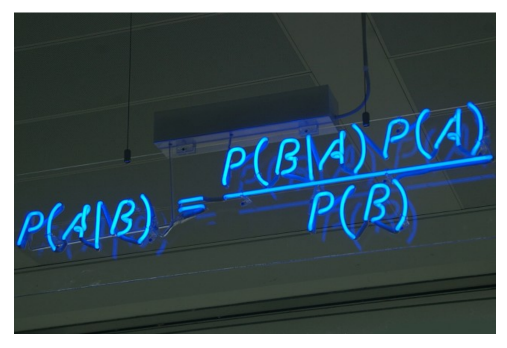
Bayesian Learning



NB

Note, we'll use another library in this notebook: GPy Installation:

```
[3]: import GPy
   import pods
```

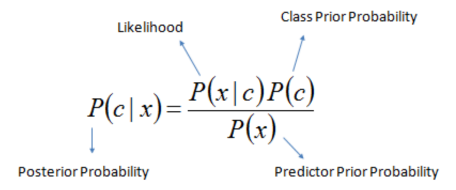
Naive Bayes

Predict the probability that a point belongs to each class, using Bayes' Theorem, assuming that the features are independent from each other.

Very fast. They work by only extracting statistics from each feature.

Naive Bayes Classifier

A Naive Bayes classifier predicts the class of each sample using Bayes' rule: P(c|x) is the posterior probability of class (target) given predictor (attribute). P(c) is the *prior* probability of class.



Bayes Rule

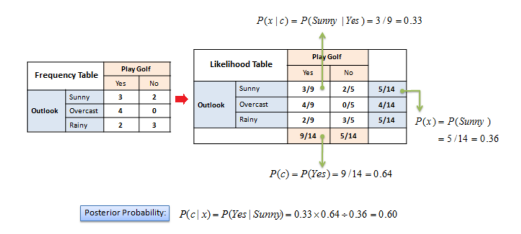
P(x|c) is the *likelihood* which is the probability of predictor given class.

P(x) is the prior probability of predictor (evidence or marginal likelihood).

Naive Bayes assumes that all features are conditionally independent from each other, in which case:

$$P(\mathbf{x}|c) = P(x_1|c) \times P(x_2|c) \times ... \times P(x_n|c)$$

Example. True or not? Players will play if weather is sunny.



nb example

Compute the posterior for every class and predict the class with highest probability

On numeric data

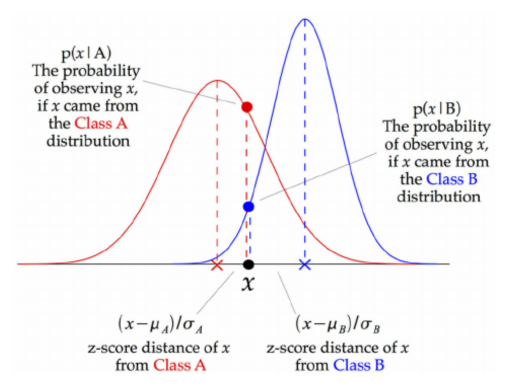
GaussianNB:

- Computes mean μ_c and standard deviation σ_c of the feature values per class
- It then fits a Gaussian distribution around the mean

$$p(x = v \mid c) = \frac{1}{\sqrt{2\pi\sigma_c^2}} e^{-\frac{(v - \mu_c)^2}{2\sigma_c^2}}$$

• Prediction are made using Bayes' theorem, by computing the joint probability given all features

$$p(c\mid \mathbf{x}) = \frac{p(c)\;p(\mathbf{x}|c)}{p(\mathbf{x})}$$
 Visualizing Naive Bayes

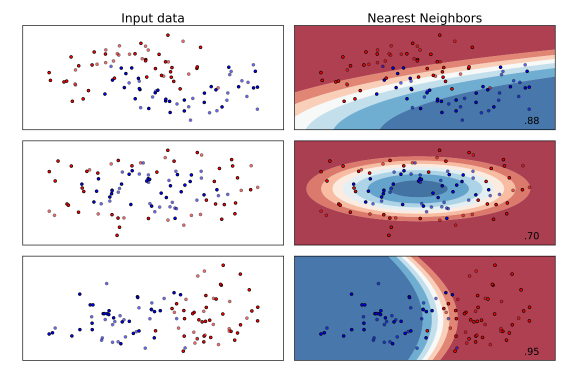


Naive Bayes image

```
[36]: from sklearn.naive_bayes import GaussianNB
    import plot_classifiers as pc

names = ["Nearest Neighbors"]
    classifiers = [GaussianNB()]

plt.rcParams.update({'font.size': 16})
    pc.plot_classifiers(names, classifiers, figuresize=(12,8))
```



Other Naive Bayes classifiers:

- BernoulliNB
 - Assumes binary data
 - Feature statistics: Number of non-zero entries per class
- MultinomialNB
 - Assumes count data
 - Feature statistics: Average value per class

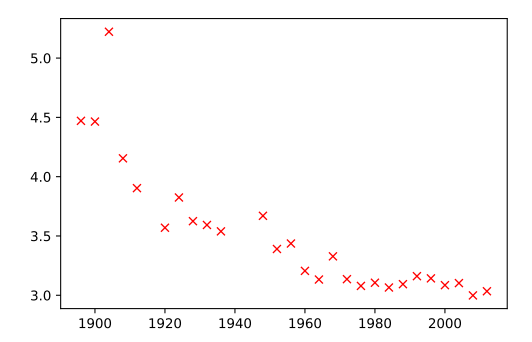
Mostly used for text classification (bag-of-words data)

