# Gaussian Processes for Probabilistic Estimates of Earthquake Ground Shaking: A 1-D Proof-of-Concept

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## Abstract

Estimates of seismic wave speeds in the Earth (seismic velocity models) are key input parameters to earthquake simulations for ground motion prediction. Owing to the non-uniqueness of the seismic inverse problem, typically many velocity models exist for any given region. The arbitrary choice of which velocity model to use in earthquake simulations impacts ground motion predictions. However, current hazard analysis methods do not account for this source of uncertainty. We present a proof-of-concept ground motion prediction workflow for incorporating uncertainties arising from inconsistencies between existing seismic velocity models. Our analysis is based on the probabilistic fusion of overlapping seismic velocity models using scalable Gaussian process (GP) regression. Specifically, we fit a GP to two synthetic 1-D velocity profiles simultaneously, and show that the predictive uncertainty accounts for the differences between the models. We subsequently draw velocity model samples from the predictive distribution and estimate peak ground displacement using acoustic wave propagation through the velocity models. The resulting distribution of possible ground motion amplitudes is much wider than would be predicted by simulating shaking using only the two input velocity models. This proof-of-concept illustrates the importance of probabilistic methods for physics-based seismic hazard analysis.

### 1 Introduction

Seismic velocity models — estimates of the Earth's seismic wave speeds — underpin earthquake ground motion prediction in seismic hazard analysis, as they are key inputs to wave equation solvers. They continue to be produced at different resolutions and scales, stemming from different methods (e.g., tomography [\[1\]](#page-5-0), reflection surveys [\[2\]](#page-5-1)). The seismic inverse problem is ill-posed as there are not enough data to constrain a unique true Earth model [\[3\]](#page-5-2). As such, many overlapping velocity models exist for a given region. Consequently, the choice of which velocity model to use in ground motion prediction is often arbitrary. Nevertheless, it has a significant impact on the results as different models have different structures, length scales, and amplitudes.

The key output of seismic hazard analysis is an estimate of peak ground motion, to assess potential infrastructure damage and inform earthquake engineering. Most commonly, empirical ground motion

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models (GMMs) [\[4,](#page-5-3) [5\]](#page-5-4) are used to predict the median and uncertainty of a ground motion parameter (e.g., peak ground displacement — PGD) for earthquake scenarios [\[6\]](#page-5-5). They make rapid predictions, but drastically simplify the underlying physical processes. Importantly, they approximate the effect of seismic velocities on ground motion, typically only using the average shear wave velocity in the uppermost 30 m [\[7\]](#page-5-6). GMMs are thus limited in accuracy and reliability. An alternative is to simulate earthquake scenarios in 3-D by solving the wave equation and extracting PGD estimates, requiring 3-D seismic velocity information as input. However, there are two issues: (i) simulating many earthquakes is computationally costly, and (ii) choices of input parameters are subjective, including the input velocity model. To address the first issue, recent advances in machine learning have begun accelerating wave propagation methods [\[8–](#page-5-7)[11\]](#page-6-0). However, current physics-based hazard analysis workflows do not consider inconsistencies between velocity models. This omits a key source of uncertainty, given that predicted ground motion can be drastically impacted by velocity structure. One possible solution is to fuse different velocity models, and use the output in earthquake simulations. Unfortunately, existing methods for velocity model fusion [e.g., [12–](#page-6-1)[14\]](#page-6-2) typically do not produce probabilistic outputs, limiting their ability to account for differences between velocity models.

In this study, we propose a workflow to account for inconsistencies between seismic velocity models in ground motion prediction. Our method is based on the probabilistic fusion of velocity models using Gaussian processes (GPs), and estimates uncertainties owing to differences between them. We then produce probabilistic ground motion predictions with respect to these uncertainties by drawing velocity model samples from the GP predictive distribution, simulating acoustic wave propagation using each sample, and extracting the PGD predictions. We illustrate that such a probabilistic method is necessary to capture the spread of possible ground motion scenarios.

