Model selection

Models (labelled by $m$) have parameters $\theta_m$ that specify the probability of data:

$$P(D|\theta_m, m).$$

If model is known, learning $\theta_m$ means finding posterior or point estimate (ML, MAP, ...).

What if we need to learn the model too?

- Could combine models into a single "supermodel", with composite parameter $(m, \theta_m)$.
  - ML learning will overfit: favours most flexible (nested) model with most parameters, even if the data actually come from a simpler one.
  - Density function on composite parameter space (union of manifolds of different dimensionalities) difficult to define $\Rightarrow$ MAP learning ill-posed.
  - Joint posterior difficult to compute — dimension of composite parameter varies [but Monte-Carlo methods may sample from much a posterior.]

$\Rightarrow$ Separate model selection step:

$$P(\theta_m, m|D) = P(\theta_m|m, D) \cdot P(m|D)$$
Model selection

Given models labeled by $m$ with parameters $\theta_m$, identify the “correct” model for data $D$. ML/MAP has no good answer: $P(D|\theta_m^\text{ML})$ is always larger for more complex (nested) models.

Neyman-Pearson hypothesis testing

- For nested models. Starting with simplest model ($m = 1$), compare (e.g. by likelihood ratio test) null hypothesis $m$ to alternative $m + 1$. Continue until $m + 1$ is rejected.
- Usually only valid asymptotically in data number.
- Conservative (N-P hypothesis tests are asymmetric).

Likelihood validation

- Partition data into disjoint training and validation data sets $D = D_t \cup D_{\text{vld}}$. Choose model with greatest $P(D_{\text{vld}}|\theta_m^\text{ML})$, with $\theta_m^\text{ML} = \arg\max P(D_t|\theta)$. [Or, better, greatest $P(D_{\text{vld}}|D_t, m)$.]
- May be biased towards simpler models; often high-variance.
- Cross-validation uses multiple partitions and averages likelihoods.

Bayesian model selection

- Choose most likely model: $\arg\max P(m|D)$.
- Principled from a probabilistic viewpoint—if true model is in set being considered—but sensitive to assumed priors etc.
- Can use posterior probabilities to weight models for combined predictions (no need to select at all).

Bayesian model selection: some terminology

A model class $m$ is a set of distributions parameterised by $\theta_m$, e.g. the set of all possible mixtures of $m$ Gaussians.

The model implies both a prior over the parameters $P(\theta_m|m)$, and a likelihood of data given parameters (which might require integrating out latent variables) $P(D|\theta_m, m)$.

The posterior distribution over parameters is

$$P(\theta_m|D, m) = \frac{P(D|\theta_m, m)P(\theta_m|m)}{P(D|m)}.$$  

The marginal probability of the data under model class $m$ is:

$$P(D|m) = \int_{\theta_m} P(D|\theta_m, m)P(\theta_m|m) \, d\theta_m.$$  

(also called the Bayesian evidence for model $m$).

The ratio of two marginal probabilities (or sometimes its log) is known as the Bayes factor:

$$\frac{P(D|m)}{P(D|m')} = \frac{P(m|D)}{P(m'|D)} \frac{p(m)}{p(m')}.$$  

The Bayesian Occam’s razor

Occam’s Razor is a principle of scientific philosophy: of two explanations adequate to explain the same set of observations, the simpler should always be preferred. Bayesian inference formalises and automatically implements a form of Occam’s Razor.

Compare model classes $m$ using their posterior probability given the data:

$$P(m|D) = \frac{P(D|m)P(m)}{P(D)}.$$  

$$P(D|m): \text{ The probability that randomly selected parameter values from the model class would generate data set } D.$$  

Model classes that are too simple are unlikely to generate the observed data set. Model classes that are too complex can generate many possible data sets, so again, they are unlikely to generate that particular data set at random. Like Goldilocks, we favour a model that is just right.

