

Filling Values Using a Model

If we want to fill missing values better, we need to predict them

We need a model, which can infer their value

- All the approaches seen so far can be considered (extremely simple) models
- ...We just need a more advanced one!

What are the desired properties of the model we seek?

Given a gap (i.e. one or more contiguous missing values), the model:

- Must be able to make a prediction about the missing values
- ...Which is consistent with all the available observations
- I.e. it should be able to interpolate the data (in generalized sense)

Most ML models cannot be used for filling (in a straightforward fashion)

Filling Values Using a Model

Density estimation does not (natively) provide predictions

To be fair, predictions can be extracted from a density estimator:

■ Given an estimator $f(\mathbf{x}, \theta)$ for $P(\mathbf{x})$ we can find the most likely value for \mathbf{x} by solving:

$$\operatorname{argmax}_{\mathbf{x}} f(\mathbf{x}, \theta)$$

- This is a Maximum A Posteriori (MAP)
- ...And its what most regressors/classifiers natively compute

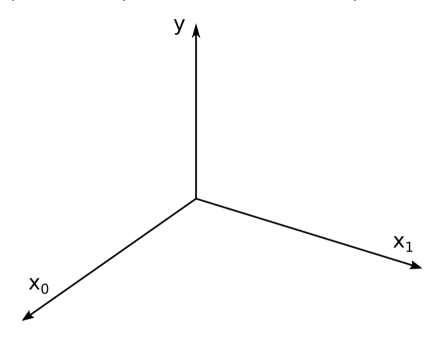
However, with a density estimator, computing the MAP can be very expensive

Auto-regressors makes use of past observations, but not the future ones

- They are designed for extrapolation (predict beyond the boundaries)
- ...And not for interpolation

One of the few viable ML models is given by Gaussian Processes (GP)

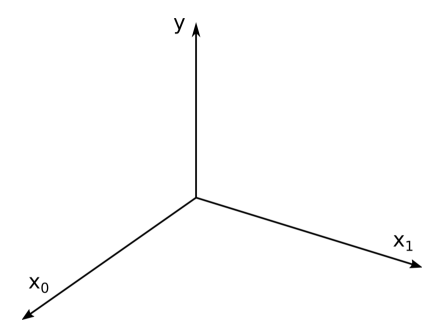
We will introduce their key concepts via an example:



- Say we want to model how rainfall changes over a stretch of land
- $y = \text{rainfall}, (x_0, x_1) = \text{position on the surface of land}$

One of the few viable ML models is given by Gaussian Processes (GP)

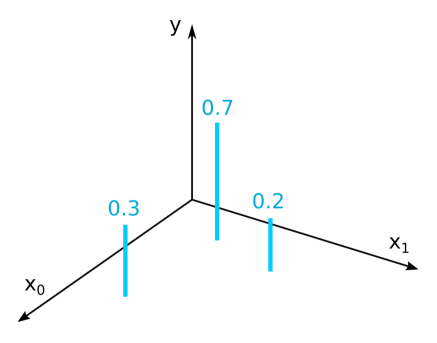
Since this is a physical phenomenon...



- lacktriangleright ...We can reasonably assume that $oldsymbol{y}$ is Normally distributed
- But unless we know more, we can say nothing else

One of the few viable ML models is given by Gaussian Processes (GP)

However, if we have a few measurements...

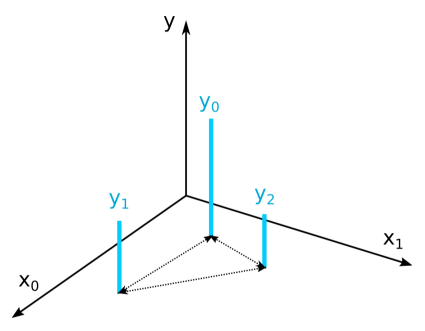


We can assume that rainfall in nearby locations is similar

- We can rely on this to estimate the chance of the measurements themselves
- ...And of rainfall at positions for which we lack measurements

One of the few viable ML models is given by Gaussian Processes (GP)

We view the measurements as components of a variable $y_X = (y_0, y_1, y_2)$



- \mathbf{y}_X will follow a multivariate Normal distribution
- ...And the covariance will depend on the distance between measurements

