specify two sets of operationally homologous landmarks $\{x^{(1)}_\ell | 1 \leq \ell \leq L\}$ and $\{x^{(2)}_\ell | 1 \leq \ell \leq L\}$ on $S_1$ and $S_2$, respectively;

(ii) Compute the distance between $S_1$ and $S_2$ by minimizing the energy functional

\[
\inf_{T \in \mathcal{E}(3)} \left( \frac{1}{L} \sum_{\ell=1}^{L} \left\| T(x^{(1)}_\ell) - x^{(2)}_\ell \right\|^2 \right)^{1/2}
\]

where $\mathcal{E}(3)$ is the group of rigid motions in $\mathbb{R}^3$.

This idea can be generalized to analyze a collection of consistently landmarked shapes, either assuming each set of landmarks on the same shape is centered at the origin so the

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1 The term “homology” in the context of evolutionary theory bears a different meaning than in modern topology; see e.g. [69, Part IV].
variational problem is defined on a product space of orthogonal groups \([31, 58, 77, 56]\), or estimate the optimal orthogonal and translation group elements jointly without the overall centering assumption \([16]\).

The key to successfully applying the Procrustes framework in statistical shape analysis is to obtain an equal number of consistent, operationally homologous landmarks on every shape in a potentially enormous collection of shapes. Consistently landmarking a collection of shapes relies crucially upon domain knowledge and tedious manual labor, and the skill to perform it “correctly” typically requires years of professional training; even then the “correctness” can be subject to debate among experts (see e.g. \([12]\) for an example on \textit{Lepilemur} teeth). In the first place, extracting a finite number of landmarks from a continuous surface inevitably loses geometric information, unless when the shapes under consideration are easily seen to be uniquely determined by the landmarks (e.g. planar polygonal shapes, as considered in \([38][21]\)), which is rarely the case for geometric morphometricians in biology; this problem of “inadequate coverage” motivated the introduction of \textit{semilandmarks}—additional points along curves containing critical curvature information about the morphology—to compensate for the loss of geometry in the landmarking process. Unfortunately, the essential arbitrariness of the semilandmarks along a curve induces additional uncertainty that needs to be quantified and reduced \([83, \S 2]\), especially in the absence of sharp anatomical features; the constraint of picking an equal number of landmarks on each shape also turns out to be far too artificial when the anatomical forms undergo complex evolutionary and developmental processes.

To mitigate both the scalability and the subjectivity issues in the existing Procrustes analysis framework, a recent trend of research in geometric morphometrics advocates \textit{automated} workflows to bypass the repetitive, laborious, and time-consuming process of manual landmark placement; see e.g. \([2, 12, 66, 44, 11, 45, 13, 22, 81, 40, 35]\) and the references therein. These techniques work directly with digitized anatomical surfaces represented as discrete triangular meshes; numerical algorithms are combined with computer graphics and geometry processing to provide high-throughput, landmark-free approaches for precise phenotyping \([65, 33, 34]\) on the discrete triangular meshes on their entirety (often consisting of thousands to millions of vertices in \(\mathbb{R}^3\)), without the manual landmarking stage to filter down the number of variables using potentially biased \textit{a priori} domain knowledge. Similar to the claim made by the proponents of landmark-based morphometrics that landmark coordinates contain more information than utilized in more traditional, measurement-based morphometrics \([83, \S 2]\), the precursors of automated geometric morphometrics believe that using whole surfaces as input passes even more information to the downstream analysis.

Despite the capability of generating high quality pairwise shape registrations, automated geometric morphometric methods suffer from interpretability problems: since all comparisons are performed merely pairwise, composing the obtained correspondences along a closed loop does not give rise to an identity map in general. This lack of \textit{transitivity} (see e.g. \([26, 23]\)) demands additional post-processing steps to translate the pairwise results into valid input for standard downstream phylogenetic analysis \([62, 27]\), which bears a strong similarity with recent studies in \textit{synchronization problems} \([16, 24]\). The loop inconsistency of pairwise correspondences also challenges the interpretability of automated geometric morphometrics, since it becomes virtually impossible to identify functionally equivalent regions across distinct anatomical structures in a consistent manner. Until fully automated geometric morphometric algorithms reach the maturity with comparable explanatory power to a human practitioner of landmark-based geometric morphometrics, deeper and more systematic
understanding of the landmarking process still seem of great interest and value.

The methodology we propose in this paper incorporates an algorithmic landmarking procedure into the automated pairwise registration algorithms. We point out that detecting morphometrically meaningful landmarks on anatomical surfaces in a completely unsupervised manner is generally a daunting task, since some of the most reliable landmarks are determined by patterns of juxtapositions of tissues—termed “Type 1 landmarks” by Bookstein [70]—which are almost always absent on the triangular meshes input to automated algorithms [10]. The selection process is further complicated by the requirement of the consistency of relative landmark positions across the data collected, as well as the specific functionality of the biological organism being studied [83]. While geometry processing algorithms (e.g. [6, 47, 15, 14, 79]) are capable of detecting sharp geometric features characterized by metric or topology (Bookstein’s “Type 2 landmarks” [70]), as well as producing high quality pairwise registrations for accurate determination of operationally homologous loci, semilandmarks and a majority of “Type 3 landmarks” in Bookstein’s typology of landmarks are marked simply for adequate and/or comprehensive coverage of the anatomical forms [70]. For instance, some Type 3 landmarks are included in the analysis for being “furthest away” from sharp geometric or functional features [83]. These observations motivated us to consider algorithmic analogies of geometric morphologists’ daily practice beyond the scope of computational geometry, shedding light upon landmark identification from the perspective of Bayesian statistics quantifying the “uncertainty” of morphometric analysis. As will be detailed in Section 4, we will first generate a set of candidate landmarks on each of the anatomical surfaces based on the “uncertainty” modeled by a Gaussian process, then apply a matching scheme that filters out non-corresponding candidate landmarks between a pair of surfaces based upon bounded conformal distortion [43].

2.2. Gaussian Processes. A Gaussian process (or Gaussian random field) on a Polish space \( M \) with mean function \( m : M \to \mathbb{R} \) and covariance function \( K : M \times M \to \mathbb{R} \) is defined as the stochastic process of which any finite marginal distribution on \( n \) fixed points \( x_1, \cdots, x_n \in M \) is a multivariate Gaussian distribution with mean vector

\[
m_n := (m(x_1), \cdots, m(x_n)) \in \mathbb{R}^n
\]

and variance-covariance matrix

\[
(2.2) \quad K_n := \begin{pmatrix} K(x_1, x_1) & \cdots & K(x_1, x_n) \\ \vdots & \ddots & \vdots \\ K(x_n, x_1) & \cdots & K(x_n, x_n) \end{pmatrix} \in \mathbb{R}^{n \times n}.
\]

A Gaussian process with mean function \( m : M \to \mathbb{R} \) and covariance function \( K : M \times M \to \mathbb{R} \) will be denoted as \( \text{GP}(m, K) \). Under model \( Y \sim \text{GP}(m, K) \), given observed values \( y_1, \cdots, y_n \) at locations \( x_1, \cdots, x_n \), the best linear predictor (BLP) [78, 71] for the random field at a new point \( x \) is given by the conditional expectation

\[
(2.3) \quad Y^*(x) := \mathbb{E}[Y(x) \mid Y(x_1) = y_1, \cdots, Y(x_n) = y_n] = m(x) + k_n(x)^\top K_n^{-1}(Y_n - m_n)
\]

where \( Y_n = (y_1, \cdots, y_n)^\top \in \mathbb{R}^n \), \( k_n(x) = (K(x, x_1), \cdots, K(x, x_n))^\top \in \mathbb{R}^n \); at any \( x \in M \), the expected squared error, or mean squared prediction error (MSPE), is
defined as

\[
\text{MSPE}(x) = \mathbb{E} \left[ (Y(x) - Y^*(x))^2 \right]
\]

(2.4)

\[
= \mathbb{E} \left[ (Y(x) - \mathbb{E}[Y(x) | Y(x_1) = y_1, \ldots, Y(x_n) = y_n])^2 \right]
\]

\[
= K(x, x) - k_n(x)^\top K_n^{-1} k_n(x)
\]

which is a function over \( M \). Here the expectation is with respect to all realizations \( Y \sim \text{GP}(m, K) \). Squared integral \( (L^2) \) or sup \( (L^\infty) \) norms of the pointwise MSPE are often used as a criterion for evaluating the prediction performance over the experimental domain. In geospatial statistics, interpolation with (2.3) and (2.4) is known as kriging.

