SigGPDE: Scaling Sparse Gaussian Processes on Sequential Data

Maud Lemercier 1  Cristopher Salvi 2  Thomas Cass 3  Edwin V. Bonilla 4  Theodoros Damoulas 1  Terry Lyons 2

Abstract

Making predictions and quantifying their uncertainty when the input data is sequential is a fundamental learning challenge, recently attracting increasing attention. We develop SigGPDE, a new scalable sparse variational inference framework for Gaussian Processes (GPs) on sequential data. Our contribution is twofold. First, we construct inducing variables underpinning the sparse approximation so that the resulting evidence lower bound (ELBO) does not require any matrix inversion. Second, we show that the gradients of the GP signature kernel are solutions of a hyperbolic partial differential equation (PDE). This theoretical insight allows us to build an efficient backpropagation algorithm to optimize the ELBO. We showcase the significant computational gains of SigGPDE compared to existing methods, while achieving state-of-the-art performance for classification tasks on large datasets of up to 1 million multivariate time series.

1. Introduction

Gaussian process (GP) models provide a sound mathematical framework for supervised learning that allows the incorporation of prior assumptions and provides uncertainty estimates when modelling unknown functions (Rasmussen & Williams, 2006). This is usually achieved by specifying a GP prior over functions with a suitable covariance (or kernel) along with a conditional likelihood. With this, the problem boils down to that of estimating the posterior over the function (values) given the observed data.

However, this posterior distribution is often analytically intractable and, even when the conditional likelihood is a Gaussian, GP models scale poorly on the number of observations N, with naïve approaches having a time complexity O(N^3). From a wide range of approximate techniques to scale inference in GP models to large datasets, “sparse” methods based on variational inference (VI) have emerged as one of the dominant approaches (Titsias, 2009). They consist in defining a family of approximate posteriors through M inducing variables, and selecting the distribution in this family that minimizes the Kullback-Leibler (KL) divergence between the approximation and the true posterior. This is achieved by minimizing the so-called evidence lower bound (ELBO). When the likelihood factorizes over datapoints, training can be done in minibatches of size N resulting in a per-iteration computational cost O(NM^2 + M^3), where the O(M^3) cost is due to the inversion of the covariance matrix of the M inducing variables. This yields significant computational savings when M ≪ N.

In the seminal work of Titsias (2009) the inducing variables correspond to evaluations of the GP at M pseudo input locations, which typically results in a dense covariance matrix to invert. Subsequently, other ways of constructing inducing variables have been introduced in order to mitigate the O(M^3) cost (Hensman et al., 2017; Burt et al., 2020b). The core idea consists in defining (almost) independent inducing variables, such that their covariance matrix is (almost) diagonal. These inducing variables correspond to projections of the GP on basis functions, such that the covariance matrix is a Gramian matrix with respect to some inner-product. Orthogonal basis functions yield diagonal Gramian matrices, hence these methods are often referred to as variational orthogonal features (VOFs). However existing VOF methods are limited to stationary kernels on \( \mathcal{X} \subset \mathbb{R}^d \) (d ∈ \( \mathbb{N} \)).

In this work we are interested in generalizing the VOF paradigm to the case where the input space \( \mathcal{X} \) is a set of sequences of vectors in \( \mathbb{R}^d \). One may be tempted to naively concatenate each vector in a sequence of length \( \ell \) to form a flat vector in \( \mathbb{R}^{d\ell} \). However in this case existing VOF methods cannot be directly applied because they are limited to low dimensional vectors, with \( d \leq 8 \) (Dutordoir et al., 2020). Thus, one needs kernel functions specifically designed for sequential data. The signature kernel (Cass et al., 2020) is a natural choice that has recently emerged as a leading machine learning tool for learning on sequential data. In particular, Toth & Oberhauser (2020) have proposed GPSig, a GP inference framework leveraging an approximation of
this covariance function (Király & Oberhauser, 2019) and achieving state-of-the-art performance on time series classification tasks. Nevertheless, as in standard sparse variational approaches to GPs, the inducing inputs they chose (so called *inducing tensors*) are additional variational parameters to optimize, and the resulting covariance matrix is dense.

Here we develop SigGPDE, a new scalable sparse variational inference framework for GP models on sequential data. After a brief recap on the general principles of variational inference (Sec. 2) we identify a set of VOFs naturally associated with the signature kernel. These inducing variables do not depend on any variational parameter as they are defined as projections of GP-samples onto an orthogonal basis for the RKHS associated to the signature kernel (Sec. 3). As a result, unlike the methods developed in Toth & Oberhauser (2020), in SigGPDE the optimization of the ELBO does not require any matrix inversion. Subsequently, we show that the gradients of the signature kernel are solutions of a hyperbolic partial differential equation (PDE). This theoretical insight allows us to build an efficient backpropagation algorithm to optimize the ELBO (Sec. 4). Our experimental evaluation shows that SigGPDE is considerably faster than GPSig, whilst retaining similar predictive performances on datasets of up to 1 million multivariate time series (Sec. 6).