Our key contributions are as follows: (i) We present a workflow for probabilistic earthquake ground motion prediction that accounts for inconsistencies between seismic velocity models. (ii) We demonstrate the capability of scalable GPs for the probabilistic fusion of different estimates of the same physical parameter, through a synthetic example using 1-D seismic velocity models. The code for this work is written in Python and is available at Scivier et al. [\[15\]](#page-6-3).

#### <span id="page-1-1"></span>2 Gaussian processes and data fusion

GPs [\[16\]](#page-6-4) are a class of non-parametric models for defining a distribution over function spaces. They are widely used for regression, providing robust uncertainty quantification and predictive performance. Unfortunately, exact GP regression is limited in scalability owing to a computational cost of  $\mathcal{O}(n^3)$ , where  $n$  is the number of data points. Despite the small datasets used in this study, seismic datasets of realistic size can have  $n \sim 10^6 - 10^7$ . To overcome this, approximate GP methods have been developed [\[17\]](#page-6-5). One popular method for scalable GP inference is the sparse variational Gaussian



<span id="page-1-0"></span>Figure 1: Comparison of SVGPR and PPGPR for the probabilistic fusion of seismic velocity models. (a) shows the input synthetic 1-D seismic velocity profiles with depth. (b) and (c) show the fusion results of SVGPR and PPGPR, and (d) shows the 200 function samples drawn from the PPGPR predictive distribution used in [Section 4.](#page-3-0) The shading in (b) and (c) show the SVGP posterior predictive distribution,  $q_{\text{SVGP}}(\mathbf{y}_*)$ , and the PPGPR latent predictive distribution,  $q_{\text{PPGPR}}(\mathbf{f}_*)$ , respectively, in terms of distance from the predictive means in standard deviations.

process (SVGP) [\[18,](#page-6-6) [19\]](#page-6-7), which applies variational inference to fit GPs. SVGPs introduce a set of inducing variables to approximate the full dataset using a smaller set of points  $m \ll n$ . Thus, the computational cost is reduced to  $\mathcal{O}(nm^2 + m^3)$ , making SVGPs practical to apply to large 2-D and 3-D datasets.

In this study, we use scalable GP regression for fusing seismic velocity models by fitting a GP to multiple datasets simultaneously. A key advantage of GPs is their modelling of covariance structure, which enables samples matching the spatial patterns of the input data to be drawn from the predictive distribution.

We aim to model inconsistencies between velocity estimates as uncertainty in the GP predictive distribution. Thus, we need to understand the form of the predictive variance in the SVGP model. Below we provide a brief summary of the key SVGP results to highlight the relevant context for our work. The interested reader is referred to Titsias [\[18\]](#page-6-6), Matthews et al. [\[19\]](#page-6-7), Jankowiak et al. [\[20\]](#page-6-8), and Murphy [\[21\]](#page-6-9) for full derivations and explanations. The inputs are the training set (coordinates of the input velocity models), X, the inducing point locations, Z, and the points at which we wish to predict,  $X_{*}$ . Then  $f_{X}$ ,  $f_{Z}$ ,  $f_{*}$  are the (unknown) velocity values that we predict at these locations; and y are the observed data (input velocity values). SVGP-based methods approximate the joint posterior as  $q(\mathbf{f}_*,\mathbf{f}_\mathbf{X},\mathbf{f}_\mathbf{Z}) = p(\mathbf{f}_*,\mathbf{f}_\mathbf{X} | \mathbf{f}_\mathbf{Z}) q(\mathbf{f}_\mathbf{Z}),$  where  $p(\mathbf{f}_*,\mathbf{f}_\mathbf{X} | \mathbf{f}_\mathbf{Z})$  is calculated exactly [\[21\]](#page-6-9). The variational distribution is  $q(f_z) = \mathcal{N}(f_z | m, S)$ , where m and S are (learned) variational parameters. The predictive distribution over the underlying function (at the target points)  $f_*$  is given by

$$
q\left(\mathbf{f}_{*}\right) = \int p\left(\mathbf{f}_{*} \mid \mathbf{f}_{\mathbf{Z}}\right) q\left(\mathbf{f}_{\mathbf{Z}}\right) d\mathbf{f}_{\mathbf{Z}}
$$
\n
$$
= \mathcal{N}\left(\mathbf{f}_{*} \mid \mu_{*}, \sigma_{\mathbf{f}}\left(\mathbf{x}_{*}\right)^{2}\right),\tag{1}
$$

