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pyGAM is a package for building Generalized Additive Models in Python, with an emphasis on modularity and performance. The API will be immediately familiar to anyone with experience of scikit-learn or scipy.
pyGAM is on pypi, and can be installed using pip:

```
pip install pygam
```

Or via conda-forge, however this is typically less up-to-date:

```
conda install -c conda-forge pyGAM
```

You can install the bleeding edge from github using flit. First clone the repo, cd into the main directory and do:

```
pip install flit
flit install
```

### 1.1 Optional

To speed up optimization on large models with constraints, it helps to have scikit-sparse installed because it contains a slightly faster, sparse version of Cholesky factorization. The import from scikit-sparse references nose, so you’ll need that too.

The easiest way is to use Conda:

```
conda install -c conda-forge scikit-sparse nose
```

More information is available in the scikit-sparse docs.
pyGAM is tested on Python 2.7 and 3.6 and depends on NumPy, SciPy, and progressbar2 (see requirements.txt for version information).

Optional: scikit-sparse.

In addition to the above dependencies, the datasets submodule relies on Pandas.
Citing pyGAM

To report an issue with pyGAM please use the issue tracker.
CHAPTER 5

License

GNU General Public License v3.0
CHAPTER 6

Getting Started

If you’re new to pyGAM, read the Tour of pyGAM for an introduction to the package.

6.1 Quick Start

This quick start will show how to do the following:

• **Install** everything needed to use pyGAM.
• **fit a regression model with custom terms**
• **search for the best smoothing parameters**
• **plot partial dependence functions**

6.1.1 Install pyGAM

**Pip**

```
pip install pygam
```

**Conda**

pyGAM is on conda-forge, however this is typically less up-to-date:

```
conda install -c conda-forge pygam
```
Bleeding edge

You can install the bleeding edge from github using flit. First clone the repo, cd into the main directory and do:

```
pip install flit
flit install
```

Get pandas and matplotlib

```
pip install pandas matplotlib
```

6.1.2 Fit a Model

Let’s get to it. First we need some data:

```
[1]: from pygam.datasets import wage
    X, y = wage()
```

Now let’s import a GAM that’s made for regression problems.

Let’s fit a spline term to the first 2 features, and a factor term to the 3rd feature.

```
[2]: from pygam import LinearGAM, s, f
    gam = LinearGAM(s(0) + s(1) + f(2)).fit(X, y)
```

Let’s take a look at the model fit:

```
[3]: gam.summary()
```

<table>
<thead>
<tr>
<th>Feature Function</th>
<th>Lambda</th>
<th>Rank</th>
<th>EDoF</th>
<th>P &gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>x</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

(continues on next page)
WARNING: Fitting splines and a linear function to a feature introduces a model
identifiability problem which can cause p-values to appear significant when they are not.

WARNING: p-values calculated in this manner behave correctly for un-penalized models
or models with known smoothing parameters, but when smoothing parameters have been estimated, the p-values
are typically lower than they should be, meaning that the tests reject the null too readily.

Even though we have 3 terms with a total of \((20 + 20 + 5) = 45\) free variables, the default smoothing penalty \((\lambda = 0.6)\) reduces the effective degrees of freedom to just \(~25\).

By default, the spline terms, \(s(\ldots)\), use 20 basis functions. This is a good starting point. The rule of thumb is to use a fairly large amount of flexibility, and then let the smoothing penalty regularize the model.

However, we can always use our expert knowledge to add flexibility where it is needed, or remove basis functions, and make fitting easier:

```python
[22]: gam = LinearGAM(s(0, n_splines=5) + s(1) + f(2)).fit(X, y)
```

### 6.1.3 Automatically tune the model

By default, spline terms, \(s()\) have a penalty on their 2nd derivative, which encourages the functions to be smoother, while factor terms, \(f()\) and linear terms \(l()\), have a l2, ie ridge penalty, which encourages them to take on smaller values.

\(\lambda\), short for \(\lambda\), controls the strength of the regularization penalty on each term. Terms can have multiple penalties, and therefore multiple \(\lambda\).

```python
[14]: print(gam.lam)
[[0.6], [0.6], [0.6]]
```

Our model has 3 \(\lambda\) parameters, currently just one per term.

Let’s perform a grid-search over multiple \(\lambda\) values to see if we can improve our model.

We will seek the model with the lowest generalized cross-validation (GCV) score.

Our search space is 3-dimensional, so we have to be conservative with the number of points we consider per dimension.
Let’s try 5 values for each smoothing parameter, resulting in a total of $5 \times 5 \times 5 = 125$ points in our grid.

```python
[15]: import numpy as np
lam = np.logspace(-3, 5, 5)
lams = [lam] * 3
gam.gridsearch(X, y, lam=lams)
gam.summary()
```

```
100% (125 of 125) |################################| Elapsed Time: 0:00:07 Time: 0:00:07

LinearGAM
===============================================
Distribution: NormalDist Effective DoF: 9.2948
Link Function: IdentityLink Log Likelihood: -24119.7277
Number of Samples: 3000 AIC: 48260.0451
 48260.1229
 1244.089
 1237.1528
 0.2915
Pseudo R-Squared:

Feature Function Lambda Rank EDoF P >

x Sig. Code

s(0) [100000.] 5 2.0 7.
s(1) [1000.] 20 3.3 1.
f(2) [0.1] 5 4.0 1.
intercept 1 0.0 1.

Significance codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

WARNING: Fitting splines and a linear function to a feature introduces a model
  identifiability problem
  which can cause p-values to appear significant when they are not.

WARNING: p-values calculated in this manner behave correctly for un-penalized models
  or models with known smoothing parameters, but when smoothing parameters have been
  estimated, the p-values are typically lower than they should be, meaning that the tests reject the
  null too readily.
```

This is quite a bit better. Even though the in-sample $R^2$ value is lower, we can expect our model to generalize better because the GCV error is lower.

We could be more rigorous by using a train/test split, and checking our model’s error on the test set. We were also
quite lazy and only tried 125 values in our hyperopt. We might find a better model if we spent more time searching across more points.

For high-dimensional search-spaces, it is sometimes a good idea to try a randomized search. We can achieve this by using numpy’s random module:

```python
[16]: lams = np.random.rand(100, 3) # random points on [0, 1], with shape (100, 3)
lams = lams * 6 - 3 # shift values to -3, 3
lams = 10 ** lams # transforms values to 1e-3, 1e3

[17]: random_gam = LinearGAM(s(0) + s(1) + f(2)).gridsearch(X, y, lam=lams)
random_gam.summary()

100% (100 of 100) |###########################| Elapsed Time: 0:00:07 Time: 0:00:07

LinearGAM
===============================================
Distribution: NormalDist Effective DoF: 15.6683
Link Function: IdentityLink Log Likelihood: -24115.6727
Number of Samples: 3000 AIC: 48264.6819
    48264.8794
    1247.2011
    1235.4817
    0.2939

Feature Function Lambda Rank EDoF P >
================================= ==================== ============ ============
    == =================
    =============== ==
s(0) [137.6336] 20 6.3 7.08e-03 **
s(1) [128.3511] 20 5.4 1.11e-16 ***
f(2) [0.3212] 5 4.0 1.11e-16 ***
intercept 1 0.0 1.11e-16 ***
```

Significance codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

WARNING: Fitting splines and a linear function to a feature introduces a model identifiability problem which can cause p-values to appear significant when they are not.

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(continues on next page)
are typically lower than they should be, meaning that the tests reject the null too readily.

In this case, our deterministic search found a better model:

```
[18]: gam.statistics_['GCV'] < random_gam.statistics_['GCV']
[18]: True
```

The `statistics_` attribute is populated after the model has been fitted. There are lots of interesting model statistics to check out, although many are automatically reported in the model summary:

```
[19]: list(gam.statistics_.keys())
[19]: ['n_samples', 'm_features', 'edof_per_coef', 'edof', 'scale', 'cov', 'se', 'AIC', 'AICC', 'pseudo_r2', 'GCV', 'UBRE', 'loglikelihood', 'deviance', 'p_values']
```

### 6.1.4 Partial Dependence Functions

One of the most attractive properties of GAMs is that we can decompose and inspect the contribution of each feature to the overall prediction.

This is done via *partial dependence* functions.

Let’s plot the partial dependence for each term in our model, along with a 95% confidence interval for the estimated function.

```
[20]: import matplotlib.pyplot as plt

[21]: for i, term in enumerate(gam.terms):
    if term.isintercept:
        continue
    XX = gam.generate_X_grid(term=i)
    pdep, confi = gam.partial_dependence(term=i, X=XX, width=0.95)

    plt.figure()
    plt.plot(XX[:, term.feature], pdep)
    plt.plot(XX[:, term.feature], confi, c='r', ls='--')
    plt.title(repr(term))
    plt.show()
```
Note: we skip the intercept term because it has nothing interesting to plot.
6.2 A Tour of pyGAM

6.2.1 Introduction

Generalized Additive Models (GAMs) are smooth semi-parametric models of the form:

\[ g(\mathbb{E}[y|X]) = \beta_0 + f_1(X_1) + f_2(X_2, X_3) + \ldots + f_M(X_N) \]

where \( X.T = [X_1, X_2, \ldots, X_N] \) are independent variables, \( y \) is the dependent variable, and \( g() \) is the link function that relates our predictor variables to the expected value of the dependent variable.

The feature functions \( f_\_i() \) are built using penalized B splines, which allow us to automatically model non-linear relationships without having to manually try out many different transformations on each variable.
GAMs extend generalized linear models by allowing non-linear functions of features while maintaining additivity. Since the model is additive, it is easy to examine the effect of each $X_i$ on $Y$ individually while holding all other predictors constant.

The result is a very flexible model, where it is easy to incorporate prior knowledge and control overfitting.

### 6.2.2 Generalized Additive Models, in general

$$y \sim \text{ExponentialFamily}(\mu|X)$$

where

$$g(\mu|X) = \beta_0 + f_1(X_1) + f_2(X_2, X_3) + \ldots + f_M(X_N)$$

So we can see that a GAM has 3 components:

- distribution from the exponential family
- link function $g(\cdot)$
- functional form with an additive structure $\beta_0 + f_1(X_1) + f_2(X_2, X_3) + \ldots + f_M(X_N)$

**Distribution:**

Specified via: `GAM(distribution='...')`

Currently you can choose from the following:

- 'normal'
- 'binomial'
- 'poisson'
- 'gamma'
- 'inv_gauss'

**Link function:**

We specify this using: `GAM(link='...')`

Link functions take the distribution mean to the linear prediction. So far, the following are available:

- 'identity'
- 'logit'
- 'inverse'
- 'log'
- 'inverse-squared'
Functional Form:

Specified in \texttt{GAM(terms=...)} or more simply \texttt{GAM(...)}

In \texttt{pyGAM}, we specify the functional form using terms:

- \texttt{l()} linear terms: for terms like \texttt{X}_i
- \texttt{s()} spline terms
- \texttt{f()} factor terms
- \texttt{te()} tensor products
- \texttt{intercept}

With these, we can quickly and compactly build models like:

\begin{verbatim}
[12]: from pygam import GAM, s, te
    GAM(s(0, n_splines=200) + te(3,1) + s(2), distribution='poisson', link='log')
[12]: GAM(callbacks=['deviance', 'diffs'], distribution='poisson',
     fit_intercept=True, link='log', max_iter=100,
     terms=s(0) + te(3, 1) + s(2), tol=0.0001, verbose=False)
\end{verbatim}

which specifies that we want a:

- spline function on feature 0, with 200 basis functions
- tensor spline interaction on features 1 and 3
- spline function on feature 2

Note:

\texttt{GAM(..., intercept=True)} so models include an intercept by default.

in Practice...

in \texttt{pyGAM} you can build custom models by specifying these 3 elements, or you can choose from \texttt{common models}:

- \texttt{LinearGAM} identity link and normal distribution
- \texttt{LogisticGAM} logit link and binomial distribution
- \texttt{PoissonGAM} log link and Poisson distribution
- \texttt{GammaGAM} log link and gamma distribution
- \texttt{InvGauss} log link and inv_gauss distribution

The benefit of the common models is that they have some extra features, apart from reducing boilerplate code.