2. Background

We begin with a general summary of variational inference for GPs. In this section, it is assumed that the input space is $\mathcal{X} \subset \mathbb{R}^d$. Standard models with zero-mean GP priors and iid conditional likelihoods can be written as follows

$$f \sim \mathcal{GP}(0, k(\cdot, \cdot)) \quad p(y|f) = \prod_{i=1}^{N} p(y_i|f(x_i)),$$

where $k(\cdot, \cdot)$ is the covariance function. The general setting for sparse GPs consists in specifying a collection of $M$ variables as well as a joint distribution with variational parameters $\mathbf{m}$ (mean vector) and $\Sigma$ (covariance matrix)

$$\mathbf{u} = \{u_m\}_{m=1}^{M}, \quad q(\mathbf{u}) = \mathcal{N}(\mathbf{m}, \Sigma).$$

These variables induce a family of approximate posteriors that are GPs with finite dimensional marginal densities of the form $q(f, \mathbf{u}) = p(f|\mathbf{u})q(\mathbf{u})$. Considering any input $x \in \mathcal{X}$, the mean and covariance functions of these GPs are

$$\mu_q(x) = C_{f, \mathbf{u}}C_{\mathbf{u}, \mathbf{u}}^{-1}\mathbf{m},$$

$$k_q(x, y) = k(x, y) - C_{f, \mathbf{u}}C_{\mathbf{u}, \mathbf{u}}^{-1}(C_{\mathbf{u}, \mathbf{u}} - \Sigma)C_{\mathbf{u}, f},$$

where the vector $C_{f, \mathbf{u}}$ and the matrix $C_{\mathbf{u}, \mathbf{u}}$ are defined as

$$[C_{f, \mathbf{u}}]_m = \mathbb{E}[u_m f(x)], \quad [C_{\mathbf{u}, \mathbf{u}}]_{m, m'} = \mathbb{E}[u_m u_{m'}].$$

Provided the inducing variables $\mathbf{u}$ are deterministic conditioned on $f$, one has the following lower bound (ELBO) on the marginal log likelihood (Matthews, 2017)

$$\log p(y) \geq \mathbb{E}_{q(f)}[\log p(y|f)] - KL[q(\mathbf{u})||p(\mathbf{u})],$$

where $p(\mathbf{u}) = \mathcal{N}(0, C_{\mathbf{u}, \mathbf{u}})$. Maximizing the right-hand-side of eq. (5) is equivalent to minimizing the KL divergence between $q(f)$ and the true posterior distribution.

The original variational inference framework outlined in Titsias (2009) consists in setting $u_m = f(z_m)$ where $z_m \in \mathcal{X}$ is a pseudo input living in the same space as $x$ that may either be fixed or optimized. The per-iteration cost of optimizing the ELBO is $O(\tilde{N}M^2 + M^3)$, where $\tilde{N}$ is the minibatch size and $M^3$ is the cost of computing $C_{\mathbf{u}, \mathbf{u}}^{-1}$ via a Cholesky decomposition.

Recently, a considerable effort has been devoted to the construction of inducing variables $\mathbf{u}$ which yield a structured covariance matrix $C_{\mathbf{u}, \mathbf{u}}$ whose inversion has a reduced computational complexity (Hensman et al., 2017). This line of work is often referred to as inter-domain sparse GPs, owing to the fact that the pseudo inputs are not constrained to live in $\mathcal{X}$ as before. In particular, Burt et al. (2020b); Dutordoir et al. (2020) have shown that provided one can find an orthogonal basis of functions for the RKHS associated with the kernel $k(\cdot, \cdot)$, it is possible to define the inducing variables as projections of the GP samples onto this basis. This construction yields a diagonal covariance matrix $C_{\mathbf{u}, \mathbf{u}}$.

3. Variational Inference with Orthogonal Signature Features

Here we present our first contribution, namely the use of *orthogonal signature features* as inducing variables for GPs on sequential data. We begin with a summary of the theoretical background needed to define GPs endowed with the signature kernel. In this section $\mathcal{X}$ is no longer a subspace of $\mathbb{R}^d$ but will be defined as a space of *paths* hereafter.

3.1. The signature

Consider a time series $\mathbf{x}$ as a collection of points $x_t \in \mathbb{R}^{d-1}$ with corresponding time-stamps $t_t \in \mathbb{R}$ such that

$$\mathbf{x} = ((t_0, x_0), (t_1, x_1), ..., (t_n, x_n))$$

with $0 = t_0 < \ldots < t_n = T$. Let $X : [0, T] \to \mathbb{R}^d$ be the piecewise linear interpolation of the data such that $X_{t_i} = (t_i, x_i)$. We denote by $\mathcal{X}$ the set of all continuous piecewise linear paths defined over the time interval $[0, T]$ and with values on $\mathbb{R}^d$.