### 2.3. Gaussian Process Landmarking.

We now provide a quick summary of the Gaussian process landmarking algorithm from [25]. The input to this algorithm is a triangular mesh \( G = (V, E, F) \). Denote the set of vertices \( V \) as \( \{x_1, \ldots, x_{|V|}\} \subset \mathbb{R}^3 \), and construct the Gaussian process landmarking (2.5) as

\[
k_{t/2} = \left( e^{-\frac{\|x_i-x_j\|^2}{\sigma^2}} \right)_{1 \leq i, j \leq |V|} =: K.
\]

We calculate the mean and Gaussian curvature functions \( \eta : V \rightarrow \mathbb{R}, \kappa : V \rightarrow \mathbb{R} \) on the triangular mesh \( (V, E, F) \) using standard algorithms in computational differential geometry [17, 3]. For any \( \lambda \in [0, 1] \) and \( \rho > 0 \), define the value of the weight function \( w = w_{\lambda, \rho} \) at each vertex \( x_i \) by

\[
w_{\lambda, \rho}(x_i) = \frac{\lambda |\kappa(x_i)|^{\rho} + (1 - \lambda) |\eta(x_i)|^{\rho}}{\sum_{k=1}^{|V|} |\kappa(x_k)|^{\rho} \nu(x_k) \sum_{k=1}^{|V|} |\eta(x_k)|^{\rho} \nu(x_k)}\]

where \( \nu(x_k) \) is the area of the Voronoi cell centered at \( x_i \) on the triangular mesh \( T \), \( W \) is a diagonal matrix of size \( n \times n \) with \( \nu(x_k) \) at its \( k \)-th diagonal entry \( (1 \leq k \leq |V|) \), and \( K \) is the discrete squared exponential kernel matrix (2.5).

We define the reweighted heat kernel \( k_{t/2}^{W_{\lambda, \rho}} \) is then defined on \( V \times V \) as

\[
k_{t/2}^{W_{\lambda, \rho}}(x_i, x_j) = \sum_{k=1}^{|V|} k_{t/2}(x_i, x_k) k_{t/2}(x_k, x_j) w_{\lambda, \rho}(x_k) \nu(x_k) =: K^T W K
\]

where the (unweighted) heat kernel \( k_{t/2} \) is calculated as in (2.5).

We now detail the greedy landmarking procedures. Until a fixed total number of landmarks are collected, at each step \( k \) the algorithm computes the uncertainty score \( \Sigma_{(k)} \) on \( V \) from the existing \( (k - 1) \) landmarks \( \xi_1, \ldots, \xi_{k-1} \) by

\[
\Sigma_{(k)}(x_i) = k_{t}^{W_{\lambda, \rho}}(x_i, x_1) - k_{t}^{W_{\lambda, \rho}}(x_i, \xi_1)^\top k_{t}^{W_{\lambda, \rho}}(\xi_1, \xi_1)^{-1} k_{t}^{W_{\lambda, \rho}}(x_i, \xi_1) \quad \forall x_i \in V
\]

where

\[
k_{t}^{W_{\lambda, \rho}}(x_i, \xi_1) := \left( k_{t}^{W_{\lambda, \rho}}(x_i, \xi_1), \ldots, k_{t}^{W_{\lambda, \rho}}(x_i, \xi_{k-1}) \right)^\top,
\]

\[
k_{t}^{W_{\lambda, \rho}}(\xi_1, \xi_1) := \left( k_{t}^{W_{\lambda, \rho}}(\xi_1, \xi_1) \ldots \right.
\]

\[
\vdots
\]

\[
\left. \vdots \right) k_{t}^{W_{\lambda, \rho}}(\xi_{k-1}, \xi_{k-1}) \right).
\]
and pick the $k$-th landmark $\xi_k$ according to the rule

$$\xi_k = \arg\max_{x_i \in V} \Sigma(k)(x_i).$$

If there exist more than one maximizers of $\Sigma^{(k)}$, we just randomly pick one; at step 1 the algorithm simply picks the vertex maximizing $x \mapsto k^w_{\lambda,\rho}(x,x)$ on $V$. See Algorithm 2.1 for a comprehensive description. More details can be found in [25, §3]; we only mention here that (2.6) and (2.7) are discretizations of

$$w_{\lambda,\rho}(x) = \frac{\lambda |\kappa(x)|^\rho}{\int_M |\kappa(\xi)|^\rho \, d\text{vol}_M(\xi)} + \frac{(1-\lambda) |\eta(x)|^\rho}{\int_M |\eta(\xi)|^\rho \, d\text{vol}_M(\xi)}, \quad \forall x \in M$$

and

$$k^w_t(x,y) = \int_M k_{t/2}(x,z) k_{t/2}(z,y) w(z) \, d\text{vol}_M(z), \quad \forall x,y \in M$$

respectively. Guided by uncertainty specified through such curvature-reweighted covariance structure, the Gaussian process landmarking often identifies landmarks of abundant biological information—for instance, the first few Gaussian process landmarks are often highly biologically informative, and demonstrate comparable level of coverage with the observer landmarks manually picked by human experts. See Figure 1 for a visual comparison between the automatically generated landmarks with the observer landmarks manually placed by evolutionary anthropologists on a different digitized fossil molar.

Algorithm 2.1 greedily picks as the next landmark the point on the manifold $M$ that maximizes the pointwise prediction error. This sequential optimization approach is reminiscent of a popular approximation scheme in entropy-based experimental design [42, 60, 71]. The key observation here is the equivalence between minimizing the conditional entropy (or maximizing the information gain) and maximizing the determinant of the marginal covariance matrix at the design points; see e.g. [71, §6.2.1], [42, §3.1], [60, §2.2]. More concretely, if we denote the maximum entropy of a Gaussian process $\text{GP}(m,K)$ on a manifold $M$ with respect to any $n$ observations as

$$\text{OPT}(n) := \max_{\{x_1, \ldots, x_n\} \subset M} \det K_n$$

where $K_n$ is the variance-covariance matrix defined in (2.2), and write $\text{GPL}(n)$ for the value of $\det K(X_1, \cdots, X_n)$ where $X_1, \cdots, X_n$ are generated using Algorithm 2.1, then by the submodularity of the entropy function [39] we conclude, using classic results [57], that