6.2.3 Terms and Interactions

\texttt{pyGAM} can also fit interactions using tensor products via \texttt{te()}
gam = PoissonGAM(s(0, n_splines=200) + te(3, 1) + s(2)).fit(X, y)

and plot a 3D surface:

```
import matplotlib.pyplot as plt
from mpl_toolkits import mplot3d
plt.ion()
plt.rcParams['figure.figsize'] = (12, 8)

XX = gam.generate_X_grid(term=1, meshgrid=True)
Z = gam.partial_dependence(term=1, X=XX, meshgrid=True)
ax = plt.axes(projection='3d')
a = ax.plot_surface(XX[0], XX[1], Z, cmap='viridis')
```

For simple interactions it is sometimes useful to add a by-variable to a term

```
from pygam import LinearGAM, s
from pygam.datasets import toy_interaction
X, y = toy_interaction(return_X_y=True)
gam = LinearGAM(s(0, by=1)).fit(X, y)
gam.summary()
```
6.2.4 Regression

For regression problems, we can use a linear GAM which models:

\[ E[y | X] = \beta_0 + f_1(X_1) + f_2(X_2, X_3) + \cdots + f_M(X_N) \]

```python
from pygam import LinearGAM, s, f
from pygam.datasets import wage

X, y = wage(return_X_y=True)

# model
gam = LinearGAM(s(0) + s(1) + f(2))
gam.gridsearch(X, y)
```

(continues on next page)
## plotting

```python
# plotting
plt.figure();
fig, axs = plt.subplots(1,3);

titles = ['year', 'age', 'education']
for i, ax in enumerate(axs):
    XX = gam.generate_X_grid(term=i)
    ax.plot(XX[:, i], gam.partial_dependence(term=i, X=XX))
    ax.plot(XX[:, i], gam.partial_dependence(term=i, X=XX, width=.95)[1], c='r', ls='--')
    if i == 0:
        ax.set_ylim(-30,30)
        ax.set_title(titles[i]);
```

Even though our model allows coefficients, our smoothing penalty reduces us to just 19 effective degrees of freedom:

```python
[4]: gam.summary()
```

```
LinearGAM
```

```
Distribution: NormalDist Effective DoF: 19.2602
```

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**pyGAM Documentation**

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<table>
<thead>
<tr>
<th>Feature Function</th>
<th>Lambda</th>
<th>Rank</th>
<th>EDoF</th>
<th>P &gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>s(0) [15.8489]</td>
<td>20</td>
<td>6.9</td>
<td>5.2e-03 **</td>
<td></td>
</tr>
<tr>
<td>s(1) [15.8489]</td>
<td>20</td>
<td>8.5</td>
<td>11e-16 ***</td>
<td></td>
</tr>
<tr>
<td>f(2) [15.8489]</td>
<td>5</td>
<td>3.8</td>
<td>11e-16 ***</td>
<td></td>
</tr>
<tr>
<td>intercept</td>
<td>0</td>
<td>1</td>
<td>0.0</td>
<td>1.</td>
</tr>
</tbody>
</table>

Significance codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '. ' 0.1 ' ' 1

**WARNING:** Fitting splines and a linear function to a feature introduces a model identifiability problem, which can cause p-values to appear significant when they are not.

**WARNING:** p-values calculated in this manner behave correctly for un-penalized models or models with known smoothing parameters, but when smoothing parameters have been estimated, the p-values are typically lower than they should be, meaning that the tests reject the null too readily.

With **LinearGAMs**, we can also check the prediction intervals:

```python
from pygam import LinearGAM
from pygam.datasets import mcycle

X, y = mcycle(return_X_y=True)

gam = LinearGAM(n_splines=25).gridsearch(X, y)
XX = gam.generate_X_grid(term=0, n=500)

plt.plot(XX, gam.predict(XX), 'r--')
plt.plot(XX, gam.prediction_intervals(XX, width=.95), color='b', ls='--')

plt.scatter(X, y, facecolor='gray', edgecolors='none')
plt.title('95% prediction interval');
```

With **LinearGAMs**, we can also check the prediction intervals:
And simulate from the posterior:

```python
# continuing last example with the mcycle dataset
for response in gam.sample(X, y, quantity='y', n_draws=50, sample_at_X=XX):
    plt.scatter(XX, response, alpha=.03, color='k')
plt.plot(XX, gam.predict(XX), 'r--')
plt.plot(XX, gam.prediction_intervals(XX, width=.95), color='b', ls='--')
plt.title('draw samples from the posterior of the coefficients')
```

100% (11 of 11) |###################################| Elapsed Time: 0:00:00 Time: 0:00:00
100% (11 of 11) |###################################| Elapsed Time: 0:00:00 Time: 0:00:00
100% (11 of 11) |###################################| Elapsed Time: 0:00:00 Time: 0:00:00
100% (11 of 11) |###################################| Elapsed Time: 0:00:00 Time: 0:00:00

[57]: Text(0.5,1,'draw samples from the posterior of the coefficients')
6.2.5 Classification

For binary classification problems, we can use a logistic GAM which models:

\[
\log \left( \frac{P(y = 1 | X)}{P(y = 0 | X)} \right) = \beta_0 + f_1(X_1) + f_2(X_2, X_3) + \cdots + f_M(X_N)
\]

[11]: from pygam import LogisticGAM, s, f
   from pygam.datasets import default

X, y = default(return_X_y=True)

gam = LogisticGAM(f(0) + s(1) + s(2)).gridsearch(X, y)

fig, axs = plt.subplots(1, 3)
titles = ['student', 'balance', 'income']

for i, ax in enumerate(axs):
    XX = gam.generate_X_grid(term=i)
    pdep, confi = gam.partial_dependence(term=i, width=.95)
    ax.plot(XX[:, i], pdep)
    ax.plot(XX[:, i], confi, c='r', ls='--')
    ax.set_title(titles[i]);
We can then check the accuracy:

```python
[8]: gam.accuracy(X, y)
```

```python
[8]: 0.9739
```

Since the scale of the Binomial distribution is known, our gridsearch minimizes the Un-Biased Risk Estimator (UBRE) objective:

```python
[9]: gam.summary()
```

LogisticGAM

-----------------------------------------------

```
Distribution: BinomialDist Effective DoF: 3.8047
Link Function: LogitLink Log Likelihood: -788.877
Number of Samples: 10000 AIC: 1585.3634
```

(continues on next page)
6.2.6 Poisson and Histogram Smoothing

We can intuitively perform **histogram smoothing** by modeling the counts in each bin as being distributed Poisson via **PoissonGAM**.

```python
from pygam import PoissonGAM
from pygam.datasets import faithful

X, y = faithful(return_X_y=True)

gam = PoissonGAM().gridsearch(X, y)

plt.hist(faithful(return_X_y=False)['eruptions'], bins=200, color='k');
plt.plot(X, gam.predict(X), color='r')
plt.title('Best Lambda: {0:.2f}'.format(gam.lam[0][0]));
```

100% (11 of 11) |##########################################################| Elapsed Time: 0:00:00 Time: 0:00:00
GAMs with a Normal distribution suffer from the limitation of an assumed constant variance. Sometimes this is not an appropriate assumption, because we’d like the variance of our error distribution to vary.

In this case we can resort to modeling the **expectiles** of a distribution.

Expectiles are intuitively similar to quantiles, but model tail expectations instead of tail mass. Although they are less interpretable, expectiles are much faster to fit, and can also be used to non-parametrically model a distribution.

```
[52]: from pygam import ExpectileGAM
from pygam.datasets import mcycle

X, y = mcycle(return_X_y=True)

# lets fit the mean model first by CV
gam50 = ExpectileGAM(expectile=0.5).gridsearch(X, y)

# and copy the smoothing to the other models
lam = gam50.lam

# now fit a few more models
gam95 = ExpectileGAM(expectile=0.95, lam=lam).fit(X, y)
gam75 = ExpectileGAM(expectile=0.75, lam=lam).fit(X, y)
gam25 = ExpectileGAM(expectile=0.25, lam=lam).fit(X, y)
gam05 = ExpectileGAM(expectile=0.05, lam=lam).fit(X, y)
```
We fit the **mean model** by cross-validation in order to find the best smoothing parameter \( \lambda \) and then copy it over to the other models.

This practice makes the expectiles less likely to cross.

### 6.2.8 Custom Models

It’s also easy to build custom models by using the base **GAM** class and specifying the **distribution** and the **link** function:

```python
from pygam import GAM
from pygam.datasets import trees

X, y = trees(return_X_y=True)
```

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pyGAM Documentation

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gam = GAM(distribution='gamma', link='log')
gam.gridsearch(X, y)

plt.scatter(y, gam.predict(X))
plt.xlabel('true volume')
plt.ylabel('predicted volume')

We can check the quality of the fit by looking at the Pseudo R-Squared:

[28]: gam.summary()

GAM

Distribution: GammaDist Effective DoF:
  25.3616
Link Function: LogLink Log Likelihood:
  -26.1673
Number of Samples: 31 AIC:
  105.0579
  501.5549
  0.0088

(continues on next page)
6.2.9 Penalties / Constraints

With GAMs we can encode prior knowledge and control overfitting by using penalties and constraints.