For any path $X \in \mathcal{X}$ and any $\alpha \in \{1, \ldots, d\}$, we will denote its $\alpha^{th}$ channel by $X^{(\alpha)}$ so that at any time $t \in [0, T]$:

$$X_t = (X_t^{(1)}, \ldots, X_t^{(d)}),$$

where $X_t^{(\alpha)} = \langle X_t \rangle^{(\alpha)}$.
as depicted on Fig. 1a with \( d = 3 \).

The signature \( S : \mathcal{X} \to H \) is a feature map defined for any path \( X \in \mathcal{X} \) as the following infinite collection of statistics

\[
S(X) = \left\{ \left( S(X)^{(\alpha)} \right) \right\}_{\alpha = 1}^{d},
\]

\[
\left\{ \left( S(X)^{(\alpha_1, \alpha_2)} \right) \right\}_{\alpha_1, \alpha_2 = 1}^{d},
\]

\[
\left\{ \left( S(X)^{(\alpha_1, \alpha_2, \alpha_3)} \right) \right\}_{\alpha_1, \alpha_2, \alpha_3 = 1}^{d}
\]

where each term is a scalar equal to the iterated integral

\[
S(X)^{(\alpha_1, \ldots, \alpha_j)} = \int_{0 < s_1 < \ldots < s_j < T} dX^{(\alpha_1)} \cdots dX^{(\alpha_j)}
\] (8)

The feature space \( H \) associated to the signature is a Hilbert space defined as the direct sum of tensor powers of \( \mathbb{R}^d \)

\[
H = \bigoplus_{k=0}^{\infty} (\mathbb{R}^d)^{\otimes k} = \mathbb{R} \oplus \mathbb{R}^d \oplus (\mathbb{R}^d)^{\otimes 2} \oplus \ldots
\] (9)

where \( \otimes \) denotes the outer product (Lyons, 1998; 2014).

**Interpretability of the signature features** An important aspect of sequential data is that the order of the observations is not commutative, in the sense that reordering the elements of a sequence can completely change its meaning. By definition the terms in the signature capture this fact. In effect, the \( j \)-fold iterated integral in eq. (8) is defined as an integral over the simplex \( 0 < s_1 < \ldots < s_j < T \) which explicitly encodes the ordering of events happening across different channels \( X^{(\alpha_1)}, \ldots, X^{(\alpha_j)} \). This provides the signature features with a natural interpretability as highlighted several times in prior work (Arribas et al., 2018; Moore et al., 2019; Lemercier et al., 2021).

### 3.2. The signature kernel

The signature kernel \( k : \mathcal{X} \times \mathcal{X} \to \mathbb{R} \) is a reproducing kernel associated to the signature feature map and defined for any pair of paths \( X, Y \in \mathcal{X} \) as the following inner product

\[
k(X, Y) = \langle S(X), S(Y) \rangle_H.
\] (10)

From the structure of \( H \) and the properties of the signature it turns out that the signature kernel can be decomposed according to the expansion (Cass et al., 2020)

\[
k(X, Y) = \sum_{j=0}^{\infty} \sum_{|\alpha| = j} S(\alpha)^{\alpha} S(\alpha)^{\alpha},
\] (11)

where the inner summation is over the set of multi-indices

\[
\{ \alpha = (\alpha_1, \ldots, \alpha_j) : \alpha_1, \ldots, \alpha_j \in \{1, \ldots, d\} \}
\] (12)

In their recent article, Cass et al. (2020) provide a kernel trick for the signature kernel by proving the relation

\[
k(X, Y) = U(T, T)
\] (13)

where the function of two variables \( U : [0, T] \times [0, T] \to \mathbb{R} \) is the solution of the following hyperbolic PDE

\[
\frac{\partial^2 U}{\partial s^2} = (X_s T Y_t) U
\] (14)

with boundary conditions \( U(0, \cdot) = 1 \) and \( U(\cdot, 0) = 1 \). This kernel trick is explained with simple arguments in the proof of Cass et al. (2020, Thm. 2.5). The sketch of the proof goes as follows: one first shows that the inner-product in eq. (10) satisfies a double integral equation which comes from the fact that the signature itself solves an integral equation. Then one uses the fundamental theorem of calculus to differentiate with respect to the two time variables to obtain the PDE.

Next, we propose a simple parametrization of this kernel.