$$\text{OPT}(n) \geq \text{GPL}(n) \geq (1 - 1/e) \text{OPT}(n).$$

In other words, the entropy of the greedy algorithm is equivalent to the maximum entropy up to a multiplicative constant. While this general submodularity-based framework justifies the information-theoretic asymptotic near-optimality of Gaussian Process Landmarking (Algorithm 2.1), in [25] we established a stronger optimality result in the context of Gaussian processes: up to a multiplicative constant, the MSPE with respect to $n$ Gaussian process landmarks coincides with the optimal MSPE attainable over all sets of $n$ points on $M$, for any positive integer $n \in \mathbb{Z}_{\geq 0}$. More details about this can be found in [25, §4].
Fig. 1: Left: Sixteen observer landmarks on a digitized fossil molar of a *Teilhardina* (one of the oldest known fossil primates closely related with living tarsiers and anthropoids [7]) identified manually by evolutionary anthropologists as ground truth, first published in [12]. Right: The first 22 landmarks picked by Gaussian Process Landmarking (Algorithm 2.1). The numbers next to each landmark indicate the order of appearance. The Gaussian process landmarks strikingly resembles the observer landmarks: the red landmarks (Number 1-5, 7, 8, 10, 11, 16, 19) signals geometric sharp features (cusps or saddle points corresponding to local maximum/minimum Gaussian curvature); the blue landmarks sit either along the curvy cusp ridges and grooves (Number 13, 18, 20, 22) or at the basin (Number 9), serving the role often played by semilandmarks; the four green landmarks (Number 6, 12, 15, 17) approximately delimit the “outline” of the tooth in occlusal view.

3. The Role of Reweighting in the Kernel Construction. An essential construction in the application of Algorithm 2.1 to geometric morphometrics is the reweighted kernel (2.10). Intuitively, the reweighting step modifies the Euclidean heat kernel (2.5) by amplifying the influence of locations with relatively high weights. To investigate the role played by the reweighted kernel in greater detail, we study the behavior of the reweighted kernel in the asymptotic regime when the number of samples increases to infinity. As will be demonstrated in Theorem 3.1, the reweighted kernel defines a diffusion process on the manifold with a backward Kolmogorov operator conjugate to the Witten Laplacian, of which the eigenfunctions has desired localization behavior near critical points of the weight function.

To motivate Theorem 3.1, let us consider i.i.d. (with respect to the standard volume measure) samples \( \{x_i\}_{i=1}^N \) on a closed Riemannian manifold \((M, g)\), as well as an arbitrary function \( f \in C^2(M) \). To simplify the discussion, assume for the moment that the samples are uniformly distributed on \( M \) with respect to the normalized volume form on \( M \). It is well known in manifold learning (see e.g. [72, 18]) that, for
Algorithm 2.1 Gaussian Process Landmarking with Reweighted Heat Kernel

1. procedure $GPL(T, L, \lambda \in [0, 1], \rho > 0, \epsilon > 0)$ \triangleright Triangular Mesh $T = (V, E)$, number of landmarks $L$
2. $\kappa, \eta \leftarrow \text{DiscreteCurvatures}(T)$ \triangleright calculate discrete Gaussian curvature $\kappa$ and mean curvature $\eta$ on $T$
3. $\nu \leftarrow \text{VoronoiAreas}(T)$ \triangleright calculate the area of Voronoi cells around each vertex $x_i$
4. $w_{\lambda, \rho} \leftarrow \text{CalculateWeight}(\kappa, \eta, \lambda, \rho, \nu)$ \triangleright calculate weight function $w_{\lambda, \rho}$ according to (2.9)
5. $W \leftarrow \left[\exp\left(-\frac{\|x_i - x_j\|^2}{\epsilon}\right)\right]_{1 \leq i, j \leq |V|} \in \mathbb{R}^{|V| \times |V|}$
6. $\Lambda \leftarrow \text{diag}(w_{\lambda, \rho}(x_1) \nu(x_1), \cdots, w_{\lambda, \rho}(x_{|V|}) \nu(x_{|V|})) \in \mathbb{R}^{|V| \times |V|}$
7. $\xi_1, \cdots, \xi_L \leftarrow \emptyset$ \triangleright initialize landmark list
8. $\Psi \leftarrow 0$
9. $\ell \leftarrow 1$
10. $K_{\text{full}} \leftarrow W^T \Lambda W \in \mathbb{R}^{|V| \times |V|}$
11. $K_{\text{trace}} \leftarrow \text{diag}(K_{\text{full}}) \in \mathbb{R}^{|V|}$
12. while $\ell < L + 1$ do
13. if $\ell = 1$ then
14. $\Sigma \leftarrow K_{\text{trace}}$
15. else
16. $\Sigma \leftarrow K_{\text{trace}} - \text{diag}(\Psi \left[ (\Psi[[\xi_1, \cdots, \xi_{\ell}, \cdot]], \cdot) \right] \in \mathbb{R}^{|V|}$ \triangleright calculate uncertainty scores by (2.8)
17. end if
18. $\xi_{\ell} \leftarrow \text{argmax} \Sigma$
19. $\Psi \leftarrow K_{\text{full}}[; [\xi_1, \cdots, \xi_{\ell}, \cdot]]$
20. $\ell \leftarrow \ell + 1$
21. end while
22. return $\xi_1, \cdots, \xi_L$
23. end procedure

any $x \in M,$

\begin{equation}
\frac{1}{N} \sum_{j=1}^{N} e^{-\frac{\|x-x_j\|^2}{\sigma^2}} f(x_j) \rightarrow \frac{1}{\text{Vol}(M)} \int_{M} e^{-\frac{\|x-y\|^2}{\sigma^2}} f(y) \, \text{dVol}(y)
\end{equation}

as $n \to \infty$. Denote the positive weight function (2.9) used in (2.10) by

\begin{equation}
w(x) = e^{-V(x)}, \quad V(x) \geq 0 \quad \forall x \in M.
\end{equation}

Note that the normalization in (2.9) ensures that the value of the weight function is bounded between 0 and 1, which is implied in (3.2), for numerical stability. Repeatedly using the “law of large numbers” argument (3.1), we have the following convergence
for the reweighted kernel:

\[
\frac{1}{N^2} \sum_{k=1}^{N} e^{-\frac{\|x-x_k\|^2}{2\epsilon}} e^{-V(x_k)} \sum_{j=1}^{N} e^{-\frac{\|x_j-x\|^2}{2\epsilon}} f(x_j) \rightarrow \frac{1}{|\text{Vol}(M)|^2} \int_M \int_M e^{-\frac{\|x-y\|^2}{2\epsilon}} e^{-V(z)} e^{-\frac{\|x-z\|^2}{2\epsilon}} f(y) \, d\text{Vol}(z) \, d\text{Vol}(y)
\]

as \(N \to \infty\).

This motivates us to consider the asymptotic behavior of the integral operator on the right hand side in the asymptotic regime \(\epsilon \to \infty\), a recurring theme in manifold learning (see e.g. [8, 18, 73, 23]). For simplicity of notation, we will denote the integral operator with respect to kernel \(K\):

\[
T_K f(x) := \frac{1}{|\text{Vol}(M)|} \int_M K(x,y) f(y) \, d\text{Vol}(y).
\]

**Theorem 3.1.** Let \(M\) be a \(d\)-dimensional closed manifold. For any smooth functions \(f, V \in C^\infty(M)\) with \(V \geq 0\), for kernel function

\[
K^\epsilon_V(x,y) := \frac{1}{|\text{Vol}(M)|} \int_M e^{-\frac{\|x-z\|^2}{2\epsilon}} e^{-V(z)} e^{-\frac{\|y-z\|^2}{2\epsilon}} \, d\text{Vol}(z)
\]

we have

\[
\frac{T_{K^\epsilon} f(x)}{T_{K^\epsilon} 1} \to f(x) + \epsilon \left( \Delta f(x) - \nabla f(x) \cdot \nabla V(x) \right) + O(\epsilon^2) \quad \text{as } \epsilon \to 0
\]

where \(1\) stands for the constant function on \(M\) taking value 1.