**Available penalties** - second derivative smoothing (default on numerical features) - L2 smoothing (default on categorical features)

**Available constraints** - monotonic increasing/decreasing smoothing - convex/concave smoothing - periodic smoothing

We can inject our intuition into our model by using monotonic and concave constraints:

```python
from pygam import LinearGAM, s
from pygam.datasets import hepatitis

X, y = hepatitis(return_X_y=True)

gam1 = LinearGAM(s(0, constraints='monotonic_inc')).fit(X, y)
gam2 = LinearGAM(s(0, constraints='concave')).fit(X, y)

fig, ax = plt.subplots(1, 2)
ax[0].plot(X, y, label='data')
ax[0].plot(X, gam1.predict(X), label='monotonic fit')
ax[0].legend()

ax[1].plot(X, y, label='data')
```

(continues on next page)
6.2.10 API

pyGAM is intuitive, modular, and adheres to a familiar API:

```python
from pygam import LogisticGAM, s, f
from pygam.datasets import toy_classification

X, y = toy_classification(return_X_y=True, n=5000)

gam = LogisticGAM(s(0) + s(1) + s(2) + s(3) + s(4) + f(5))
gam.fit(X, y)
```

Since GAMs are additive, it is also super easy to visualize each individual feature function, \( f_i(X_i) \). These feature functions describe the effect of each \( X_i \) on \( y \) individually while marginalizing out all other predictors:
6.2.11 Current Features

Models

pyGAM comes with many models out-of-the-box:

- GAM (base class for constructing custom models)
- LinearGAM
- LogisticGAM
- GammaGAM
- PoissonGAM
- InvGaussGAM
- ExpectileGAM
Terms

• l() linear terms
• s() spline terms
• f() factor terms
• te() tensor products
• intercept

Distributions

• Normal
• Binomial
• Gamma
• Poisson
• Inverse Gaussian

Link Functions

Link functions take the distribution mean to the linear prediction. These are the canonical link functions for the above distributions:

• Identity
• Logit
• Inverse
• Log
• Inverse-squared

Callbacks

Callbacks are performed during each optimization iteration. It’s also easy to write your own.

• deviance - model deviance
• diffs - differences of coefficient norm
• accuracy - model accuracy for LogisticGAM
• coef - coefficient logging

You can check a callback by inspecting:

```python
[32]: _ = plt.plot(gam.logs_['deviance'])
```
Linear Extrapolation

```
[33]: from pygam import LinearGAM
   from pygam.datasets import mcycle
   X, y = mcycle()
   gam = LinearGAM()
   gam.gridsearch(X, y)
   XX = gam.generate_X_grid(term=0)
   m = X.min()
   M = X.max()
   XX = np.linspace(m - 10, M + 10, 500)
   Xl = np.linspace(m - 10, m, 50)
   Xr = np.linspace(M, M + 10, 50)
   plt.figure()
   plt.plot(XX, gam.predict(XX), 'k')
   plt.plot(Xl, gam.confidence_intervals(Xl), color='b', ls='--')
   plt.plot(Xr, gam.confidence_intervals(Xr), color='b', ls='--')
   _ = plt.plot(X, gam.confidence_intervals(X), color='r', ls='--')
```

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6.2.12 References

   Generalized Additive Models: an introduction with R

2. Hastie, Tibshirani, Friedman
   The Elements of Statistical Learning

3. James, Witten, Hastie and Tibshirani
   An Introduction to Statistical Learning

4. Paul Eilers & Brian Marx, 1996 Flexible Smoothing with B-splines and Penalties

5. Kim Larsen, 2015
   GAM: The Predictive Modeling Silver Bullet

   UCI Machine Learning: Notes on IRLS

7. Paul Eilers & Brian Marx, 2015
   International Biometric Society: A Crash Course on P-splines
6.3 User API

6.3.1 Generalized Additive Model Classes

GAM

```python
class pygam.pygam.GAM(terms='auto', max_iter=100, tol=0.0001, distribution='normal',
    link='identity', callbacks=['deviance', 'diffs'], fit_intercept=True, verbose=False, **kwargs)
```

Bases: pygam.core.Core, pygam.terms.MetaTermMixin

Generalized Additive Model

**Parameters**

- `terms (expression specifying terms to model, optional.)` – By default a univariate spline term will be allocated for each feature.

  For example:

  ```
  >>> GAM(s(0) + l(1) + f(2) + te(3, 4))
  ```

  will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

- `callbacks (list of str or list of CallBack objects, optional)` – Names of callback objects to call during the optimization loop.

- `distribution (str or Distribution object, optional)` – Distribution to use in the model.

- `link (str or Link object, optional)` – Link function to use in the model.

- `fit_intercept (bool, optional)` – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.

- `max_iter (int, optional)` – Maximum number of iterations allowed for the solver to converge.

- `tol (float, optional)` – Tolerance for stopping criteria.

- `verbose (bool, optional)` – whether to show pyGAM warnings.

**coef_**

Coefficient of the features in the decision function. If fit_intercept is True, then self.coef_[0] will contain the bias.

Type array, shape (n_classes, m_features)

**statistics_**

Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

Type dict
logs_
Dictionary containing the outputs of any callbacks at each optimization loop.

The logs are structured as {callback: [...]}  
  Type  dict

References

Hastie, Tibshirani, Friedman The Elements of Statistical Learning  
Paul Eilers & Brian Marx, 2015 International Biometric Society: A Crash Course on P-splines  

confidence_intervals (X, width=0.95, quantiles=None)  
estimate confidence intervals for the model.

Parameters

• X (array-like of shape (n_samples, m_features)) – Input data matrix
• width (float on [0,1], optional) –
• quantiles (array-like of floats in (0, 1), optional) – Instead of specifying the prediction width, one can specify the quantiles. So width=.95 is equivalent to quantiles=[.025, .975]

Returns intervals

Return type np.array of shape (n_samples, 2 or len(quantiles))

Notes

Wood 2006, section 4.9 Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.

deviance_residuals (X, y, weights=None, scaled=False)  
method to compute the deviance residuals of the model
these are analogous to the residuals of an OLS.

Parameters

• X (array-like) – Input data array of shape (n_samples, m_features)
• y (array-like) – Output data vector of shape (n_samples,)
• weights (array-like shape (n_samples,) or None, optional) – Sample weights. If None, defaults to array of ones
• scaled (bool, optional) – whether to scale the deviance by the (estimated) distribution scale

Returns deviance_residuals – with shape (n_samples,)

Return type np.array

fit (X, y, weights=None)  
Fit the generalized additive model.
Parameters

- \(X\) (array-like, shape (n_samples, m_features)) – Training vectors.
- \(y\) (array-like, shape (n_samples,)) – Target values, i.e. integers in classification, real numbers in regression.
- \(weights\) (array-like shape (n_samples,) or None, optional) – Sample weights. If None, defaults to array of ones.

Returns `self` – Returns fitted GAM object

Return type object

`generate_X_grid(term, n=100, meshgrid=False)`
create a nice grid of X data
array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong if term is >= 0, we generate n samples per feature, which results in n^deg samples, where deg is the degree of the interaction of the term

Parameters

- \(term\) (int,) – Which term to process.
- \(n\) (int, optional) – Number of data points to create.
- \(meshgrid\) (bool, optional) – Whether to return a meshgrid (useful for 3D plotting) or a feature matrix (useful for inference like partial predictions).

Returns

- If \(meshgrid\) is False – np.array of shape (n, n_features) where m is the number of (sub)terms in the requested (tensor)term.
- Else – tuple of length m, where m is the number of (sub)terms in the requested (tensor)term. Each element in the tuple contains a np.ndarray of size \(n^m\).

Raises `ValueError` – If the term requested is an intercept since it does not make sense to process the intercept term.

`gridsearch(X, y, weights=None, return_scores=False, keep_best=True, objective='auto', progress=True, **param_grids)`
Performs a grid search over a space of parameters for a given objective

Warning: `gridsearch` is lazy and will not remove useless combinations from the search space, e.g.

```
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```
will result in 10 loops, of which 5 are equivalent because `fit_splines = False`

Also, it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

Parameters

- \(X\) (array-like) – Input data of shape (n_samples, m_features)
- \(y\) (array-like) – Label data of shape (n_samples,)
- \(weights\) (array-like shape (n_samples,), optional) – Sample weights
**return_scores** (*boolean, optional*) – whether to return the hyperparameters and score for each element in the grid

**keep_best** (*boolean, optional*) – whether to keep the best GAM as self.

**objective** (*{'auto', 'AIC', 'AICC', 'GCV', 'UBRE'}, optional*) – Metric to optimize. If *auto*, then grid search will optimize *GCV* for models with unknown scale and *UBRE* for models with known scale.

**progress** (*bool, optional*) – whether to display a progress bar

****kwargs** – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.

If no parameter are specified, lam=np.logspace(-3, 3, 11) is used. This results in a 11 points, placed diagonally across lam space.

If grid is iterable of iterables of floats, the outer iterable must have length m.features. the cartesian product of the subgrids in the grid will be tested.

If grid is a 2d numpy array, each row of the array will be tested.

The method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

**Returns**

- if return_scores=True – model_scores: dict containing each fitted model as keys and corresponding objective scores as values
- else – self: ie possibly the newly fitted model

**Examples**

For a model with 4 terms, and where we expect 4 lam values, our search space for lam must have 4 dimensions.

We can search the space in 3 ways:

1. via cartesian product by specifying the grid as a list. our grid search will consider 11 ** 4 points:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = [lam] * 4
>>> gam.gridsearch(X, y, lam=lams)
```

2. directly by specifying the grid as a np.ndarray. This is useful for when the dimensionality of the search space is very large, and we would prefer to execute a randomized search:

```python
>>> lams = np.exp(np.random.random(50, 4) * 6 - 3)
>>> gam.gridsearch(X, y, lam=lams)
```

3. copying grids for parameters with multiple dimensions. if we specify a 1D np.ndarray for lam, we are implicitly testing the space where all points have the same value

```python
>>> gam.gridsearch(lam=np.logspace(-3, 3, 11))
```