### 3.3. Parametrization of the signature kernel

In many real-world problems the input path \( X \) contains a large number \( d \) of different channels, only some of which are relevant. For any coordinate \( \alpha \in \{1, \ldots, d\} \) and time index \( t \in [0, T] \) one can rescale each channel \( X^{(\alpha)} \) by a scalar hyperparameter \( \theta_{\alpha} \) yielding the rescaled path

\[
X^{(\alpha)}_t := (\theta_1 X^{(1)}_t, \ldots, \theta_d X^{(d)}_t).
\] (15)

From eq. (8) it is straightforward to see that the corresponding rescaled signature satisfies the following relation

\[
S_{\theta}(X)^{(\alpha_1, \ldots, \alpha_j)} := S(X^{(\theta)^{(\alpha_1, \ldots, \alpha_j)}})
\] (16)

\[
= \theta_{\alpha_1} \cdots \theta_{\alpha_j} S(X)^{(\alpha_1, \ldots, \alpha_j)}
\] (17)

for any \( \alpha_1, \ldots, \alpha_j \in \{1, \ldots, d\} \). As a result, akin to an automatic relevance determination (ARD) parametrization, the signature kernel of eq. (11) can be reparametrized as

\[
k_{\theta}(X, Y) = \sum_{j=0}^{\infty} \sum_{|\alpha| = j} S_{\theta}(X)^{\alpha} S_{\theta}(Y)^{\alpha}.
\] (18)

### 3.4. Variational Orthogonal Signature Features

In the sequel we build on the results from the previous sections to define the orthogonal signature features underlying our sparse variational inference framework for GPs on sequential data.

By Steinwart & Christmann (2008, Thm. 4.21), the reproducing kernel Hilbert space (RKHS) \( \mathcal{H} \) associated to the parametrized signature kernel \( k_{\theta} \) can be defined as

\[
\mathcal{H} = \{ g : X \mapsto \langle S_{\theta}(X), h \rangle_H \}, \quad h \in H.
\] (19)
Besides, for any two functions $g_1, g_2 \in \mathcal{H}$ such that,
\begin{align}
  g_1 : X \mapsto \langle S_\theta(X), h_1 \rangle_H, \\
  g_2 : X \mapsto \langle S_\theta(X), h_2 \rangle_H,
\end{align}
the inner product $\langle \cdot, \cdot \rangle_H$ induces the inner product on $\mathcal{H}$
\begin{equation}
  \langle g_1, g_2 \rangle_H = \langle h_1, h_2 \rangle_H. 
\end{equation}

This result relies on the unicity of $h_1$ and $h_2$ in the decomposition of $g_1$ and $g_2$, which follows from Diehl & Reizenstein (2019, Lemma 3.4) and Xu & Zhang (2007, Lemma 5).1

The key to our setup is that the set of signature features
\begin{equation}
  S^\perp = \{ S_\theta(\cdot)^\alpha : X \mapsto S_\theta(X)^\alpha \}_{\alpha=\alpha_1,\ldots,\alpha_j} 
\end{equation}
forms an orthonormal basis for the RKHS $\mathcal{H}$, i.e.
\begin{equation}
  \langle S_\theta(\cdot)^\alpha, S_\theta(\cdot)^{\alpha'} \rangle_H = \delta_{\alpha,\alpha'},
\end{equation}

An important property of the orthonormal basis $S^\perp$ is that its elements are naturally ordered. This ordering is due to the property that for any path $X \in \mathcal{X}$ the terms of the signature decay factorially (Lyons et al., 2007)
\begin{equation}
  |S(X)^\alpha| = O \left( \frac{1}{|\alpha|!} \right),
\end{equation}
as shown in Fig. 1. Hence to index the signature orthogonal features $S_\theta^0, S_\theta^1, S_\theta^2, \ldots$ we order first by increasing level $j$, and then by sorting the multi-indices $\alpha$ within a level. From eqs. (24) and (25) we define our inducing variables as orthogonal projections2 of the GP onto the first $M$

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Illustration of the first terms of the signature $S(X)$ for a 3-dimensional path $X$. Each blue circle corresponds to a signature feature $S(X)^\alpha$ with $\alpha = (\alpha_1, \ldots, \alpha_j)$. The size of the circle reflects the feature importance according to the property $|S(X)^\alpha| = O(1/|\alpha|!)$. The first feature $S^{(0)}$ which is always equal to 1 is omitted in this schematic.}
\end{figure}

\begin{equation}
  u_m = \langle f, S_\theta^m \rangle_H, \quad 1 \leq m \leq M. 
\end{equation}

With this choice of inducing variables we easily deduce the following covariances (Hensman et al., 2017)
\begin{equation}
  \mathbb{E}[u_m f(X)] = S_\theta^m(X) \quad \text{and} \quad \mathbb{E}[u_m u_m'] = \delta_{m,m'}, 
\end{equation}
which implies that the covariance matrix $C_u$ is the identity. For any path $X \in \mathcal{X}$ we use the convenient vector notation
\begin{equation}
  S_M(X) := [S_\theta^0(X), \ldots, S_\theta^M(X)] \in \mathbb{R}^M, 
\end{equation}
to obtain the approximate posterior $\mathcal{GP}(\mu, \nu)$ with mean and covariance functions defined by the following equations
\begin{equation}
  \mu(X) = S_M(X)^T \mathbf{m} \quad \text{and} \quad \nu(X,Y) = k_\theta(X,Y) - S_M(X)^T (I_M - \Sigma) S_M(Y).
\end{equation}

We note that the signature and the signature kernel can be easily computed on real time series using existing python libraries (Lyons, 2010; Reizenstein & Graham, 2018).