Probabilistic interpretation of regression

Let's look at a the following regression problem

```
[107]: data = pods.datasets.olympic_marathon_men()
    x = data['X']
    y = data['Y']
    plt.plot(x, y, 'rx')

[<matplotlib.lines.Line2D at 0x11ac37cc0>]
```



Let's first try to fit a linear model

$$f(\mathbf{x}_i) = \mathbf{x}_i^\top \mathbf{w}$$

We can solve this via linear algebra by making a design matrix of the data, which includes the $x_0 = 1$ column, to represent the bias. Hence, each vector \mathbf{x}_i is given by appending a 1 onto the original vector

$$\mathbf{x}_i = \begin{bmatrix} 1 \\ x_i \end{bmatrix}$$

We can do this for the entire data set to form a design matrix X,

$$\mathbf{X} = \begin{bmatrix} \mathbf{x}_1^\top \\ \mathbf{x}_2^\top \\ \vdots \\ \mathbf{x}_N^\top \end{bmatrix} = \begin{bmatrix} 1 & x_1 \\ 1 & x_2 \\ \vdots & \vdots \\ 1 & x_N \end{bmatrix},$$

y = mx + c

which in numpy is done and solved with the following commands:

```
[11]: X = np.hstack((np.ones_like(x), x)) # [ones(size(x)) x]
    w = np.linalg.solve(np.dot(X.T, X), np.dot(X.T, y))
    print(w)

[[ 28.895]
    [ -0.013]]
```

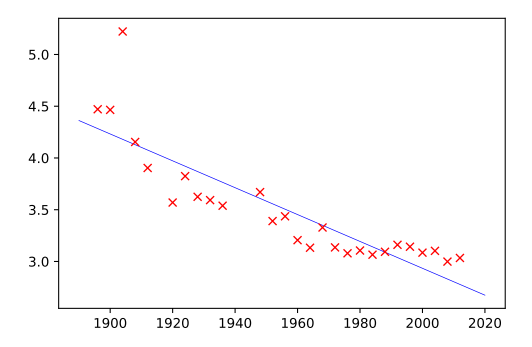
We can now fit the function

```
[14]: m = w[1]; c=w[0]
    x_test = np.linspace(1890, 2020, 130)[:, None]

    f_test = m*x_test + c

    plt.plot(x_test, f_test, 'b-')
    plt.plot(x, y, 'rx')
```

[<matplotlib.lines.Line2D at 0x111529b70>]



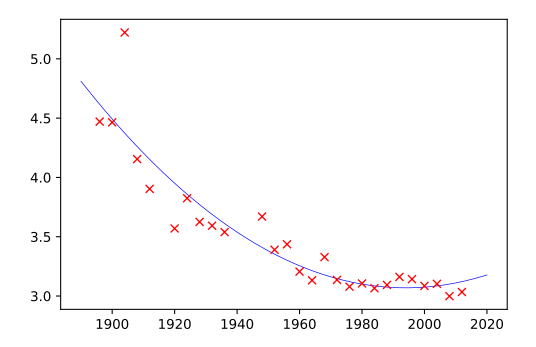
Now we will fit a quadratic model using more *basis functions*. Firstly, we need to create a new design matrix that contains the quadratic basis,

$$\mathbf{\Phi} = egin{bmatrix} \mathbf{1} & \mathbf{x} & \mathbf{x}^2 \end{bmatrix}$$

```
[23]: Phi = np.hstack([np.ones(x.shape), x, x**2])
    w = np.linalg.solve(np.dot(Phi.T, Phi), np.dot(Phi.T, y))
    print(w)
    f_test = w[2]*x_test**2 + w[1]*x_test + w[0]
    plt.plot(x_test, f_test, 'b-')
    plt.plot(x, y, 'rx')

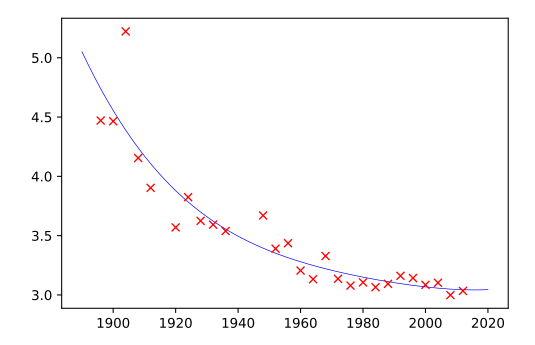
[[ 643.642]
    [ -0.643]
    [ 0. ]]
```

[<matplotlib.lines.Line2D at 0x11296f4a8>]



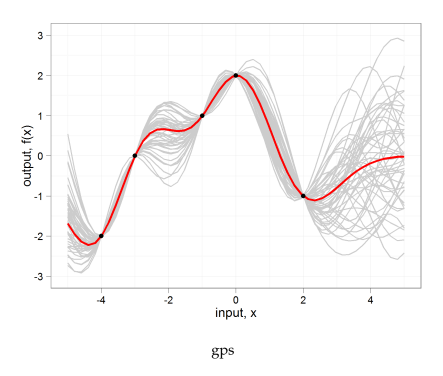
[33]: Phi = np.hstack([np.ones(x.shape), x, x**2, x**3, x**4, x**5, x**6])
 w = np.linalg.solve(np.dot(Phi.T, Phi), np.dot(Phi.T, y))
 f_test = w[6]*x_test**6 + w[5]*x_test**5 + w[4]*x_test**4 + w[3]*x_test**3
 plt.plot(x_test, f_test, 'b-')
 plt.plot(x, y, 'rx')

[<matplotlib.lines.Line2D at 0x116f4a470>]



Gaussian processes

Processes where the probabilities of possible base functions are learned/updated based on new data.