is equivalent to:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = np.array([lam] * 4)
>>> gam.gridsearch(X, y, lam=lams)
```
**loglikelihood** \((X, y, \text{weights}=\text{None})\)

compute the log-likelihood of the dataset using the current model

**Parameters**

- **X** *(array-like of shape \((n_{samples}, m\text{-features})\)) – containing the input dataset*
- **y** *(array-like of shape \((n,)\)) – containing target values*
- **weights** *(array-like of shape \((n,), \text{optional}\)) – containing sample weights*

**Returns** *log-likelihood – containing log-likelihood scores*

**Return type** *np.array of shape \((n,)\)*

**partial_dependence** *(term, X=\text{None}, width=\text{None}, quantiles=\text{None}, meshgrid=\text{False})*

Computes the term functions for the GAM and possibly their confidence intervals.

if both width=\text{None} and quantiles=\text{None}, then no confidence intervals are computed

**Parameters**

- **term** *(int, \text{optional}) – Term for which to compute the partial dependence functions.*
- **X** *(array-like with input data, \text{optional}) – if meshgrid=\text{False}, then \(X\) should be an array-like of shape \((n_{samples}, m\text{-features})\).*

  - if meshgrid=\text{True}, then \(X\) should be a tuple containing an array for each feature in the term.

  - if None, an equally spaced grid of points is generated.
- **width** *(float on \((0, 1), \text{optional}) – Width of the confidence interval.*
- **quantiles** *(array-like of floats on \((0, 1), \text{optional}) – instead of specifying the prediction width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]. if None, defaults to width.*
- **meshgrid** *(bool, \text{whether to return and accept meshgrids.}) – Useful for creating outputs that are suitable for 3D plotting.*

  - Note, for simple terms with no interactions, the output of this function will be the same for meshgrid=\text{True} and meshgrid=\text{False}, but the inputs will need to be different.

**Returns**

- **pdeps** *(np.array of shape \((n_{samples},))\)*
- **conf_intervals** *(list of length \(\text{len(term)}\)) – containing np.arrays of shape \((n_{samples}, 2 \text{ or } \text{len(quantiles)})\)*

**Raises** *ValueError : – If the term requested is an intercept since it does not make sense to process the intercept term.*

**See also:**

- **generate_X_grid()** for help creating meshgrids.

**predict** *(\(X\))*

predict expected value of target given model and input \(X\) often this is done via expected value of GAM given input \(X\)
**predict(x)**

**Parameters**

- **X** *(array-like of shape (n_samples, m_features))** – containing the input dataset

**Returns**

- **y** – containing predicted values under the model

**Return type**

np.array of shape (n_samples,)

**predict_mu(x)**

**Parameters**

- **X** *(array-like of shape (n_samples, m_features),) – containing the input dataset

**Returns**

- **y** – containing expected values under the model

**Return type**

np.array of shape (n_samples,)

**sample(x, y, quantity='y', sample_at_X=None, weights=None, n_draws=100, n_bootstraps=5, objective='auto')**

Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with **gridsearch** (with the parameter **keep_best=True**) before calling **sample**, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. **n_bootstraps** many “bootstrap samples” of the response (**y**) are simulated by drawing random samples from the model’s distribution evaluated at the expected values (**mu**) for each sample in **x**.
2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter **lam** given the response data **y**.
3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.

**Notes**

A **gridsearch** is done **n_bootstraps** many times, so keep **n_bootstraps** small. Make **n_bootstraps < n_draws** to take advantage of the expensive bootstrap samples of the smoothing parameters.

**Parameters**

- **X** *(array of shape (n_samples, m_features))** – empirical input data
- **y** *(array of shape (n_samples,))** – empirical response vector
- **quantity** (**{'y', 'coef', 'mu'}, default: 'y'**) – What quantity to return pseudorandom samples of. If **sample_at_X** is not None and **quantity** is either ‘y’ or ‘mu’, then samples are drawn at the values of **X** specified in **sample_at_X**.
- **sample_at_X** *(array of shape (n_samples_to_simulate, m_features) or)** –
  - **optional (None, )** – Input data at which to draw new samples.

Only applies for **quantity** equal to ‘y’ or to ‘mu’. If **None**, then **sample_at_X** is replaced by **X**.
• **weights** (*np.array of shape (n_samples,))*) – sample weights

• **n_draws** (*positive int, optional (default=100))*) – The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters

• **n_bootstraps** (*positive int, optional (default=5))*) – The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If *n_bootstraps* is 1, then only the already fitted model’s smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.

• **objective** (*string, optional (default='auto')*) – metric to optimize in grid search. must be in ['AIC', 'AICc', 'GCV', 'UBRE', 'auto'] if 'auto', then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

Returns

draws – Simulations of the given quantity using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

If quantity == ‘coef’, then the number of columns of draws is the number of coefficients (len(self.coef_)).

Otherwise, the number of columns of draws is the number of rows of sample_at_X if sample_at_X is not None or else the number of rows of X.

Return type 2D array of length n_draws

References


score (*X, y, weights=None*)

method to compute the explained deviance for a trained model for a given X data and y labels

Parameters

• **X** (*array-like*) – Input data array of shape (n_samples, m_features)

• **y** (*array-like*) – Output data vector of shape (n_samples,)

• **weights** (*array-like shape (n_samples,) or None, optional*) – Sample weights. if None, defaults to array of ones

Returns explained deviance score

Return type np.array (n_samples,)

summary ()

produce a summary of the model statistics

Parameters None –

Returns

Return type None
LinearGAM

class pygam.pygam.LinearGAM(terms='auto', max_iter=100, tol=0.0001, scale=None, callbacks=['deviance', 'difs'], fit_intercept=True, verbose=False, **kwargs)

Bases: pygam.pygam.GAM

Linear GAM

This is a GAM with a Normal error distribution, and an identity link.

Parameters

• **terms** *(expression specifying terms to model, optional.)* – By default a univariate spline term will be allocated for each feature.

  For example:

  ```
  >>> GAM(s(0) + l(1) + f(2) + te(3, 4))
  ```

  will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

• **callbacks** *(list of str or list of CallBack objects, optional)* – Names of callback objects to call during the optimization loop.

• **fit_intercept** *(bool, optional)* – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.

• **max_iter** *(int, optional)* – Maximum number of iterations allowed for the solver to converge.

• **tol** *(float, optional)* – Tolerance for stopping criteria.

• **verbose** *(bool, optional)* – Whether to show pyGAM warnings.

coef_

  Coefficient of the features in the decision function. If fit_intercept is True, then self.coef_[0] will contain the bias.

  Type array, shape (n_classes, m_features)

statistics_

  Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

  Type dict

logs_

  Dictionary containing the outputs of any callbacks at each optimization loop.

  The logs are structured as `{callback: [...]}`

  Type dict

References


confidence_intervals(X, width=0.95, quantiles=None)
estimate confidence intervals for the model.

Parameters

- X (array-like of shape (n_samples, m_features)) – Input data matrix
- width (float on [0,1], optional) –
- quantiles (array-like of floats in (0, 1), optional) – Instead of specifying the prediction width, one can specify the quantiles. So width=.95 is equivalent to quantiles=[.025, .975]

Returns intervals

Return type np.array of shape (n_samples, 2 or len(quantiles))

Notes

Wood 2006, section 4.9 Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.

deviance_residuals(X, y, weights=None, scaled=False)
method to compute the deviance residuals of the model
these are analogous to the residuals of an OLS.

Parameters

- X (array-like) – Input data array of shape (n_samples, m_features)
- y (array-like) – Output data vector of shape (n_samples,)
- weights (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones
- scaled (bool, optional) – whether to scale the deviance by the (estimated) distribution scale

Returns deviance_residuals – with shape (n_samples,)

Return type np.array

fit(X, y, weights=None)
Fit the generalized additive model.

Parameters

- X (array-like, shape (n_samples, m_features)) – Training vectors.
- y (array-like, shape (n_samples,)) – Target values, ie integers in classification, real numbers in regression
- weights (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones

Returns self – Returns fitted GAM object

Return type object
**generate_X_grid** *(term, n=100, meshgrid=False)*

create a nice grid of X data

array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong

if term is >= 0, we generate n samples per feature, which results in n^deg samples, where deg is the degree of the interaction of the term

**Parameters**

- **term** *(int, )* – Which term to process.
- **n** *(int, optional)* – number of data points to create
- **meshgrid** *(bool, optional)* – Whether to return a meshgrid (useful for 3d plotting) or a feature matrix (useful for inference like partial predictions)

**Returns**

- *if meshgrid is False* – np.array of shape (n, n_features) where m is the number of (sub)terms in the requested (tensor)term.
- *else* – tuple of len m, where m is the number of (sub)terms in the requested (tensor)term.
  each element in the tuple contains a np.ndarray of size (n)^m

**Raises** `ValueError`: – If the term requested is an intercept since it does not make sense to process the intercept term.

**get_params** *(deep=False)*

returns a dict of all of the object's user-facing parameters

**Parameters**

- **deep** *(boolean, default: False)* – when True, also gets non-user-facing parameters

**Returns**

**Return type** `dict`

**gridsearch** *(X, y, weights=None, return_scores=False, keep_best=True, objective='auto', progress=True, **param_grids)*

Performs a grid search over a space of parameters for a given objective

**Warning:** gridsearch is lazy and will not remove useless combinations from the search space, eg.

```
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```

will result in 10 loops, of which 5 are equivalent because fit_splines = False

Also, it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

**Parameters**

- **X** *(array-like)* – input data of shape (n_samples, m_features)
- **y** *(array-like)* – label data of shape (n_samples,)
- **weights** *(array-like shape (n_samples,), optional)* – sample weights
- **return_scores** *(boolean, optional)* – whether to return the hyperpamaters and score for each element in the grid
- **keep_best** *(boolean, optional)* – whether to keep the best GAM as self.
• **objective** (\{'auto\', 'AIC', 'AICc', 'GCV', 'UBRE'\}, optional) – Metric to optimize. If *auto*, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

• **progress** (bool, optional) – whether to display a progress bar

• **kwargs** – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.

If no parameter are specified, \(\text{lam} = \text{np.logspace}(-3, 3, 11)\) is used. This results in a 11 points, placed diagonally across lam space.

If grid is iterable of iterables of floats, the outer iterable must have length \(m_{\text{features}}\). the cartesian product of the subgrids in the grid will be tested.

If grid is a 2d numpy array, each row of the array will be tested.

The method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

**Returns**

• if `return_scores=True` – model Scores: dict containing each fitted model as keys and corresponding objective scores as values

• else – self: ie possibly the newly fitted model

**Examples**

For a model with 4 terms, and where we expect 4 lam values, our search space for lam must have 4 dimensions.

We can search the space in 3 ways:

1. via cartesian product by specifying the grid as a list. our grid search will consider \(11 \times 4\) points:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = [lam] * 4
>>> gam.gridsearch(X, y, lam=lams)
```

2. directly by specifying the grid as a np.ndarray. This is useful for when the dimensionality of the search space is very large, and we would prefer to execute a randomized search:

```python
>>> lams = np.exp(np.random.random(50, 4) * 6 - 3)
>>> gam.gridsearch(X, y, lam=lams)
```

3. copying grids for parameters with multiple dimensions. if we specify a 1D np.ndarray for lam, we are implicitly testing the space where all points have the same value

```python
>>> gam.gridsearch(lam=np.logspace(-3, 3, 11))
```

is equivalent to:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = np.array([lam] * 4)
>>> gam.gridsearch(X, y, lam=lams)
```

**loglikelihood** \((X, y, weights=None)\)
compute the log-likelihood of the dataset using the current model

**Parameters**
• \textbf{X} (array-like of shape \((n\_samples, m\_features)\)) – containing the input dataset
• \textbf{y} (array-like of shape \((n,)\)) – containing target values
• \textbf{weights} (array-like of shape \((n,)\), optional) – containing sample weights

Returns \textbf{log-likelihood} – containing log-likelihood scores

Return type \textbf{np.array of shape \((n,)\)}

\texttt{partial\_dependence(} \texttt{term, X=None, width=None, quantiles=None, meshgrid=False)\texttt{)}

Computes the term functions for the GAM and possibly their confidence intervals.

if both width=None and quantiles=None, then no confidence intervals are computed

Parameters

• \textbf{term} (int, optional) – Term for which to compute the partial dependence functions.
• \textbf{X} (array-like with input data, optional) – if \texttt{meshgrid=False}, then \textbf{X} should be an array-like of shape \((n\_samples, m\_features)\).
  if \texttt{meshgrid=True}, then \textbf{X} should be a tuple containing an array for each feature in the term.
  if None, an equally spaced grid of points is generated.
• \textbf{width} (float on \((0, 1)\), optional) – Width of the confidence interval.
• \textbf{quantiles} (array-like of floats on \((0, 1)\), optional) – instead of specifying the prediction width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]. if None, defaults to width.
• \textbf{meshgrid} (bool, whether to return and accept meshgrids.) – Useful for creating outputs that are suitable for 3D plotting.