Bayesian model comparison: Occam’s razor at work

Compare the model classes using their posterior probability:

$$P(m|D) = \frac{P(D|m)P(m)}{P(D)}.$$  

Model classes that are too simple are unlikely to generate the observed data set. Model classes that are too complex can generate many possible data sets, so again, they are unlikely to generate that particular data set at random. Like Goldilocks, we favour a model that is just right.
Conjugate-exponential families (recap)

Can we compute \( P(D|m) \)?

Sometimes.

Suppose \( P(D|\theta_m, m) \) is a member of the exponential family:

\[
P(D|\theta_m, m) = \prod_{i=1}^{N} P(x_i|\theta_m, m) = \prod_{i=1}^{N} e^{g(x_i)\theta_m - A(\theta_m)}.
\]

If our prior on \( \theta_m \) is conjugate:

\[
P(\theta_m|m) = e^{s_0(\theta_m - \theta_m^0)/Z(s_0, n_0)}
\]

then the joint is in the same family:

\[
P(D, \theta_m|m) = e^{\left(\sum_i s(x_i) + s_0\right)\theta_m - (N + n_0)A(\theta_m)/Z(s_0, p)}
\]

and so:

\[
P(D|m) = \int d\theta_m P(D, \theta_m|m) = Z(\sum_i s(x_i) + s_0, N + n_0)/Z(s_0, p)
\]

But this is a special case. In general, we need to approximate . . .

Laplace approximation

We want to find \( P(D|m) \) = \( \int P(D, \theta_m|m) \) \( d\theta_m \).

As data size \( N \) grows (relative to parameter count \( d \)), \( \theta_m \) becomes more constrained.

\( \Rightarrow P(D, \theta_m|m) \propto P(\theta_m|m) \) becomes concentrated on posterior mode \( \theta_m^* \).

Idea: approximate \( \log P(D, \theta_m|m) \) to second-order around \( \theta^* \):

\[
\int P(D, \theta_m|m) d\theta_m = \int e^{\log P(D, \theta_m|m)} d\theta_m
= \int e^{\log P(D, \theta_m^*|m) + \sum \log P(D, \theta_m^*|m) (\theta_m - \theta_m^*) + \frac{1}{2} (\theta_m - \theta_m^*) A(\theta_m - \theta_m^*)} d\theta_m
= \int P(D, \theta_m^*|m) e^{-\frac{1}{2}(\theta_m - \theta_m^* A(\theta_m - \theta_m^*)} d\theta_m
= P(D|\theta_m^*|m)P(\theta_m^*|m)(2\pi)^{-\frac{d}{2}}|A|^{-\frac{1}{2}}
\]

\( A = \nabla^2 \log P(D, \theta^*|m) \) is the negative Hessian of \( \log P(D, \theta|m) \) evaluated at \( \theta_m^* \).

This is equivalent to approximating the posterior by a Gaussian: an approximation which is asymptotically correct.

Practical Bayesian approaches

- Laplace approximation:
  - Approximate posterior by a Gaussian centred at the maximum a posteriori parameter estimate.

- Bayesian Information Criterion (BIC)
  - an asymptotic \((N \to \infty)\) approximation.

- Variational Bayes
  - Lower bound on the marginal probability.
  - Biased estimate.
  - Easy and fast, and often better than Laplace or BIC.

- Monte Carlo methods:
  - Importance sampling: estimate evidence using samples \( \theta^{(i)} \) from arbitrary \( f(\theta) \):
    \[
    \sum_{i} P(D(\theta^{(i)}), m)P(\theta^{(i)}|m) \to \int d\theta f(\theta) P(D, \theta|m) = P(D|m)
    \]
    *Reversible jump* Markov Chain Monte Carlo: sample from posterior on composite \((m, \theta_m)\).
    - If samples for each \( m \propto p(m|D) \).
    - Both exact in the limit of infinite samples, but may have high variance with finite samples.

Not an exhaustive list (Bethe approximations, Expectation propagation, . . . )

We will discuss Laplace and BIC now, leaving the rest for the second half of course.