Probabilistic interpretation or regression

When there are more observations than unknowns (overdetermined systems), we cannot perfectly fit

$$y = mx + c$$

This issue can be solved by assuming that the data in inherently uncertain, and model it explictly by introducing a type of slack variable, ϵ_i , known as noise.

For each observation we now have the equation

$$y_i = mx_i + c + \epsilon_i$$
.

The slack variable represented the difference between our actual prediction and the true observation. This is also known as the *residual*.

We now have an additional n variables to estimate, one for each data point, $\{\epsilon_i\}$. With the original m and c we now have n+2 parameters to be estimated from n observations (underdetermined system).

We can however make assumptions about the noise distribution, i.e. that the slack variables are distributed according to a probability density. One often assumes Gaussian noise:

$$\epsilon_i \sim \mathcal{N}(0, \sigma^2),$$

with zero mean and variance σ^2 .

In the Bayesian approach, we also assume a *prior distribution* for the parameters, w:

$$\mathbf{w} \sim \mathcal{N}(\mathbf{0}, \alpha \mathbf{I})$$

I.e, each element of the parameters vector, w_i , was drawn from a Gaussian density with variance α

$$w_i \sim \mathcal{N}(0, \alpha)$$

Gaussian process model parameters:

- parameters of the prior
- parameters of the basis functions
- noise level

```
[104]: # set prior variance on w
    alpha = 4.
    # set the order of the polynomial basis set
    degree = 5
    # set the noise variance
    sigma2 = 0.01
```

Now we have the variance, we can sample from the prior distribution to see what form we are imposing on the functions *a priori*.

```
[108]: # Build the basis matrices (on Olypics data)
    def polynomial(x, degree, loc, scale):
        degrees = np.arange(degree+1)
        return ((x-loc)/scale)**degrees

scale = np.max(x) - np.min(x)
    loc = np.min(x) + 0.5*scale

num_data = x.shape[0]
    num_pred_data = 100 # how many points to use for plotting predictions
    x_pred = np.linspace(1880, 2030, num_pred_data)[:, None] # input location
    Phi_pred = polynomial(x_pred, degree=degree, loc=loc, scale=scale)
    Phi = polynomial(x, degree=degree, loc=loc, scale=scale)
```

Weight Space View

To generate typical functional predictions from the model, we need a set of model parameters. We assume that the parameters are drawn independently from a Gaussian density,

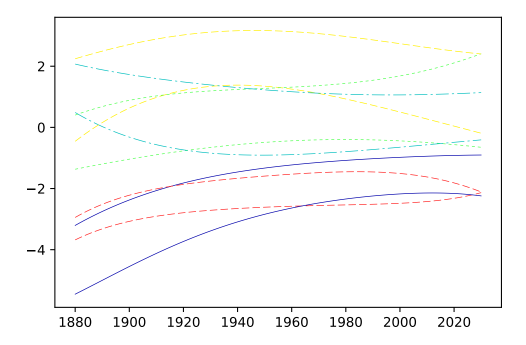
$$\mathbf{w} \sim \mathcal{N}(\mathbf{0}, \alpha \mathbf{I}),$$

then we can combine this with the definition of our prediction function $f(\mathbf{x})$,

$$f(\mathbf{x}) = \mathbf{w}^{\top} \boldsymbol{\phi}(\mathbf{x}).$$

We can now sample from the prior density to obtain a vector \mathbf{w} using the function np.random.normal and combine these parameters with our basis to create some samples of what $f(\mathbf{x})$ looks like,

```
[109]: num_samples = 10
   K = degree+1
   for i in range(num_samples):
        z_vec = np.random.normal(size=(K, 1))
        w_sample = z_vec*np.sqrt(alpha)
        f_sample = np.dot(Phi_pred,w_sample)
        plt.plot(x_pred, f_sample)
```



Function space view

We can use standard properties of multivariate Gaussians to write down the probability density that is implied over **f**.

We know that if \mathbf{w} is sampled from a multivariate Gaussian with covariance $\alpha \mathbf{I}$ and zero mean, then assuming that Φ is a deterministic matrix (i.e. it is not sampled from a probability density) then the vector \mathbf{f} will also be distributed according to a zero mean multivariate normal as follows,

$$\mathbf{f} \sim \mathcal{N}(\mathbf{0}, \alpha \mathbf{\Phi} \mathbf{\Phi}^{\top}).$$

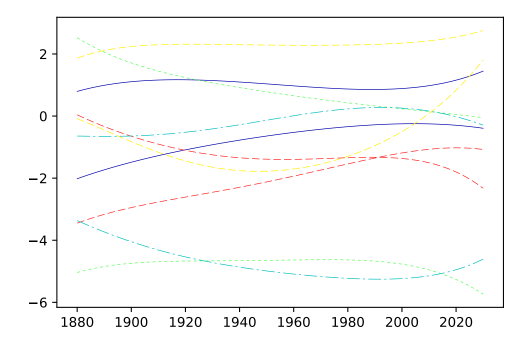
What happens if we sample f directly from this density, rather than first sampling w and then multiplying by Φ .

