Introduction to Gaussian Processes - RAW

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Abstract

The Gaussian processes have proven to be a powerfull framework for robust estimation and a flexible model for non-linear regression, case which will be the main object of this work, with some implementations of real situations.

Applications in disease mapping

The model 1.1

There's several applications using GP and here we'll resume an example for disease mapping presented by [Vanhatalo et al., 2010]. Then let's assume that our phenomenon is ruled by an function f. But, we interested in the distribution of them, considering the approach presented in this work. So, we may say that we evaluated each observation y_i is Poisson distributed and its mean is given by an unknown function f_i . With this we assume that our observations and our functions are independent and then we can evaluate our joint distribution for the likehood by the product of each one [Vanhatalo, 2010].

$$\begin{cases} y_1, y_2, \dots, y_n \sim \prod_{i=1}^n Poisson\left(e_i \exp(f_i)\right) \\ f(\mathbf{x})|\theta \sim \mathcal{GP}\left(m(\mathbf{x}), k(\mathbf{x}, \mathbf{x}'|\theta)\right) \\ \theta \sim \text{half-t}(\nu, A)^* \end{cases}$$
(1.1a)

$$f(\mathbf{x})|\theta \sim \mathcal{GP}(m(\mathbf{x}), k(\mathbf{x}, \mathbf{x}'|\theta))$$
 (1.1b)

$$\theta \sim \text{half-t}(\nu, A)^*$$
 (1.1c)

In this case, we used the Poisson distribution for the likelihood because the nature of the process. The phenomenon here is the relative risk of death μ in a region of the country. So, if we consider y the counting of deaths on this region, we can model the phenomenon with a Poisson process [Vanhatalo et al., 2010].

For numerical reasons, we transform $f = \log(\mu)$. Finally, we assume an uncertainty over the parameters θ of the kernel functions too, then, our hierarchical model stays for the posterior distribution as

$$p(\mathbf{f}|\mathbf{y}, \mathbf{x}) \propto \int p(\mathbf{y}|\mathbf{f}) \mathcal{GP}(\mathbf{f}|m(\mathbf{x}), k(\mathbf{x}, \mathbf{x}'|\theta)) p(\theta) d\theta.$$
 (1.2)

This function isn't analytically tractable because of the Poisson process, but it is possible its evaluation with approximation methods.

^{*}The values ν and A are not arbitrary, but deterministic [Vanhatalo et al., 2010].

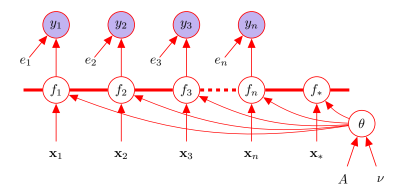


Figure 1: Graphical model for the GP for regression. Colored circles represent observed variables and whited ones represent the unknowns. The thick horizontal bar in f_i node represents a set of fully connected nodes of the Gaussian field. Note that an observation y_i is conditionally independent of all other nodes given the corresponding latent variable, f_i . Because of the marginalization property of GPs addition of further inputs, \mathbf{x} , latent variables, f, and unobserved targets, y_* , does not change the distribution of any other variables.

References

[Vanhatalo, 2010] Vanhatalo, J. (2010). Speeding up the inference in Gaussian process models. PhD thesis, Aalto University School of Science and Technology, Faculty of Information and Natural Sciences, Department of Biomedical Engineering and Computational Science, Aalto.

[Vanhatalo et al., 2010] Vanhatalo, J., Pietiläinen, V., and Vehtari, A. (2010). Approximate inference for disease mapping with sparse gaussian processes. <u>Statistics in medicine</u>, 29:1580–607.