Bayesian Information Criterion (BIC)

BIC can be obtained from the Laplace approximation:

\[
\log P(D|m) \approx \log P(\theta_m^*|m) + \log P(D|\theta_m^*, m) + \frac{d}{2} \log 2\pi - \frac{1}{2} \log |A|
\]

We have

\[
A = \nabla^2 \log P(D, \theta^*|m) = \nabla^2 \log P(D|\theta^*, m) + \nabla^2 \log P(\theta^*|m)
\]

So as the number of iid data \( N \to \infty \), \( A \) grows as \( NA_0 + \text{constant for a fixed matrix } A_0 \).

\( \Rightarrow \log |A| \to \log |NA_0| = \log(N^d |A_0|) = d \log N + \log |A_0| \).

Retaining only terms that grow with \( N \) we get:

\[
\log P(D|m) \approx \log P(D|\theta_m^*, m) - \frac{d}{2} \log N
\]

Properties:

- Quick and easy to compute.
- Does not depend on prior.
- We can use the ML estimate of \( \theta \) instead of the MAP estimate (= as \( N \to \infty \)).
- Related to the "Minimum Description Length" (MDL) criterion.
- Assumes that in the large sample limit, all the parameters are well-determined (i.e. the model is identifiable; otherwise, \( d \) should be the number of well-determined parameters).
- Neglects multiple modes (e.g. permutations in a MoG).
Hyperparameters and Evidence optimisation

In some cases, we need to choose between a family of continuously parameterised models.

\[ P(D|\eta) = \int P(D|\theta)P(\theta|\eta) \, d\theta \]

This choice can be made by ascending the gradient in:
- the exact evidence (if tractable).
- the approximated evidence (Laplace, EP, Bethe, . . . )
- a free-energy bound on the evidence (Variational Bayes)

or by placing a hyperprior on the hyperparameters \( \eta \), and sampling from the posterior

\[ P(\eta|D) = \frac{P(D|\eta)P(\eta)}{P(D)} \]

using Markov chain Monte Carlo sampling.

The evidence for linear regression

- The posterior on \( \mathbf{w} \) is normal: \( \Sigma_w = \left( \frac{X^T X}{\alpha} + C^{-1} \right)^{-1}; \bar{\mathbf{w}} = \Sigma_w \frac{X^T Y}{\alpha} \).
  Note: \( X \) is a matrix where columns are input vectors, and \( Y \) is a row vector of corresponding predicted outputs.

- The evidence, \( \mathcal{E}(C, \sigma^2) = \int P(Y|X, \mathbf{w}, \sigma^2)P(\mathbf{w}|C) \, d\mathbf{w} \), is given by:

\[ \mathcal{E}(C, \sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} \left( \frac{1}{2} \left( \frac{Y}{\sigma^2} - \frac{X^T \Sigma_w X}{\sigma^4} \right) Y^T \right) \]

- For optimization, general forms for the gradients are available. If \( \theta \) is a parameter in \( C \):

\[ \frac{\partial}{\partial \theta} \log \mathcal{E}(C, \sigma^2) = \frac{1}{2} \text{Tr} \left[ (C - \Sigma_w - \bar{\mathbf{w}} \bar{\mathbf{w}}^T) \frac{\partial}{\partial \theta} C^{-1} \right] \]

\[ \frac{\partial}{\partial \sigma^2} \log \mathcal{E}(C, \sigma^2) = \frac{1}{\sigma^2} \left[ -N + \text{Tr} \left( I - \Sigma_w C^{-1} \right) + \frac{1}{\sigma^2} (Y - \bar{\mathbf{w}} \bar{\mathbf{w}}^T)(Y - \bar{\mathbf{w}} \bar{\mathbf{w}}^T)^T \right] \]

Evidence optimisation in linear regression

Consider simple linear regression:

\[ \mathbf{w} \sim \mathcal{N}(\mathbf{0}, C) \]

\[ y_i \sim \mathcal{N}(\mathbf{\bar{w}}^T \mathbf{x}_i, \sigma^2) \]