Note, for simple terms with no interactions, the output of this function will be the same for \texttt{meshgrid=True} and \texttt{meshgrid=False}, but the inputs will need to be different.

Returns

• \textbf{pdeps} (np.array of shape \((n\_samples,)\))
• \textbf{conf\_intervals} (list of length \texttt{len(term)}) – containing np.arrays of shape \((n\_samples, 2 \text{ or len(quantiles)})\)

Raises \texttt{ValueError} – If the term requested is an intercept since it does not make sense to process the intercept term.

See also:

\texttt{generate\_X\_grid()} for help creating meshgrids.

\texttt{predict(X)}

\texttt{predict expected value of target given model and input X often this is done via expected value of GAM given input X}

Parameters \textbf{X} (array-like of shape \((n\_samples, m\_features)\)) – containing the input dataset

Returns \textbf{y} – containing predicted values under the model
Return type np.array of shape (n_samples,)

**predict_mu** (*X*)

predict expected value of target given model and input *X*

**Parameters**

- `X` ([array-like of shape (n_samples, m_features),]) – containing the input dataset

**Returns**

- `y` – containing expected values under the model

Return type np.array of shape (n_samples,)

**prediction_intervals** (*X*, *width*=0.95, *quantiles*=None)

estimate prediction intervals for LinearGAM

**Parameters**

- `X` ([array-like of shape (n_samples, m_features)]) – input data matrix

- `width` (float on [0,1], optional (default=0.95)) –

- `quantiles` ([array-like of floats in [0, 1], default: None]) – instead of specifying the prediction width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]

**Returns**

- `intervals` np.array of shape (n_samples, 2 or len(quantiles))

**sample** (*X*, *y*, *quantity*='y', *sample_at_X*=None, *weights*=None, *n_draws*=100, *n_bootstraps*=5, *objective*='auto')

Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with `gridsearch` (with the parameter `keep_best=True`) before calling `sample`, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. *n_bootstraps* many “bootstrap samples” of the response (*y*) are simulated by drawing random samples from the model’s distribution evaluated at the expected values (*mu*) for each sample in *X*.

2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter *lam* given the response data *y*.

3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.

**Notes**

A `gridsearch` is done *n_bootstraps* many times, so keep *n_bootstraps* small. Make *n_bootstraps < n_draws* to take advantage of the expensive bootstrap samples of the smoothing parameters.

**Parameters**

- `X` ([array of shape (n_samples, m_features)]) – empirical input data

- `y` ([array of shape (n_samples,))] – empirical response vector
• **quantity** (\{'y', 'coef', 'mu'\}, default: 'y') – What quantity to return pseudorandom samples of. If `sample_at_X` is not None and `quantity` is either ‘y’ or ‘mu’, then samples are drawn at the values of X specified in `sample_at_X`.

• **sample_at_X** (array of shape (n_samples_to_simulate, m_features) or) –

• **optional** (None, ) – Input data at which to draw new samples.

  Only applies for `quantity` equal to ‘y’ or to ‘mu’. If `None`, then `sample_at_X` is replaced by X.

• **weights** (np.array of shape (n_samples,)) – sample weights

• **n_draws** (positive int, optional (default=100)) – The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters

• **n_bootstraps** (positive int, optional (default=5)) – The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If `n_bootstraps` is 1, then only the already fitted model’s smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.

• **objective** (string, optional (default='auto') – metric to optimize in grid search. must be in ['AIC', 'AICc', 'GCV', 'UBRE', 'auto'] if `auto`, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

**Returns**

`draws` – Simulations of the given `quantity` using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

If `quantity == 'coef'`, then the number of columns of `draws` is the number of coefficients (len(self.coef_)).

Otherwise, the number of columns of `draws` is the number of rows of `sample_at_X` if `sample_at_X` is not `None` or else the number of rows of X.

**Return type** 2D array of length n_draws

**References**


```python
def score(self, X, y, weights=None):
    # method to compute the explained deviance for a trained model for a given X data and y labels
    # Parameters
    # • X (array-like) – Input data array of shape (n_samples, m_features)
    # • y (array-like) – Output data vector of shape (n_samples,)
    # • weights (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones
    
    # Returns explained deviance score
    
    # Return type np.array() (n_samples,)
```
set_params (deep=False, force=False, **parameters)
sets an object’s parameters

Parameters

- **deep** (boolean, default: False) – when True, also sets non-user-facing parameters
- **force** (boolean, default: False) – when True, also sets parameters that the object does not already have
- **parameters** (parameters to set)

Returns

Return type self

summary()
produce a summary of the model statistics

Parameters None –

Returns

Return type None

GammaGAM

class pygam.pygam.GammaGAM (terms='auto', max_iter=100, tol=0.0001, scale=None, callbacks=['deviance', 'diffs'], fit_intercept=True, verbose=False, **kwargs)
Bases: pygam.pygam.GAM
Gamma GAM
This is a GAM with a Gamma error distribution, and a log link.

NB Although canonical link function for the Gamma GLM is the inverse link, this function can create problems for numerical software because it becomes difficult to enforce the requirement that the mean of the Gamma distribution be positive. The log link guarantees this.

If you need to use the inverse link function, simply construct a custom GAM:

```python
>>> from pygam import GAM
>>> gam = GAM(distribution='gamma', link='inverse')
```

Parameters

- **terms** (expression specifying terms to model, optional.) – By default a univariate spline term will be allocated for each feature.
  For example:

  ```python
  >>> GAM(s(0) + l(1) + f(2) + te(3, 4))
  ```

  will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

- **callbacks** (list of str or list of CallBack objects, optional) – Names of callback objects to call during the optimization loop.

- **fit_intercept** (bool, optional) – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.
pyGAM Documentation

- **max_iter** (*int, optional*) – Maximum number of iterations allowed for the solver to converge.
- **tol** (*float, optional*) – Tolerance for stopping criteria.
- **verbose** (*bool, optional*) – Whether to show pyGAM warnings.

**coef_**
Coefficient of the features in the decision function. If fit_intercept is True, then self.coef_[0] will contain the bias.

Type: array, shape (n_classes, m_features)

**statistics_**
Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

Type: dict

**logs_**
Dictionary containing the outputs of any callbacks at each optimization loop.

The logs are structured as `{callback: [...]}`

Type: dict

References

**confidence_intervals** (*X, width=0.95, quantiles=None*)
Estimate confidence intervals for the model.

Parameters
- **X** (*array-like of shape (n_samples, m_features]*) – Input data matrix
- **width** (*float on [0,1], optional*) –
- **quantiles** (*array-like of floats in (0, 1), optional*) – Instead of specifying the prediciton width, one can specify the quantiles. So `width=.95` is equivalent to `quantiles=[.025, .975]`

Returns intervals

Return type: np.array of shape (n_samples, 2 or len(quantiles))

Notes
Wood 2006, section 4.9 Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.
**deviance_residuals** *(X, y, weights=None, scaled=False)*

method to compute the deviance residuals of the model

these are analogous to the residuals of an OLS.

**Parameters**

- **X** *(array-like)* – Input data array of shape (n_samples, m_features)
- **y** *(array-like)* – Output data vector of shape (n_samples,)
- **weights** *(array-like shape (n_samples,) or None, optional)* – Sample weights. if None, defaults to array of ones
- **scaled** *(bool, optional)* – whether to scale the deviance by the (estimated) distribution scale

**Returns** `deviance_residuals` – with shape (n_samples,)

**Return type** `np.array`

**fit** *(X, y, weights=None)*

Fit the generalized additive model.

**Parameters**

- **X** *(array-like, shape (n_samples, m_features))* – Training vectors.
- **y** *(array-like, shape (n_samples,))* – Target values, ie integers in classification, real numbers in regression
- **weights** *(array-like shape (n_samples,) or None, optional)* – Sample weights. if None, defaults to array of ones

**Returns** `self` – Returns fitted GAM object

**Return type** `object`

**generate_X_grid** *(term, n=100, meshgrid=False)*

create a nice grid of X data

array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong

if term is >= 0, we generate n samples per feature, which results in n^deg samples, where deg is the degree of the interaction of the term

**Parameters**

- **term** *(int,)* – Which term to process.
- **n** *(int, optional)* – number of data points to create
- **meshgrid** *(bool, optional)* – Whether to return a meshgrid (useful for 3d plotting) or a feature matrix (useful for inference like partial predictions)

**Returns**

- **if meshgrid is False** – np.array of shape (n, n_features) where m is the number of (sub)terms in the requested (tensor)term.
- **else** – tuple of len m, where m is the number of (sub)terms in the requested (tensor)term. each element in the tuple contains a np.ndarray of size (n)^m

**Raises** `ValueError`: – If the term requested is an intercept since it does not make sense to process the intercept term.
**get_params** *(deep=False)*
returns a dict of all of the object’s user-facing parameters

Parameters **deep** *(boolean, default: False)* – when True, also gets non-user-facing parameters