The proof of Theorem 3.1 can be found in Section SM1. Theorem 3.1 indicates that a proper normalization of the reweighted kernel (2.10) gives rise to an approximation to the heat kernel of the backward Kolmogorov operator

\[
L = -\Delta + \nabla V \cdot \nabla.
\]

Note that this operator is also the infinitesimal generator of the diffusion process determined by the stochastic differential equation

\[
dX_t = -\nabla V(X_t) \, dt + \sqrt{2} \, dW_t
\]

where \(W_t\) is the standard Wiener process defined on \(M\). In particular, Theorem 3.1 suggests that our construction of the reweighted kernel encodes information from the weight function into the dynamics of the stochastic process on \(M\).

**Remark 3.2.** It is interesting to compare the result of Theorem 3.1 with the related but different kernel construction in diffusion maps [18, 55]. The integral operator in (3.4) is a properly normalized version of integrating a smooth function against the kernel

\[
\tilde{K}^\epsilon_V(x,y) := \int_M e^{-\frac{\|x-z\|^2}{2\epsilon}} e^{-V(z)} e^{-\frac{\|y-z\|^2}{2\epsilon}} \, d\text{Vol}(z)
\]

which is obtained from sandwiching \(\exp[-V(\cdot)]\) with the squared exponential kernel. If we pick a different order of sandwiching, namely, construct the kernel as

\[
K^\epsilon_V(x,y) := e^{-V(x)} e^{-\frac{\|x-y\|^2}{2\epsilon}} e^{-V(y)},
\]
then a quick computation in the same spirit as [9, Theorem 1] leads to

$$\frac{T_{K_V} f(x)}{T_{K_V} 1} \to f(x) + \epsilon \left( \frac{1}{2} \Delta f(x) - \nabla f(x) \cdot \nabla V(x) \right) + O(\epsilon^2)$$

as $\epsilon \to 0$.

In other words, the infinitesimal generator differs from the one calculated in (3.4) only by a multiplicative factor in front of the Laplace-Beltrami operator. We actually observe very similar numerical results when replacing $\tilde{K}_V$ with $K_V$ in Algorithm 2.1, when appropriate bandwidth parameters are chosen. Empirically, we find the bandwidth parameter easier to tune for kernel $\tilde{K}_V$ than for $K_V$.

With these preparation, we now illustrate the effect of kernel reweighting by replacing (3.2) with

$$w_\epsilon(x) = e^{-V(x)/\epsilon}, \quad V(x) \geq 0 \quad \forall x \in M.$$  

Note that we can always redefine the potential function $V$ so that (3.2) is written in the form of (3.5), for any fixed $\epsilon > 0$; in particular, replacing $V(x)$ by $V(x)/\epsilon$ does not change its critical points. An almost identical calculation as in the proof of Theorem 3.1 leads to

**Corollary 3.3.** Under the same assumptions as in Theorem 3.1,

$$\frac{T_{K_{1/\epsilon} V} f(x)}{T_{K_{1/\epsilon} V} 1} \to f(x) + \epsilon \left( \Delta f(x) - \nabla f(x) \cdot \frac{1}{\epsilon} \nabla V(x) \right) + O(\epsilon^2)$$

as $\epsilon \to 0$

where $K_{1/\epsilon}$ is obtained from (3.3) by replacing the potential $V$ with $1/\epsilon V$.

Similar to Theorem 3.1, Corollary 3.3 shows that a proper normalization of the kernel reweighted by (3.5) approximates the heat kernel of the operator

$$L_\epsilon := -\Delta + \frac{1}{\epsilon} \nabla V \cdot \nabla.$$

The dependence on $\epsilon$ is of particular interest. It is well known in the literature of semi-classical analysis that $\epsilon L_\epsilon$ is conjugate to the semiclassical Witten Laplacian [82] on 0-forms:

$$\Delta_{V,\epsilon} = \epsilon e^{-V/2\epsilon} (\epsilon L_\epsilon) e^{V/2\epsilon},$$

where

$$\Delta_{V,\epsilon} = -\epsilon^2 \Delta + \frac{1}{4} |\nabla V|^2 - \frac{\epsilon}{2} \Delta V.$$  

Since the eigenfunctions of $\Delta_{V,\epsilon}$ corresponding to the leading small eigenvalues concentrate near the critical points of the potential function $V$ for sufficiently small $\epsilon > 0$ (see e.g. [36] or [64, Theorem 3.9]), the eigenfunctions corresponding to the leading small eigenvalues of $L_\epsilon$ also concentrate near the critical points of $V$ after being multiplied by $e^{-V/2\epsilon}$, which, by the proof of Theorem 3.1, can be approximated by the square root of the denominator of (3.6). Note that the matrix $K_{\text{full}}$ in Algorithm 2.1 corresponds to the integral kernel in the numerator of (3.6) if we choose $\Lambda$ to be the diagonal matrix with $\exp [-V(v_i)/\epsilon]$ at its $i$-th diagonal entry, where $v_i$ is the $i$-th vertex on the triangular mesh; setting $D$ to be the diagonal matrix with the $i$-th
row sum of $K_{\text{full}}$ at its $i$-th diagonal entry, the Witten Laplacian $\Delta_{V, \epsilon}$ can then be approximated using

$$D^{-1/2}K_{\text{full}}D^{-1/2}$$

up to a scalar multiplication by the bandwidth $\epsilon$. We would thus expect the first few eigenfunctions corresponding to the smallest eigenvalues of $D^{-1/2}K_{\text{full}}D^{-1/2}$ to concentrate near the critical points of $V$. This can be easily verified with numerical experiments; see Figure 2 for an example.

**Fig. 2:** Concentration of the eigenfunctions of the Witten Laplacian with respect to a potential function $V$. The left boxed subplot illustrates the potential function $V$ using a heat map; we constructed this potential to have 4 wells centered around 4 cusps manually picked on a *Lorisidae* mandibular molar. The 4 eigenfunctions corresponding to the 4 smallest eigenvalues of the Witten Laplacian with potential $V$ are depicted on the $2 \times 2$ panel on the right; it can be read off from the colorbars that the support of each of these 4 eigenfunctions are concentrated around critical points of $V$.

**Remark 3.4.** The localization effects on eigenfunctions can also be achieved by adding a diagonal matrix, which represents a potential function on the domain of interest, to the kernel matrix; this idea has found applications in biomedical data analysis [19] and computer graphics [50]. The insight we gained from relating the reweighted kernel to Witten Laplacian reveals that the localization can be equivalently realized through a simple nonlinear procedure. This phenomenon can be explained as follows: while adding a diagonal potential to the kernel provides a direct discretization of the Schrödinger operator $L_V = -\Delta + V$, the reweighted kernel (2.10) can be thought of as a discretization of the heat kernel of $L_V$ by taking only “one-hop” paths in the Feynmann-Kac formula for $e^{-tL_V}$. An extensive discussion along this direction is beyond the scope of the current paper; interested readers may find useful the works [75, 76, 32] and the references therein.