- Maximize

\[ P(\mathbf{y}_1, \ldots, \mathbf{y}_N|\mathbf{x}_1, \ldots, \mathbf{x}_N, C, \sigma^2) = \int P(\mathbf{y}_1, \ldots, \mathbf{y}_N|\mathbf{x}_1, \ldots, \mathbf{x}_N, \mathbf{w}, \sigma^2)P(\mathbf{w}|C) \, d\mathbf{w} \]

to find optimal values of \( C, \sigma \).
- Compute the posterior \( P(\mathbf{w}|\mathbf{y}_1, \ldots, \mathbf{y}_N, \mathbf{x}_1, \ldots, \mathbf{x}_N, C, \sigma^2) \) given these optimal values.

Automatic Relevance Determination

The most common form of evidence optimization for regression (due to MacKay and Neal) takes \( C^{-1} = \text{diag}(\alpha) \) (i.e. \( \mathbf{w} \sim \mathcal{N}(0, \alpha^{-1}) \)) and then optimizes the precisions \( \{\alpha_i\} \).

Setting the gradients to 0 and solving gives

\[ \alpha_i^{\text{new}} = \frac{1 - \alpha_i \Sigma_w |_{i}}{\bar{\mathbf{w}}_i^T} \]

\[ (\sigma^2)^{\text{new}} = \frac{(Y - \bar{\mathbf{w}} X)(Y - \bar{\mathbf{w}}^T X)^T}{N - \Sigma_i (1 - \Sigma_w |_{i})} \]

During optimization the \( \alpha_i \)'s meet one of two fates

\[ \alpha_i \to \infty \Rightarrow w_i = 0 \quad \text{irrelevant input} \ x_i \]

\[ \alpha_i \text{ finite} \Rightarrow w_i = \text{argmax} P (w_i | X, Y, \alpha_i) \quad \text{relevant input} \ x_i \]

This procedure, Automatic Relevance Determination (ARD), yields sparse solutions that improve on ML regression. (cf. L1-regression or LASSO).

Evidence optimisation is also called maximum marginal likelihood or ML-2 (Type 2 maximum likelihood).
### Prediction averaging

![Diagram](image)

Linear regression predicts output $y$ given input vector $x$ by:

$$ y \sim \mathcal{N}(w^T x, \sigma^2) $$

Posterior over $w$ is Gaussian with covariance $\Sigma_w = (\frac{1}{\tau^2}XX^T + \frac{1}{\sigma^2}I)^{-1}$ and mean $\bar{w} = \frac{1}{\tau^2}XX^T$ (where $X$ is matrix with columns being input vectors, $Y$ is row vector of outputs).

Given a new input vector $x'$, the predicted output $y'$ is (integrating out $w$):

$$ y' | x' \sim \mathcal{N}(\bar{w}^T x', \bar{w}^T \Sigma_w x' + \sigma^2) $$

the additional variance term $x'^T \Sigma_w x'$ comes from the posterior uncertainty in $w$.

### Predictions with marginalised linear regression

Now, include the test input vector $x'$ and test output $y'$:

$$ \begin{bmatrix} y' \\ y \end{bmatrix} | X, x' \sim \mathcal{N}( \begin{bmatrix} 0 \\ 0 \end{bmatrix}, \begin{bmatrix} \tau^2 X^T X + \sigma^2 I & \tau^2 X^T x' \\ \tau^2 x'^T X & \tau^2 x'^T x' + \sigma^2 \end{bmatrix}) $$

We can find $y'|Y$ by the standard multivariate Gaussian result:

$$ a \sim \mathcal{N}( \begin{bmatrix} 0 \\ 0 \end{bmatrix}, \begin{bmatrix} A & C \\ C^T & B \end{bmatrix}) \Rightarrow b|a \sim \mathcal{N}(A^T b - C^T a, B - C^T A^{-1} C) $$