4. Reverse-mode automatic differentiation for the signature kernel

In order to optimize the ELBO with respect to the parameters $\theta$ one needs to take derivatives of the signature kernel $k_\theta$ of eq. (29) with respect to each of its input paths. Given that $k_\theta$ solves the PDE (14) it can be computed using appropriate PDE numerical solvers. Therefore, in theory the differentiation could be carried out by leveraging the automatic differentiation tools of modern deep learning libraries (Tensorflow, PyTorch etc.). However, backpropagating through the operations of the PDE solver can be inefficient.

Here we show that the gradients of $k_\theta$ can be computed efficiently without backpropagating through the operations
of the PDE solver as they are the solutions of a second PDE analogous to eq. (14). The ability not to rely on automatic differentiation allows for an efficient fitting of SigGPDE both in the terms of time complexity and memory cost.

4.1. Differentiating the signature kernel along the direction of a path

Consider a time series \( x \) as a collection of points \( x_i \in \mathbb{R}^d \) with corresponding time-stamps \( s_i \in \mathbb{R} \) such that
\[
x = ( (s_0, x_0), (s_1, x_1), \ldots, (s_d, x_d) )
\]
(30)
with \( s_0 < \ldots < s_d \). Every vector \( x_i \) in the sequence can be written with respect to the canonical basis of \( \mathbb{R}^d \) as
\[
x_i = \sum_{j=1}^{d} x_{i,j} e_j
\]
(31)

Let \( X : [0, T] \rightarrow \mathbb{R}^d \) be the piecewise linear interpolation of the data such that \( X_i = (t_i, x_i) \). Similarly for a second time series \( Y \) and resulting piecewise linear interpolation \( \gamma \). Recall the definition of signature kernel as
\[
k_\theta(X, Y) = k(X^\theta, Y^\theta),
\]
(32)
where \( X^\theta \) and \( Y^\theta \) are the rescaled paths of eq. (15).

By the chain rule one has that
\[
\frac{\partial k_\theta}{\partial \theta} = \frac{\partial k}{\partial X^\theta} \frac{\partial X^\theta}{\partial \theta} + \frac{\partial k}{\partial Y^\theta} \frac{\partial Y^\theta}{\partial \theta}
\]
(33)
Hence, to formulate a backpropagation algorithm in a rigorous way compatible with the TensorFlow library used in this work, we need to give meaningful to the following gradients
\[
\{ \frac{\partial}{\partial x_{i,j}} k(X, Y) \}_{i,j=1}^{\ell,d}
\]
(34)

The technical difficulty here consists in reconciling the discontinuous nature of the input path \( X \) and the discrete nature of the locations \( x_{i,j} \) where one wants to compute the gradients and given by the knots of the time series \( x \).

Next we introduce a collection of localised impulses and define the concept of directional derivative of the signature kernel along a path in order to make sense of the gradients in eq. (34). These definitions will be followed by the main result of this section, namely that the directional derivative of \( k \) solves another PDE similar to eq. (14) for the signature kernel, for which we derive an explicit solution via the technique of variation of parameters (Thm. 4.1).

Definition 1. For any \( i = 1, \ldots, \ell \) and any \( j = 1, \ldots, d \) define the localised impulse \( \gamma_{i,j} : [0, T] \rightarrow \mathbb{R}^d \) as the solution of the following ordinary differential equation (ODE)
\[
\dot{\gamma}_{i,j} = \frac{1}{\ell} e_j \mathbb{1}_{\{t \in [(i-1)/\ell, i/\ell)\}}, \quad \gamma_{i,j}(0) = 0
\]
(35)

Definition 2. For any path \( \gamma \in \mathcal{X} \) the directional derivative of the signature kernel \( k \) along \( \gamma \) is defined as
\[
k_\gamma(X, Y) := \frac{\partial}{\partial \epsilon} k(X + \epsilon \gamma, Y) \big|_{\epsilon=0}
\]
(36)
Each gradient of the signature kernel \( k_\theta \) at the knot \( x_{i,j} \) reported in eq. (34) can be identified with the directional derivative of \( k_\theta \) along the localised impulse \( \gamma_{i,j} \) of Def. 1
\[
\frac{\partial}{\partial x_{i,j}} k(X, Y) := k_{\gamma_{i,j}}(X, Y)
\]
(37)