The connection between the reweighted kernel (2.10) and the semi-classical analysis of the Witten Laplacian provides insights for the behavior of the Gaussian process landmarks generated from Algorithm 2.1. Given a Gaussian process GP $(m, K)$
defined on a manifold $M$, the eigenfunctions of $K$—properly reweighted by their corresponding eigenvalues—give rise to the Karhunen-Loève basis for $\text{GP}(m, K)$, with respect to which the sample paths of $\text{GP}(m, K)$ adopt expansions with i.i.d. standard normal coefficients. If the low-frequency components of these expansions tend to concentrate at certain regions on $M$, when fitting an unknown function in $\text{GP}(m, K)$ using an active learning procedure, it could be much more efficient if one begins with the inquiry for the function value at those regions of concentration. After information at regions of concentration are collected to some extent, it becomes more beneficial to also incorporate ambient information that resides in the complement of these regions in $M$. Therefore, spreading landmarks on $M$ using a Gaussian process with reweighted kernel balances out the information prioritized by the weight function and the space-filling experimental design strategies. The main reason of not selecting landmarks solely based on the weight function is that the weight function can be too spurious to produce reliable, semantically meaningful landmarks, at least for our application in geometric morphometrics; see Figure 3 for an example for such a comparison.

Fig. 3: An equal number (20) of landmark points on the same surface generated using three different strategies. **Left:** Gaussian process landmarks generated using Algorithm 2.1 with a exponential squared kernel function reweighted by weight function $w$; **Middle:** The local maxima of the weight function used in the reweighed kernel, which appears semantically less meaningful from the perspective of geometric morphometrics; **Right:** Points produced by geodesic farthest point sampling (see e.g. [28, 53]), a greedy algorithm commonly used for generating uniformly sampled (or approximately space-filling) points on a triangular mesh. Comparing these three sampling approaches, Gaussian process landmarks has the advantage of tending to fill up the manifold while prioritizing the choice of semantically meaningful features for geometric morphometrics.

4. Gaussian Process Landmarking for Automated Geometric Morphometrics. This section is divided into three parts. **Subsection 4.1** compares Gaussian process landmarks generated from Algorithm 2.1 with “ground truth” landmarks manually picked by comparative biologists on a real dataset of anatomical surfaces, demonstrating comparable levels of coverage of biologically significant features. The results presented in this section provide quantitative evidence that Gaussian process landmarks are capable of capturing geometric features encoding important information for comparative biologists on individual anatomical surfaces. **Subsection 4.2** adapts the image feature matching algorithm of [46] for the registration of pairs of anatomical surfaces via matching Gaussian process landmarks computed on each individual surface. We compare the resulting shape correspondence maps with a baseline obtained using previously developed continuous Procrustes analysis in [2, 12]; the results suggest that, though Gaussian process landmarks are generated on each indi-
vidual shape separately, they implicitly encode operationally homologous features that can be compared and contrasted across shapes, and such pairwise comparison results are comparable with those obtained from standard Procrustes shape analysis based on “ground truth” observer landmarks. The pairwise shape distances induced from matching landmarks are turned into ordination plots (two-dimensional embeddings of distance matrices, commonly employed for visualizing the “morphospace” characterizing shape variances) in Section SM3; it turns out that Gaussian process landmarks lead to a favorable ordination, which is in better agreement with observations made by comparative biologists and paleontologists in existing literature.

4.1. Unsupervised Landmarking on Individual Anatomical Surfaces. Gaussian process landmarks are generated on each anatomical surface individually, regardless of the total size of the shape collection; the landmarks manually selected by human experts (observers), however, may well depend on the information gradually exposed to the human expert as he/she moves through a collection of surfaces. It is thus surprising that the individually generated, “local” Gaussian process landmarks bear striking similarity with the observer landmarks selected with certain extent of collection-wise or “global” knowledge, as illustrated in [25, Figure 2] through an example fossil molar. This subsection is devoted to a more thorough and quantitative comparison between Gaussian process landmarks and “ground truth” observer landmarks placed by human experts.

We begin by collecting in Figure 4 results obtained by applying the Gaussian Process Landmarking algorithm to several different types of anatomical surfaces, including a subset of the second mandibular molars of primates (first published in [12]) and a subset of the astragalus and calcaneus bones of tarsiers first published in [13]. It can be recognized from Figure 4 that the algorithm is capable of consistently capturing both geometric features (“Type 3 landmarks” [70]) and semilandmarks (delineating ridges and grooves).

To quantitatively validate the biological informativeness of Gaussian process landmarks, we use a dataset of second mandibular molars on which the “observer landmarks,” or the landmarks manually selected by experienced comparative biologists, are readily available (see [12]). We calculate, on each anatomical surface, the median geodesic distance from an observer landmark to its closest Gaussian process landmark. This median geodesic distance can obviously be calculated for any other type of landmarks in place of the Gaussian process landmarks; we shall refer to it as the median observer-to-automatic landmark distance, for the sake of simplicity. We compute the median observer-to-automatic landmark distances for a varying number of automatic landmarks, obtaining a curve that encodes the rate with which the median observer-to-automatic landmark distances decay to zero as the number of automatic landmarks increases. Comparing curves obtained from different automatic landmark generation methods then provides us with a way to evaluate how closely each type of automation “mimics” the observer landmarks. When we perform such curve comparisons for a large collection of surfaces in a dataset, one can in principle construct statistics (e.g. “mean” or “standard deviation” in the “space of curves”) along the lines of Functional Data Analysis (FDA, see e.g. [68, 67] and the references therein), but we will have to defer such a statistically systematic treatment to future work.

Using this strategy, we compare Gaussian process landmarks with landmarks obtained from Geodesic Farthest Point Sampling (GFPS) [28, 53], a popular downsampling technique in automated geometric morphometrics, on a dataset of 116 second mandibular molars of prosimian primates and closely related non-primates first pub-
Fig. 4: Gaussian process landmarks on several different types of anatomical surfaces. All triangular meshes are acquired from μCT scans. **Top Row:** Six second mandibular molars of prosimian primates and non-primate close relatives, from a dataset of 116 molars first published in [12]. **Middle Row:** Six astragalus bones of tarsiers from a dataset of 40 astragali first published in [13]. **Bottom Row:** Six calcaneus bones of tarsiers from a dataset of 40 calcanei first published in [13]. On all three types of anatomical surfaces, Gaussian process landmarks tend to play the roles of both landmarks and semilandmarks (c.f. [83, §2]): the curvature-reweighted kernel promotes automatically selecting sharp peaks or saddle points on the anatomical surface; after most of the prominent geometric features—normally recognized as Type 2 landmarks [83]—are captured, the uncertainty-based criterion encourages the identification of semilandmark-type points along ridges and grooves.
GAUSSIAN PROCESS LANDMARKING FOR 3D GM

4.2. Surface Registration via Matching Gaussian Process Landmarks.

We demonstrate in this subsection the benefit of using Gaussian process landmarks for establishing correspondences between pairs of surfaces. In particular, although Gaussian process landmarks are obtained independently on each surface, they turn out to encode geometrically significant features — shared more often among similar or related shapes — that are recognized as “operational homologous” [13] loci by comparative biologists. Statistical analysis also suggest that these correspondence maps reach comparable explanatory power to observer landmarks placed by human experts, in terms of induced shape distances. The comparison between the morphospaces characterized by these shape distances is deferred to Section SM3.