Returns

Return type dict

**gridsearch** *(X, y, weights=None, return_scores=False, keep_best=True, objective='auto', progress=True, **param_grids)*
Performs a grid search over a space of parameters for a given objective

```python
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```
will result in 10 loops, of which 5 are equivalent because fit_splines = False

Also, it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

Warning: gridsearch is lazy and will not remove useless combinations from the search space, eg.

Parameters

- **X** *(array-like)* – input data of shape (n_samples, m_features)
- **y** *(array-like)* – label data of shape (n_samples,)
- **weights** *(array-like shape (n_samples,), optional)* – sample weights
- **return_scores** *(boolean, optional)* – whether to return the hyperpamaters and score for each element in the grid
- **keep_best** *(boolean, optional)* – whether to keep the best GAM as self.
- **objective** *({'auto', 'AIC', 'AICc', 'GCV', 'UBRE'}, optional)* – Metric to optimize. If auto, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.
- **progress** *(bool, optional)* – whether to display a progress bar
- ****kwargs** – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.

If no parameter are specified, lam=np.logspace(-3, 3, 11) is used. This results in a 11 points, placed diagonally across lam space.

If grid is iterable of iterables of floats, the outer iterable must have length m_features. the cartesian product of the subgrids in the grid will be tested.

If grid is a 2d numpy array, each row of the array will be tested.

The method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

Returns

- if return_scores=True – model_scores: dict containing each fitted model as keys and corresponding objective scores as values
- else – self: ie possibly the newly fitted model
Examples

For a model with 4 terms, and where we expect 4 lam values, our search space for lam must have 4 dimensions.

We can search the space in 3 ways:

1. via cartesian product by specifying the grid as a list. our grid search will consider \(11^4\) points:

   ```
   >>> lam = np.logspace(-3, 3, 11)
   >>> lams = [lam] * 4
   >>> gam.gridsearch(X, y, lam=lams)
   ```

2. directly by specifying the grid as a np.ndarray. This is useful for when the dimensionality of the search space is very large, and we would prefer to execute a randomized search:

   ```
   >>> lams = np.exp(np.random.random(50, 4) * 6 - 3)
   >>> gam.gridsearch(X, y, lam=lams)
   ```

3. copying grids for parameters with multiple dimensions. if we specify a 1D np.ndarray for lam, we are implicitly testing the space where all points have the same value

   ```
   >>> gam.gridsearch(lam=np.logspace(-3, 3, 11))
   ```

   is equivalent to:

   ```
   >>> lam = np.logspace(-3, 3, 11)
   >>> lams = np.array([lam] * 4)
   >>> gam.gridsearch(X, y, lam=lams)
   ```

loglikelihood \((X, y, \text{weights}=\text{None})\)

compute the log-likelihood of the dataset using the current model

Parameters

- **X** *(array-like of shape \((n\text{-samples, m\_features})\)) – containing the input dataset*
- **y** *(array-like of shape \((n,)) – containing target values*
- **weights** *(array-like of shape \((n,)), optional) – containing sample weights*

Returns **log-likelihood** – containing log-likelihood scores

Return type **np.array of shape \((n,)\)**

partial_dependence \((\text{term, X=\text{None, width=\text{None, quantiles=\text{None, meshgrid=\text{False}}})}\)

Computes the term functions for the GAM and possibly their confidence intervals.

if both width=\text{None} and quantiles=\text{None}, then no confidence intervals are computed

Parameters

- **term** *(int, optional) – Term for which to compute the partial dependence functions.*
- **X** *(array-like with input data, optional) – if meshgrid=False, then X should be an array-like of shape \((n\_samples, m\_features).*
  
  if meshgrid=True, then X should be a tuple containing an array for each feature in the term.
if None, an equally spaced grid of points is generated.

• width (float on (0, 1), optional) – Width of the confidence interval.

• quantiles (array-like of floats on (0, 1), optional) – instead of specifying the prediction width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]. if None, defaults to width.

• meshgrid (bool, whether to return and accept meshgrids.) – Useful for creating outputs that are suitable for 3D plotting.

Note, for simple terms with no interactions, the output of this function will be the same for meshgrid=True and meshgrid=False, but the inputs will need to be different.

Returns

• pdeps (np.array of shape (n_samples,))

• conf_intervals (list of length len(term)) – containing np.arrays of shape (n_samples, 2 or len(quantiles))

Raises ValueError – If the term requested is an intercept since it does not make sense to process the intercept term.

See also:

generate_X_grid() for help creating meshgrids.

predict (X)

Predict expected value of target given model and input X often this is done via expected value of GAM given input X

Parameters X (array-like of shape (n_samples, m_features)) – containing the input dataset

Returns y – containing predicted values under the model

Return type np.array of shape (n_samples,)

predict_mu (X)

Predict expected value of target given model and input X

Parameters X (array-like of shape (n_samples, m_features)), – containing the input dataset

Returns y – containing expected values under the model

Return type np.array of shape (n_samples,)

sample (X, y, quantity='y', sample_at_X=None, weights=None, n_draws=100, n_bootstraps=5, objective='auto')

Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with gridsearch (with the parameter keep_best=True) before calling sample, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. n_bootstraps many “bootstrap samples” of the response (y) are simulated by drawing random samples from the model’s distribution evaluated at the expected values (mu) for each sample in X.
2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter $\text{lam}$ given the response data $y$.

3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.

**Notes**

A gridsearch is done $n_{\text{bootstraps}}$ many times, so keep $n_{\text{bootstraps}}$ small. Make $n_{\text{bootstraps}} < n_{\text{draws}}$ to take advantage of the expensive bootstrap samples of the smoothing parameters.

**Parameters**

- $X$ (array of shape (n_samples, m_features)) – empirical input data
- $y$ (array of shape (n_samples,)) – empirical response vector
- $\text{quantity}$ ('y', 'coef', 'mu'), default: 'y') – What quantity to return pseudorandom samples of. If $\text{sample}_\text{at}_X$ is not None and $\text{quantity}$ is either 'y' or 'mu', then samples are drawn at the values of $X$ specified in $\text{sample}_\text{at}_X$.
- $\text{sample}_\text{at}_X$ (array of shape (n_samples_to_simulate, m_features) or) –
  - optional (None,) – Input data at which to draw new samples.
    Only applies for $\text{quantity}$ equal to 'y' or 'mu'. If None, then $\text{sample}_\text{at}_X$ is replaced by $X$.
- $\text{weights}$ (np.array of shape (n_samples,)) – sample weights
- $n_{\text{draws}}$ (positive int, optional (default=100)) – The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters
- $n_{\text{bootstraps}}$ (positive int, optional (default=5)) – The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If $n_{\text{bootstraps}}$ is 1, then only the already fitted model's smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.
- $\text{objective}$ (string, optional (default='auto') – metric to optimize in grid search. must be in ['AIC', 'AICC', 'GCV', 'UBRE', 'auto'] if 'auto', then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

**Returns**

draws – Simulations of the given $\text{quantity}$ using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

If $\text{quantity} == \text{\textquoteleft\textquoteleft coef\textquoteright\textquoteright}$, then the number of columns of $\text{draws}$ is the number of coefficients (len(self.coef_)).

Otherwise, the number of columns of $\text{draws}$ is the number of rows of $\text{sample}_\text{at}_X$ if $\text{sample}_\text{at}_X$ is not None or else the number of rows of $X$.

**Return type** 2D array of length n_draws
References


score(X, y, weights=None)
method to compute the explained deviance for a trained model for a given X data and y labels

Parameters

• X (array-like) – Input data array of shape (n_samples, m_features)
• y (array-like) – Output data vector of shape (n_samples,)
• weights (array-like shape (n_samples,) or None, optional) – Sample weights. If None, defaults to array of ones

Returns explained deviance score

Return type np.array() (n_samples,)

set_params(deep=False, force=False, **parameters)
sets an object’s parameters

Parameters

• deep (boolean, default: False) – when True, also sets non-user-facing parameters
• force (boolean, default: False) – when True, also sets parameters that the object does not already have
• **parameters (parameters to set) –

Returns

Return type self

summary()
produce a summary of the model statistics

Parameters None –

Returns

Return type None

InvGaussGAM
class pygam.pygam.InvGaussGAM(terms='auto', max_iter=100, tol=0.0001, scale=None, call-backs=['deviance', 'difffs'], fit_intercept=True, verbose=False, **kwargs)

Bases: pygam.pygam.GAM

Inverse Gaussian GAM

This is a GAM with a Inverse Gaussian error distribution, and a log link.

NB Although canonical link function for the Inverse Gaussian GLM is the inverse squared link, this function can create problems for numerical software because it becomes difficult to enforce the requirement that the mean of the Inverse Gaussian distribution be positive. The log link guarantees this.

If you need to use the inverse squared link function, simply construct a custom GAM:
```python
>>> from pygam import GAM
>>> gam = GAM(distribution='inv_gauss', link='inv_squared')
```

### Parameters

- **terms** *(expression specifying terms to model, optional)* – By default a univariate spline term will be allocated for each feature.

  For example:

  ```python
  >>> GAM(s(0) + l(1) + f(2) + te(3, 4))
  ```

  will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

- **callbacks** *(list of str or list of CallBack objects, optional)* – Names of callback objects to call during the optimization loop.

- **fit_intercept** *(bool, optional)* – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.

- **max_iter** *(int, optional)* – Maximum number of iterations allowed for the solver to converge.

- **tol** *(float, optional)* – Tolerance for stopping criteria.

- **verbose** *(bool, optional)* – Whether to show pyGAM warnings.

**coef_**

Coefficient of the features in the decision function. If `fit_intercept` is True, then `self.coef_[0]` will contain the bias.

Type array, shape (n_classes, m_features)

**statistics_**

Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

Type dict

**logs_**

Dictionary containing the outputs of any callbacks at each optimization loop.

The logs are structured as `{callback: [...]}`

Type dict

### References


**confidence_intervals** *(X, width=0.95, quantiles=None)*

Estimate confidence intervals for the model.

**Parameters**
**X** (array-like of shape (n_samples, m_features)) – Input data matrix

**width** (float on [0,1], optional) –

**quantiles** (array-like of floats in (0, 1), optional) – Instead of specifying the prediction width, one can specify the quantiles. So width=.95 is equivalent to quantiles=[.025, .975]

**Returns intervals**

Return type np.array of shape (n_samples, 2 or len(quantiles))

### Notes

**Wood 2006, section 4.9** Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.

device_resids (X, y, weights=None, scaled=False)
method to compute the deviance residuals of the model
these are analogous to the residuals of an OLS.

**Parameters**

- **X** (array-like) – Input data array of shape (n_samples, m_features)
- **y** (array-like) – Output data vector of shape (n_samples,)
- **weights** (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones
- **scaled** (bool, optional) – whether to scale the deviance by the (estimated) distribution scale

**Returns device_resids** – with shape (n_samples,)

Return type np.array

**fit** (X, y, weights=None)
Fit the generalized additive model.

**Parameters**

- **X** (array-like, shape (n_samples, m_features)) – Training vectors.
- **y** (array-like, shape (n_samples,)) – Target values, i.e. integers in classification, real numbers in regression
- **weights** (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones

**Returns self** – Returns fitted GAM object

Return type object

generate_X_grid (term, n=100, meshgrid=False)
create a nice grid of X data
array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong if term is >= 0, we generate n samples per feature, which results in n^deg samples, where deg is the degree of the interaction of the term

**Parameters**
• **term** (*int,*) – Which term to process.
• **n** (*int, optional*) – Number of data points to create
• **meshgrid** (*bool, optional*) – Whether to return a meshgrid (useful for 3d plotting) or a feature matrix (useful for inference like partial predictions)

**Returns**

- **if meshgrid is False** – np.array of shape (n, n_features) where m is the number of (sub)terms in the requested (tensor)term.
- **else** – tuple of len m, where m is the number of (sub)terms in the requested (tensor)term.
  each element in the tuple contains a np.ndarray of size (n)^m

**Raises** *ValueError* – If the term requested is an intercept since it does not make sense to process the intercept term.

**get_params** (*deep=False*)
returns a dict of all of the object’s user-facing parameters

**Parameters**

- **deep** (*boolean, default: False*) – when True, also gets non-user-facing parameters

**Returns**

**Return type** *dict*

**gridsearch** (*X, y, weights=None, return_scores=False, keep_best=True, objective='auto', progress=True, **param_grids*)
Performs a grid search over a space of parameters for a given objective

**Warning:** *gridsearch* is lazy and will not remove useless combinations from the search space, eg.