where,

$$
\mu_* = \mathbf{K}_{*,\mathbf{Z}} \mathbf{K}_{\mathbf{Z},\mathbf{Z}} \mathbf{K}_{\mathbf{Z}} \mathbf{K}_{\mathbf{Z}} \mathbf{K}_{\mathbf{Z}} \mathbf{K}_{\mathbf{Z}} \mathbf{K}_{\mathbf{Z} \mathbf{Z}} \mathbf{K}_{\mathbf{Z
$$

with e.g.,  $\mathbf{K}_{*,\mathbf{Z}} = k(\mathbf{X}_*,\mathbf{Z})$ , and  $k(\cdot,\cdot)$  is the (chosen) covariance function. Assuming a Gaussian likelihood, measurements (at the target points)  $y_*$  are related to the underlying function (at the target points)  $f_*$  as  $p(y_* | f_*, \sigma_y^2) = \mathcal{N}(y_* | f_*, \sigma_y^2 I_*)$ , where  $\sigma_y^2$  is the (learned) observational noise variance. Thus, the predictive distribution over  $y_*$  is,

<span id="page-2-0"></span>
$$
q(\mathbf{y}_{*}) = \int p(\mathbf{y}_{*} | \mathbf{f}_{*}, \sigma_{y}^{2}) q(\mathbf{f}_{*}) d\mathbf{f}_{*}
$$
  
=  $\mathcal{N}\left(\mathbf{y}_{*} | \mu_{*}, \sigma_{\mathbf{f}}(\mathbf{x}_{*})^{2} + \sigma_{y}^{2} \mathbb{I}_{*}\right).$  (2)

The predictive variance at a target point  $x_{*,i}$  is thus the sum of input-dependent variance over the underlying function,  $\sigma_f(x_{*,i})^2$ , and observational noise,  $\sigma_y^2$ : Var  $(x_{*,i}) = \sigma_f(x_{*,i})^2 + \sigma_y^2$ . Despite this symmetry in the predictive variance, Jankowiak et al. [\[20\]](#page-6-8) highlight that the typical SVGP objective function (variational ELBO) targets only large  $\sigma_y^2$  – often resulting in  $\sigma_y^2 \gg$  $\sigma_f(\mathbf{x}_{*,i})^2$ , which we see from the data-fit term:  $\mathcal{L}_{\text{SVGP}} \supset -\frac{1}{2\sigma_y^2} |\mathbf{y}_i - \mu_{\mathbf{X},i}|^2$ . This means we would model disagreements between different velocity models as observational noise. Given the degree of disagreement varies spatially,  $\sigma_y$  would need to be input-dependent [e.g., [22,](#page-6-10) [23\]](#page-6-11). However, this would result in noisy samples in the predictive distribution (see [Fig. 1b](#page-1-0)) — making them useless for downstream tasks.

We instead wish to model inconsistencies between velocity models as input-dependent uncertainty in the underlying physical process (i.e.,  $\sigma_f(\mathbf{x})$ ). To enable this, we use the parametric predictive GP regression (PPGPR) model [\[20\]](#page-6-8). PPGPR is a variation of SVGP with an objective function that directly targets the predictive distribution [\(Eq. \(2\)\)](#page-2-0). Notably, the PPGPR objective encourages large  $\sigma_f(\mathbf{x}_*)^2$ , as seen from the data-fit term:  $\mathcal{L}_{\text{PPGPR}} \supset -\frac{1}{2} \frac{1}{\sigma_y^2 + \sigma_f(\mathbf{x}_i)^2} |\mathbf{y}_i - \mu_{\mathbf{X},i}|^2$ . Contrary to SVGP,

this typically results in  $\sigma_f(x_{*,i})^2 \gg \sigma_y^2$ . Therefore, we can choose to ignore  $\sigma_y^2$  and use the PPGPR latent predictive distribution,  $q(f_*)$ .