4.2.1. Experimental Setup and Methodology.

**Bounded Distortion Gaussian Process Landmark Matching.** Let $S_1$, $S_2$ be two-dimensional disk-type surfaces (conformally equivalent to planar disks by the Uniformization Theorem; see e.g. [2, 44, 45]) and let $\xi^{(1)}_1, \ldots, \xi^{(1)}_{L_1} \in S_1$ and $\xi^{(2)}_1, \ldots, \xi^{(2)}_{L_2} \in S_2$ be two sets of Gaussian process landmarks computed using Algorithm 2.1 on $S_1$ and $S_2$, respectively. Note that the algorithm we present in this subsection works equally well for $L_1 \neq L_2$, though we choose $L_1 = L_2 = 40$ throughout this paper to simplify the discussion. Adopting the approach suggested in [46] for feature-based image matching, we devise the following two-step approach for establishing geometrically-consistent matchings between the two sets of Gaussian process landmarks:

1) **Parametrization:** For each surface $S_j$ ($j = 1, 2$) we follow [74, 41] to compute an as-isometric-as-possible (AIAP) two-dimensional parametrization, which is a diffeomorphism $\Phi_j : S_j \rightarrow \Omega_j \subset \mathbb{R}^2$ from $S_j$ to a connected planar domain $\Omega_j$
minimizing the (discretization of the) isometric distortion energy

\[ F(\phi) := \int_{S_j} \left( |\nabla \phi(x)|^2 + |\nabla \phi^{-1}(x)|^2 \right) \, dv_{M}(x). \]

Each landmark \( \xi^{(j)}_\ell \) is mapped to a unique corresponding point \( \zeta^{(j)}_\ell = \Phi_j \left( \xi^{(j)}_\ell \right) \in \Omega_j \subset \mathbb{R}^2 \), where

\[ \Phi_j := \arg\min_{\phi: S_j \to \mathbb{R}^2} F(\phi). \]

(2) **Bounded Distortion Matching:** Following [46], we search within the set of planar diffeomorphisms between \( \Omega_1 \) and \( \Omega_2 \) with conformal distortion [43] bounded by a pre-fixed constant \( K \geq 1 \). This algorithm strives to find a maximal subset of geometrically-consistent correspondences within an initial set of candidate matches. In the extreme case of \( K = 1 \), the search is constrained within the set of strictly angle-preserving (conformal) maps for the continuous isometric distortion energy and the set of planar rigid transformations for the discretized isometric distortion energy; we select \( K = 1.5 \) in this experiment to slightly enlarge the search space of candidate maps.

As input to this matching algorithm, for each Gaussian process landmark \( \zeta^{(1)}_\ell \) on \( S_1 \), we choose \( T \geq 2 \) Gaussian process landmarks \( \zeta^{(2)}_{t=1}, \ldots, \zeta^{(2)}_{t=T} \) as initial putative candidate matches from the Gaussian process landmarks \( \{ \zeta^{(2)}_j \mid 1 \leq j \leq L_2 \} \) on \( S_2 \); the algorithm then searches for the bounded distortion map \( \Psi : \Omega_1 \to \Omega_2 \) that minimizes the mismatch count

\[ \sum_{\ell=1}^{L_1} \sum_{k=1}^{T} \left\| \Psi \left( \zeta^{(1)}_\ell \right) - \zeta^{(2)}_{t=k} \right\|^0, \]

where, following the notations of [46], \( \| \cdot \|^0 \) denotes the mixed \((2, 0)\)-norm

\[ \left\| \Psi \left( \zeta^{(1)}_\ell \right) - \zeta^{(2)}_{t=k} \right\|^0 = \begin{cases} 1 & \text{if } \Psi \left( \zeta^{(1)}_\ell \right) \neq \zeta^{(2)}_{t=k}, \\ 0 & \text{otherwise.} \end{cases} \]

The initial candidate matches in our experiments are generated by comparing the Wave Kernel Signature (WKS) [5] of the Gaussian process landmarks — the \( T \) Gaussian process landmarks on \( S_2 \) with most similar WKS’s (measured in Euclidean distances) with that of each \( \zeta^{(1)}_t \in S_1 \) are selected as \( \{ \zeta^{(2)}_{t \rightarrow t} \mid 1 \leq t \in T \} \).

Ideally, “incorrect” initial matches, which potentially lead to large conformal distortions, will be filtered out by minimizing (4.2) under the bounded conformal distortion constraint. Our implementation uses \( T = 2 \); see Figure 6(b) for an illustrative example.

The pairwise registration algorithm outputs a subset of \( 1 \leq L \leq \min \{ L_1, L_2 \} \) one-to-one correspondences \( \zeta^{(1)}_t \leftrightarrow \zeta^{(2)}_t, \ t = 1, \ldots, L, \) from the initial \( L_1T \) candidate Gaussian process landmark matches; the number of matched landmarks, \( L \), is automatically determined by the final bounded distortion map and could be vary between different pairs of surfaces. See Figure 6(c)(d) for an illustrative example. A final step in this algorithm pipeline “interpolates” the \( L \) pairs of matched Gaussian process landmarks to obtain a diffeomorphism between the surfaces \( S_1 \) and \( S_2 \); see Figure 6(d) for
Fig. 6: The outline of our approach for landmark matching and surface registration. (a) Gaussian process landmarks computed using Algorithm 2.1 on two surfaces; (b) planar parameterizations of the surfaces that minimize (4.1), overlaid with putative landmark matches (indicated by purple lines); (c) Bounded-Distortion correspondences: a subset of geometrically-consistent matches computed by approximately minimizing (4.2); (d) Pairs of corresponding Gaussian process landmarks found by bounded distortion matching, illustrated by spheres with matching colors; (e) Texture-map visualization of an inter-surface map interpolating the landmark correspondences depicted in (d).

an illustrative example. For this purpose, we use the technique developed in [74, 41] which computes a map $\Psi : \Omega_1 \to \Omega_2$ that minimizes the isometric distortion energy (4.1) subject to the $L$ linear equality constraints $\Psi(\tilde{\zeta}_\ell^{(1)}) = \tilde{\zeta}_\ell^{(2)} (1 \leq \ell \leq L)$ representing the “sparse” correspondences between Gaussian process landmarks. The composition $f_{12} = \Phi_2^{-1} \circ \Psi \circ \Phi_1$ finally produces the desired map $f : S_1 \to S_2$ that approximately minimizes the mismatch count (4.2). This last interpolation step is indispensable for the purpose of interpretability (producing full-surface registrations for visual comparisons) as well as evaluation (inducing shape Procrustes distances for ordination, see Section SM3). Figure 6 outlines the complete workflow of landmark matching and surface registration for a pair of molars surfaces from [12]; a few examples are shown in Figure 7.

**Evaluation Metrics and the Baseline.** We demonstrate the rich taxonomic information captured by the maps output from the proposed bounded distortion Gaussian process landmark matching (GP-BD) algorithm, by comparing the shape Procrustes distances induced by maps computed from GP-BD with those obtained by the same feature matching methodology but alternative choices of landmarking schemes. For any inter-surface map $f : S_1 \to S_2$, we define the Procrustes distance induced by $f$ as

$$d_P(f) = \left( \min_{R \in \mathbb{E}_3} \int_{S_1} \| f(x) - R(x) \|^2 \, d\text{vol}_{S_1}(x) \right)^{\frac{1}{2}},$$

where $\mathbb{E}_3$ stands for the rigid motion group in $\mathbb{R}^3$. The distance $d_P(f)$ measures the spatial registration error induced by the map $f$ between the two surfaces. To ensure that the Procrustes distances induced by the various methods listed in Table 1 is
Abbreviation | Description of the pairwise surface registration method
--- | ---
GP-BD | BD-filtering for GP landmarks computed with Algorithm 2.1
GP\textsubscript{Euc}-BD | BD-filtering for GP landmarks with the standard Euclidean heat kernel (2.5)
GP\textsubscript{nW}-BD | BD-filtering for GP landmarks with non-weighted kernel (\(w \equiv 1\) in (2.10))
GT-BD | BD-filtering for GT landmarks
GT\textsuperscript{2}-BD | BD-filtering for GT landmarks, also using the ground truth correspondences as candidate initial matches
CPM | Continuous Procrustes Maps (CPM) computed using the method of [12, 2]

Table 1: The pairwise surface registration methods compared in Subsection 4.2 and Section SM3. BD-filtering stands for \textit{bounded distortion maps based feature matching filtering}; GP stands for “Gaussian process”; GT stands for “Ground Truth,” i.e. those landmarks placed by experienced comparative biologists.

comparable, we normalize each surface to have unit surface area. When the map \(f\) is the continuous Procrustes map produced from the algorithm presented in [2], \(d_P\) gives exactly the \textit{continuous Procrustes distance} between \(S_1\) and \(S_2\).