Formally:

A GP is a stochastic process, i.e. a collection of indexed random variables

- lacktriangle Each variable y_x is indexed via a tuple x (e.g. location, time...)
- The index is continuous and the collection infinite
- lacktriangle Every finite subset of y_x variables follows a Multivariate Normal Distribution Some examples:
- \mathbf{y}_{x} could be the rainfall rate at location \mathbf{x}
- \mathbf{y}_{x} could be the twitter volume at time x
- \mathbf{y}_{x} could be the traffic volume at time x

In general y_x is the value of a (stochastic) function for input x

Multivariate Normal Distritbuion

The multivariate normal distribution?

- It works for many real world phenomena
- It has a <u>closed-form density function</u> that is (relatively) easy to compute
- lacksquare The PDF is defined via a (vector) mean μ and a covariance matrix Σ
- lacksquare ...And often we can assume $\mu=0$, so knowing Σ is enough

So, given a set of indexes/input values of interest X

Then, if we manage to know Σ , we can compute:

- lacksquare The probability density $f(\hat{\mathbf{y}}_{\mathbf{X}})$ of some given observations
 - I.e. the probability of a dataset
- lacksquare The conditional density $f(\hat{y}_x \mid \hat{\mathbf{y}}_{\mathbf{X}})$ of a new observation
 - lacksquare I.e. a prediction w.r.t. a set of know observations \hat{y}_X

Defining the Covariance Matrix

How do we define Σ ?

We assume that the covariance depends on x (and not on the y)

- lacksquare Given two variables y_{x_i} and y_{x_j} , their covariance is given by $K(x_i,x_j)$
- \blacksquare Where K is called a kernel function (and is user chosen)

Given any finite set of variables $\{y_{x_1}, \dots y_{x_n}\}$, the covariance matrix is:

$$\Sigma = \begin{pmatrix} K(x_1, x_1) & K(x_1, x_2) & \cdots & K(x_1, x_n) \\ K(x_2, x_1) & K(x_2, x_2) & \cdots & K(x_2, x_n) \\ \vdots & \vdots & \vdots & \vdots \\ K(x_n, x_1) & K(x_n, x_2) & \cdots & K(x_n, x_n) \end{pmatrix}$$

■ I.e. it's entirely specified via the kernel

Unfortunately, choosing the kernel completely by hand would still be too difficult

Fitting a Gaussian Process

In practice, to define the kernel we:

- lacksquare Pick a parameterized kernel function $K_{ heta}(x_i,x_j)$, where heta = parameter vector
- lacksquare Collect training observations $\hat{oldsymbol{y}}_{oldsymbol{X}}$

Then we choose θ so as to maximize the likelihood of the training data, i.e.:

$$\operatorname{argmax}_{\theta} f(\mathbf{\hat{y}_X}, \theta)$$

■ Where $f(\hat{\mathbf{y}}_{\mathbf{X}}, \theta)$ is the estimated probability density of the observations

The training problem

- Is a (possibly challenging) numerical optimization problem
- ...Which is typically solved to local optimality (e.g. via gradient descent)

Let's see how to use Gaussian Processes in scikit-learn

First, let us choose a target function:

```
In [2]: f = lambda x: x * np.sin(2*np.pi*x) + x # target function
x = np.linspace(0, 3, 1000)
y = pd.Series(index=x, data=f(x))
nab.plot_gp(target=y, figsize=figsize)
```

A Jupyter widget could not be displayed because the widget state could not be found. This could happen if the kernel storing the widget is no longer available, or if the widget state was not saved in the notebook. You may be able to create the widget by running the appropriate cells.