Let's try this. First of all we define the covariance (joined variability between 2 variables) as

$$\mathbf{K} = \alpha \mathbf{\Phi} \mathbf{\Phi}^{\mathsf{T}}.$$

```
[111]: K = alpha*np.dot(Phi_pred, Phi_pred.T)
```

We can use $np.random.multivariate_normal$ for sampling from a multivariate normal with covariance given by K and zero mean,

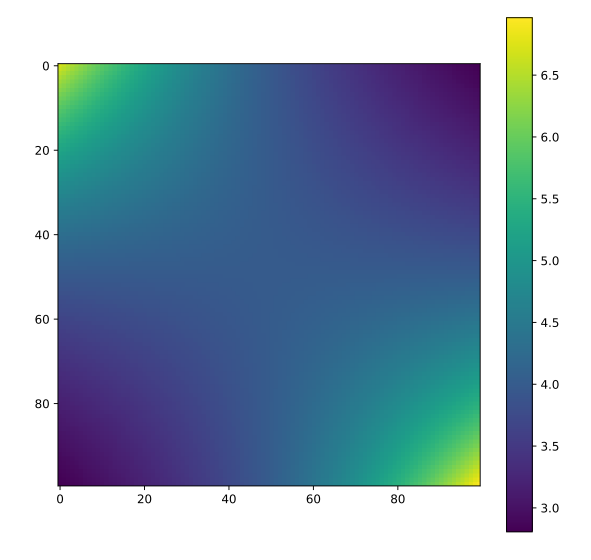


These look very similar! Indeed, they are effectively drawn from the same mutivariate normal density.

When sampling ${\bf f}$ directly we created the covariance for ${\bf f}$. We can visualise the covariance matrix:

```
[114]: fig, ax = plt.subplots(figsize=(8,8))
    im = ax.imshow(K, interpolation='none')
    fig.colorbar(im)

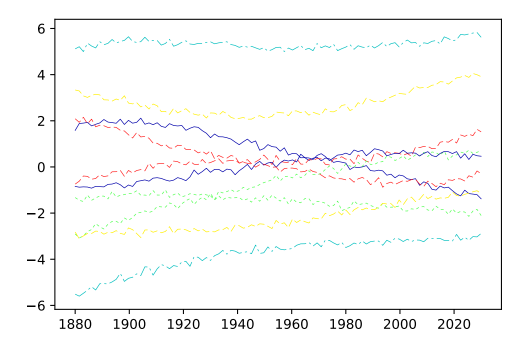
<matplotlib.colorbar.Colorbar at 0x117488908>
```



Noisy functions

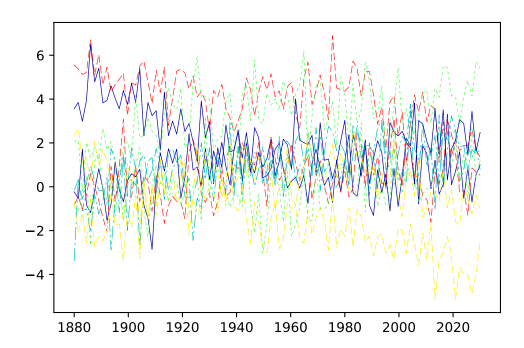
We normally add Gaussian noise to obtain our observations:

$$y = f + \epsilon$$



We can also increase the variance of the noise

```
[119]: sigma2 = 1.
    K = alpha*np.dot(Phi_pred, Phi_pred.T) + sigma2*np.eye(x_pred.size)
    for i in range(10):
        y_sample = np.random.multivariate_normal(mean=np.zeros(x_pred.size),
        plt.plot(x_pred.flatten(), y_sample.flatten())
```



Gaussian Process

In a Gaussian process you specify the *covariance function* directly, rather than *implicitly* through a basis matrix and a prior over parameters.

Gaussian processes have the advantage that they can be *nonparametric*: they can have *infinite* basis functions.

The RBF (Gaussian) covariance function is specified by

$$k(\mathbf{x}, \mathbf{x}') = \alpha \exp\left(-\frac{\|\mathbf{x} - \mathbf{x}'\|^2}{2\ell^2}\right).$$

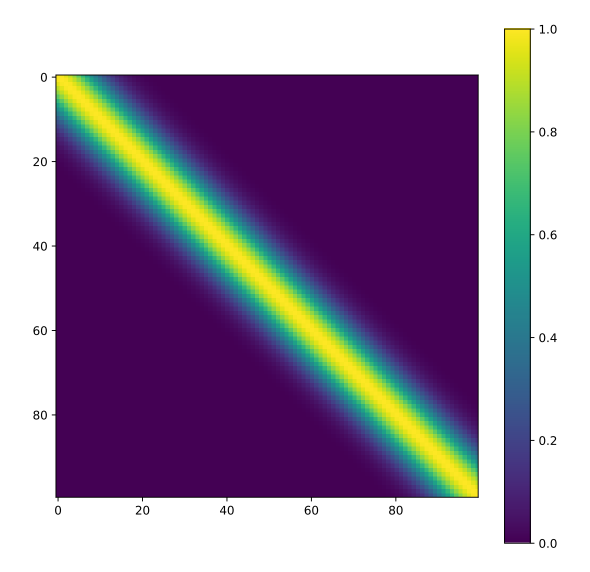
where $\|\mathbf{x} - \mathbf{x}'\|^2$ is the squared distance between the two input vectors

$$\|\mathbf{x} - \mathbf{x}'\|^2 = (\mathbf{x} - \mathbf{x}')^{\top} (\mathbf{x} - \mathbf{x}')$$

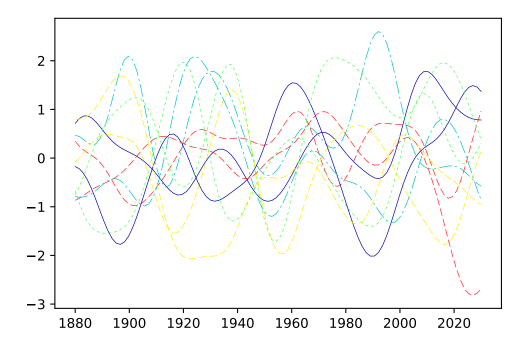
Let's build a covariance matrix based on this function.