The alternatives considered in this section include two other different types of Gaussian process landmarks (GP\textsubscript{nW}-BD and GP\textsubscript{Euc}-BD) with alternative kernel functions (different from (2.7)), two different strategies of utilizing the “ground truth” user-placed observer landmarks (GT-BD and GT\textsuperscript{2}-BD) as “oracles” representing the “ideal” landmarks for feature matching, as well as the baseline \textit{continuous Procrustes maps} (CPM) reported in [2, 12]; Table 1 provides a summary of these variants. Specifically, GP\textsubscript{Euc}-BD and GP\textsubscript{nW}-BD are based on Gaussian processes with the standard squared exponential kernel (2.5) and the trivially weighted kernel (setting \(w \equiv 1\) in (2.10)), respectively. GT-BD simply replaces the GP landmarks with “ground truth” (GT) landmarks, which then follows through with candidate selection via WKS and bounded distortion maps based feature matching filtering (BD-filtering); GT\textsuperscript{2}-BD also takes ground truth landmarks as inputs, but skips the candidate selection by setting the ground truth correspondences between observer landmarks as candidate matches, though BD-filtering still applies and prunes out potential “geometrically incompatible” correspondences leading to large conformal distortions. A detailed description of observer landmark acquisition can be found in [12].

The shape distances will be compared in several different ways. We will first compare cumulative distributions of the pairwise shape distance values, followed by two statistical tests addressing (i) the correlation between each automatic shape distance and the observer-determined landmarks Procrustes distance (ODLP), and (ii) the capability of each shape distance at distinguishing taxonomic groups. A qualitative but more intuitive comparison of the morphospaces [52] characterized by these shape distances will be deferred to supplementary materials Section SM3, in the form of ordination plots (two dimensional embedding of the shape distance matrices as visual representations for shape variances across species groups). The discussion in Section SM3 is however oriented slightly more towards readers with some background in comparative biology.

\subsection*{4.2.2. Comparison Results.} We compared the pairwise surface registration methods listed in Table 1 on three different dataset of anatomical surfaces: (i) 116 second mandibular molars of prosimian primates and closely related non-primates; (ii) 57 proximal first metatarsals of prosimian primates, New and Old World monkeys; (iii) 45 distal radii of apes and humans. Detailed descriptions of all 3 datasets can be
Fig. 7: Inter-surface maps and induced Procrustes distances: texture overlaid on the Source surface (top row) is mapped by $f : S_1 \rightarrow S_2$ so as to visualize correspondence. The second row (GP-BD) shows maps induced by correspondences established between Gaussian process landmarks computed with Algorithm 2.1. The bottom row (CPM) compares the baseline continuous procrustes maps. GP-BD outperforms CPM in examples (a)-(c), wherein the map it induces establishes a better correspondence between morphological features of the surfaces; example (d) shows a case in which GP-BD is outperformed by CPM. The inset values are the the Procrustes distance $d_P$ associated with each map.

found in [12]. We computed correspondences between each pair of surfaces within this dataset, totaling in over 13,000 inter-surface registrations for dataset (i), over 1,500 for dataset (ii), and nearly 1,000 for dataset (iii). Figure 7 shows example pairs of surfaces, visualizing inter-surface maps maps induced by GP-BD correspondences as well as the baseline CPM maps. These examples illustrate typical differences between maps computed from GP-BD and CPM along with their Procrustes distances; GP-BD maps often offer an improvement over CPM in terms of both visual quality and their ability to relate biologically meaningful and operationally homologous regions.

Comparing Cumulative Distributions of Distance Values. We first provide a crude comparison across the various methods listed in Table 1. Figure 8 plots the cumulative distributions of Procrustes distances induced by each type of pairwise registration method in Table 1, which serves as a direct comparison of the proportions of pairwise inter-surface correspondences for which the Procrustes distance $d_P$ (4.3) is below a
given threshold. The figure further includes a curve for the Observer-Determined Landmarks Procrustes (ODLP) distances, computed using the standard Procrustes analysis between sets of corresponding landmarks manually placed by human experts [12]. Recall from Subsection 4.2.1 that the surface areas are normalized so as to ensure that Procrustes distances induced by different shape correspondences are comparable.

Noticeably, the cumulative distribution of GP-BD distances most closely resembles that of GT-BD, obtained by replacing the Gaussian process landmarks with the ground-truth observer landmarks (but otherwise using exactly the same algorithm involving WKS and BD-filtering). Also, GP-BD outperforms GP_{\text{Euc}}-BD (Gaussian process with the standard Euclidean heat kernel). This comparison suggests that the Gaussian process landmarks computed with Algorithm 2.1 provide a good proxy for geometrically significant features needed to determine meaningful correspondences between surfaces.

Expectedly, GP-BD falls short compared to distances that rely on both the ground-truth observer landmarks and their true correspondences (i.e. GT^2-BD and ODLP). Comparing GP-BD to the baseline continuous Procrustes maps (CPM) in terms of distance distributions is equivocal as the two curves in Figure 8 cross each other when the Procrustes distance threshold is about 0.15. Nonetheless, comparing GP-BD to CPM indicates that the former produces more pairwise correspondences with shape distances less than 0.1 (at which the vertical gap between the two curves reaches its maximum). This, along with the correlation reported in [26] between smaller continuous Procrustes (cP) distances and better morphometric interpretability of the associated maps, implies that Gaussian process landmarks potentially lead to more stable and interpretable comparative biological analysis if combined with other globally transitive geometric morphometric methods.

**Statistical Tests Comparing the Distance Matrices.** We now provide a more thorough comparison, using nonparametric statistical tests, for the shape distance matrices induced by the various methods listed in Table 1. Treating the OPLD (Observer-Determined Landmarks Procrustes distance) matrix as a proxy for the “ground truth” accepted among geometric morphometricians (c.f. [12]), we first compare the correlation between each automatically computed shape distance matrix with ODLP using the Mantel correlation test [49]; the explanatory power of the shape distances (in terms of the ability to distinguish taxonomic groups) are then compared using a variant of the multivariate analysis of variance (MANOVA) for distance matrices.