#### 3 Fusing synthetic seismic velocity models

We present a proof-of-concept demonstrating the applicability of PPGPR for the probabilistic fusion of two synthetic 1-D seismic velocity models. We note that we do not consider uncertainties attached to input velocity models in this study (i.e., the input models themselves are not probability distributions). Two datasets,  $s_1$  and  $s_2$ , are sampled ( $n = 25$  data points, each) from a GP prior, using radial basis function (RBF) kernels with different length scales. The samples have different coordinates in an overlapping region. The first model is set as the first sample,  $m_1 = s_1$ . The second model is a weighted superposition of the two samples,  $m_2 = \frac{2}{3}s_1 + \frac{1}{3}s_2$ , to create larger-scale similarities and smaller-scale differences — typical of different seismic velocity models. [Fig. 1a](#page-1-0) shows the input velocity models, which are 1-D profiles with respect to depth.

At training time, the input models are concatenated and used to condition the GP as a single dataset. For regression, we employ both PPGPR and SVGPR for comparison, using scaled RBF kernels,  $m = 20$  inducing points (with learned locations), and Gaussian likelihoods. The models are trained using the Adam optimiser [\[24\]](#page-6-12) and identical hyperparameters (i.e., learning rate and number of iterations). Hyperparameters are chosen through trial-and-error, and full hyperparameter details are provided in the code [\[15\]](#page-6-3). Training the models takes one minute for each method, on a laptop using an NVIDIA T500 2GB GDDR6 GPU.

[Fig. 1b](#page-1-0) and c show the results of SVGPR and PPGPR, respectively, on the two velocity models. As discussed in [Section 2,](#page-1-1) optimising the SVGP objective results in  $\sigma_{\text{obs}}^2 \gg \sigma_f(x)^2$ , making it unsuitable for this task owing to a lack of input-dependence on the predictive variance and noisy function samples. On the other hand, PPGPR performs well, with predictive samples appearing to reflect the spatial patterns of the input models. In an ideal case of maximum likelihood estimation for fitting a univariate Gaussian distribution to two observations,  $y_1$  and  $y_2$ , the resulting distribution is  $\mathcal{N}\left(\mu=\frac{1}{2}(y_1+y_2),\sigma^2=\left(\frac{y_1-y_2}{2}\right)^2\right)$  [\[25\]](#page-7-0). In our example, we therefore expect the  $\pm 1\sigma$  contours to approximately follow each of the input velocity models. We interpolated  $m_1$  and  $m_2$  at the test points using cubic splines and calculated the root mean square error (RMSE) of the SVGP and PPGPR predictive means and variances with respect to the above ideal result. The RMSEs on  $\mu_{\text{SVGP}}$ and  $\sigma_{\text{SVGP}}^2$  were 0.243 and 0.098 (in wave speed units), respectively, while for  $\mu_{\text{PPGPR}}$  and  $\sigma_{\text{PPGPR}}^2$ the RMSEs were 0.045 and 0.012 (in wave speed units). The PPGPR predictive distribution thus appropriately quantifies the uncertainty on the knowledge of seismic velocities in the region. Most importantly for our application, the covariance structure of the data is modelled. This enables the drawing of samples from the predictive distribution that match the spatial patterns of the input data. Despite only fusing two velocity models here, our approach is generally applicable for fusing any number of input datasets.

## <span id="page-3-0"></span>4 Probabilistic ground motion prediction

We propose a proof-of-concept workflow for propagating the predicted uncertainty on seismic velocities through simulations of the acoustic wave equation, to produce probabilistic ground motion predictions. First, we draw 200 function samples from the PPGPR latent predictive distribution [\(Fig. 1d](#page-1-0)). Then for each sample, we simulate the 1-D acoustic wave equation for displacement,  $\bf{u}$ , with a Ricker wavelet as the earthquake source, using a finite difference scheme (6 s for 200 simulations on a laptop — vectorised over velocity models). At the surface, we implement a free-surface boundary condition (i.e., the acoustic pressure  $p = 0$ ). At depth, we implement an absorbing boundary layer according to Chern [\[26\]](#page-7-1). [Fig. 2a](#page-4-0)–f shows snapshots of the displacement field at various time steps in one of the simulations. For each simulation, we record the peak ground displacement at the surface (i.e., PGD; depth  $= 0$ ), producing one PGD estimate per simulation (i.e., per sample velocity model). [Fig. 2g](#page-4-0) shows a histogram of the recorded PGD measurements from the simulations. Additionally, we ran simulations using interpolated versions of  $m_1$  and  $m_2$  and marked the resulting PGD measurements in [Fig. 2g](#page-4-0), to investigate how much information is gained by running simulations for many velocity model samples. Clearly, the PGD measurements for  $m_1$  and  $m_2$  do not account for the spread of possible ground motions, given the degree of knowledge of seismic velocities in this example. Despite being 1-D, our work already shows that it is not possible to approximate the full distribution of possible ground motions using only two velocity models.