4.2. A PDE for the gradients of the signature kernel

Recall that the signature kernel \( k_\theta \) solves the following PDE
\[
\frac{\partial^2 U}{\partial s \partial t} = (\dot{X}_s \dot{Y}_t) U
\]
(38)
Integrating both sides with respect to \( s \) and \( t \) one obtains
\[
U(s, t) = 1 + \int_{u=0}^{s} \int_{v=0}^{t} U(u, v)(\dot{X}_u \dot{Y}_v) du dv
\]
(39)
Let’s denote by \( U_\gamma : [0, T] \times [0, T] \rightarrow \mathbb{R} \) the directional derivative \( k_\gamma \) evaluated at the restricted paths \( X_{|0,s], Y_{|0,t]} \)
\[
U_\gamma(s, t) := k_\gamma(X_{|0,s], Y_{|0,t]})
\]
(40)
The combination of eqs. (39) and (40) yields the relation
\[
U_\gamma(s, t) = \frac{\partial}{\partial \epsilon} \left( k((X + \epsilon \gamma)_{|0,s], Y_{|0,t]}}) \big|_{\epsilon=0}
\]
\[
= \frac{\partial}{\partial \epsilon} \left( \int_{0}^{s} \int_{0}^{t} U(u, v) \left( \dot{X}_u + \epsilon \dot{\gamma}_u \right)^T \dot{Y}_v du dv \right) \bigg|_{\epsilon=0}
\]
\[
= \int_{0}^{s} \int_{0}^{t} \left( U_\gamma(u, v) \dot{X}_u^T \dot{Y}_v + U(u, v) \dot{\gamma}_u^T \dot{Y}_v \right) du dv
\]

Hence, differentiating the last equation first with respect to \( t \) and then \( s \) we get that the directional derivative \( k_\gamma \) of the signature kernel along the path \( \gamma \) solves the following PDE
\[
\frac{\partial^2 U_\gamma}{\partial s \partial t} = (\dot{X}_s^T \dot{Y}_t) U_\gamma + (\dot{\gamma}_s^T \dot{Y}_t) U
\]
(41)
with boundary conditions
\[
U_\gamma(0, \cdot) = 0, \quad U_\gamma(\cdot, 0) = 0
\]
(42)
As a result, the gradients in eq. (34) of the signature kernel with respect to each of its input paths can be computed in a single call to a PDE solver, which concatenates the original state and the partial derivatives (41) into a single vector. Each partial derivative follows the dynamics of (41) where one replaces the direction \( \gamma \) by the relevant localised impulse \( \gamma_{i,j}, \tau_{i,j} \) for \( X \) and \( Y \) respectively. We outline the resulting procedure in Alg. 1, where the concatenated partial derivatives are denoted by \( U_\gamma(s, t) \). Note that to optimize the ELBO we only need to differentiate \( k(X, X) \), which is the case presented in the algorithm. The generalization to the case \( k(X, Y) \) is straightforward using the chain rule.
Algorithm 1 Backpropagation for $k_{\theta}(X,X)$ via PDE (41)

1: **Input:** Path $X$, localised impulses $\gamma = \{\gamma_{i,j}\}$ fully determined by the time series $x$.
2: $u_{0,0} : [1,0,\ldots,0]$, $u_{0,0} = [1,0,\ldots,0]$ \hfill // Boundary conditions for the augmented state
3: \textbf{def} aug_dynamics$([U(s,t), U_\gamma(s,t)]$, $s,t$):
4: \textbf{return} $\begin{bmatrix} X_T^s X_t U(s,t), X_T^s X_t U_\gamma(s,t) + \hat{\gamma}_s X_t U(s,t) \end{bmatrix}$ \hfill // Dynamics for the augmented state
5: $[U(T,T), U_\gamma(T,T)] = \text{PDESolve}(u_{0,0}, u_{0,0}, \text{aug\_dynamic}, T,T)$ \hfill \// Keep the solutions at each $(s,t)$
6: **Output:** $2 \cdot U_\gamma(T,T)$ \hfill // Gradients of the kernel at the knots of $X$

Algorithm 2 Backpropagation for $k_{\theta}(X,X)$ via variation of parameters (Thm. 4.1)

1: **Input:** Path $X$, localised impulses $\gamma = \{\gamma_{i,j}\}$ fully determined by the time series $x$
2: $u_{0,0} : [1,0,\ldots,1]$, $u_{0,0} = [1,0,\ldots,1]$ \hfill // Boundary conditions for the augmented state
3: \textbf{def} aug_dynamics$([U(s,t), \tilde{U}(s,t)]$, $s,t$):
4: \textbf{return} $\begin{bmatrix} X_T^s X_t U(s,t), X_T^s X_t \tilde{X}_{T-t} \tilde{U}(s,t) \end{bmatrix}$ \hfill // Dynamics for the augmented state
5: $[U, \tilde{U}] = \text{PDESolve}(u_{0,0}, u_{0,0}, \text{aug\_dynamic}, T,T)$ \hfill \// Simple final TensorFlow operations
6: $U_\gamma = \text{tf.sum}(U \cdot \tilde{U} \cdot \gamma X)$ \hfill // Gradients of the kernel at the knots of $X$
7: **Output:** $2 \cdot U_\gamma$

4.3. An explicit solution by variation of parameters

From this second PDE (41) we derive the following theorem (proved in Appendix A), that allows to compute the directional derivative $k_\gamma$ of the signature kernel directly from its evaluations at $X$, $Y$ and at $\overline{X}$, $\overline{Y}$, where $\overline{X}$, $\overline{Y}$ are respectively the paths $X$, $Y$ reversed in time.