```python
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```

will result in 10 loops, of which 5 are equivalent because `fit_splines = False`

Also, it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

**Parameters**

- **X** (*array-like*) – input data of shape (n_samples, m_features)
- **y** (*array-like*) – label data of shape (n_samples,)
- **weights** (*array-like shape (n_samples,), optional*) – sample weights
- **return_scores** (*boolean, optional*) – whether to return the hyperpamaters and score for each element in the grid
- **keep_best** (*boolean, optional*) – whether to keep the best GAM as self.
- **objective** (*{'auto', 'AIC', 'AICc', 'GCV', 'UBRE'}, optional*) – Metric to optimize. If *auto*, then grid search will optimize *GCV* for models with unknown scale and *UBRE* for models with known scale.
- **progress** (*bool, optional*) – whether to display a progress bar
- ****kwargs – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.
If no parameter are specified, `lam=np.logspace(-3, 3, 11)` is used. This results in a 11 points, placed diagonally across lam space.

If grid is iterable of iterables of floats, the outer iterable must have length `m_features`. the cartesian product of the subgrids in the grid will be tested.

If grid is a 2d numpy array, each row of the array will be tested.

The method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

**Returns**

- if `return_scores=True` – `model_scores`: dict containing each fitted model as keys and corresponding objective scores as values
- else – `self`: ie possibly the newly fitted model

**Examples**

For a model with 4 terms, and where we expect 4 lam values, our search space for lam must have 4 dimensions.

We can search the space in 3 ways:

1. via cartesian product by specifying the grid as a list. our grid search will consider $11^{**4}$ points:

   ```python
   >>> lam = np.logspace(-3, 3, 11)
   >>> lams = [lam] * 4
   >>> gam.gridsearch(X, y, lam=lams)
   ```

2. directly by specifying the grid as a np.ndarray. This is useful for when the dimensionality of the search space is very large, and we would prefer to execute a randomized search:

   ```python
   >>> lams = np.exp(np.random.random(50, 4) * 6 - 3)
   >>> gam.gridsearch(X, y, lam=lams)
   ```

3. copying grids for parameters with multiple dimensions. if we specify a 1D np.ndarray for lam, we are implicitly testing the space where all points have the same value

   ```python
   >>> gam.gridsearch(lam=np.logspace(-3, 3, 11))
   ```

   is equivalent to:

   ```python
   >>> lam = np.logspace(-3, 3, 11)
   >>> lams = np.array([lam] * 4)
   >>> gam.gridsearch(X, y, lam=lams)
   ```

**loglikelihood** *(X, y, weights=None)*

compute the log-likelihood of the dataset using the current model

**Parameters**

- `X` *(array-like of shape (n_samples, m_features)) – containing the input dataset*
- `y` *(array-like of shape (n,)) – containing target values*
- `weights` *(array-like of shape (n,), optional) – containing sample weights*
Returns log-likelihood – containing log-likelihood scores

Return type np.array of shape (n,)

partial_dependence(term, X=None, width=None, quantiles=None, meshgrid=False)
Computes the term functions for the GAM and possibly their confidence intervals.
if both width=None and quantiles=None, then no confidence intervals are computed

Parameters

- **term** (int, optional) – Term for which to compute the partial dependence functions.
- **X** (array-like with input data, optional) – if meshgrid=False, then X should be an array-like of shape (n_samples, m_features).
  if meshgrid=True, then X should be a tuple containing an array for each feature in the term.
  if None, an equally spaced grid of points is generated.
- **width** (float on (0, 1), optional) – Width of the confidence interval.
- **quantiles** (array-like of floats on (0, 1), optional) – instead of specifying the prediction width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]. if None, defaults to width.
- **meshgrid** (bool, whether to return and accept meshgrids.) – Useful for creating outputs that are suitable for 3D plotting.

Note, for simple terms with no interactions, the output of this function will be the same for meshgrid=True and meshgrid=False, but the inputs will need to be different.

Returns

- **pdeps** (np.array of shape (n_samples,))
- **conf_intervals** (list of length len(term)) – containing np.arrays of shape (n_samples, 2 or len(quantiles))

Raises ValueError: – If the term requested is an intercept since it does not make sense to process the intercept term.

See also:

generate_X_grid() for help creating meshgrids.

predict(X)
predict expected value of target given model and input X often this is done via expected value of GAM given input X

Parameters X (array-like of shape (n_samples, m_features)) – containing the input dataset

Returns y – containing predicted values under the model

Return type np.array of shape (n_samples,)

predict_mu(X)
predict expected value of target given model and input X

Parameters X (array-like of shape (n_samples, m_features),) – containing the input dataset
Returns \( y \) – containing expected values under the model

Return type np.array of shape (n_samples,)

**sample** \((X, y, quantity='y', sample_at_X=None, weights=None, n_draws=100, n_bootstraps=5, objective='auto')\)

Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with gridsearch (with the parameter keep_best=True) before calling **sample**, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. \( n_{\text{bootstraps}} \) many “bootstrap samples” of the response \((y)\) are simulated by drawing random samples from the model’s distribution evaluated at the expected values \((\mu)\) for each sample in \(X\).
2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter \(\lambda\) given the response data \(y\).
3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.

**Notes**

A gridsearch is done \( n_{\text{bootstraps}} \) many times, so keep \( n_{\text{bootstraps}} \) small. Make \( n_{\text{bootstraps}} < n_{\text{draws}} \) to take advantage of the expensive bootstrap samples of the smoothing parameters.

**Parameters**

- **X** (array of shape \((n_{\text{samples}}, m_{\text{features}})\)) – empirical input data
- **y** (array of shape \((n_{\text{samples}},)\)) – empirical response vector
- **quantity** (\([''y'', 'coef', 'mu'], default: 'y')\) – What quantity to return pseudorandom samples of. If \(\text{sample}_at_{\text{X}}\) is not None and \(\text{quantity}\) is either ‘y’ or ‘mu’, then samples are drawn at the values of \(X\) specified in \(\text{sample}_at_{\text{X}}\).
- **sample_at_X** (array of shape \((n_{\text{samples_to_simulate}}, m_{\text{features}})\)) – Input data at which to draw new samples. Only applies for \(\text{quantity}\) equal to ‘y’ or to ‘mu’. If None, then \(\text{sample}_at_{\text{X}}\) is replaced by \(X\).
- **weights** (np.array of shape \((n_{\text{samples}},)\)) – sample weights
- **n_draws** (positive int, optional (default=100)) – The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters
- **n_bootstraps** (positive int, optional (default=5)) – The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If \(n_{\text{bootstraps}}\) is 1, then only the already fitted model’s smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.
**objective** (string, optional (default='auto') – metric to optimize in grid search. must be in ['AIC', 'AICc', 'GCV', 'UBRE', 'auto'] if `auto' then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

Returns

draws – Simulations of the given quantity using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

If quantity == 'coef', then the number of columns of draws is the number of coefficients (len(self.coef_)).

Otherwise, the number of columns of draws is the number of rows of sample_at_X if sample_at_X is not None or else the number of rows of X.

Return type 2D array of length n_draws

References


`score` (X, y, weights=None)
method to compute the explained deviance for a trained model for a given X data and y labels

Parameters

• X (array-like) – Input data array of shape (n_samples, m_features)

• y (array-like) – Output data vector of shape (n_samples,)

• weights (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones

Returns explained deviance score

Return type np.array() (n_samples,)

`set_params` (deep=False, force=False, **parameters)
sets an object’s parameters

Parameters

• deep (boolean, default: False) – when True, also sets non-user-facing parameters

• force (boolean, default: False) – when True, also sets parameters that the object does not already have

• **parameters (parameters to set)–

Returns

Return type self

`summary` ()
produce a summary of the model statistics

Parameters None –

Returns

Return type None
LogisticGAM

class pygam.pygam.LogisticGAM(terms='auto', max_iter=100, tol=0.0001, callbacks=['deviance', 'diffs', 'accuracy'], fit_intercept=True, verbose=False, **kwargs)

Bases: pygam.pygam.GAM

Logistic GAM

This is a GAM with a Binomial error distribution, and a logit link.

Parameters

- **terms** *(expression specifying terms to model, optional.)* – By default a univariate spline term will be allocated for each feature.

  For example:

  >>> GAM(s(0) + l(1) + f(2) + te(3, 4))

  will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

- **callbacks** *(list of str or list of CallBack objects, optional)* – Names of callback objects to call during the optimization loop.

- **fit_intercept** *(bool, optional)* – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.

- **max_iter** *(int, optional)* – Maximum number of iterations allowed for the solver to converge.

- **tol** *(float, optional)* – Tolerance for stopping criteria.

- **verbose** *(bool, optional)* – whether to show pyGAM warnings.

**coef_**

Coefficient of the features in the decision function. If fit_intercept is True, then self.coef_[0] will contain the bias.

  Type array, shape (n_classes, m_features)

**statistics_**

Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

  Type dict

**logs_**

Dictionary containing the outputs of any callbacks at each optimization loop.

  The logs are structured as {callback: [...]}

  Type dict

References


6.3. User API

**accuracy** \((X=None, y=None, mu=None)\)

computes the accuracy of the LogisticGAM

**Parameters**

- **note** \((X \text{ or } mu \text{ must be defined. defaults to } mu)\)
- **X** \((\text{array-like of shape } (n\_samples, m\_features), \text{ optional} (\text{default}=\text{None}))\) – containing input data
- **y** \((\text{array-like of shape } (n,))\) – containing target data
- **mu** \((\text{array-like of shape } (n\_samples,)), \text{ optional} (\text{default}=\text{None})\) – expected value of the targets given the model and inputs

**Returns**

**Return type** float in \([0, 1]\)

**confidnce_intervals** \((X, \text{width}=0.95, \text{quantiles}=\text{None})\)

estimate confidence intervals for the model.

**Parameters**

- **X** \((\text{array-like of shape } (n\_samples, m\_features))\) – Input data matrix
- **width** \((\text{float on } [0,1], \text{ optional})\) –
- **quantiles** \((\text{array-like of floats in } (0, 1), \text{ optional})\) – Instead of specifying the prediction width, one can specify the quantiles. So \(\text{width}=.95\) is equivalent to \(\text{quantiles}=[.025, .975]\)

**Returns** intervals

**Return type** np.array of shape \((n\_samples, 2 \text{ or len(quantiles)})\)

**Notes**

**Wood 2006, section 4.9** Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.

**deviance_residuals** \((X, y, \text{weights}=\text{None}, \text{scaled}=\text{False})\)

method to compute the deviance residuals of the model

these are analogous to the residuals of an OLS.

**Parameters**

- **X** \((\text{array-like})\) – Input data array of shape \((n\_samples, m\_features)\)
- **y** \((\text{array-like})\) – Output data vector of shape \((n\_samples,)\)
- **weights** \((\text{array-like shape } (n\_samples,)) \text{ or } \text{None, optional})\) – Sample weights. if None, defaults to array of ones
- **scaled** \((\text{bool, optional})\) – whether to scale the deviance by the (estimated) distribution scale

**Returns** deviance_residuals – with shape \((n\_samples,)\)

**Return type** np.array
**fit** *(X, y, weights=None)*

Fit the generalized additive model.