Let's see how to use Gaussian Processes in scikit-learn

Then we build a small training set:

```
In [3]: np.random.seed(42)
    n_tr = 15
    x_tr = np.linspace(0.2, 2.8, n_tr) + 0.2*np.random.rand(n_tr)
    x_tr.sort()
    y_tr = pd.Series(index=x_tr, data=f(x_tr) + 0.2*np.random.rand(n_tr))
    nab.plot_gp(target=y, samples=y_tr, figsize=figsize)
```

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Let's see how to use Gaussian Processes in scikit-learn

Then we need to choose a kernel

- There are <u>many available options</u>
- We will start with a simple Radial Basis Function (i.e. Gaussian) kernel

$$K(x_i, x_j) = e^{-\frac{d(x_i, x_j)^2}{2l}}$$

The correlation decreases with the (Euclidean) distance $d(x_i, x_j)$:

- Intuitively, the closer the points, the higher the correlation
- lacktriangle The l parameter (scale) control the rate of the reduction

We have 15 indexes, so the kernel will define a 15×15 covariance matrix

Here's how to use an RBF kernel in scikit-learn

```
In [4]: from sklearn.gaussian_process.kernels import RBF
kernel = RBF(1, (1e-3, 1e3))
```

The RBF kernel has a single parameter, representing its scale

The extra (tuple) parameter represents a pair of bounds

- During training only parameter values within the boundaries will be considered
- Bounds can be very useful for controlling the training process
- ...Based on the available domain information

Now we can train a Gaussian Process

```
In [5]: from sklearn.gaussian_process import GaussianProcessRegressor
    gp = GaussianProcessRegressor(kernel=kernel, n_restarts_optimizer=9)
    gp.fit(y_tr.index.values.reshape(-1,1), y_tr.values) # needs 2D input
    gp.kernel_
Out[5]: RBF(length_scale=0.229)
```

- Training uses Gradient Descent...
- ...To maximize the likelihood (density) of the training data
- Restarts are needed to mitigate issues due to local optima

And then we can obtain the predictions:

```
In [6]: xp, std = gp.predict(x.reshape(-1,1), return_std=True)
xp = pd.Series(index=y.index, data=xp)
std = pd.Series(index=y.index, data=std)
```

We can now plot the predictions

```
In [9]: nab.plot_gp(target=y, samples=y_tr, pred=xp, std=std, figsize=figsize)
```

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- We get both a point estimate (the blue line)
- ...And a confidence interval (the light blue areas)

But how did we manage that?

There are two tricks at work here:

1) The training examples $\hat{\mathbf{y}}_{\mathbf{X}}$ are part of the GP parameters

- This is similar to what we have (e.g.) in KDE
- ...And quite different from other ML methods (e.g. NN, DTs, SVMs...)

Given an new input value x, the GP output corresponds to:

$$f\left(y_{x}\mid\hat{\mathbf{y}}_{\mathbf{X}}\right)$$

- I.e. the probability of y_x conditioned on the known examples
- 2) For computing Σ , we only need to know the input values (i.e. X and x)
- lacktriangle This is by construction: all our kernels $K(x_i,x_j)$ are built this way
- lacksquare Which means that we can compute $f\left(y_x \mid \mathbf{\hat{y}_X}\right)$ via a closed-form expression

Thanks to this, for $f\left(y_{x}\mid\mathbf{\hat{y}_{X}}\right)$ we can get

The conditional mean (which is also the MAP):

$$\arg\max_{y_x} f\left(y_x \mid \mathbf{\hat{y}_X}\right)$$

■ This will be our "prediction"

The conditional standard deviation

$$\sqrt{\operatorname{Var}\left[f\left(y_{x}\mid\hat{\mathbf{y}}_{\mathbf{X}}\right)\right]}$$

■ This is used to define the confidence intervals

In practice, we GP output is a probability distribution

A Numeric Example

As an example, say we want a prediction for x=2.5, i.e. $y_{2.5}$

...And let's assume that our training set contains only one example

- We will consider separately the tenth and first example in our dataset
- lacksquare We have: $(\hat{x}_9,\hat{y}_{\hat{x}_9})\simeq (2.01,2.27)$ and $(\hat{x}_0,\hat{y}_{\hat{x}_0})\simeq (0.27,0.58)$

The covariance matrix in the two cases is therefore:

$$\Sigma_{y_x, \hat{y}_{\hat{x}_9}} = \begin{pmatrix} K(2.01, 2.01) & K(2.01, 2.5) \\ K(2.5, 2.01) & K(2.5, 2.5) \end{pmatrix}$$

$$\Sigma_{y_x, \hat{y}_{\hat{x}_0}} = \begin{pmatrix} K(0.27, 0.27) & K(0.27, 2.5) \\ K(2.5, 0.27) & K(2.5, 2.5) \end{pmatrix}$$