So

$$ y'|Y, X, x' \sim \mathcal{N} \left( \tau^2 x'^T X + \sigma^2 \right)^{-1} Y', \tau^2 x'^T x' + \sigma^2 - \tau^2 x'^T X \left( \tau^2 X^T X + \sigma^2 \right)^{-1} \tau^2 x'^T x' \right) $$

$$ \sim \mathcal{N} \left( \frac{1}{\tau^2} x'^T \Sigma X Y^T, x'^T \Sigma x' + \sigma^2 \right) \quad \Sigma = \left( \frac{1}{\tau^2} XX^T + \frac{1}{\sigma^2} I \right)^{-1} $$

- Same answer as obtained by integrating wrt posterior over $w$.
- Evidence $P(Y|X)$ is just probability under joint Gaussian; also reduces to expression found previously.
- Thus, Bayesian linear regression can be derived from a joint, parameter-free distribution on all the outputs conditioned on all the inputs.

### Marginalised linear regression

![Diagram](image)

Integrate out $w$: the joint distribution of $y_1, \ldots, y_N$ given $x_1, \ldots, x_N$ is Gaussian. The means and covariances are:

$$ E[y_i] = E[w^T x_i] = 0, \quad x_i = 0 $$

$$ E[(y_i - \bar{y})^2] = E[(\bar{w}^T w)(w^T x_i)] + \sigma^2 = \tau^2 x_i^T x_i + \sigma^2 $$

$$ E[(y_i - \bar{y})(y_j - \bar{y})] = E[(\bar{w}^T w)(w^T x_i)] = \tau^2 x_i^T x_j $$

$$ y_1, y_2, \ldots, y_N \sim \mathcal{N} \left( \begin{bmatrix} 0 \\ 0 \\ \vdots \\ 0 \end{bmatrix}, \begin{bmatrix} \tau^2 x_1^T x_1 + \sigma^2 & \tau^2 x_1^T x_2 & \ldots & \tau^2 x_1^T x_N \\ \tau^2 x_2^T x_1 & \tau^2 x_2^T x_2 + \sigma^2 & \ldots & \tau^2 x_2^T x_N \\ \vdots & \vdots & \ddots & \vdots \\ \tau^2 x_N^T x_1 & \tau^2 x_N^T x_2 & \ldots & \tau^2 x_N^T x_N + \sigma^2 \end{bmatrix} \right) $$

### Nonlinear regression

![Diagram](image)

We can also introduce a nonlinear mapping $x \mapsto \phi(x)$. Each element of $\phi(x)$ is a (nonlinear) feature extracted from $x$. May be many more features than elements on $x$.

The regression function $f(x) = w^T \phi(x)$ is nonlinear, but outputs $Y$ still jointly Gaussian:

$$ Y^T | X \sim \mathcal{N}(0, \tau^2 \Phi^T \Phi + \sigma^2 I) $$

where the $i^{th}$ column of matrix $\Phi$ is $\phi_i(x)$.

Proceeding as before, the predictive distribution over $y'$ for a test input $x'$ is:

$$ y'|x', Y, X \sim \mathcal{N} \left( \tau^2 \phi(x')^T K^{-1} Y', \tau^2 \phi(x')^T \Phi K^{-1} \phi(x') + \sigma^2 - \tau^4 \phi(x')^T \Phi K^{-1} \Phi^T \phi(x') \right) $$

$$ K = \tau^2 \Phi^T \Phi + \sigma^2 I $$
The Gaussian process

The covariance kernel

\[ Y^T X \sim \mathcal{N} \left( 0_N, \Sigma + \sigma^2 I_N \right) \]

The covariance of the output vector \( Y \) plays a central role in the development of the theory of Gaussian processes.

Define the covariance kernel function \( K : \mathbb{X} \times \mathbb{X} \to \mathbb{R} \) such that if \( x, x' \in \mathbb{X} \) are two input vectors with corresponding outputs \( y, y' \), then

\[ K(x, x') = \text{Cov}[y, y'] = E[y y'] - E[y]E[y'] \]

In the nonlinear regression example we have \( K(x, x') = \tau^2 \phi(x)^T \phi(x') + \sigma^2 \delta_{x=x'} \).