**Theorem 4.1.** For any $\gamma \in \mathcal{X}$ the directional derivative $k_\gamma(X,Y)$ of the signature kernel along the path $\gamma$ satisfies the following relation

$$k_\gamma(X,Y) = \int_0^T \int_0^T U(s,t)\tilde{U}(T-s, T-t)(\gamma_s^T \gamma_t)dsdt$$

where $\tilde{U}(s,t) = k(\overline{X}|_{[0,s]}, \overline{Y}|_{[0,t]})$ and where $\overline{X}$, $\overline{Y}$ are respectively the paths $X$, $Y$ reversed in time.

The full backpropagation procedure is described in Alg. 2.

5. Related work

In this section we expand on the material presented in Sec. 2, focusing on the most recent approaches to scalable GPs on $\mathbb{R}^d$ with VOFs and on sparse GPs for sequential data.

**Variational Fourier Features** In Hensman et al. (2017) the inducing variables are defined for scalar input $\mathcal{X} = \mathbb{R}$ as projections of the GP-sample onto the truncated Fourier basis. This type of inducing variables can be constructed for GPs with Matérn-type kernels. Although the resulting covariance matrix of the inducing variables is not diagonal, it can be decomposed into the sum of a diagonal matrix and rank one matrices. As a result it can be inverted using the Woodbury identity, which makes it possible to scale GP inference on $\mathbb{R}$. The generalization to GPs on $\mathbb{R}^d$ is done by taking the outer product of the Fourier basis on $\mathbb{R}$.

**Eigenfunction inducing features** Closest to our work is the eigenfunction inducing features developed by Burt et al. (2020a), where the inducing variables are also defined as projections of the GP-sample onto an orthogonal basis of functions for the RKHS associated with the GP kernel. This relies on a Mercer’s expansion of the kernel. From here one identifies this orthogonal basis functions by solving an eigendecomposition problem. For example Dutordoir et al. (2020) map the input data to the hypersphere $S^{d-1} \subset \mathbb{R}^d$ and then show that spherical harmonics form an orthogonal basis for RKHS associated to zonal kernels defined on $S^{d-1}$.

**GPs with signature covariances** Toth & Oberhauser (2020) propose a different sparse GP inference framework for sequential data with signature covariances (GPSig). In this work the inducing variables are either taken to be inducing sequences (IS) in the original input space (GPSig-IS) of sequences or inducing tensors (IT) in the corresponding feature space (GPSig-IT). The chosen covariance function is an approximation of the signature kernel based on truncating the signature to a finite level $n$. For GPSig-IT, this truncation makes the feature space finite dimensional and allows to optimize inducing tensors defined over such truncated space. Unlike our method, the inducing tensors are additional variational parameters to optimize. The covariance matrix $C_{uu}$ is dense and its inversion incurs an additional $O(M^3)$ cost. In Table 1 we compare the computational complexities of GPSig-IT, GPSig-IS and SigGPDE. A similar table for the memory complexity can be found in Appendix C.
6. Experiments

In this last section, we benchmark SigGPDE against GPSig-IT and GPSig-IS from Toth & Oberhauser (2020) on various multivariate time series classification tasks. For GPSig-IS, we use inducing sequences of length $\ell = 5$ as recommended in Toth & Oberhauser (2020). We highlight how SigGPDE performs competitively in terms of accuracy and uncertainty quantification but with a significant speed-up in the fitting compared to the other baselines.

We use a mixture of UEA & UCR time series datasets (timeseriesclassification.com) and real world data for the final example. In the latter we discuss how the predictions provided by SigGPDE can be interpreted in a natural way via the interpretability of the interated integrals defining the signature and discussed in Sec. 3.1.

We measure the classification accuracy on the test set, assess the uncertainty quantification with mean negative log-predictive probabilities (NLPP) and report the runtime per iteration. For each dataset all models are trained 3 times using a random training-validation split. The validation split is used to monitor the NLPP when optimizing the hyper-parameters of the models. Further details on the training procedure can be found in Appendix B. All code is written in TensorFlow using GPFlow (De G. Matthews et al., 2017).

6.1. Classifying digits in sequential MNIST

We start with a handwritten digit classification task, where writers were asked to draw the digits from 0 to 9. The instances are made up of 2-d trajectories of the pen traced across a digital screen. The trajectories are of length $\ell = 8$. The training and test sets are of size 7,494 and 3,498 respectively. We made use of $M = 500$ inducing features. In the results reported in Table 2, SigGPDE achieves even better accuracy and NLPP than the GPSig baselines, whilst being almost twice as fast than GPSig-IT.