■ Each matrix defines a multivariate Normal distribution

Let's actually build the matrices in Python

- Note: scikit-learn kernels are not designed to be used on individual points
- So, for this we will rely on basic numpy methods

We start with \hat{x}_9 and x, which are close to each other

Then we do the same for \hat{x}_0 and x, which are far apart

\hat{x}_9 and x are close to each other, so $\hat{y}_{\hat{x}_9}$ and y_x are strongly correlated

```
In [12]: yr = np.linspace(-5, 5, 100)
  nab.plot_distribution_2D(f_9x, yr, yr, figsize=figsize)
  plt.xlabel('y_{x_9}'); plt.ylabel('y_{x}'); plt.tight_layout()
```

A Jupyter widget could not be displayed because the widget state could not be found. This could happen if the kernel storing the widget is no longer available, or if the widget state was not saved in the notebook. You may be able to create the widget by running the appropriate cells.

lacksquare Still, if we know neither $\hat{y}_{\hat{x}_0}$ nor y_x , we can only say that they are likely both

zero

But we do know $\hat{y}_{\hat{y}_9}$! So, we can use this information_

```
In [13]: nab.plot_distribution_2D(f_9x, yr, yr, figsize=figsize)
  plt.axvline(10*(y_tr[x_tr[9]] + 5), color='tab:orange');
  plt.xlabel('y_{x_9}'); plt.ylabel('y_{x}'); plt.tight_layout()
```

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lacktriangle Given the observation, the most likely value for y_x changes considerably

\hat{x}_0 and x are far apart, so $\hat{y}_{\hat{x}_0}$ and y_x are loosely correlated

```
In [14]: nab.plot_distribution_2D(f_0x, yr, yr, figsize=figsize)
   plt.axvline(10*(y_tr[x_tr[0]] + 5), color='tab:orange');
   plt.xlabel('y_{x_0}'); plt.ylabel('y_{x}'); plt.tight_layout()
```

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lacksquare Knowing $\hat{y}_{\hat{x}_0}$ is not going to be of much help here

So, a few key insight to keep in mind:

- Superficially, GPs behave like functions that output probability distribution
- Internally, this is enabled by two components:
 - The kernel, defining how all the points are correlated
 - A set of observations, use to obtain conditional distributions

In scikit-learn:

When we call the fit method:

- The optimizer adjusts the kernel parameters
- lacksquare ...And the observations $\hat{y}_{\hat{x}}$ are stored

When we call the predict method:

- The covariance matrix is built
- The model computes the conditional distributions

Now, let's try to improve our model

We need to choose a kernel appropriate to our dataset

```
In [15]: nab.plot_gp(target=y, samples=y_tr, figsize=figsize)
```

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■ We a bit of noise, a period, and a trend

So, let us deal with the noise first

```
In [16]: from sklearn.gaussian_process.kernels import WhiteKernel
    kernel = WhiteKernel(0.1, (1e-3, 1e3))
    kernel += RBF(1, (1e-2, 1e2))
```

Whitekernel captures the presence of noise in the data

$$K(x_i, x_j) = \sigma^2$$
 iff $x_i = x_j$, 0 otherwise

- lacktriangle The only parameter of <code>whitekernel</code> represents the noise level $oldsymbol{\sigma}^2$
- A small noise level prevent overfitting the data
- ...But too much noise leads to useless predictions!

It's often a good idea to have magnitude parameters in the kernel

```
In [17]: from sklearn.gaussian_process.kernels import ConstantKernel
    kernel = WhiteKernel(0.1, (1e-2, 1e2))
    kernel += ConstantKernel(1, (1e-2, 1e2)) * RBF(1, (1e-2, 1e2))
```

constantkernel is a constant factor (in this case a relative weight)