<span id="page-4-0"></span>Figure 2: Wavefield snapshots and probabilistic ground motion prediction. (a)–(f) shows wavefield displacement snapshots at increasing time steps for one simulation. Each panel includes the source location (yellow star), the maximum PGD up to that time step (yellow dot), and the underlying velocity model of the simulation. The shaded region indicates where an absorbing boundary layer is applied  $[26]$ .  $(g)$  shows a histogram of the PGD measurements from the simulations, and highlights the median and middle 70% of predictions. Also shown are the PGD measurements resulting from simulations using just  $m_1$  or  $m_2$  as input.

## 5 Limitations

This work is a proof-of-concept and can be extended in several ways. For example, we do not account for data with varying length scales or structure, or address kernel design or choice, which would be required for dealing with real seismic datasets. For real-world applicability, it will also be important to extend our workflow from 1-D to 2-D and 3-D, and to solve the elastic wave equation instead of the acoustic wave equation. Working with 3-D velocity models would add complexity, but in principle it would consist of changing the GP coordinate space from 1-D to 3-D. There are many optimised 3-D seismic wave propagation codes that could then be used for the simulation component of the workflow [e.g., [11,](#page-6-0) [27](#page-7-2)[–29\]](#page-7-3). If the input velocity models had different spatial densities of data points, the objective function would be weighted towards one of them, and the result would be skewed. This can be readily solved by weighting their contributions to the objective. Additionally in this work, the synthetic input velocity models do not have uncertainties, which we plan to incorporate in the future (i.e., the input velocity models would themselves be probability distributions).

In this study, we were unable to compare our method with existing methods for velocity model fusion. Current methods are generally designed for enhancing larger-scale velocity models using smaller-scale models — and are thus not applicable when the models occupy the same domain and/or have similar spatial data density, as in our case. Additionally, existing methods typically do not produce probabilistic outputs, meaning it is not possible to compare them in the ground motion prediction component of the study [\(Section 4\)](#page-3-0).

## 6 Conclusion

Seismic velocity models underpin predictions of earthquake ground motion. Current methods for physics-based seismic hazard analysis do not account for inconsistencies between existing velocity models. In this study, we present a proof-of-concept workflow for probabilistic ground motion prediction that takes this source of uncertainty into account. Firstly, we demonstrate the applicability of scalable GP regression to the probabilistic fusion of input velocity models, showing that inconsistencies between velocity models can be modelled as predictive uncertainty. This provides access to any number of plausible velocity models for the region by drawing samples from the predictive distribution. Secondly, we build up a distribution of possible ground motion scenarios for the family of possible velocity models according to the GP predictive distribution. Our results show a much wider spread of possible peak ground motions than would be predicted by simulating earthquake scenarios using just the input velocity models themselves. We thus highlight the value of using probabilistic methods, such as the one presented here, in physics-based seismic hazard analysis to account for differences between velocity models.

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Reproducibility The code used to produce the results and figures in this study is available at Scivier et al. [\[15\]](#page-6-3).

Software The code for this work was written in Python, and used the open-source software libaries JUPYTER NOTEBOOKS [\[30\]](#page-7-4), BINDER [\[31\]](#page-7-5), NUMPY v2.1.3 [\[32\]](#page-7-6), SCIPY v1.14.1 [\[33\]](#page-7-7), MATPLOTLIB v3.9.2 [\[34\]](#page-7-8), PYTORCH v2.5.1 [\[35\]](#page-7-9), and GPYTORCH v1.13 [\[36\]](#page-7-10).

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