The covariance kernel has two properties:

- Symmetric: \( K(x, x') = K(x', x) \) for all \( x, x' \).
- Positive semidefinite: the matrix \( [K(x_i, x_j)] \) formed by any finite set of input vectors \( x_1, \ldots, x_n \) is positive semidefinite.

Theorem: A covariance kernel \( K : \mathbb{X} \times \mathbb{X} \to \mathbb{R} \) is symmetric and positive semidefinite if and only if there is a feature map \( \phi : \mathbb{X} \to \mathbb{H} \) such that

\[ K(x, x') = \phi(x)^T \phi(x') \]

The feature space \( \mathbb{H} \) can potentially be infinite dimensional.

The Gaussian process

A Gaussian process (GP) is a collection of random variables, any finite number of which have (consistent) Gaussian distributions.

In our regression setting, for each input vector \( x \) we have an output \( f(x) \). Given \( X = \{x_1, \ldots, x_n\} \), the joint distribution of the outputs \( F = \{f(x_1), \ldots, f(x_n)\} \) is:

\[ F|X, K \sim \mathcal{N}(0, K(X, X)) \]

Thus the random function \( f(x) \) (as a collection of random variables, one \( f(x) \) for each \( x \)) is a Gaussian process.

In general, a Gaussian process is parametrized by a mean function \( m(x) \) and covariance kernel \( K(x, x') \), and we write

\[ f(\cdot) \sim \mathcal{GP}(m(\cdot), K(\cdot, \cdot)) \]

Posterior Gaussian process: on observing \( X \) and \( F \), the conditional joint distribution of \( F' = \{f(x_1'), \ldots, f(x_n')\} \) on another set of input vectors \( x_1', \ldots, x_n' \) is still Gaussian:

\[ F'|X', X, F \sim \mathcal{N}(K(X', X)K(X, X)^{-1}F, K(X', X') - K(X', X)K(X, X)^{-1}K(X, X')) \]

thus the posterior over functions \( f(\cdot)|X, F \) is still a Gaussian process.

Regression using the covariance kernel

For non-linear regression, all operations depended on \( K(x, x') \) rather than explicitly on \( \phi(x) \).

So we can define the joint in terms of \( K \) implicitly using a (potentially infinite-dimensional) feature map \( \phi(x) \).

\[ Y|X, K \sim \mathcal{N}(0_N, K(X, X)) \]

where the \( i, j \) entry in the covariance matrix \( K(X, X) \) is \( K(x_i, x_j) \).

This is called the kernel trick.

Prediction: compute the predictive distribution of \( y' \) conditioned on \( Y \):

\[ y'|X, Y, K \sim \mathcal{N}(K(x', X)K(X, X)^{-1}Y, K(x', x') - K(x', X)K(X, X)^{-1}K(X, x')) \]

Evidence: this is just the Gaussian likelihood:

\[ P(Y|X, K) = \left| 2\pi K(X, X) \right|^{-\frac{1}{2}} e^{-\frac{1}{2}y^T \left( \Sigma + \sigma^2 I \right)^{-1}y} \]

Evidence optimisation: the covariance kernel \( K \) often has parameters, and these can be optimized by gradient ascent in \( \log P(Y|X, K) \).

Regression with Gaussian processes

We seek to learn the function that maps inputs \( x_1, \ldots, x_N \) to outputs \( y_1, \ldots, y_N \).

Instead of assuming a specific form, we consider a random function drawn from a GP prior:

\[ f(\cdot) \sim \mathcal{GP}(0, K(\cdot, \cdot)) \]

Any function is possible (no restriction on support) but some are (much) more likely \( a \text{ priori} \).