■ ...And allows the optimizer to tune the magnitude of the RBF kernel

Let's repeat training again:

```
In [18]: gp = GaussianProcessRegressor(kernel=kernel, n_restarts_optimizer=9)
    gp.fit(y_tr.index.values.reshape(-1,1), y_tr.values) # needs 2D input
    print(gp.kernel_)

WhiteKernel(noise_level=0.01) + 2.21**2 * RBF(length_scale=0.321)

/usr/local/lib/python3.9/site-packages/sklearn/gaussian_process/kernels.py:402: ConvergenceWar
    ning: The optimal value found for dimension 0 of parameter kl_noise_level is close to the spe
    cified lower bound 0.01. Decreasing the bound and calling fit again may find a better value.
    warnings.warn("The optimal value found for "
```

Let us see the new predictions

```
In [19]: xp, std = gp.predict(x.reshape(-1,1), return_std=True)
xp = pd.Series(index=y.index, data=xp)
std = pd.Series(index=y.index, data=std)
nab.plot_gp(target=y, samples=y_tr, pred=xp, std=std, figsize=figsize)
```

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Better, but we are still not exploiting the period and the trend

So, let us take them into account, starting with the period

```
In [20]: from sklearn.gaussian_process.kernels import ExpSineSquared
   kernel = WhiteKernel(0.1, (1e-2, 1e2))
   kernel += ConstantKernel(1, (1e-2, 1e2)) * RBF(1, (1e-2, 1e2))
   kernel += ExpSineSquared(1, 1, (1e-2, 1e2), (1e-2, 1e2))
```

ExpSineSquared captures the period:

$$K(x_i, x_j) = e^{-2\frac{\sin^2\left(\pi \frac{d(x_i, x_j)}{p}\right)}{l^2}}$$

- lacksquare The correlation grows is the distance is close to a multiple of the period p
- lacktriangle The scale parameter l control the rate of decrease/increase
- lacksquare In the implementation, the first parameter is l and the second p

Now, let's try to capture the trend

```
In [21]: from sklearn.gaussian_process.kernels import DotProduct
    kernel = WhiteKernel(0.1, (1e-2, 1e2))
    kernel += ConstantKernel(1, (1e-2, 1e2)) * RBF(1, (1e-2, 1e2))
    kernel += ExpSineSquared(1, 1, (1e-2, 1e2), (1e-2, 1e2))
    kernel += DotProduct(1, (1e-2, 1e2))
```

DotProduct (somewhat) captures the trend:

$$K(x_i, x_j) = \sigma^2 + x_i x_j$$

- lacktriangle The larger the x values, the larger the correlation
- This allows the distance from the mean (which is zero) to grow
- lacktriangle The $oldsymbol{\sigma}$ parameter controls the base level of correlation
- Unlike all kernels so far DotProduct is not translation-invariant (we sa!

The new predictions are a bit better at the edges of the plot

```
In [22]: gp = GaussianProcessRegressor(kernel=kernel, n_restarts_optimizer=9)
         gp.fit(y tr.index.values.reshape(-1,1), y tr.values) # needs 2D input
         print(gp.kernel )
         xp, std = qp.predict(x.reshape(-1,1), return std=True)
         xp = pd.Series(index=y.index, data=xp)
         std = pd.Series(index=y.index, data=std)
         nab.plot gp(target=y, samples=y tr, pred=xp, std=std, figsize=figsize)
         WhiteKernel(noise level=0.01) + 1.17**2 * RBF(length scale=0.305) + ExpSineSquared(length scal
         e=2.17, periodicity=0.939) + DotProduct(sigma 0=0.0197)
         /usr/local/lib/python3.9/site-packages/sklearn/gaussian process/kernels.py:402: ConvergenceWar
         ning: The optimal value found for dimension 0 of parameter k1 k1 k1 noise level is close to
         the specified lower bound 0.01. Decreasing the bound and calling fit again may find a better v
         alue.
           warnings.warn("The optimal value found for "
                                                 Figure 1
                       target
                       predictions
                       95% C.I.
                       samples
```

Considerations

Gaussian Processes are a very flexible ML technique

- They can be used to making predictions
- ...Together with their confidence intervals
- ...But also for (conditional) density estimation
- ...And for generating data

They are non-trivial to use:

- In particular, choosing a kernel requires some practice and some understanding
- Automating the process is possible, but complex (grid search is likely not enough)

Gaussian processes tend to perform better: