USING GAUSSIAN PROCESS REGRESSION TO DENOISE IMAGES AND REMOVE ARTEFACTS FROM MICROARRAY DATA

by

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A thesis submitted in conformity with the requirements for the degree of Master of Science Graduate Department of Computer Science University of Toronto

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Abstract

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Natural image denoising is a well-studied problem of computer vision, but still eludes sufficiently good solutions. Removing spatial artefacts from DNA microarray data is of significant practical importance to biologists, but overly simple methods are commonly used. We show that the two seemingly disparate problems of image denoising and spatial artefact removal from DNA microarrays can be solved using similar algorithms when the problems are viewed as a decomposition of the data into spatially-independent and spatially-dependent components. We discuss previously used methods and how Gaussian processes for regression (GPR) may be used to solve both problems. We present a new method employing a novel covariance function for use in GPR in the special case of twodimensional grid data. Finally, we show that our new method compares well with popular previous methods in both image denoising and microarray artefact removal applications.

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Chapter 1

Introduction

1.1 Noisy grid data

This thesis primarily concerns itself with two-dimensional grid data $\{(x_i, y_i)\}_{i=1}^N \subset \mathbb{R}^2 \times \mathbb{R}$, where x_i describes the location of the *i*th data point on a finite, uniformly-spaced grid, and y_i is its value. We also assume the grid is *complete* in that every point of the grid has a data point in our data set. In the spatial statistics literature, this is sometimes called regular lattice data [4]. A common example of complete grid data is a rectangular image, where the locations are the pixel row and column indices and the values are the pixel intensities. Our goal will be to explain each y_i as a decomposition into two components, with one of them, n_i , being zero-mean, independent, and identically distributed

$$y_i = s_i + n_i \tag{1.1}$$

Of course, further assumptions on the behaviour of s and n are necessary to make the problem meaningful. In our work the s component will have some dependence on location, whereas the n component will be independent of location. We refrain from the terms "signal" and "noise" when referring to these components because which component is actually the noise depends on the context. Instead we think of s and n as short for *spatial* (or dependent) and *independent*, respectively. Although there are potentially many applications, we will focus on two:

- 1. DNA microarray spatial artefact removal: y_i is the (log of the) expression level for probe *i* on a DNA microarray, s_i is the contribution from spatial artefacts, and n_i is the true expression level of the probe. x_i is the position of probe *i* on the array. Here n_i is the signal (the independence from location comes from the assumption that the probes are placed on the array in random order).
- 2. Natural image denoising: y_i is the gray-level of a noisy image at pixel *i*, s_i is the contribution of the true image pixel, and n_i is pixel-independent noise. x_i is the position of pixel *i* in the image. Here s_i is the signal.

We show schematically an example of each problem in Figure 1.1.

Given the nature of these interpretations of s, informally we expect s to exhibit spatial regularity or correlations; however, we neither make differentiability nor continuity (as a function of the spatial variable x) a requirement.

Historically, these two problems have been attacked separately, although the main difference between them is that in the first case we are interested in the spatially independent term and in the second in the spatially-dependent term. Because we usually expect the signal-component to have greater energy than the noise-component, we expect the independent component to dominate in the case of microarray artefact removal, and the dependent component to dominate in the case of image denoising.

More importantly, it seems methods designed for either application can be used for the other with minor modifications. It may be that certain methods really are better for one application, or that one is best for both. In this thesis we investigate this possibility by adapting algorithms from both the bioinformatics and image processing communities to both applications and comparing to our own novel approach using Gaussian process regression.

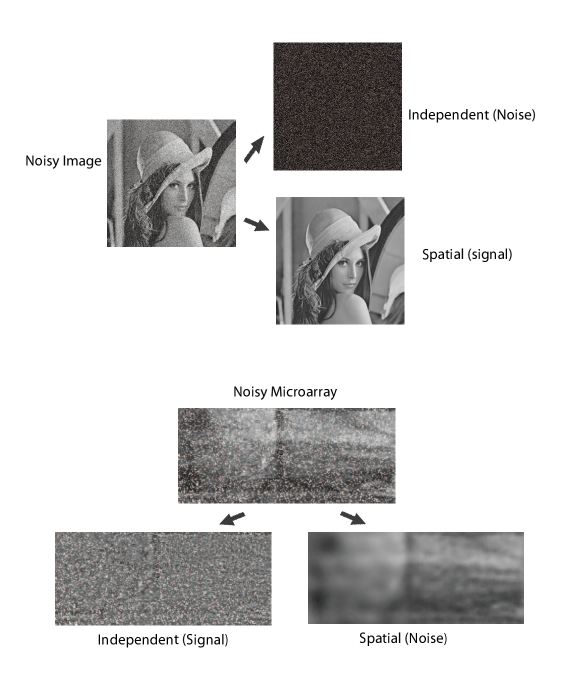


Figure 1.1: The two decomposition problems we will discuss. Top, a noisy image decomposed into noise and the original image. Bottom, a noisy microarray decomposed into probe expression and spatial artefacts.

1.2 Outline of thesis

The work described in this thesis began as an application of Gaussian processes to the problem of denoising microarray image data; however, we realized that the methods developed for this application could easily be viewed also as natural image denoising methods. Since there may be interest in our approach from both bioinformatics and image processing communities, who may have widely differing backgrounds, an attempt has been made to structure the thesis so that a reader from either community may read it and learn about our method without learning about the other field. To this end, in Chapter 2, we present previous methods used to solve either problem. In Chapter 3, we present a brief introduction to Gaussian processes for regression and a description of our method. In Chapter 4, we describe the problem of microarray artefact removal and present some experimental results, using the methods discussed. In Chapter 5, we discuss the application of these methods to image denoising and present experimental results in that domain. In Chapter 6, we summarize our contributions and discuss possible future work. A dependency chart for the chapters is shown in Figure 1.2.

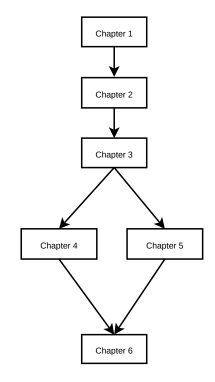


Figure 1.2: Chapter dependency chart.

Chapter 2

Related Work

The methods we will discuss in this section were designed for only one of image or microarray denoising applications, but may be used for either in principle. Before describing related methods it is useful to have more definitions.

Recall our dataset $\mathcal{D} = \{(x_i, y_i)\}_{i=1}^N$ is in the form of a *complete grid* isomorphic to a Euclidean rectangular plane, which for visualization purposes could be viewed as an image. For simplicity, in this chapter we refer to each data point as a *pixel* and $p_i = (x_i, y_i)$ as the *i*th pixel.

Define the k-neighbourhood (for positive integer k) of a pixel p, $N_k(p)$, as the set of pixel-values within k pixels of p not including itself. As a slight abuse of notation, if p = (x, y), we may also denote $N_k(p)$, by $N_k(x)$. More precisely, $N_1(p)$ is the set of values of pixels adjacent to p. For k > 1, $N_k(p)$ is $N_{k-1}(p)$ and their adjacent pixel-values except p. We show $N_k(p)$, for k = 1, 2 in Figure 2.1. Typically, we would like for $N_k(p)$ to have the same cardinality of $4k^2 + 4k$ for all pixels ; to this end we augment the original image by adding artificial pixels to the boundaries. The values assigned to these pixels will depend on the method, as is discussed below.

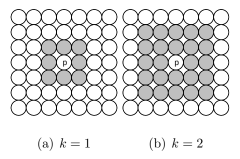


Figure 2.1: k-neighbourhood pixels for pixel p are shown in gray.

2.1 Median filtering

The median filter is a simple, commonly used non-linear filtering method which attempts to predict the spatial component of a pixel by taking the median of its k-neighbourhood. In increasing k there is a trade-off between maintaining the fine structure of the spatial component and removing the independent component. It is ideal for removing isolated "specks" of very high or low intensity (perhaps due to dropout), called *salt and pepper* noise in the image processing literature, where neighbouring pixels are expected to have very similar spatial components (see an example in Figure 2.2). It is likely less ideal for our case where the independent component is present in all pixels, but the hope is that the median would be a similar pixel with a independent component near zero.



Figure 2.2: An image corrupted by salt and pepper noise. (Source: http://homepages.inf.ed.ac.uk/rbf/HIPR2/noise.htm)

2.2 Mean and Gaussian Filtering

For our problem, we can observe that $E[y_i] = E[s_i]$, and might aim to "average-out" the zero-mean independent component by averaging the values of similar (defined by the method) pixels. We will see that similarity effectively induces a set of weights $\{w_{ij}\}_{j=1}^N$ to yield a weighted-average predictor (WAP) of the spatial component at pixel i, $\hat{s}_i = \sum_{j=1}^N w_{ij}y_j$. The simplest such method is like the median filter except we take the mean instead of the median. In our notation,

$$w_{ij} \propto \begin{cases} 1, & \text{if } y_j \in N_k(x_i) \\ 0, & \text{otherwise} \end{cases}$$
(2.1)

$$\sum_{j=1}^{N} w_{ij} = 1 \tag{2.2}$$

In fact this coincides with nearest neighbours algorithm for grid data where the number of neighbours is the size of the neighbourhood.

Gaussian filtering is a commonly used image filtering technique which is a WAP with weights defined as

$$w_{ij} \propto \exp\left(-\frac{\|x_i - x_j\|_2^2}{h^2}\right), i \neq j$$
(2.3)

$$w_{ii} = 0 \tag{2.4}$$

$$\sum_{i=1}^{N} w_{ij} = 1 \tag{2.5}$$

where $\|\cdot\|_2$ is the L2 norm. Because of the rapid decay of w_{ij} as a function of the distance $\|x_i - x_j\|_2$, Gaussian smoothing is effectively a local filtering method.

The parameter h can be selected in a variety of ways. For example, the Spatial Trend Removal (STR) method [9], which was developed to remove spatial trends in microarray data, optimizes h using gradient-based optimization on the mean squared error between the original and filtered datasets:

$$\frac{1}{N}\sum_{i=1}^{N}(y_i - \hat{s}_i)^2 \tag{2.6}$$

If viewed from a machine learning perspective, this error may be interpreted as the leave-one-out cross-validation error in the task of predicting the original dataset.

In order to increase the smoothness of the estimated true image, STR ignores the pixels which differ the most from an initial true image estimated by a median filter (7x7 window). The authors report that ignoring 10 % of pixels with the greatest difference works well on their microarray data (this can be viewed as removing outliers). However, this is an adjustable parameter.

As an image denoising algorithm, Gaussian filtering is well-known to over-smooth images, resulting in the loss of significant detail, especially edge sharpness.

2.3 Non-local Means

In STR, similarity between pixels is a function of the distance of the locations of pixels. In Non-local Means [3], the distance between pixels is mostly ignored, and similarity is defined as the distance between k-neighbourhoods. Thus, the pixels it chooses to average may very well be nonlocal. It was designed to denoise natural images, and the motivation behind it is that natural images (here, the spatial component) usually exhibit significant structural redundancy. The selection of pixels to average should therefore be guided by this structure in the image. For example, the pixels in a constant-coloured region or forming an edge would all have very similar k-neighbourhoods even though pixels at opposite ends of the region or edge are distant from each other in pixel-location space. Figure 2.3 shows the weight distributions for the centre pixel in various images. The algorithm is perhaps ideally suited for images with significant periodicity such as image (e) in Figure 2.3.

We may write the spatial component predictions produced in the form $\hat{s}_i = \sum_{j=1}^N w_{ij} y_j$, but here w_{ij} depends on all pixel values, not just the locations; therefore, we do not have a linear predictor as in the WAP methods previously discussed. More precisely, for each pixel, p = (x, y), we may arbitrarily enumerate its k-neighbourhood, $n_1, \ldots, n_{|N_k(x)|}$, and form a neighbourhood-vector denoted, $V_k(x) = (n_1, \ldots, n_{|N_k(x)|})$. Then we define the weights as

$$w_{ij} \propto \exp\left(-\frac{\|V_k(x_i) - V_k(x_j)\|_{2,a}^2}{h^2}\right), i \neq j$$
 (2.7)

$$w_{ii} = 0 \tag{2.8}$$

$$\sum_{j=1}^{N} w_{ij} = 1 \tag{2.9}$$

where $\|\cdot\|_{2,a}$ is the weighted L2 norm, where the weights come from a Gaussian kernel with variance a^2 .

The parameters to this algorithm are h and k. Although the authors of [3] do not provide a methodology to choosing them, they suggest $h \in [10\sigma, 15\sigma]$, where σ is an estimate of the standard deviation of the independent component, and k = 7.

The time complexity is dominated by the pairwise neighbourhood-distance calculations $\mathcal{O}(kN^2)$. In practice, one might restrict attention to the pixels within a large window centred on the pixel of interest to speed-up the algorithm and also force a local bias in the weights. A new parameter M specifies the search window size. The time for distance calculations is reduced to $\mathcal{O}(kNM^2)$ (note $M^2 \leq N$). The authors suggest M = 21 works well in their experience.

2.4 Other methods not compared

There are a myriad other classes of methods which we do not review nor include in performance comparisons in this thesis. Among them methods based on *wavelet thresholding* ([5]) from image processing are worth mentioning. Briefly, after performing a wavelet transform, wavelet coefficients below a threshold (associated with high frequency noise) are set to zero. These methods are fast, but it is unclear what the optimal set of wavelets are. Although their algorithms are not open source, many state-of-the-art commercial

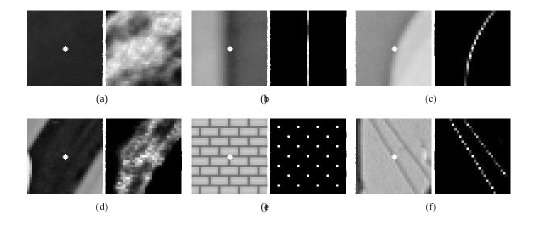


Figure 2.3: In each pair of images: the left shows an image with the centre pixel highlighted; the right shows the weight distribution for the centre pixel. Source: [3].

image denoising software packages such as Noise Ninja and NeatImage are based on this technology. A comparison of the NL means algorithm with non-state-of-the-art wavelet thresholding algorithms is done in [2].

Chapter 3

Gaussian process method

3.1 Using Gaussian processes for noisy 2-dimensional interpolation

Recall the decomposition of the observations into spatial and independent components we are striving for

$$y_i = s_i + n_i \tag{3.1}$$

In order for Gaussian processes to be used efficiently we assume $\{n_i\}_{i=1}^N$ are Gaussian in addition to being independent, identically distributed with zero mean. We will see that the variance of n_i will be estimated from the data automatically.

Initially, we will also assume $s_i = f(x_i)$ for some unknown (latent) function of the two coordinate inputs x; hence, n_i will denote the departures of y_i from $f(x_i)$ and our task can be seen as determining the value of f at all data point locations, what one may call the *two-dimensional noisy interpolation problem*.

The approach of parametric methods is to parameterize a class of functions conjectured to contain the right f and use the data to estimate its parameters. Coming up with the right parametric form is generally difficult. By contrast, Gaussian process regression discovers f without explicitly stating a parametric form but rather by vaguely specifying its expected behaviour through a "distribution over functions".

3.1.1 Gaussian process priors

A stochastic process is a collection of random variables $\{Y_x : x \in I\}$ indexed by some set I, which often denotes time, but in general, as in our case, is multidimensional. Note that x is not a random variable, but merely an index for random variables. We may also write Y_x as Y(x) to emphasize that x is typically viewed as the input and Y(x) as the output (we use indices, inputs interchangeably). The Kolmogorov extension theorem guarantees the existence of a stochastic process with consistent joint distributions for any finite collection of random variables from the process [15]. Consistency here means that the marginal probability distributions should agree with those derived from higher dimensional joint distributions. Gaussian processes are stochastic processes where these joint distributions are Gaussian.

A sample function of a stochastic process is a single realization of each of its random variables. It can then be viewed as a function of the index set, I. For example, in our case of grid data, the index set is a finite subset of a two-dimensional plane; hence, we may visualize a sample function as a surface in \mathbb{R}^3 . An interesting property of stochastic processes is that the joint distributions defining it induce a distribution over sample functions. In general, we do not know the analytic form of sample functions but, in the case of Gaussian processes, can obtain the value of the function at an arbitrary number of indices (inputs). We give a procedure for doing this in Section 3.1.2. With a notion of "distribution over functions" in hand, we may talk about priors and posteriors over functions induced by Gaussian processes. A distribution over functions can also be achieved by parameterizing a class of functions and setting prior distributions over the parameters, but the Gaussian process prior maybe preferred if no prior knowledge justifies choosing a particular parametric form. The mean function, $M : I \to \mathbb{R}$, of a stochastic process specifies the mean of each random variable $M(x) = E[Y_x]$. The covariance function of a stochastic process, C : $I \times I \to \mathbb{R}$, specifies the covariance of any two random variables, $C(x, x') = Cov(Y_x, Y_{x'})$ and determines the smoothness properties of the process. As a Gaussian distribution is completely specified by its mean and covariance, a mean function, M, and a covariance function, C, completely specify a Gaussian process, which we denote $\mathcal{GP}(M, C)$.

If we have some strong prior knowledge of the behaviour of the mean function, we could specify a parametric form – say of a quadratic polynomial $M(x) = ax^2 + bx + c$ (with hyperparameters a, b, c) – but typically our knowledge is vague so we set M(x) to a constant. By pre-processing our data we can make the most appropriate value for this constant equal to zero.

Specifying the covariance function requires more thought, but as an illustration consider the *squared exponential covariance* function:

$$C(x, x') = \sigma_a^2 \exp\left(-\left\|\frac{x - x'}{\ell}\right\|^2\right)$$
(3.2)

where σ_a^2 and ℓ are constants. The interpretation of these constants is elaborated in 3.1.4, but we can immediately see that $Cov(Y_x, Y_{x'})$ is maximized when x and x' are close, encoding the prior belief that the sample functions from the corresponding Gaussian process are continuous. In fact, with this covariance function they are differentiable [14], which is not always a good assumption. If this were the only possible covariance function, Gaussian process priors would be insufficiently expressive of our beliefs; luckily, there is an infinite supply of covariance functions.

A real, symmetric function C on $I \times I$ is said to be *semi-positive definite* if for any $N, x_1, \ldots, x_N \in I$, and $v \in \mathbb{R}^N$, we have $v^T A v \ge 0$ where $A_{ij} = C(x_i, x_j)$. Given a semipositive definite function C, the Kolmogorov extension theorem guarantees that there exists a (unique) zero-mean Gaussian process with covariance function E[y(x)y(x')] = C(x, x') [20]. That is, there exists a zero-mean Gaussian process, $\mathcal{GP}(0, C)$, for every such "kernel" function, C. In practice most of the work in specifying a Gaussian process prior is spent choosing the covariance function. This is the topic of Section 3.1.4.

3.1.2 Sampling functions from a Gaussian process

Although examining the mathematical structure of a covariance function gives clues into its properties, these can also be determined by looking at sample functions. It is easy to generate sample functions from Gaussian processes, since although we may not know explicitly the functional form of the sample function, we can obtain the values of the function at any finite number of indices. Let us use the shorthand $f \sim \mathcal{GP}(M, C)$ to signify that f is a sample function of $\mathcal{GP}(M, C)$. One possible procedure to obtain the values of a sample function $f \sim \mathcal{GP}(0, C)$ at n given indices of interest, $x_1, \ldots, x_n \in I$, is as follows:

- 1. Compute the covariance matrix $K_{ij} = C(x_i, x_j)$, for i, j = 1, ..., n.
- 2. Compute the Cholesky Decomposition of K, which is the lower triangular matrix L such that $K = LL^{T}$.
- 3. Draw *n* independent samples $v_1, \ldots, v_n \sim \mathcal{N}(0, 1)$ and form the vector $\mathbf{v} = (v_1, \ldots, v_n)^T$.
- 4. Compute values of the sample function at the *n* input points by $f(x_i) = y_i$ where $\mathbf{y} = (y_1, \dots, y_n)^T = Lv.$

It is easy to show that $\langle \mathbf{y} \rangle = 0$ and $\langle \mathbf{y} \mathbf{y}^T \rangle = K$. That is we have generated $\mathbf{y} \sim \mathcal{N}(0, K)$. By choosing a large enough number of indices, we can get an arbitrarily good idea of the sample function. In Figure 3.1, we show the values of sample functions drawn from two different Gaussian processes over a region of the input space (an interval).

¹called the Gram matrix in the kernel-methods terminology

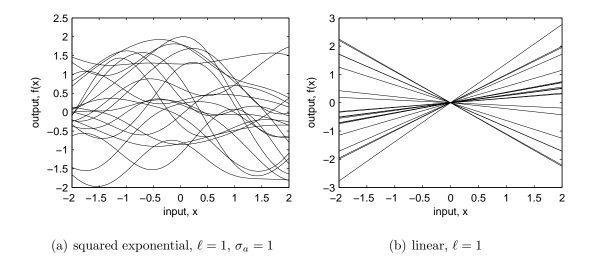


Figure 3.1: 20 sample functions from zero-mean Gaussian Processes with squaredexponential and linear $(C(x, x') = x^T x')$ covariance functions. Note that at each location, x, the mean value over all functions is about zero.

3.1.3 Gaussian process regression

One possible way to approach the noisy interpolation problem is through Gaussian process Regression (GPR). Although a form of GPR has long been practised by spatial statisticians with a technique called *kriging* [4], renewed interest from the machine learning community began after it was re-derived in the Bayesian machine learning framework and given a new probabilistic interpretation. Neal showed that the prior distribution encoded by certain neural networks converges to a Gaussian process in the limit of infinite hidden units [11], and Rasmussen and Williams [21] investigated it as a general machine learning method.

Suppose we want to have a prior "distribution" over functions without restricting ourselves to a parametric form, and that we want to be able to compute a posterior over functions having seen noisy data. GPR provides one way of doing so in a principled (Bayesian) way. If we further assume a Gaussian noise model we can do this using efficient matrix operations. The noise model and prior over functions are

$$y_i | f(x_i) \sim \mathcal{N}(f(x_i), \sigma_{\varepsilon}^2)$$
 (3.3)

$$f \sim \mathcal{GP}(0, C) \tag{3.4}$$

where σ_{ε}^2) is the noise variance (to be estimated), and *C* is the covariance function of the Gaussian process. The use of a zero-mean function for the Gaussian process encodes the belief that at each location x, f(x) has equal probability of being positive or negative; it does not say that sample functions themselves need have zero-mean. In order to make this prior belief reasonable, we normalize the data to have (sample) mean of zero and standard deviation of 1. We perform the same normalization to the inputs to simplify the choice of initial hyperparameters when optimizing the marginal likelihood (see Section 3.1.5).

First, we consider the noise-less case, $y_i = f(x_i)$. We learned how to sample functions from the prior in the previous section, but really what we want to do in the context of regression is to predict the values of f(x) for x not in the training data, i.e. test data. Let us consider a single test input x^* and predict its output y^* . Now the prior distribution for y_1, \ldots, y_n, y^* is multivariate Gaussian, so the conditional distribution $y^*|y_1, \ldots, y_n$ is univariate Gaussian. Some elementary manipulations using Gaussian identities yields

$$E[y^*|x^*, \{(x,y)\}_{i=1}^N] = k^T K^{-1}[y_1, \dots, y_n]^T$$
(3.5)

$$Var(y^*|x^*, \{(x,y)\}_{i=1}^N) = Cov(x^*, x^*) - k^T K^{-1} k$$
(3.6)

where
$$k \in \mathbb{R}^N, k_i = Cov(x^*, x_i)$$
 (3.7)

We see that the mean is a linear combination of training outputs, which led spatial statisticians to call it a linear predictor. The uncertainty in the value of the function at the test input x^* has been reduced, since $k^T K^{-1} k \ge 0$, but the magnitude of this reduction does not depend on the training outputs. One of the advantages of GPR is we have a full predictive distribution rather than just a point prediction. If a point prediction is required we can determine the optimal guess for our loss function from the predictive distribution; often we use the squared error loss function, in which case we guess the mean of the distribution.

In the noise-free case, the predictive mean interpolates the training outputs exactly and there is zero uncertainty (variance) in the output at the training inputs (for example see figure 3.2). More realistically, we are interested in the case with noisy outputs:

$$y = f(x) + \varepsilon \tag{3.8}$$

$$\varepsilon \sim \mathcal{N}(0, \sigma_{\varepsilon}^2)$$
 (3.9)

which we have seen before in equation 3.4. Conveniently, the additive noise assumption means the covariance between outputs becomes

$$Cov(y_i, y_j) = C(x_i, x_j) + \delta_{ij}\sigma_{\varepsilon}^2$$
(3.10)

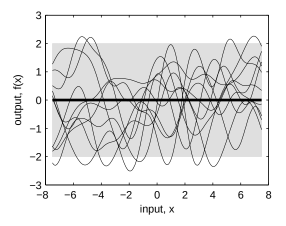
where δ_{ij} is 1 if i = j and 0 otherwise. That is, adding noise changes the covariance function from C to $(C + \delta \sigma_e^2)$. To obtain the covariance matrix, we simply add the diagonal matrix $\sigma_{\varepsilon}^2 I$ to the old noise-free covariance matrix. We then can obtain analogous predictive distribution equations for the noise-free output at x^* , denoted by f^* :

$$E[f^*|x^*, \{(x,y)\}_{i=1}^N] = k^T (K + \sigma_{\varepsilon}^2 I)^{-1} [y_1, \dots, y_n]^T$$
(3.11)

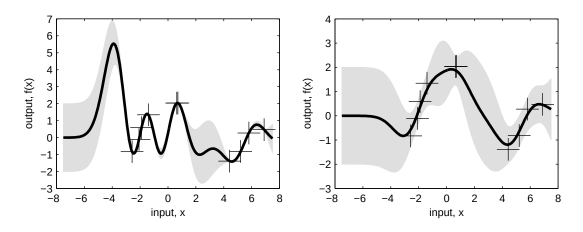
$$Var(f^*|x^*, \{(x,y)\}_{i=1}^N) = Cov(x^*, x^*) - k^T (K + \sigma_{\varepsilon}^2 I)k$$
(3.12)

where
$$k \in \mathbb{R}^N, k_i = Cov(x^*, x_i)$$
 (3.13)

One notable difference between the noise-free and noisy cases, is that in the latter case, the predictive mean of the noise-free outputs at the training input locations is in general not equal to the training output (which has noise added). Indeed, we will use the predictive mean as an estimate for the spatial component in the spatial-independent decomposition in equation 3.1.



(a) 10 samples functions from a GP prior



(b) Predictive distribution with noise-less ob- (c) Predictive distribution with noisy observaservations tions

Figure 3.2: Experiments with the squared exponential covariance function ($\ell = 1, \sigma_a = 1, \sigma_{\varepsilon}^2 = 0.2$). The predictive distributions are shown before and after seeing some data points. The mean function is shown by the dark solid line; the two standard deviation region at each location x is shown in gray. We see that in regions of sparse data, the predictive distribution is unchanged from the prior and in data dense regions, the uncertainty in the value of the latent function is smaller than in the prior.

3.1.4 Selecting the covariance function

Since we assume a mean function equal to 0 (in the absence of knowledge), all we really need to infer the latent function, f, is an appropriate covariance function that encodes our prior beliefs of its behaviour. This is therefore where most of the tweaking/tuning occurs in Gaussian process methods.

The approach we take is to decide on an appropriate class of covariance functions indexed by hyperparameters (we call them hyperparameters since they parameterize our prior for the latent function, not the latent function itself). We set these hyperparameters to maximize an objective function called *the evidence* or *marginal likelihood*, which automatically finds a compromise between model complexity and data fit. This is described further in section 3.1.5.

Next, we describe a few common classes of covariance functions. Let D be the dimensionality of x.

1. Linear:

$$C(x, x') = \sum_{i=1}^{D} \frac{x_i x'_i}{\ell_i^2}$$
(3.14)

Hyperparameters: $\ell_1, \ldots, \ell_D > 0$. The sample functions are linear in the inputs x. Having different ℓ_i values, also known as the *lengthscale* for the i^{th} input dimension, adjusts the relative importance of different inputs. For example, certain input features may be effectively "ignored" by setting their corresponding ℓ_i , to be very large relative to other lengthscales.

A Gaussian process with a linear covariance function is equivalent to Bayesian linear regression, with normally distributed weights:

$$y = w \cdot x \tag{3.15}$$

$$w_i \sim \mathcal{N}(0, \ell_i^{-2}) \tag{3.16}$$

Inferring w in this model is more efficient than the GP approach when D < N, since it involves inverting a D by D matrix rather than an N by N one. However, it is useful to combine the linear term with another covariance function through the operations of addition or multiplication to be discussed in the next section.

2. Power Exponential:

$$C(r) = \sigma_a^2 \exp\left(-\left\|\frac{x-x'}{\ell}\right\|^P\right)$$
(3.17)

Hyperparameters: σ_a^2 determines the variance in the outputs; ℓ determines the characteristic lengthscale of the input space. Although P could be made a hyperparameter, normally it is fixed ($0 < P \leq 2$, to ensure semi-positive definiteness) to ease the optimization process. For any choice of P, the sample functions will be continuous, but with varying degrees of smoothness.

When P = 2, we obtain the popular "squared exponential" which has the property that sample functions are infinitely continuously differentiable. It is interesting to note that a Gaussian process using this covariance function is equivalent to a radial basis network with an infinite number of basis functions (one centred on each point of the input space) [14]. If this were attempted directly, an infinite amount of computation would be required to manipulate infinite-dimensional matrices; the Gaussian process approach is able to do the same efficiently, in a finite amount of time.

When P < 2, we lose differentiability. Roughly speaking, lower values of P result in more erratic sample functions. For example, we obtain functions related to the well-known "Brownian motion" with P = 1. Figure 3.3, shows the effect of changing P while other hyperparameters are fixed.

As for the linear covariance function, we may wish to scale each input dimension with its own lengthscale. In doing so, we may arrive at the following covariance

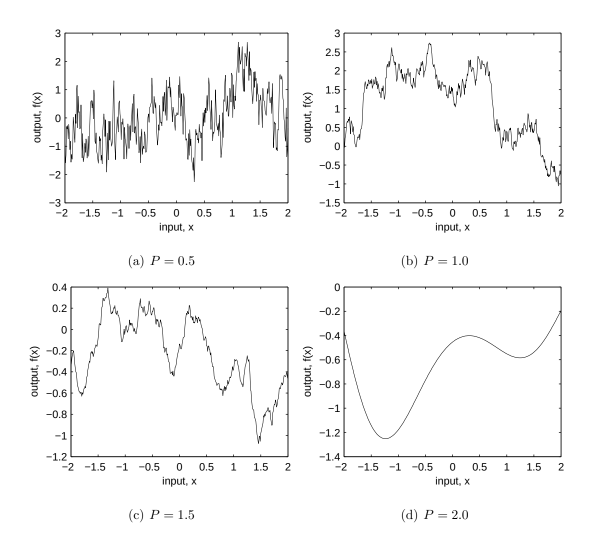


Figure 3.3: One sample function from a Gaussian process with power exponential covariance function, for various values of P (other hyperparameters are fixed at $\ell = 1$, $\sigma_a = 1$).

function:

$$C(x, x') = \sigma_a^2 \exp\left(-\left(\sum_{i=1}^D \left(\frac{x_i - x'_i}{\ell_i}\right)^2\right)^{P/2}\right)$$
(3.18)

Here, instead of single ℓ , we have ℓ_i for input dimension *i*.

3. Constant:

$$C(x, x') = \alpha^2 \tag{3.19}$$

Sample functions correspond to a constant function whose value has prior variance of α^2 and mean zero. This covariance function has little use on its own, and is usually combined with others by addition (see 3.1.4), in which case it models the prior variance for the constant part (or bias) of the latent function. The α may be treated as a hyperparameter but in practice it is better to fix it to a reasonable value. A reasonable value can be determined from the range of possible values the bias could take on, which can be determined from the data. The danger in leaving it as a hyperparameter is that it may be set to an unreasonably large value, which can cause numerical issues when computing the Cholesky decomposition of the covariance matrix.

There are many more covariance functions, but those listed above are the ones we will use in our work and includes arguably the most common, the squared exponential covariance function. For a compendium of covariance functions, the read is advised to consult the technical report by Abrahamsen [1].

Combining covariance functions

It is in general non-trivial to design reasonable legal (semi-positive definite) covariance functions, except through the addition and multiplication of known covariance functions. The sum or product of any number of semi-positive definite kernel functions is still semipositive definite; hence, adding or multiplying two covariance functions yields another. Thus,

$$C_{add}(x, x') = C_1(x, x') + C_2(x, x')$$
(3.20)

$$C_{prod}(x, x') = C_1(x, x') \cdot C_2(x, x')$$
(3.21)

are valid covariance functions, if C_1 and C_2 are too.

Sample functions drawn from a GP with covariance C_{add} exhibit the properties of functions drawn separately from C_1 and C_2 and then added. The interpretation of the product operation is less obvious but we may observe that while in the case of addition, a large contribution from either C_1 or C_2 will result in a large overall covariance, in the case of product, both C_1 and C_2 need to be large for the overall covariance to be large. Another way to make use of these combination operations is to have C_1 and C_2 depend on different subsets of the inputs; in fact, we take this approach in section 3.2. As can be imagined, a great variety of covariance functions can be produced through these operations based on small set of base covariance functions.

Other ways of combining covariance functions into new ones are discussed in [17].

3.1.5 Selecting hyperparameters

Bayesian approach

In the Bayesian formalism, all parameters and hyperparameters have prior distributions encoding our beliefs. Because Gaussian process regression is a non-parametric method, we only have to worry about hyperparameters, which we denote by θ_C (having fixed our covariance function C). The predictive distribution for a new test output y_{N+1} (given dataset, $D = \{(x_i, y_i)\}_{i=1}^N$) is obtained by averaging predictions from different hyperparameters weighted by their posterior probability:

$$p(y_{N+1}|x_{N+1},\mathcal{D}) = \int p(y_{N+1}|x_{N+1},\mathcal{D},\theta_C)p(\theta_C|x_{N+1},\mathcal{D})d\theta_C$$
(3.22)

In general, equation 3.22 cannot be evaluated analytically, and we need to resort to numerical methods. The investigation of these methods is beyond the scope of this thesis;

for a full treatment see the work of Neal ([12],[10]), and Rasmussen and Williams [14]. One property of these methods is that they require significant additional computation time, which we avoid in order to maximize the practicality of our algorithm.

Marginal likelihood / evidence maximization

Instead of averaging predictions from all hyperparameter sets, as an approximation we may use the prediction from the single set with maximum posterior probability, given by

$$p(\theta_C | \mathcal{D}) \propto p(\mathbf{y} | \{x\}_{i=1}^N, \theta_C) p(\theta_C)$$
(3.23)

where $\mathbf{y} = (y_1, \ldots, y_N)$, $p(\mathbf{y} | \{x\}_{i=1}^N, \theta_C)$ is the probability of the data or (*the evidence*, or *marginal likelihood*) and $p(\theta_C)$ is the prior distribution on the hyperparameters. For simplicity, this prior is often ignored (i.e. set to improper uniform), although using a proper prior may help rule out unreasonable values of θ_C ; therefore, maximization of the posterior probability of the hyperparameters reduces to maximization of the evidence.

The maximization problem is non-convex in general and it is carried out through a gradient-based optimization technique such as conjugate-gradients, or L-BFGS. Numerically, it is easier to perform this maximization in the log domain. Since we know that $\mathbf{y} | \{\mathbf{x}_i\}_{i=1}^N, \theta \sim \mathcal{N}(0, K)$, we have

$$\log(p(\mathbf{y}|\{\mathbf{x}_i\}_{i=1}^N, \theta)) = -\frac{N}{2}\log(2\pi) - \frac{1}{2}\log(|K|) - \frac{1}{2}\mathbf{y}^T K^{-1}\mathbf{y}$$
(3.24)

from which we can obtain the gradient with respect to θ

$$\frac{\partial \log(p(\mathbf{y} | \{\mathbf{x}_i\}_{i=1}^N, \theta))}{\partial \theta} = -\frac{1}{2} \operatorname{tr}(\mathbf{K}^{-1} \frac{\partial \mathbf{K}}{\partial \theta}) + \frac{1}{2} \mathbf{y}^T \mathbf{K}^{-1} \frac{\partial \mathbf{K}}{\partial \theta} \mathbf{K}^{-1} \mathbf{y}$$
(3.25)

The asymptotic time complexity is dominated by the inversion of K, the covariance matrix, which takes $\mathcal{O}(N^3)$ time. As with all non-convex optimization problems, local optima can be a problem, although perhaps not too serious a one in practice, since the number of hyperparameters is usually small relative to the number of data points [8]. In our experiments, we have chosen to use the approach of evidence maximization rather than the true Bayesian approach, in order to make the algorithm more practical in terms of time complexity and avoiding the difficulty of choosing appropriate prior distributions for hyperparameters.

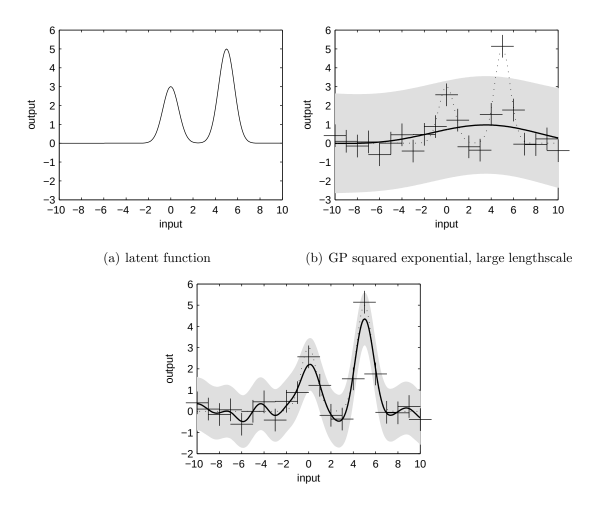
3.1.6 Limitations of the 2-d interpolation approach

Except for the linear covariance function, the basic covariance functions we discussed all can be written as a function of the separation vector x - x', between two inputs xand x'. Covariance functions that have this property are called *stationary*. It is readily seen that the operations of addition and multiplication of covariance functions preserve stationarity. Unfortunately, stationarity implies certain reasonable functions are highly improbable such as functions which vary rapidly in one part of the input space and slowly in another. This corresponds to a process with a input-dependent (i.e. non-stationary) lengthscale. A concrete 1-dimensional example of this is shown in figure 3.4, where a rapid change in the function only occurs at two isolated parts of the input space and is unchanging elsewhere.

Although the linear covariance function is non-stationary, it only has the power to add a linear trend to sample functions. Dealing with reasonable non-stationary covariance functions is somewhat involved and beyond the scope of this thesis, but it is examined in the work of Paciorek [13]. In the next section, we propose a novel way to extend the modelling power of stationary covariance functions in the special case of grid data.

3.2 Adding pseudo-inputs to grid data

Recall that in two-dimensional grid data, the data points are uniformly spaced in a rectangle. Each point has a k-neighbourhood as defined in Chapter 2 (boundary cases are treated specially). Our aim is to incorporate local neighbourhood information into the



(c) GP squared exponential, short lengthscale

Figure 3.4: A function that is badly modelled by stationary Gaussian processes. We added a small amount of noise, sampled a few data points, then modelled it with a squared exponential GP. We show the predictive distributions of two different sets of parameters corresponding to different local maxima in the marginal likelihood. In (b) the predictive model is too simple; in (c) it is too complex.

Gaussian process previously discussed for 2-dimensional noisy interpolation, effectively making the covariance vary throughout the grid even if the distance between points is fixed while still using a stationary covariance function. We achieve this by adding extra inputs to the data points.

Previously our input x was simply the two Cartesian coordinates encoding the location of our data points. Let us now call this x_{coord} , and consider augmenting our input vector with neighbourhood information, $x_{neigh} = V_k(x_{coord})$ (we find k = 1 works best)², and define our new, augmented input vector, $x_{aug} = (x_{coord}, x_{neigh})$. It is perhaps incorrect to call x_{neigh} extra inputs since they correspond to the noisy outputs for other data points; for this reason we use the term *pseudo-inputs* for x_{neigh} . However, in the following we treat them as if they were regular inputs.