Observations \( y_i \) usually taken to be noisy versions of latent \( f(x) \):

\[ y_i|f(x_i), K \sim \mathcal{N}(f(x_i), \sigma_0^2) \]

Evidence: given by the multivariate Gaussian likelihood:

\[ P(Y|X, K) = \left| 2\pi (K(X, X) + \sigma_0^2 I) \right|^{-\frac{1}{2}} e^{-\frac{1}{2}y^T (K(X, X) + \sigma_0^2 I)^{-1}y} \]

Posterior: also a GP:

\[ f(\cdot)|X, Y \sim \mathcal{GP}(K(\cdot, \cdot)(K(X, X) + \sigma_0^2 I)^{-1}Y, K(\cdot, \cdot) - K(\cdot, X)(K(X, X) + \sigma_0^2 I)^{-1}K(X, \cdot)) \]

Predictions: posterior on \( f \), plus observation noise:

\[ y'|X, Y, f \sim \mathcal{N}(E[f(\cdot)|X, Y], \text{Var}[f(\cdot)|X, Y] + \sigma_0^2) \]

Evidence Optimisation: gradient ascent in \( \log P(Y|X) \).
Samples from a Gaussian process

We can draw sample functions from a GP by fixing a set of input vectors \( x_1, \ldots, x_N \), and drawing a sample \( f(x_1), \ldots, f(x_N) \) from the corresponding multivariate Gaussian. This can then be plotted.

Example prior and posterior GPs:

Another approach is to

- sample \( f(x_1) \) first,
- then \( f(x_2) | f(x_1) \),
- and generally \( f(x_n) | f(x_1), \ldots, f(x_{n-1}) \) for \( n = 1, 2, \ldots \).

Examples of covariance kernels

- Polynomial:
  \[
  K(x, x') = (1 + x^T x')^m \quad m = 1, 2, \ldots
  \]

- Squared-exponential (or exponentiated-quadratic):
  \[
  K(x, x') = \theta^2 e^{-\frac{\|x - x'\|^2}{2\eta^2}}
  \]

- Periodic (exp-sine):
  \[
  K(x, x') = \theta^2 e^{-2 \sin^2 \left( \frac{\pi (x - x')}{\tau} \right)}
  \]

- Rational Quadratic:
  \[
  K(x, x') = \left( 1 + \frac{\|x - x'\|^2}{2\alpha \eta^2} \right)^{-\alpha} \quad \alpha > 0
  \]

Forms of kernels

If \( K_1 \) and \( K_2 \) are covariance kernels, then so are:

- Rescaling: \( \alpha K_1 \) for \( \alpha > 0 \).
- Addition: \( K_1 + K_2 \)
- Elementwise product: \( K_1 K_2 \)
- Mapping: \( K_1(\phi(x), \phi(x')) \) for some function \( \phi \).

A covariance kernel is translation-invariant if

\[
K(x, x') = h(|x - x'|)
\]

A GP with a translation-invariant covariance kernel is stationary: if \( f(\cdot) \sim \GP(0, K) \), then so is \( f(\cdot - x) \sim \GP(0, K) \) for each \( x \).

A covariance kernel is radial or radially symmetric if

\[
K(x, x') = h(||x - x'||)
\]

A GP with a radial covariance kernel is stationary with respect to translations, rotations, and reflections of the input space.
Overparameterised models can overfit. In the GP, the parameter is the function \( f(x) \) which can be infinite-dimensional.

However, the Bayesian treatment integrates over these parameters: we never identify a single “best fit” \( f \), just a posterior (and posterior mean). So \( f \) cannot be adjusted to overfit the data.

The GP is an example of the larger class of nonparametric Bayesian models.

- Infinite number of parameters.
- Often constructed as the infinite limit of a nested family of finite models (sometimes equivalent to infinite model averaging).
- Parameters integrated out, so effective number of parameters to overfit is zero or small (hyperparameters).
- No need for model selection. Bayesian posterior on parameters will concentrate on “submodel” with largest integral automatically.
- No explicit need for Occam’s razor, validation or added regularisation penalty.