<matplotlib.colorbar.Colorbar at 0x11bfabef0>

```
[121]: # Exponentiated quadratic is another name for RBF
      def exponentiated_quadratic(x, x_prime, variance, lengthscale):
           squared\_distance = ((x-x\_prime) **2).sum()
           return variance*np.exp((-0.5*squared_distance)/lengthscale**2)
       # Compute covariances directly
      def compute_kernel(X, X2, kernel, **kwargs):
          K = np.zeros((X.shape[0], X2.shape[0]))
           for i in np.arange(X.shape[0]):
               for j in np.arange(X2.shape[0]):
                   K[i, j] = kernel(X[i, :], X2[j, :], **kwargs)
           return K
       # Visualize
      K = compute_kernel(x_pred, x_pred, exponentiated_quadratic, variance=1.,
      fig, ax = plt.subplots(figsize=(8,8))
      im = ax.imshow(K, interpolation='none')
      fig.colorbar(im)
```



Finally, we can sample functions with this kernel (covariance matrix)



Gaussian process optimization

The Gaussian process perspective takes the marginal likelihood of the data to be a joint Gaussian density with a covariance given by \mathbf{K} .

The model likelihood is of the form,

$$p(\mathbf{y}|\mathbf{X}) = \frac{1}{(2\pi)^{\frac{n}{2}}|\mathbf{K}|^{\frac{1}{2}}} \exp\left(-\frac{1}{2}\mathbf{y}^{\top} \left(\mathbf{K} + \sigma^{2}\mathbf{I}\right)^{-1}\mathbf{y}\right)$$

where the input data, \mathbf{X} , influences the density through the covariance matrix, \mathbf{K} whose elements are computed through the covariance function, $k(\mathbf{x}, \mathbf{x}')$.

Hence, the negative log likelihood (the objective function) is given by,

$$E(\boldsymbol{\theta}) = \frac{1}{2} \log |\mathbf{K}| + \frac{1}{2} \mathbf{y}^{\top} \left(\mathbf{K} + \sigma^2 \mathbf{I} \right)^{-1} \mathbf{y}$$

where the *parameters* of the model are also embedded in the covariance function, they include the parameters of the kernel (such as lengthscale and variance), and the noise variance, σ^2 .

```
[127]: class GP():
    def __init__(self, X, y, sigma2, kernel, **kwargs):
        self.K = compute_kernel(X, X, kernel, **kwargs)
        self.X = X
        self.y = y
        self.sigma2 = sigma2
        self.kernel = kernel
        self.kernel_args = kwargs
        self.update_inverse()

def update_inverse(self):

# Preompute the inverse covariance and some quantities of interes
```

```
# the log determinant of the covariance matrix.
self.logdetK = np.linalg.det(self.K+self.sigma2*np.eye(self.K.sha
# The matrix inner product of the inverse covariance
self.Kinvy = np.dot(self.Kinv, self.y)
self.yKinvy = (self.y*self.Kinvy).sum()

def log_likelihood(self):
    # use the pre-computes to return the likelihood
    return -0.5*(self.K.shape[0]*np.log(2*np.pi) + self.logdetK + sel

def objective(self):
    # use the pre-computes to return the objective function
    return -self.log_likelihood()
```

NOTE: This is not the correct *numerical* way to compute this!
self.Kinv = np.linalg.inv(self.K+self.sigma2*np.eye(self.K.shape[

Making predictions

The model makes predictions for f that are unaffected by future values of f^* .

If we think of f^* as test points, we can still write down a joint probability density over the training observations, f and the test observations, f^* .

This joint probability density will be Gaussian, with a covariance matrix given by our covariance function, $k(\mathbf{x}_i, \mathbf{x}_j)$.

$$\begin{bmatrix} \mathbf{f} \\ \mathbf{f}^* \end{bmatrix} \sim \mathcal{N} \left(\mathbf{0}, \begin{bmatrix} \mathbf{K} & \mathbf{K}_* \\ \mathbf{K}_*^\top & \mathbf{K}_{*,*} \end{bmatrix} \right)$$

where K is the covariance computed between all the training points,

 \mathbf{K}_* is the covariance matrix computed between the training points and the test points, $\mathbf{K}_{*,*}$ is the covariance matrix computed between all the tests points and themselves.

Conditional Density

Just as in naive Bayes, we defined the joint density (although there it was over both the labels and the inputs, p(y, X) and now we need to define *conditional* distributions that answer particular questions of interest.

We will need the conditional density for making predictions.

$$\mathbf{f}^* | \mathbf{y} \sim \mathcal{N}(\boldsymbol{\mu}_f, \mathbf{C}_f)$$

with a mean given by

$$\boldsymbol{\mu}_f = \mathbf{K}_*^{\top} \left[\mathbf{K} + \sigma^2 \mathbf{I} \right]^{-1} \mathbf{y}$$

and a covariance given by

$$\mathbf{C}_f = \mathbf{K}_{*,*} - \mathbf{K}_*^{\top} \left[\mathbf{K} + \sigma^2 \mathbf{I} \right]^{-1} \mathbf{K}_*.$$

Let's compute what those posterior predictions are for the olympic marathon data.

```
[130]: # set covariance function parameters
    variance = 16.0
    lengthscale = 32
    # set noise variance
    sigma2 = 0.05
```

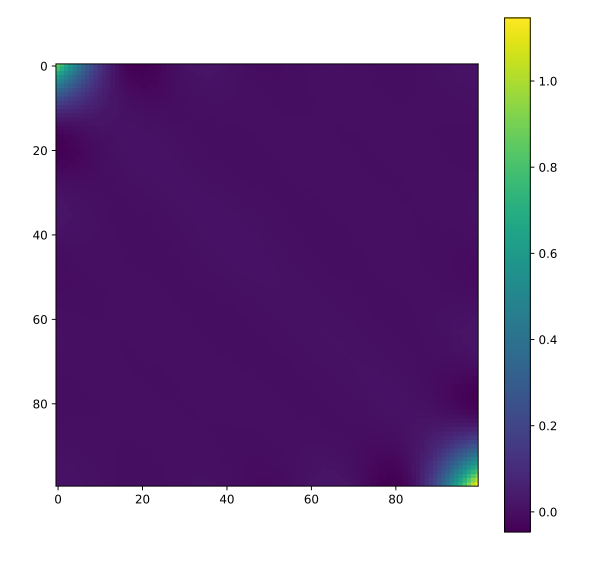
```
def posterior_f(self, X_test):
    K_star = compute_kernel(self.X, X_test, self.kernel, **self.kernel_ar
    K_starstar = compute_kernel(X_test, X_test, self.kernel, **self.kernel
    A = np.dot(self.Kinv, K_star)
    mu_f = np.dot(A.T, y)
    C_f = K_starstar - np.dot(A.T, K_star)
    return mu_f, C_f

# attach the new method to class GP():
GP.posterior_f = posterior_f

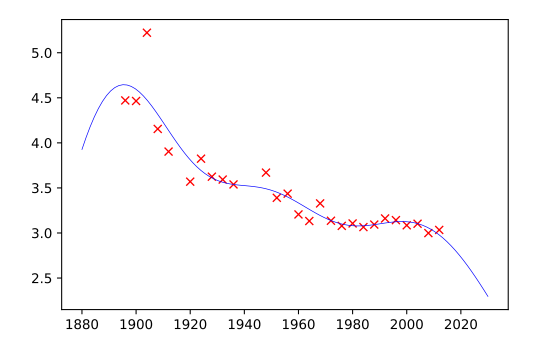
model = GP(x, y, sigma2, exponentiated_quadratic, variance=variance, leng
mu_f, C_f = model.posterior_f(x_pred)

fig, ax = plt.subplots(figsize=(8,8))
im = ax.imshow(C_f, interpolation='none')
fig.colorbar(im)
```

<matplotlib.colorbar.Colorbar at 0x117e81898>



We can now plot the mean

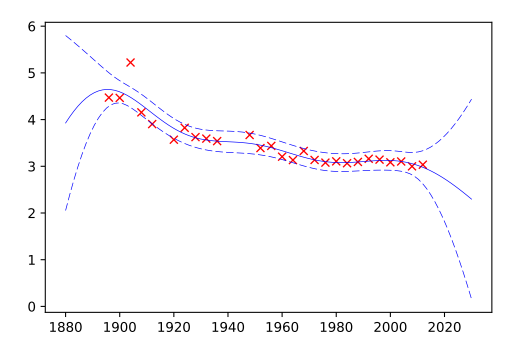


as well as the associated error bars

```
[132]: var_f = np.diag(C_f)[:, None]
    std_f = np.sqrt(var_f)

    plt.plot(x, y, 'rx')
    plt.plot(x_pred, mu_f, 'b-')
    plt.plot(x_pred, mu_f+2*std_f, 'b--')
    plt.plot(x_pred, mu_f-2*std_f, 'b--')

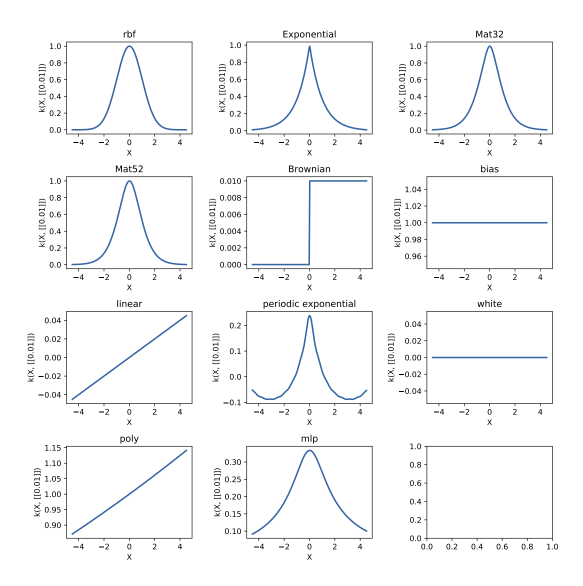
[<matplotlib.lines.Line2D at 0x11aa42550>]
```



Gaussian Processes with GPy

- GPyRegression
- Generate a kernel first
 - State the dimensionality of your input data
 - Variance and lengthscale are optional, default = 1
 kernel = GPy.kern.RBF(input_dim=1, variance=1., lengthscale=1.)
 - Other kernels:

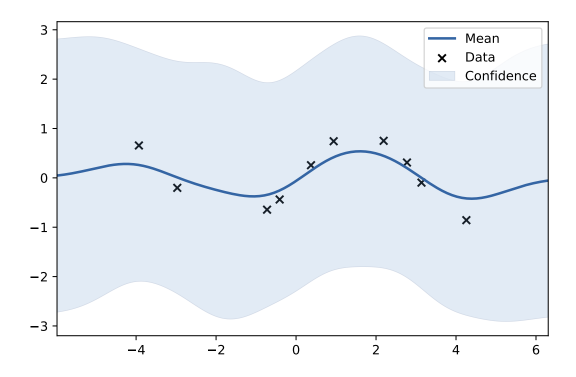
 GPy.kern.BasisFuncKernel?
- Build model:



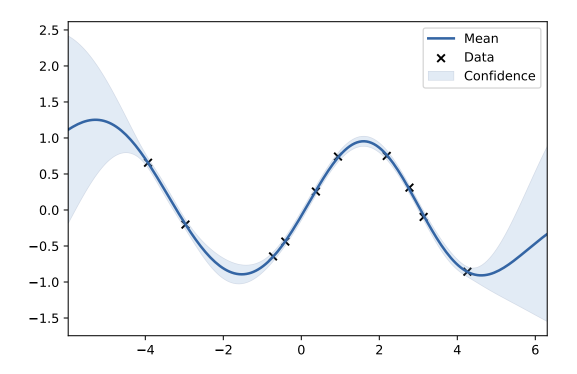
Matern is a generalized RBF kernel that can scale between RBF and Exponential Build the untrained GP. The shaded region corresponds to ~95% confidence intervals (i.e. +/- 2 standard deviation)

```
[72]: # Generate noisy sine data
   X = np.random.uniform(-5.,5.,(10,1))
   Y = np.sin(X) + np.random.randn(10,1)*0.05

# Build untrained model
   kernel = GPy.kern.RBF(input_dim=1, variance=1., lengthscale=1.)
   m = GPy.models.GPRegression(X,Y,kernel)
   fig = m.plot()
```

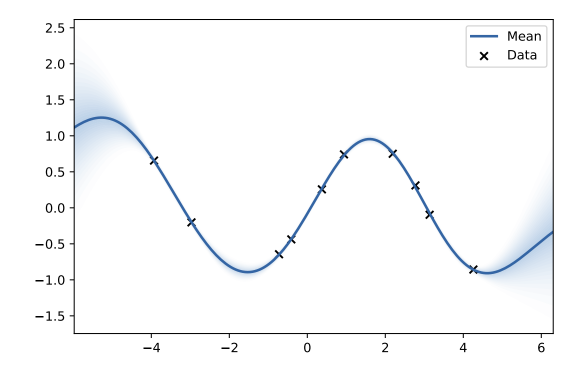


Train the model (optimize the parameters): maximize the likelihood of the data. Best to optimize with a few restarts: the optimizer may converges to the high-noise solution. The optimizer is then restarted with a few random initialization of the parameter values.



You can also plot densities

[74]: fig = m.plot(plot_density=True)



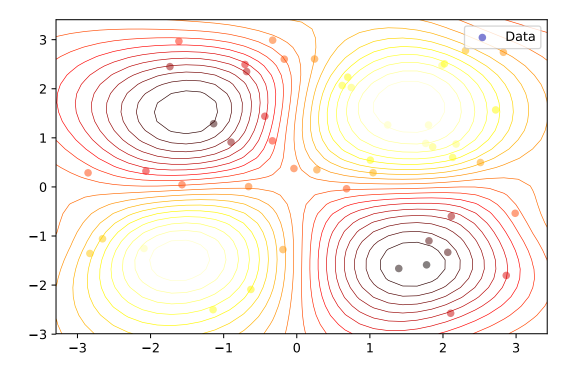
You can also show results in 2D

```
[80]: # sample inputs and outputs
X = np.random.uniform(-3.,3.,(50,2))
Y = np.sin(X[:,0:1]) * np.sin(X[:,1:2])+np.random.randn(50,1)*0.05

# define kernel
ker = GPy.kern.Matern52(2,ARD=True) + GPy.kern.White(2)

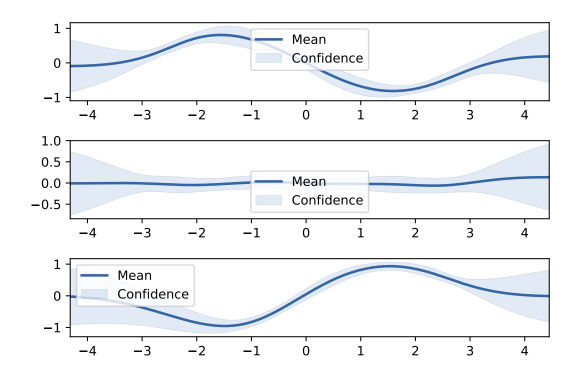
# create simple GP model
m = GPy.models.GPRegression(X,Y,ker)

# optimize and plot
m.optimize(max_f_eval = 1000)
fig = m.plot()
```



We can plot 2D slices using the fixed_inputs argument to the plot function. fixed_inputs is a list of tuples containing which of the inputs to fix, and to which value.

```
[89]: slices = [-1, 0, 1.5]
    figure = GPy.plotting.plotting_library().figure(3, 1)
    for i, y in zip(range(3), slices):
        canvas = m.plot(figure=figure, fixed_inputs=[(1,y)], row=(i+1), plot_d
```



For vertical slices, simply fix the other input: fixed_inputs=[(0,y)]

1

0

-1 -

Mean

Confidence

<u>-</u>2

```
[92]: slices = [-1, 0, 1.5]
      figure = GPy.plotting.plotting_library().figure(3, 1)
      for i, y in zip(range(3), slices):
          canvas = m.plot(figure=figure, fixed_inputs=[(0,y)], row=(i+1), plot_d
      1
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                                                           3
                                                                  4
```

ò

1

-1

2

3

4

Gaussian Processes with scikit-learn

- GaussianProcessRegressor
- Hyperparameters:
 - kernel: kernel specifying the covariance function of the GP
 - * Default: "1.0 * RBF(1.0)"
 - * Typically leave at default. Will be optimized during fitting
 - alpha: regularization parameter
 - * Tikhonov regularization of the assumed covariance between the training points.
 - * Adds a (small) value to the diagonal of the kernel matrix during fitting.
 - * Larger values:
 - · correspond to increased noise level in the observations
 - · also reduce potential numerical issues during fitting
 - * Default: 1e-10
 - n_restarts_optimizer: number of restarts of the optimizer
 - * Default: 0. Best to do at least a few iterations.
 - * Optimizer finds the kernel's parameters which maximize the log-marginal likelihood
- Retrieve predictions and confidence interval after fitting:

```
y_pred, sigma = gp.predict(x, return_std=True)
```

Example

gp.fit(X, y)

```
[36]: from sklearn.gaussian_process import GaussianProcessRegressor
    from sklearn.gaussian_process.kernels import RBF, ConstantKernel as C

def f(x):
        """The function to predict."""
        return x * np.sin(x)

X = np.atleast_2d([1., 3., 5., 6., 7., 8.]).T

# Observations
y = f(X).ravel()

# Mesh the input space for evaluations of the real function, the prediction
# its MSE
x = np.atleast_2d(np.linspace(0, 10, 1000)).T

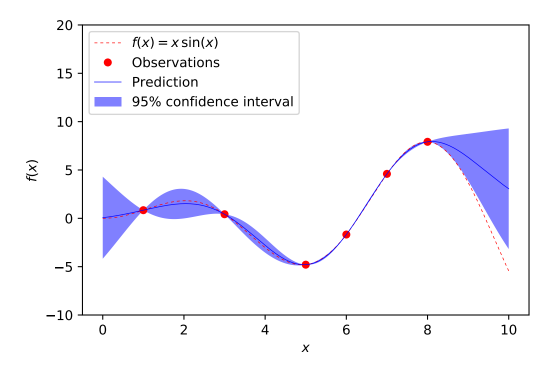
# Instanciate a Gaussian Process model
kernel = C(1.0, (1e-3, 1e3)) * RBF(10, (1e-2, 1e2))
gp = GaussianProcessRegressor(kernel=kernel, n_restarts_optimizer=9)

# Fit to data using Maximum Likelihood Estimation of the parameters
```

Make the prediction on the meshed x-axis (ask for MSE as well)

<matplotlib.legend.Legend at 0x1169aaf98>

plt.legend(loc='upper left')



Example with noisy data

```
[39]: X = np.linspace(0.1, 9.9, 20)
    X = np.atleast_2d(X).T

# Observations and noise
    y = f(X).ravel()
    dy = 0.5 + 1.0 * np.random.random(y.shape)
    noise = np.random.normal(0, dy)
    y += noise
```

```
# Instanciate a Gaussian Process model
gp = GaussianProcessRegressor(kernel=kernel, alpha=(dy / y) ** 2,
                                n_restarts_optimizer=10)
# Fit to data using Maximum Likelihood Estimation of the parameters
gp.fit(X, y)
# Make the prediction on the meshed x-axis (ask for MSE as well)
y_pred, sigma = gp.predict(x, return_std=True)
# Plot the function, the prediction and the 95% confidence interval based
# the MSE
fig = plt.figure()
plt.plot(x, f(x), 'r:', label=u'\$f(x) = x\,\\\sin(x)\$')
plt.errorbar(X.ravel(), y, dy, fmt='r.', markersize=10, label=u'Observation
plt.plot(x, y_pred, 'b-', label=u'Prediction')
plt.fill(np.concatenate([x, x[::-1]]),
          np.concatenate([y_pred - 1.9600 * sigma,
                          (y_pred + 1.9600 * sigma)[::-1]]),
          alpha=.5, fc='b', ec='None', label='95% confidence interval')
plt.xlabel('$x$')
plt.ylabel('$f(x)$')
plt.ylim(-10, 20)
plt.legend(loc='upper left')
plt.show()
   20
           - f(x) = x \sin(x)
             Prediction
   15
             95% confidence interval
             Observations
   10
(x)
    5
    0
   -5
  -10
                  2
         0
                            4
                                      6
                                                8
                                                         10
```

Х

Gaussian processes

The advantages of Gaussian processes are: - The prediction interpolates the observations (at least for regular kernels). - The prediction is probabilistic (Gaussian) so that one can compute empirical confidence intervals. - Versatile: different kernels can be specified.

The disadvantages of Gaussian processes include: - They are not sparse, i.e., they use the whole samples/features information to perform the prediction. - They lose efficiency in high dimensional spaces – namely when the number of features exceeds a few dozens.