Due to the dependence of entries in a distance matrix (e.g. constrained by the triangular inequality), assessments of correlation between distance matrices, or between a distance matrix and a (continuous or categorical) variable, often involves repeatedly permuting the rows and columns of the distance matrix (see e.g. [63, 59, 29] and the references therein). Table 2 demonstrates the results of Mantel correlation analysis [49] between the distance matrices computed in this paper against the ODLP distance matrix. In all mantel tests in this paper, we set the number of permutations to 9999 since it is recommended in [4] that at least 5000 permutations should be done for tests with an \( \alpha \)-level of 0.01. GT-BD and GT^2-BD correlate best with ODLP, which is as expected since they both rely directly on the same set of observer-determined landmarks used for computing the ODLP distances; their high correlation with ODLP justifies the use of bounded distortion maps for capturing biologically meaningful and corresponding geometric features. GP-BD outperforms CPM, which is consistent with our observation in Figure 8 that a majority (over 80\%) of the distance values computed from GP-BD is more similar in distribution to the distances computed with ground truth landmarks. The advantageously higher correlation of GP-BD over GP_{\text{Euc}} and
Fig. 8: Comparing GP-BD to the alternative methods listed in Table 1. Each curve represents the proportion of correspondences, among a collection of 116 surfaces, for which the Procrustes distance $d_P$ falls below a given threshold. The cumulative distribution of GP-BD distances most closely resembles that of GT-BD which utilizes the ground truth observer landmarks. It falls short of GT²-BD and the Observer-Determined Landmarks Procrustes (ODLP), both of which rely additionally on the “ground truth” correspondences between observer landmarks. In terms of producing more relatively smaller distances between the shape pairs (which is favorable by the correlation between smaller distances and enhanced morphometric interpretability reported in [26]; also suggested by the cumulative distribution curves of ODLP, GT²-BD, and GT-BD), GP-BD outperforms the baseline continuous Procrustes maps (CPM), as well as variants of Gaussian process landmarks with alternative kernel constructions, GPₙW-BD (Gaussian process landmarks with a non-weighted kernel) and GPₑuc-BD (Gaussian process landmarks with a standard Euclidean heat kernel).

|                | GT   | GT²  | CPM  | GP   | GPₑuc | GPₙW |
|----------------|------|------|------|------|-------|------|
| Molars         | 0.7042 | 0.7563 | 0.6647 | 0.6870 | 0.6135 | 0.6257 |
| First Metatarsals | 0.7371 | 0.8326 | 0.4887 | 0.7318 | 0.6607 | 0.7117 |
| Radii          | 0.3273 | 0.4909 | 0.1775 | 0.3231 | 0.2510 | 0.2746 |

Table 2: Correlation coefficients output from Mantel correlation analysis for the various distances computed in this paper versus ODLP distances, for three different datasets. See Table 1 for the list of abbreviations; note that we omitted “-BD” for the sake of space. The relatively high correlations of GT-BD and GT²-BD versus ODLP is not surprising due to their direct dependence on the observer-determined landmarks; the first 2 columns can thus be viewed as performance upper bounds when “oracle landmark correspondences” are provided. For all three datasets, the distance matrix computed from GP-BD correlates better with ODLP than CPM, as well as almost all of its variants, which is consistent with the CDF plot Figure 8 and the ordination plots in Figure SM5. The $P$-values of all results are < 0.01.

In addition, we perform Permutational Multivariate Analysis of Variance (PERMANOVA) [4] for the distance matrices computed in this paper, with the taxonomic groups shown in the ordination plots (Figure SM5, see detailed explanations in Section SM3) as the treatment effects for the molar dataset; for the first metatarsal and radius datasets, we use family and species groups as treatment effects, respectively.
The results are presented in Table 3. The purpose of this test is to quantitatively compare the power of differentiating taxonomic groups for these distance matrices. The pseudo $F$-statistics in PERMANOVA is a properly normalized ratio between among-group and within-group sum of squared distances. The statistical significance is then calculated from the fraction—among sufficiently many shuffles of the rows and columns of the distance matrix—of the permutation instances that produce a higher pseudo $F$-ratio. While all distance matrices demonstrate statistically significant pseudo $F$-ratios in this test, the GT-BD distance matrix leads the board of pseudo $F$-ratios, indicating its superior ability of separating taxonomic groups; the GP-BD distance matrix and its two variants consistently outperform CPM in terms of pseudo $F$-ratios, verifying again the improved quality of pairwise anatomical surface registrations as well as the induced shape Procrustes distances.

![Table 3](image)

Table 3: Pseudo $F$-ratios output from PERMANOVA for various distance matrices computed in this paper for three different datasets. See Table 1 for the list of abbreviations; note that we omitted “-BD” for the sake of space. The number of groups for the molar dataset is 30, equaling to the number of polygonal regions in the ordination plot in Figure SM5; the numbers of groups for first metatarsals and radius are 8 (families) and 4 (genuses), respectively. The first column refers to observer-determined landmarks Procrustes (ODLP) distances calculated in [12] as a baseline; the first 3 columns can thus be viewed as performance upper bounds when “oracle landmark correspondences” are provided. For all three datasets, the pseudo-$F$ ratio of the GP-BD distance matrix is better or at least comparable to the oracles. All distance matrices are statistically significant in terms of their powers of separating species groups: the $P$-values of all results are $< 0.01$.

5. **Discussions.** In this paper, we apply the uncertainty-based landmark generating algorithm proposed in [25] to three-dimensional geometric morphometrics. The algorithmically produced landmarks resemble biologists' landmarks selected with expert knowledge, providing adequate coverage for both geometric features and semantically “uncertain” regions on the anatomical surfaces of practical interest. We tested the applicability of this landmarking procedure for various tasks on real datasets.

The Gaussian process landmarking algorithm presented in this paper takes one anatomical surface as input at one time, which is not exactly consistent with the methodology of geometric morphometricians. In fact, these biologists do not simply place landmark on individual surface; rather, they are trained to take into consideration an entire collection of shapes, so as to consistently place landmarks that are in joint correspondence. In standard practice, landmarking a new anatomical surface typically involve repeated comparisons with all the other surfaces in the collection, and the already placed landmarks on a surface are still subject to change upon future knowledge acquired from landmarking more surfaces. It would be highly interesting to adapt our algorithm to accommodate for this type of group-wise comparison strategies as well.

Though geometric approaches in general lack the ability to recognize Type 1 landmarks (recall Bookstein’s typology reviewed in Subsection 2.1), we think that
the similarities between Gaussian process landmarks and user selected Type 2 and Type 3 landmarks call into question user-based landmarks. In particular, we found the ordination plots to be as successful, and in some ways superior, to the user-based landmarks in reflecting previous ideas about shape affinities. While $F$-ratios were highest in the user-determined sample, we note that it is highly likely that, for any underdetermined landmarks (e.g. the Type 1 landmarks, Type 3 landmarks, or when trying to place Type 2 landmarks on eroded features like blunted cusp tips) biologists will tend to minimize variance within species. In other words, if there is not enough geometry to allow consistent placement of a point, users are likely to unconsciously choose a positioning that visually maximizes similarity to other members of the species. Given the likely bias towards minimizing within group error by biologists during landmarking, it is quite remarkable that the Gaussian process approach comes so close to the “ground truth”, and exceeds by so much the other automated methods in terms of $F$-ratio. In user-placed landmarks, there is also the question of how many and which landmarks were chosen, not to mention whether different users are as accurate in placing the same landmarks. Finally, there is the limitation that any traditional landmark used must be present in every specimen of the sample. Given the challenges with the traditional user based approach, and the demonstrated ability of Gaussian process to emulate qualitatively and quantitatively Type 2 and Type 3 landmarks, we believe the Gaussian process landmarking algorithm has great potential to be an alternative, automated approach for such landmarks.

Software. MATLAB code for the surface registration algorithm is publicly available at https://github.com/shaharkov/GPLmkBDMatch.

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