The covariance function we use to combine the two input sets in a reasonable way is:

$$C(x_{aug}, x'_{aug}) = \sigma_a^2 C_{coord}(x_{coord}, x'_{coord}) C_{neigh}(x_{neigh}, x'_{neigh})$$
(3.26)

$$C_{coord}(x_{aug}, x'_{aug}) = \exp\left(-\frac{|x_{coord} - x'_{coord}|^2}{l_{coord}^2}\right)$$
(3.27)

$$C_{neigh}(x_{aug}, x'_{aug}) = \exp\left(-\frac{|x_{neigh} - x'_{neigh}|^2}{l_{neigh}^2}\right)$$
(3.28)

or

$$C(x_{aug}, x'_{aug}) = \sigma_a^2 \exp\left(-\frac{|x_{coord} - x'_{coord}|^2}{l_{coord}^2} - \frac{|x_{neigh} - x'_{neigh}|^2}{l_{neigh}^2}\right)$$
(3.29)

There are two characteristic length-scales, one for the location and one for the neighbourhood. This allows the model to automatically determine which of the two "super features" of distance in actual space and distance in neighbourhood space is more relevant. Fixing l_{coord} and letting $l_{neigh} \rightarrow \infty$, we get the 2-d interpolation scheme introduced in section 3.1; while fixing l_{neigh} and letting $l_{coord} \rightarrow \infty$ we get a covariance only depending on neighbourhood as in the NL-means algorithm. Having these as hyperparameters

²Recall the definition from Chapter 2: $V_k(x)$ is the set of k-neighbourhood values centred at x, put in a vector.

allows for the evidence maximization mechanism to decide how to trade-off between the two extremes of using mostly local and mostly non-local information to calculate the covariance between outputs.

The resulting covariance function encodes our prior belief that data points both near each other and with similar neighbourhood structures should have similar outputs. Although the covariance functions used are all stationary, the predictions produced clearly cannot come from a stationary process depending on only location inputs.

One consequence of using a higher-dimensional input space is we have lost continuity with respect to x_{coord} , i.e. two data points which are close in the grid may be far apart in the new input space, therefore freeing the outputs from being close. However, if the locations and the neighbourhoods of two data points are close (which is not uncommon, e.g. a constant region), the covariance will be large and the outputs close. This behaviour is actually desirable and preferable in our situation since the type of surfaces we wish to model are not globally continuous. For example, consider modelling an edge in a natural image. Points on either side of the edge are close in location but have unrelated outputs (discontinuous), whereas points on the edge have similar outputs (continuous). As previously mentioned in section 2.3, the points on the edge have similar k-neighbourhoods, different from those on either side of the edge. In our augmented input space this subtlety can be learned by the model.

We have also lost the ability to generate datasets, since our pseudo-inputs require knowning the dataset beforehand! However, given a noisy dataset, we can generate the noise-free (or spatial) component corresponding to the dataset, which is where our interest primarily lies.

3.2.1 Novel general algorithm for denoising grid data

Our algorithm for discovering the decomposition of noisy grid data is to use GPR with a zero-mean function and a covariance function which is a sum of covariance functions we have discussed:

$$C(x, x') = \alpha + C_1(x, x') + \delta(x, x')\sigma_{\varepsilon}^2$$
(3.30)

$$C_1(x, x') = \sigma_a^2 \exp\left(-\frac{|x_{coord} - x'_{coord}|^2}{l_{coord}^2} - \frac{|x_{neigh} - x'_{neigh}|^2}{l_{neigh}^2}\right)$$
(3.31)

The hyperparameters of the covariance function are $\sigma_a, l_{coord}, l_{neigh}, \sigma_{\varepsilon}$, and α . Usually α can be set manually based on the data (see 3.1.4) but the rest are learned (in the absence of other knowledge) by maximizing the evidence. It suffices to initialize the hyperparameters to reasonable values.

After the hyperparameters are selected, the mean of the predictive distribution at each grid location is used as the prediction for the spatial component. The difference of this and the actual value is the prediction for the independent component.

For large grids, it may be better to have different sets of hyperparameters for different regions of the grid. We consider a scheme for achieving this in the next section.

3.3 Managing the $\mathcal{O}(N^3)$ time complexity

The space complexity of Gaussian process Regression is only $\mathcal{O}(N^2)$, required to store the covariance matrix, so it is mainly the $\mathcal{O}(N^3)$ time complexity which poses a problem. For modern computers, datasets larger than 10^4 are infeasible and limiting N to 10^3 or less is desirable.

We overcome this difficulty by partitioning the grid into fixed-sized, overlapping patches and running our algorithm on each patch separately. By fixing the size of a patch to N_{part} pixels, the time complexity is now $\mathcal{O}(N_{part}^3 P)$, where P is the number of patches. Since the number of patches grows linearly with the number of pixels, asymptotically our algorithm is now linear in the number of pixels, $\mathcal{O}(N)$, with a constant factor corresponding to the time to process a single patch.

The partitioning scheme also adds flexibility to the model by allowing the hyperparameters of our covariance function to be different in different patches. This allows our method to better model widely differing features throughout the image. The patch size should be small enough so that the GP model can explain the within-patch variability, but not too small so as not to have enough data points for learning / pattern discovery. A partitioning of the standard "Lena" image which works well is found in figure 3.5. The patches are made to overlap so as to increase the continuity of the predictions when moving between patches. Predictions in the overlapping regions are calculated by averaging the predictions from individual patches.

Our approach of partitioning the space is similar to splines, where, say, a cubic polynomial may not be able capture the underlying function well, but may approximate it well in a small interval; the use of multiple cubic polynomials with different parameters in each partition of space is analogous to what we do. Often, splines are constrained to be continuous at the *knots* (boundaries of partitions), but we don't explicitly force this. In chapter 4, we also investigate reducing the dataset by summarizing blocks of data by



(a) Lena image

(b) with partitions

Figure 3.5: A possible partitioning of the standard 512x512 image of "Lena". The patches are mostly 29x29 in size.

their median and then using the block medians as data values.

3.4 Details of marginal likelihood optimization step

In our implementation, we normalize the inputs and outputs to have mean 0 and variance 1 in a pre-processing step. Initially, we initialize the hyperparameters σ_a , l_{coord} , $l_{neigh} = 1$ and initialize σ_{ε}^2 to an estimate of the independent-component variance. We fix $\alpha = 36$ rather than leave it as a hyperparameter. The hyperparameter search is carried out using conjugate-gradients as a first-order gradient-based method when maximizing the marginal likelihood. With this initialization, we obtained consistently good local maxima, making additional random restarts unnecessary.

Processing a single 30 x 30 patch in our MATLAB implementation takes approximately 4 minutes with a 3 GHz CPU.

Chapter 4

Microarray denoising

Our first application of the methods discussed is for removing spatial artefacts in data from DNA microarray experiments. Readers interested only in the domain of image de-noising may skip this chapter and move on to the next one without loss of continuity.

4.1 Description of the problem

Often biologists or medical researchers are interested in the relative expression levels of segments of deoxyribonucleic acid (DNA) – usually genes – in various experimental conditions (e.g. brain/liver tissue). By expression we mean the process through which DNA is "transcribed" to produce RNA, which may serve a regulatory purpose in the cell or be "translated" into a protein in the case of genes. A common way of measuring the expression level is to measure the abundance of a product of transcription, the RNA transcript. Even in the case in which transcription of DNA does not result in a protein, the RNA transcript abundance is still of interest as it may serve a regulatory role in the cell.

A common tool used to measure the abundance of RNA in a cell is the DNA microarray. A sensor with thousands of spots or probes, each designed to bind to some sequence of bases from the set A,C,G,T¹. The probes are arranged in a uniformly spaced two-dimensional grid, or array. Microarrays can be constructed economically and rapidly and allow parallel, high-throughput generation of data. Typically, the spacing between probes is small, allowing for thousands of probes to be placed on a glass slide (also a called microarray chip) roughly the size of a finger tip.

In a process called *hybridization*, a (tissue) sample of interest is dyed with a fluorophore then washed onto the slide to allow binding. In the imaging step, a laser is shone on the array, causing the dye to fluoresce. The amount of fluorescence at each spot is related to the amount of binding and in turn to the RNA abundance in the sample. From each spot, the imaging software derives an intensity value. These values are then normalized [18].

In practice it is possible to hybridize two differently dyed (using Cy3 or Cy5) samples to a single microarray. In spotted microarrays, this technique may be used to measure the relative expression levels in control and patient samples; in oligonucleuotide arrays (e.g. the data we discuss in this chapter), the signals are of sufficiently high quality that it is possible to measure the absolute abundance of RNA for two samples at each spot. For more details on the technology of microarrays, the reader may consult Schena's book ([16]).

By design, the order in which the probes are placed on the microarray slide is usually randomized (certainly in the dataset we use); hence, we expect very little correlation between position and expression after hybridization. This assumption is crucial to our denoising approach since the presence of spatial structure in the expression data can then be attributed to *spatial artefact noise* which arises from imperfect experimental procedures. There is a multitude of noise sources which we do not attempt to exhaustively examine, but they may include

¹Adenosine (A), Guanine (G), Thymine (T), Cytonsine (C) – the four nucleic acids of DNA and the building blocks of the genetic code

- stray dust particles on the slide or imaging lens [18];
- touching of the slide with fingers or other objects
- non-uniform "washing" in the hybridization process,

In other words, a great many possible things can contribute spatial artefacts and it is imprudent to place many restrictions on the shapes of artefacts we expect. Instead, we assume that the diversity of shapes is similar to that found in natural images, which tend to have a significant amount of continuity, along with some sharp edges.

There are other sources of noise in microarray data, such as that produced by crosshybridization, in which probes bind to the wrong segment of DNA [7], but we ignore these.

4.2 Data used in our experiments

We test our methods on mouse gene expression microarray data created by Zhang et al. [23] for gene function prediction. The microarrays they use are of the oligonucleotide variety, allowing for two different simultaneous hybridizations in the Cy3 and Cy5 channels in a process called *fluor reversal*. The result is we have two microarray images per slide (which we will call green and red), but we treat the data as if it were created from two separate slides. Conveniently, the dataset contains replicates (one from each channel, green and red) for each experiment; i.e. for each microarray experiment, there are two hybridizations on different slides. We take advantage of these replicates in our performance metric, as discussed in Section 4.4.

There are N = 21939 spots arranged in a 213-by-103 grid. Henceforth, the direction along the longer length will be called the vertical direction and the other will be called the horizontal direction.

We informally observed that there is a large diversity of spatial artefact shapes across

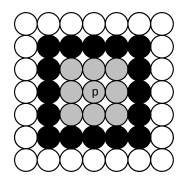
all slides, but not so much within a slide. This observation (along with preliminary experiments) influenced our decision to use only two partitions (see 3.3) for each slide. Remember, having finer partitions allows us to have more sets of hyperparameters in our Gaussian process tailored to different parts of the slide. The second reason for having finer partitions is that it reduces the computational complexity of processing the entire slide; however, we address this aspect by learning the hyperparameters on a reduced dataset, described in the next section.

4.3 GP algorithm for microarray artefact removal

Our approach to microarray denoising is to model the (log of the) intensity at each probe as the sum of spatially-independent and spatially-dependent components, seen as the contributions from the true expression level of the probe and the microarray's spatial artefacts, respectively. The spatial artefact component is also called the local/spatial trend or background, and its removal is called detrending. Once the background over the entire slide has been estimated, it can be removed by subtraction. In our algorithm, the spatial trend is a latent function inferred using Gaussian process regression and the true expression levels are modelled by IID Gaussian noise². The covariance function we use was described in Section 3.2.1.

The Gaussian model for the expression levels is not quite appropriate, since the actual distribution has a heavy right tail [9]. To make the data more amenable to our model, we derive a new dataset where the outputs are the medians of 3x3 non-overlapping blocks from the grid data, and the inputs are the centre (two) coordinates of the block and the 1-neighbourhood of the block, which is defined as the values of the data points directly adjacent to the block but not part of the block. Note that the 1-neighbourhood of a data point corresponds to the 1-neighbourhood of a block of size 1x1. Figure 4.1

 $^{^{2}}$ Note that the data is positive. As mentioned in Chapter 3, to make a zero-mean gaussian process reasonable, we pre-process the data so that it has zero mean



shows a 3x3 block centred at a point p along with its 1-neighbourhood. The hope in

Figure 4.1: Shows the 1-neighbourhood of a 3x3 block centred at p. The gray coloured circles are part of the 3x3 block and the black coloured circles form the 1-neighbourhood of the block. The white circles are other grid data points.

this pre-processing step is that the spatial artefacts over a 3x3 block are approximately constant, and hence removing the extreme data values from consideration by taking medians indirectly removes the extreme independent-component values. Figure 4.2 shows the effect of running one microarray image through our median scheme. We see that the distribution of values changes from very right-skewed to a more symmetric, more Gaussian distribution.

This pre-processing step also has a major side benefit of reducing the number of training points to 1/9 of the number. As we saw in the previous section, learning the hyperparameters involves several inversions of the covariance matrix, taking time that is $O(N^3)$ in the number of training points, which asymptotically dominates the overall time complexity of the algorithm. In the dataset used here, the slide is a 213 x 103 grid of probe positions, so N = 21939. Taking the medians of 3x3 blocks gives a new reduced grid of 71 x 34, with N = 2414, which is much more manageable on a modern-day computer. The task is reformulated as predicting the median of the spatial artefact values of a 3x3 block centred at x or (x_{coord}) . Note that even though we train on the centres of non-overlapping blocks, we may still make predictions of the medians of blocks

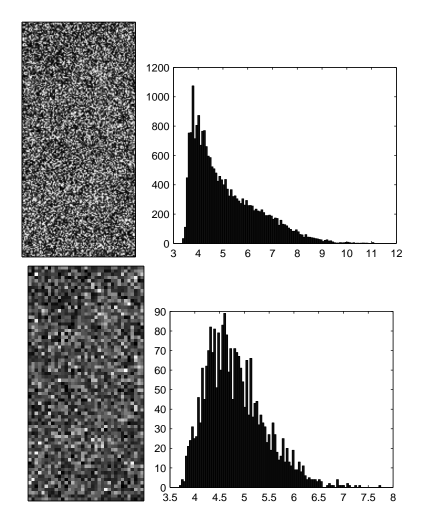


Figure 4.2: Top: A microarray slide (log domain) and its histogram of values. Bottom: The derived median version and its histogram of values.

centred at all 21939 (test) locations on the slide. This step is linear in the number of test points once the covariance matrix is inverted (taking time $O(N^3)$ in the training points).

In representing the original microarray image with 1/9 as much data, there is obviously a loss in resolution resulting in the loss of some fine spatial artefacts (say a 1 pixel line). However, such fine artefact structure is rare and the overall effect of ignoring it is small when using the correlation performance metric described in the next section.

Our medians-scheme is by no means the only way to increase the Gaussianity of data. Snelson et al. ([19]) investigates learning the parameters of a data transformation from a class (which includes log-like functions) simultaneously with the hyperparameters of the covariance function. In their Gaussian smoothing algorithm, Shai et al. ([9]) ignore data points whose values exceed a certain departure from a preliminary estimate of the local trend. However, only our scheme reduces the data set size while increasing its Gaussianity.

In our MATLAB implementation, processing each of the two partitions per chip takes approximately 4 minutes on a 3 GHz CPU computer.

4.4 Experimental methodology

We compare our own Gaussian Process method ("GP") with Gaussian filtering / smoothing ("STR", see 2.2), and a median filter with a 1-neighbourhood ("Medians", see 2.1). We do not compare with the NL-means algorithm since according to its developers, the choice of h (controls degree of filtering, see 2.3) should depend on the standard deviation of the independent component, σ_{ε} , which is unknown here. We attempted to estimate it by looking at the average (absolute) difference between adjacent values divided by two, but found that this was only accurate when the independent-component was small compared to the dependent-component, which is not the case in microarray data.

The performance metric we will use is the increase in correlation between the two

replicates of a microarray experiment after processing by the algorithm:

$$\rho(\hat{N}_{green}, \hat{N}_{red}) - \rho(Y_{green}, Y_{red}) \tag{4.1}$$

where Y_{green}, Y_{red} are the original green and red microarray image data, respectively, and $\hat{N}_{green}, \hat{N}_{red}$ are the estimates of their independent parts made by the algorithm whose performance we are measuring, and ρ is the sample correlation between two sets of measurements. It is important to emphasize that the algorithms process each replicate independently, and do not depend on the existence of replicates; the replicate information is used here only to measure performance. In general, replicates are not expected for the algorithms we discuss.

This metric is more appropriate for our purposes than mean squared error since a ground truth is unavailable, but we have replicates; furthermore, we are not concerned with the actual values of the expression, just the relative values within the slide. Under perfect experimental conditions, we expect the correlation to be very close to 1; in reality, this often is not the case. An ideal microarray denoising algorithm would bring the correlation between the two replicates as close to 1 as possible without destroying the signal. The histogram in Figure 4.3, shows the distribution of correlations between replicate-pairs before processing by the algorithms (there are 114 experiments in the Zhang data). Although most replicate-pairs already have a high-correlation, a significant proportion (29% of 114 replicate pairs) have correlation under 0.9.

Note it is possible to cheat by setting $s_i = y_i$ for all *i*, resulting in a detrended result of all zeros in both replicates, and a correlation of 1. In order to safeguard against such pathological algorithms, one can check the change in correlation between pairs after only processing one replicate in the pair; a large systematic decrease in the correlation is strong evidence of a pathological algorithm. Our observation is that the algorithms whose results we report appear not to be of this type.

In addition to the Zhang dataset already described we also conduct experiments with a semi-artificial dataset, whose results are easier to analyze. To construct this dataset we

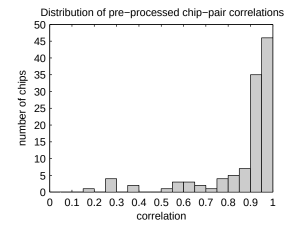


Figure 4.3: histogram of correlations between replicate pair experiments

took the highest correlated (before processing) replicate-pair out of the 114 and added artificial artefacts to one of the replicates (green). Before adding artefacts, this pair had a correlation of .98, which is close to one, and hence was apparently free of artefacts and noise. We created 10 semi-artificial microarray slide datasets by adding 10 sets of artefacts (shown in figure 4.4) to the green slide. The advantage of this dataset is we can compare the estimated artefacts with the actual artefacts, unlike in the Zhang dataset. Thus as a performance metric we may take the correlation between the green data with artefacts added after processing and original green or red slides. There is no possibility of cheating here since only one slide will be processed in a pair.

4.5 Results

4.5.1 Artificial artefacts results

Figure 4.5 shows the spatial components or (local) trend estimated by each of the three methods, for each semi-artificial microarray slide. Comparing to Figure 4.4, the Medians estimates appears to be the least successful in separating out the spatial trend. The STR and GP look more successful, but it is difficult to make strong claims about their relative

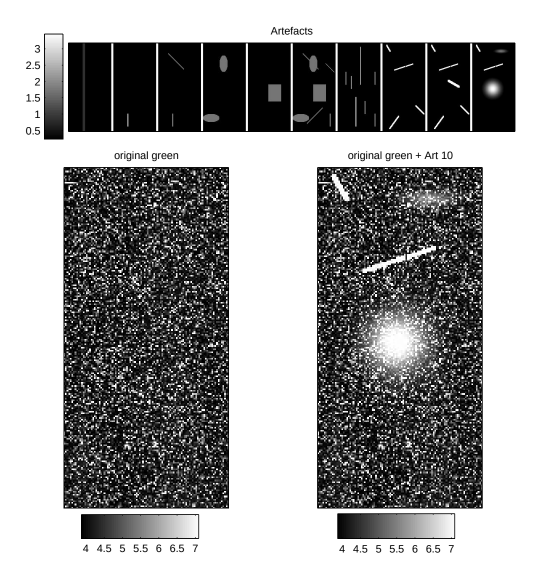


Figure 4.4: Top row: Ten sets of artificial artefacts (we label them $1, \ldots, 10$ from left to right) to be added to a microarray data slide. Bottom row: the original slide to which artefacts are added and the result of adding the 10^{th} set of artefacts

performance just from Figure 4.5. However, we may observe that the GP trends more accurately capture the thin, sharp line segments found in artefact cases 7-10.

Stronger, more objective statements can be made by looking at the change in correlation between each artificially corrupted slide and its uncorrupted replicate. These results are shown in Figure 4.6. As expected, in most cases the Medians method does worst, and in cases 1-3 actually decreases the correlation, so that not processing at all would have done better. Agreeing with our visual remarks in the previous paragraph, the GP algorithm performs better than STR in the line-segment-dominated cases of 7-9. The performance gap is less severe in case 10, which has a mix of smooth and line-segment artefacts. In all cases, the GP method performs at least as well as the STR method and never performs worse than doing nothing.

4.5.2 Real artefacts

Figure 4.7 shows a few examples of the decomposition of microarrays from the Zhang dataset into their spatially-independent (denoised microarray expression) and spatially-dependent (microarray spatial artefacts) components. These examples illustrate the diversity of spatial artefacts found in the dataset and the versatility of our method at estimating spatial artefacts/trends.

Figure 4.8 shows a scatter plot of the complete results for the Zhang dataset for the three methods under consideration. We see that GP and STR consistently beat Medians and the gap is wider for replicate-pairs with lower correlation. Another notable difference between GP/STR and medians is that GP/STR rarely decrease the correlation. For the subset of pairs which had a pre-processing correlation above 0.9, it is unclear which of GP and STR performs better, but when the pre-processing correlation<0.9, GP always performs at least as well as STR and GP is never worse than doing nothing.

Figures 4.9 and 4.10 are perhaps easier to interpret. The first figure shows that the average percentage improvement (in correlation) of GP over STR is comparable to that of

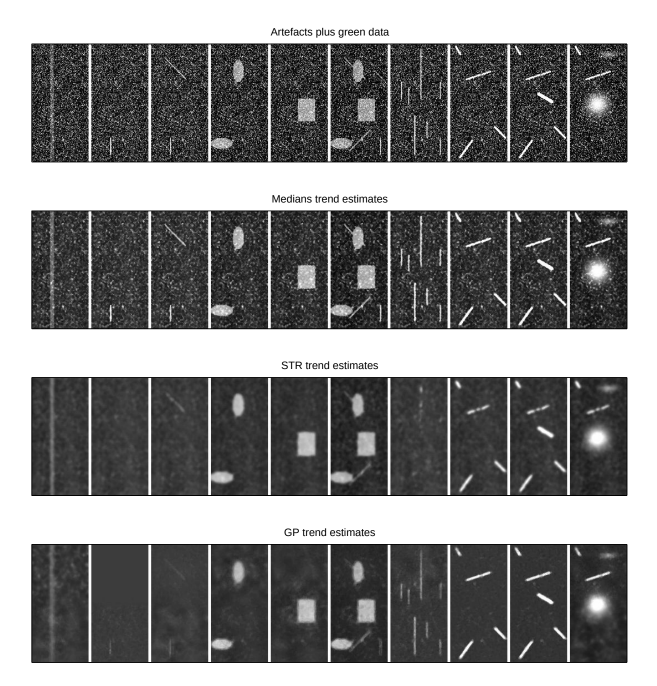


Figure 4.5: Row 1: Data to detrend used as input to the methods. Row 2: Corresponding trends estimated by the Medians algorithm. Row 3: Trends estimated by the STR algorithm. Row 4: Trends estimated by the GP algorithm.

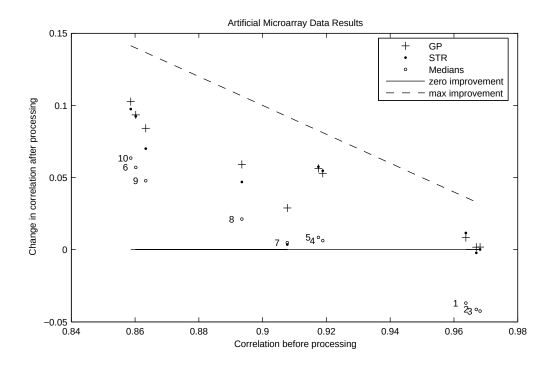


Figure 4.6: The correlation improvement results for the three methods on the artificial dataset. Each Medians result is labelled with the artefact case number (1-10).

STR over medians and the magnitude of this improvement increases as the pre-processing correlation decreases. The second figure shows a similar result using negative percentage change in 1-correlation as a measure of improvement.

4.6 Discussion

We have shown that our GP algorithm performs at least as well as and often better than STR in both our semi-artificial and the real, Zhang dataset on low PPC replicate-pairs. In the real data case, the percentage improvement of GP over STR is comparable to that of STR over medians.

With Gaussian process regression methods, one has to be careful with the time complexity, but our median-scheme combined with partitioning each slide into two has made the total processing time per chip comparable to STR^3 – on the order of 10 minutes with a 3 GHz CPU (doing each partition sequentially). Our algorithm can scale linearly to larger microarrays by fixing the partition size while increasing the number of partitions. Median filtering is still much faster but its performance is likely unacceptable.

³In the STR algorithm, determining the Gaussian kernel parameter by gradient-based optimization dominates the time complexity.

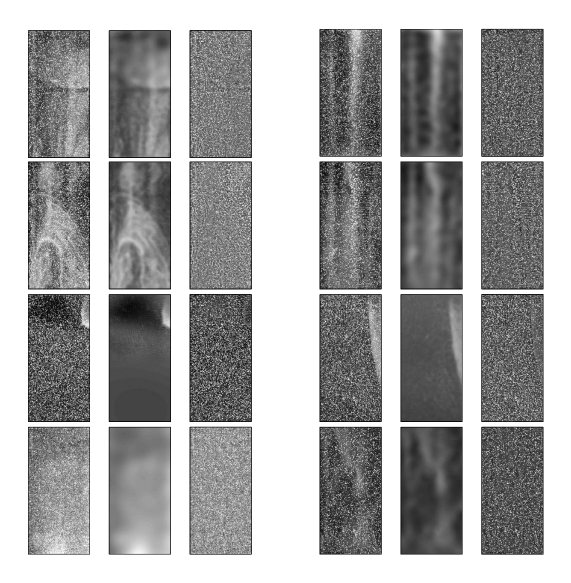


Figure 4.7: Eight examples of microarray decompositions by the our method into spatial and independent components. In each triplet of images, the left one is the original microarray slide, the centre is the estimated spatial component, and the right is the estimated independent component.

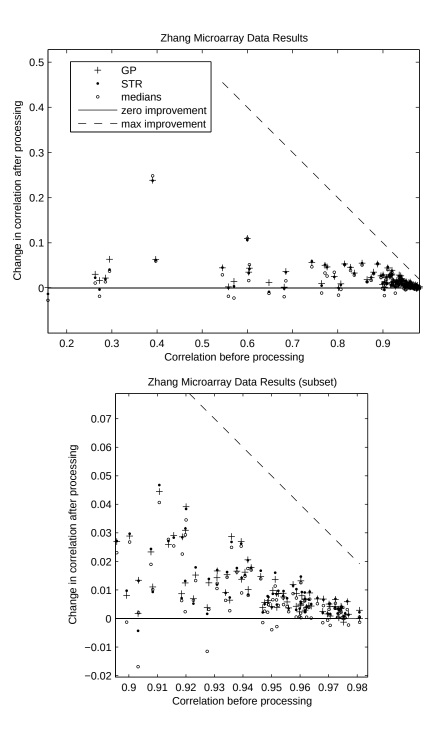


Figure 4.8: Top: Scatter plot of improvement results for all replicate pairs in the Zhang dataset. Bottom: Improvement results for replicate pairs with pre-processing correlation of at least 0.9.

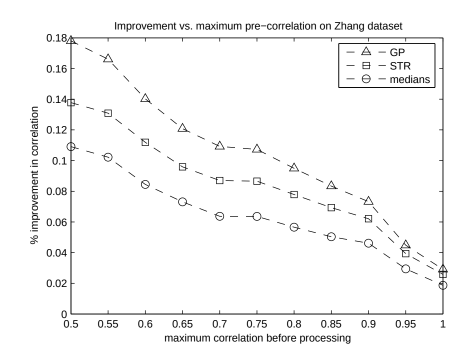


Figure 4.9: Shows the average % increase in correlation between replicate pairs after processing by the 3 algorithms considering only those pairs with a correlation (before processing) below a threshold (found on the x-axis).

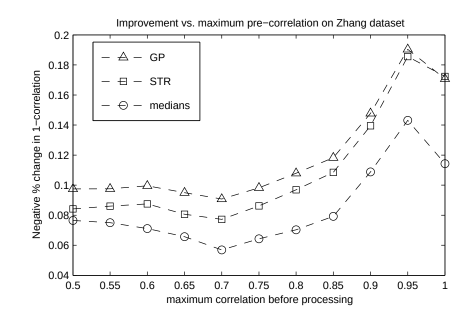


Figure 4.10: Shows the average negative % change in 1-correlation between replicate pairs after processing by the 3 algorithms considering only those pairs with a correlation (before processing) below a threshold (found on the x-axis).

Chapter 5

Image denoising

5.1 Description of Problem

Images captured by digital cameras or scanners are rarely noise-free. The nature of the noise artefacts in images can vary significantly, but here we focus on the unavoidable shot noise of the charge-coupled device (CCD), the main image sensor in most imaging devices. The CCD is essentially a two-dimensional array partitioned into small, equalsized, (usually) square regions called pixels, each of which has sensors designed to estimate the number of photons arriving in its region in some time interval (exposure time). In the extreme cases of low-light or fast-action photography, the number of photons is small enough that the sampling error is detrimental; in most other cases, the effect is not as noticeable but still undesirable. The random process of photon counting is usually thought to be a Poisson process, where we expect the number of photons, P, to have standard deviation \sqrt{P} [2]; that is, the noise is intensity dependent. However, as a simplification, most noise models in the image processing literature assume the noise is independent, identically distributed (IID) Gaussian. When the mean number of photons

Usually image data is received in the form of a matrix specifying the grey-level or

RGB vector for a pixel in row *i* and column *j*. Without loss of generality¹, we restrict ourselves to grey-level images; after re-labelling the data and allowing **x** to be the (row, column) of a pixel and *y* to be the grey-level of the image, we see we have two-dimensional grid data $\{(x_i, y_i)\}_{i=1}^N \subset \mathbb{R}^2 \times \mathbb{R}$. If we let the true image be the spatial component and the shot noise be the independent component, we arrive at the decomposition problem discussed in Chapter 1.

5.2 GP algorithm for denoising images

Our image denoising algorithm first divides an arbitrary natural image into approximately 30x30 pixel patches, which overlap with adjacent patches by 6 pixels on all sides (see 3.3). For each patch, we use Gaussian process regression using the covariance function described in Equation 3.31 to obtain the predictive mean of the noise-free image, which in this case is the spatial component. Once predictions are obtained for each patch, we combine them to produce the complete denoised image. Multiple predictions at the same pixel are averaged.

Recall, that in our covariance function, the hyperparameter σ_{ε}^2 controls the estimate of the variance in the independent component (IID Gaussian). This can be learned along with the other hyperparameters, or can be fixed to an estimate obtained from the user. In commercial image denoising software, the estimate is often obtained with the aid of the user, who provides a bounding box around a region of the image with no detail. In our experiments, we assume this estimate is available and fix σ_{ε}^2 accordingly. This increases the robustness of the hyperparameter learning by avoiding certain bad local maxima in the marginal likelihood.

¹to denoise a colour image, denoise the R,G, and B channels separately

5.3 Experimental methodology

To compare image denoising algorithms, we will take three relatively noise-free images, add various amounts of IID Gaussian noise artificially, attempt to remove the noise using the algorithms, and compare the denoised results with the original images.

5.3.1 Adding artificial noise to images

Let σ_{image}^2 be the sample variance of an image before adding noise. Define the signal-tonoise ratio (SNR) of the image with artificially added white noise $N(0, \sigma_{noise}^2)$ as

$$\sigma_{image}/\sigma_{noise}$$
 (5.1)

A lower SNR corresponds to a noisier image, or in our earlier terminology, an image with a larger independent component contribution to each pixel. In practice, an image with SNR of 10 has barely noticeable noise, and one with an SNR of less than 2 is unacceptable to most people. For our experiments, we add noise such that the SNR is 2, 5, and 10 (identical noise patterns, but different noise magnitudes).

5.3.2 Performance metrics

We focus on two algorithm performance metrics. One is quantitative and objective, the other is qualitative and subjective, but perhaps more natural.

Because we are adding noise ourselves to a ground truth image, we can calculate the mean squared error (MSE) between the denoised image and true image:

$$\frac{1}{N}\sum_{i=1}^{N} (\tilde{s}_i - \hat{s}_i)^2 \tag{5.2}$$

where \tilde{s}_i is the true value of the i^{th} pixel, and \hat{s}_i is its estimated value from the noisy data. This error measure has the advantage of being objective and is often used in the image processing literature to compare algorithms.

The other performance metric is simply a visual comparison of true and denoised images by people. We present these results by showing the actual images and discussing their visual qualities. An attempt will be made to make only obvious statements, which the reader presumably will find non-controversial.

In certain pathological cases, the MSE metric and human-visual metric produce very different rankings. A simple example is an algorithm which perfectly denoises an image and translates it 1 pixel to the left. Calculating only the MSE would lead one to believe that the algorithm is very bad; whereas, most humans would not notice the translation and believe the algorithm is very good based only on visual inspection. Apart from such cases, we find that the MSE metric often agrees with visual inspection, but visual inspection can offer insight unobtainable from just the MSE numbers.

5.3.3 Image data

We use three commonly used images in the image denoising literature. Lena, Barbara and Boat.² The three images are shown in Figure 5.1. Each image is a 512x512 pixel, 8-bit graylevel image exhibiting a large range of intensity and detail. The Lena image has a mix of very low detail in the skin and the background with very high detail in the hat. The Barbara image has less extreme range of detail but also has periodic properties in the tablecloth and head covering. The Boat image has a very high amount of detail throughout most of the image, except in some regions of the sky.

5.3.4 Methods selected for comparison

As baselines in the MSE results, we use median filtering ("medians") with a 1-neighbourhood (see Section 2.1), and Gaussian filtering using STR (see Section 2.2) without outlier de-

 $^{^{2}}$ Grayscale Lena obtained by averaging RGB channels of color version found at http://www.cs.cmu.edu/~chuck/lennapg/lenna.shtml. Barbara and Boat images obtained from http://decsai.ugr.es/~javier/denoise/test_images/index.htm.



(a) Lena







Figure 5.1: The original images before adding noise.

tection. We also report the MSE of doing nothing to the noisy image ("original+noise"). The main comparison will be between our Gaussian process algorithm ("GP") introduced in Chapter 3 and the Nonlocal means algorithm ("NL means") of Buades et al. [3], with $h = 10\sigma_{noise}$, a neighbourhood size of 7x7, and search window size of 21x21, settings recommended by the authors. It is possible that the parameters suggested by the authors are not optimal for all images and noise levels and that some sort of search over parameter space would improve the performance of the NL means algorithm; however, we do not perform such a search. A comprehensive comparison between the NL means algorithm and several standard algorithms including Gaussian smoothing is found in [2], which claims NL means has state-of-the-art performance.

5.4 Results

	SNR 2	SNR 5	SNR 10
original+noise	455.55	73.02	18.36
medians	108.85	40.30	29.54
Gauss Filter	98.88	91.40	90.39
NL means	47.74	21.91	14.63
GP	50.60	20.87	10.90

5.4.1 Lena

Table 5.1: Mean squared errors for the Lena experiments.

The mean squared errors for the Lena experiments are found in table 5.1; the visual results can be inspected in Figure 5.3. We can see from the MSE table that both the NL-means and GP algorithms beat the two baselines by a large margin but perform similarly at various SNR levels. The trend in the difference in MSEs between NL-means and GP is that GP performs better at low noise-levels whereas NL-means does slightly



Figure 5.2: A closer look at Lena's hat after denoising in the case of SNR=5. The left image is the image before adding noise; the center is the result from the NL-means method; the right image is the result of the GP method.

better at higher noise-levels.

From the visual point of view, the NL-means and GP methods compare similarly, but the GP method preserves more of the high frequency detail in the image. A striking example of this is obtained from a closer inspection of Lena's hat at the medium noiselevel of SNR=5 where both methods perform similarly on the MSE metric (see Figure 5.2).

	SNR 2	SNR 5	SNR 10
original+noise	742.70	118.91	29.78
medians	388.87	281.25	263.92
Gauss Filter	332.68	319.79	318.06
NL means	87.70	33.40	20.31
GP	120.50	38.75	16.67

5.4.2 Barbara

Table 5.2: Mean squared errors for the Barbara experiments.

The mean squared errors for the Barbara experiments are found in table 5.2; the visual results can be inspected in Figure 5.4.

The MSE results for the Barbara image show similar trends as the Lena image except the NL-means algorithm performs significantly better at the high noise-level of SNR=2. This is not so surprising considering the high level of periodicity in the image (as in the table cloth, head covering, and pants), which is the case in which NL-means excels.

Examining the visual results, we see that the NL-means method is better able to denoise the periodic regions. However, the GP method is able to better preserve high frequency details in the non-periodic regions such as the face.

	SNR 2	SNR 5	SNR 10
original+noise	542.67	86.91	21.78
medians	175.15	89.58	75.88
Gauss Filter	212.96	204.21	203.07
NL means	89.46	41.94	26.91
GP	95.87	36.97	16.73

5.4.3 Boat

Table 5.3: Mean squared errors for the boat experiments.

The mean squared errors for the Boat experiments are found in table 5.3; the visual results can be inspected in Figure 5.5.

The rankings from the MSE results are similar to those obtained in the Lena image and again visually we see that the GP method is better at preserving high-frequency detail in the image. However, since the Boat image has the highest non-periodic detail of the three images, the effect is more dramatic.

5.5 Discussion

We have shown our GP method performs favourably – especially at moderate to low noise levels – against the NL-means algorithm, using the program settings recommended by the authors in [3]. We make this claim with respect to both the MSE and visual inspection (the ultimate) performance metrics. However, we note that in the visual results we saw that the GP method is significantly better at preserving non-periodic, high-frequency detail.

Since most real applications of image denoising occur at the medium-to-low noise levels, our GP method is likely better than the other methods in most practical situations.

Although our algorithm scales linearly in the number of pixels when the patch (partition) size is fixed, the time taken for each patch (30x30 pixel) partition is on the order of 5 minutes. For the test images used, this translated to a total time for denoising an image on the order of several hours in a MATLAB implementation run on a 3 GHz CPU doing one patch at a time, whereas a good implementation of NL-means would take a few minutes. Although, the time complexity of our method is within the limits of feasibility, future work in accelerating our method would increase its general usefulness.



(a) noise added



(b) NL means



(c) GP method

Figure 5.3: Lena image results. SNR left to right: 2, 5, 10.



(a) noise added



(b) NL means



(c) GP method

Figure 5.4: Barbara image results. SNR left to right: 2, 5, 10.



(a) noise added



(b) NL means



(c) GP method

Figure 5.5: Boat image results. SNR left to right: 2, 5, 10.

Chapter 6

Concluding remarks

6.1 Summary of thesis

In this thesis, we have treated microarray spatial artefact/trend removal and image denoising as essentially the same problem – the decomposition of data into spatially independent and dependent components. We showed that both problems can be solved using very similar methods. We presented a method using Gaussian processes for regression with a new type of covariance function, which can be used with any grid dataset. Through experiments in both application domains, we have shown that our method performs favourably compared to well-established methods in the both literatures.

However, we find that our method performs best in the case of medium-to-low energy in the spatially-independent component relative to the dependent component; this corresponds to the case of (more realistic) medium-to-low level of noise in image denoising. The performance improvement over other methods is less impressive for microarrays with low-energy spatial artefacts and images with high noise levels, where the independent component plays a more significant role in the data. It may be that the problem is simply too difficult, and that little can be done beyond small improvements, which are achievable by simple algorithms. One can justify this intuition by asking probably the best image denoiser, the human mind, how well it can distinguish the image from noise in the Lena image with an SNR=1 (see Figure 6.1).



Figure 6.1: The Lena image with an SNR=1. The image is arguably difficult to visually disentangle from the noise.

6.2 Future work

One possible improvement to our Gaussian process method may reduce the pixelation effects experienced at high-noise levels in image denoising resulting from discontinuities between the predictions of adjacent patches. When processing each patch/partition of an image (or microarray), instead of ignoring all the data outside of the patch, one could incorporate certain statistics computed from all other patches. Presumably, global trends (in the spatially-dependent component) extending beyond single patches could then be more easily inferred, without increasing the amount of training data significantly. Pixelation might be reduced as continuity across patches might be more encouraged. One possibly useful statistic would be the mean of each other patch. The covariance between means and single grid points needs to be handled slightly differently than between two grid points.

Speeding up our algorithm for large image/microarray datasets is of considerable practical importance. There are a number of directions one could take to achieve this. One is to approximate the inverse of the covariance matrix, whose computation is very time-consuming when learning the hyperparameters, the dominant step with respect to time complexity. To this end, one may use a low-rank approximation to the covariance matrix as in the Nystrom approximation [22], or approximate the Cholesky factorization as in [6]. A second direction is to use a subset of the data selected by some reasonable criteria to train the hyperaparameters; we achieved a similar reduction in training data using our medians scheme when applying our algorithm to microarray data, but more general methods exist. A review of these methods can be found in Chapter 8 of Rasmussen and Williams' book ([14]).

On the theoretical side, further analysis of our use of pseudo-inputs in our covariance function can be conducted. Our pseudo-inputs in fact are a subset of the noisy-outputs. Having the covariance between outputs depend on other (noisy) outputs may be viewed with suspicion. However, the fact that our method performs well in experiments suggests something has been gained. A new interpretation of what we are doing may lead to a more principled framework and allay our anxieties.

Finally, a comparision of the image denoising performance with state-of-the-art commercial applications such as Noise Ninja and Neat Image would be interesting.

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