<table>
<thead>
<tr>
<th>Model</th>
<th>Mean Acc.</th>
<th>NLPP</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPSig-IS</td>
<td>97.42 ± 0.17</td>
<td>0.096 ± 0.005</td>
<td>0.186 (s/iter)</td>
</tr>
<tr>
<td>GPSig-IT</td>
<td>96.66 ± 0.59</td>
<td>0.115 ± 0.018</td>
<td>0.036 (s/iter)</td>
</tr>
<tr>
<td>SigGPDE</td>
<td>97.73 ± 0.13</td>
<td>0.085 ± 0.001</td>
<td>0.022 (s/iter)</td>
</tr>
</tbody>
</table>

6.2. Detecting whale call signals

In this example the task is to classify audio signals and distinguish one emitted from right whales from noise. The dataset (called RightWhaleCalls in the UEA archive) contains 10,934 train cases and 5,885 test cases. The signals are one-dimensional, sampled at 2kHz over 2 seconds, hence of length 4,000. We tackle this problem as a multivariate time series classification task, by taking the spectrogram of the univariate audio signal. The resulting streams are made of 29 channels corresponding to selected frequencies and are 30 time steps long. The results in Table 3 are obtained with $M = 700$ and show the significant speed-up of SigGPDE by almost one order of magnitude compared to GPSig. This speed-up is compensated by a minimal decrease in performance both in terms of accuracy and NLPP.

Table 1. Comparison of time complexities. $M$ is the number of inducing variables, $N$ the batch size, $d$ the number of channels in the time series, $\ell$ the length of the sequences, $n$ the truncation level (for GPSig-IT and GPSig-IS) and $\tilde{\ell}$ the length of the inducing sequences. The last line of the table corresponds to linear algebra operations including matrix multiplication and matrix inversion.
### Table 3. Classification for whale call signals

<table>
<thead>
<tr>
<th>Model</th>
<th>Mean Acc.</th>
<th>NLPP</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPSig-IS</td>
<td>86.97 ± 0.11</td>
<td>0.367 ± 0.005</td>
<td>0.438 (s/iter)</td>
</tr>
<tr>
<td>GPSig-IT</td>
<td>87.70 ± 0.42</td>
<td>0.357 ± 0.003</td>
<td>0.048 (s/iter)</td>
</tr>
<tr>
<td>SigGPDE</td>
<td>86.76 ± 0.36</td>
<td>0.382 ± 0.002</td>
<td>0.008 (s/iter)</td>
</tr>
</tbody>
</table>

#### 6.3. Large scale classification of satellite time series

This is our large scale classification example on 1 million time series. The time series in this dataset represent a vegetation index, calculated from remote sensing spectral data. The 24 classes represent different land cover types (Petitjean et al., 2012). The aim in classifying these time series of length $\ell = 46$ is to map different vegetation profiles to different types of crops and forested areas. Due to the sheer size of this dataset we only compare SigGPDE to GPSig-IT as GPSig-IS is not scalable to such large dataset. In Fig. 3 we report the accuracy, time per iteration and ELBO by progressively increasing the number of inducing variables. Compared to SigGPDE, GPSig-IT has additional variational parameters, namely the inducing tensors. This extra flexibility explains the better performances of GPSig-IT when few inducing variables are used. However, as the number of inducing features increases, SigGPDE catches up and outperforms its competitor in all monitored metrics.

#### 6.4. Weather forecast

In this last example we will be using a dataset of climatic variables recorded by the Max Planck Institute for Biogeochemistry in the weather stations of WS Beutenberg and WS Saaleauve from 2004-2020. The data consists of 7-dimensional time series recorded once per 10 minutes where each channel represents a weather feature such as temperature, pressure, humidity etc. The goal is to predict whether it will rain over the next hour from the trajectory of all other features in the preceding 6 hours. To obtain binary labels for the classification task we set the label to 1 if the precipitation is larger than 1mm and to 0 otherwise. The inference mechanism is depicted on Fig. 2.

A key feature proper to our model SigPDE is its interpretability. Looking at the variational mean vector $m$ in eq. (29), we can extract the terms with highest relevance learned by the model. As discussed in Sec. 3.1, thanks to the corresponding signature features it is possible to infer which signature features used by the GP are more responsible for the produced outcome. The most relevant predictive features for this weather forecast experiment are represented in Fig. 4.

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3https://www.bgc-jena.mpg.de/wetter/
7. Conclusion

In this paper we have developed SigGPDE, a framework to perform variational inference for GP models on sequential data with orthogonal signature features. Firstly, we constructed inducing variables so that their covariance matrix is diagonal. Secondly, we showed that the gradients of the signature kernel are solutions of a hyperbolic PDE. As a result the ELBO is cheap to evaluate as gradient descent does not require backpropagating through the operations of the PDE solver. We benchmarked SigGPDE against the state-of-the-art GPSig on different time series classification tasks, showing a significant speed up and similar performance.

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