**Parameters**

- **X** *(array-like, shape (n_samples, m_features)) –* Training vectors.
- **y** *(array-like, shape (n_samples,)) –* Target values, i.e., integers in classification, real numbers in regression.
- **weights** *(array-like shape (n_samples,) or None, optional) –* Sample weights. If None, defaults to array of ones.

**Returns**

self – Returns fitted GAM object

**Return type**

object

**generate_X_grid** *(term, n=100, meshgrid=False)*

Create a nice grid of X data.

- array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong.
- if term is >= 0, we generate n samples per feature, which results in n^deg samples, where deg is the degree of the interaction of the term.

**Parameters**

- **term** *(int,) –* Which term to process.
- **n** *(int, optional) –* number of data points to create.
- **meshgrid** *(bool, optional) –* Whether to return a meshgrid (useful for 3d plotting) or a feature matrix (useful for inference like partial predictions).

**Returns**

- if meshgrid is False – np.array of shape (n, n_features) where m is the number of (sub)terms in the requested (tensor)term.
- else – tuple of len m, where m is the number of (sub)terms in the requested (tensor)term. Each element in the tuple contains a np.ndarray of size (n)^m.

**Raises**

ValueError: – If the term requested is an intercept since it does not make sense to process the intercept term.

**get_params** *(deep=False)*

Returns a dict of all of the object’s user-facing parameters.

**Parameters**

deep *(boolean, default: False) –* When True, also gets non-user-facing parameters.

**Returns**

**Return type**

dict

**gridsearch** *(X, y, weights=None, return_scores=False, keep_best=True, objective='auto', progress=True, **param_grids)*

Performs a grid search over a space of parameters for a given objective.

**Warning:** gridsearch is lazy and will not remove useless combinations from the search space, eg.

```python
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```
will result in 10 loops, of which 5 are equivalent because `fit_splines = False`.

Also, it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

**Parameters**

- **X** (*array-like*) – input data of shape (n_samples, m_features)
- **y** (*array-like*) – label data of shape (n_samples,)
- **weights** (*array-like shape (n_samples,), optional*) – sample weights
- **return_scores** (*boolean, optional*) – whether to return the hyperparameters and score for each element in the grid
- **keep_best** (*boolean, optional*) – whether to keep the best GAM as self.
- **objective** (*{'auto', 'AIC', 'AICc', 'GCV', 'UBRE'}, optional*) – Metric to optimize. If `auto`, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.
- **progress** (*bool, optional*) – whether to display a progress bar
- ****kwargs** – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.

If no parameter are specified, lam=np.logspace(-3, 3, 11) is used. This results in a 11 points, placed diagonally across lam space.

If grid is iterable of iterables of floats, the outer iterable must have length m_features.

The cartesian product of the subgrids in the grid will be tested.

If grid is a 2d numpy array, each row of the array will be tested.

The method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

**Returns**

- if `return_scores=True` – model_scores: dict containing each fitted model as keys and corresponding objective scores as values
- else – self: ie possibly the newly fitted model

**Examples**

For a model with 4 terms, and where we expect 4 lam values, our search space for lam must have 4 dimensions.

We can search the space in 3 ways:

1. via cartesian product by specifying the grid as a list. our grid search will consider $11^4$ points:

   ```
   >>> lam = np.logspace(-3, 3, 11)
   >>> lams = [lam] * 4
   >>> gam.gridsearch(X, y, lam=lams)
   ```

2. directly by specifying the grid as a np.ndarray. This is useful for when the dimensionality of the search space is very large, and we would prefer to execute a randomized search:
```python
>>> lams = np.exp(np.random.random(50, 4) * 6 - 3)
>>> gam.gridsearch(X, y, lam=lams)
```

3. copying grids for parameters with multiple dimensions. if we specify a 1D np.ndarray for lam, we are implicitly testing the space where all points have the same value

```python
>>> gam.gridsearch(lam=np.logspace(-3, 3, 11))
```

is equivalent to:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = np.array([lam] * 4)
>>> gam.gridsearch(X, y, lam=lams)
```

loglikelihood(X, y, weights=None)

- compute the log-likelihood of the dataset using the current model

**Parameters**

- X (array-like of shape (n_samples, m_features)) – containing the input dataset
- y (array-like of shape (n,)) – containing target values
- weights (array-like of shape (n,), optional) – containing sample weights

**Returns**

- log-likelihood – containing log-likelihood scores

**Return type**

np.array of shape (n,)

partial_dependence(term, X=None, width=None, quantiles=None, meshgrid=False)

- Computes the term functions for the GAM and possibly their confidence intervals.

if both width=None and quantiles=None, then no confidence intervals are computed

**Parameters**

- term (int, optional) – Term for which to compute the partial dependence functions.
- X (array-like with input data, optional) – if meshgrid=False, then X should be an array-like of shape (n_samples, m_features).
  
  if meshgrid=True, then X should be a tuple containing an array for each feature in the term.

  if None, an equally spaced grid of points is generated.
- width (float on (0, 1), optional) – Width of the confidence interval.
- quantiles (array-like of floats on (0, 1), optional) – instead of specifying the predicton width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]. if None, defaults to width.
- meshgrid (bool, whether to return and accept meshgrids.) – Useful for creating outputs that are suitable for 3D plotting.

**Note**

- for simple terms with no interactions, the output of this function will be the same for meshgrid=True and meshgrid=False, but the inputs will need to be different.

**Returns**

- pdeps (np.array of shape (n_samples,))
• **conf_intervals** *(list of length \( \text{len(term)} \)) – containing np.arrays of shape \((\text{n_samples}, 2 \text{ or } \text{len(quantiles)})\)*

**Raises** *ValueError* : – If the term requested is an intercept since it does not make sense to process the intercept term.

See also:

*`generate_X_grid()`* for help creating meshgrids.

**predict** *(\(X\))*

Predict binary targets given model and input \(X\)

**Parameters**

\(X\) *(array-like of shape \((\text{n_samples}, m_{features}), \text{optional } (default=\text{None})\)) – containing the input dataset*

**Returns**

\(y\) – containing binary targets under the model

**Return type**

np.array of shape \((\text{n_samples},)\)

**predict_mu** *(\(X\))*

Predict expected value of target given model and input \(X\)

**Parameters**

\(X\) *(array-like of shape \((\text{n_samples}, m_{features}), \text{containing the input dataset}\)) – containing*

**Returns**

\(y\) – containing expected values under the model

**Return type**

np.array of shape \((\text{n_samples},)\)

**predict_proba** *(\(X\))*

Predict targets given model and input \(X\)

**Parameters**

\(X\) *(array-like of shape \((\text{n_samples}, m_{features}), \text{optional } (default=\text{None})\)) – containing the input dataset*

**Returns**

\(y\) – containing expected values under the model

**Return type**

np.array of shape \((\text{n_samples},)\)

**sample** *(\(X, y, \text{quantity=’y’, sample_at_X=’None’, weights=’None’, n_draws=100, n_bootstraps=5, objective=’auto’}’))*

Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with *gridsearch* (with the parameter *keep_best=True*) before calling *sample*, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. \(n_{bootstraps}\) many “bootstrap samples” of the response \((y)\) are simulated by drawing random samples from the model’s distribution evaluated at the expected values \((\mu)\) for each sample in \(X\).

2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter \(\lambda_m\) given the response data \(y\).

3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.
Notes

A gridsearch is done \( n_{\text{bootstraps}} \) many times, so keep \( n_{\text{bootstraps}} \) small. Make \( n_{\text{bootstraps}} < n_{\text{draws}} \) to take advantage of the expensive bootstrap samples of the smoothing parameters.

Parameters

- \( X \) (array of shape (\( n_{\text{samples}}, m_{\text{features}} \))): empirical input data
- \( y \) (array of shape (\( n_{\text{samples}}, \))): empirical response vector
- \( \text{quantity} \) (\{‘y’, ‘coef’, ‘mu’\}, default: ‘y’): What quantity to return pseudorandom samples of. If \( \text{sample}_\text{at}_X \) is not None and \( \text{quantity} \) is either ‘y’ or ‘mu’, then samples are drawn at the values of \( X \) specified in \( \text{sample}_\text{at}_X \).
- \( \text{sample}_\text{at}_X \) (array of shape (\( n_{\text{samples}_\text{to}_\text{simulate}}, m_{\text{features}} \)) or): optional (None): Input data at which to draw new samples. Only applies for \( \text{quantity} \) equal to ‘y’ or to ‘mu’. If None, then \( \text{sample}_\text{at}_X \) is replaced by \( X \).
- \( \text{weights} \) (np.array of shape (\( n_{\text{samples}}, \))): sample weights
- \( n_{\text{draws}} \) (positive int, optional (default=100)): The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters
- \( n_{\text{bootstraps}} \) (positive int, optional (default=5)): The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If \( n_{\text{bootstraps}} \) is 1, then only the already fitted model’s smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.
- \( \text{objective} \) (string, optional (default=’auto’)): metric to optimize in grid search. must be in [‘AIC’, ‘AICc’, ‘GCV’, ‘UBRE’, ‘auto’] if ‘auto’, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

Returns

- \( \text{draws} \): Simulations of the given \( \text{quantity} \) using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

If \( \text{quantity} == \text{‘coef’} \), then the number of columns of \( \text{draws} \) is the number of coefficients (len(self.coef_)).

Otherwise, the number of columns of \( \text{draws} \) is the number of rows of \( \text{sample}_\text{at}_X \) if \( \text{sample}_\text{at}_X \) is not None or else the number of rows of \( X \).

Return type 2D array of length \( n_{\text{draws}} \)

References


\( \text{score} \) (\( X, y \))
method to compute the accuracy for a trained model for a given \( X \) data and \( y \) labels
Parameters

- \(X\) (array-like) – Input data array of shape \((n_{\text{samples}}, m_{\text{features}})\)
- \(y\) (array-like) – Output data vector of shape \((n_{\text{samples}},)\)

Returns accuracy score

Return type np.array() \((n_{\text{samples}},)\)

\textbf{set_params} (\textit{deep=False}, \textit{force=False}, **\textit{parameters})

sets an object’s parameters

Parameters

- \textit{deep} (boolean, default: \textit{False}) – when True, also sets non-user-facing parameters
- \textit{force} (boolean, default: \textit{False}) – when True, also sets parameters that the object does not already have
- **\textit{parameters} (parameters to set)

Returns

Return type self

\textbf{summary}()

produce a summary of the model statistics

Parameters None –

Returns

Return type None

\textbf{PoissonGAM}

class \texttt{pygam.pygam.PoissonGAM}(\textit{terms='auto', max_iter=100, tol=0.0001, callbacks=['deviance', 'diffs'], fit_intercept=True, verbose=False, **\textit{kwargs}})

Bases: \texttt{pygam.pygam.GAM}

Poisson GAM

This is a GAM with a Poisson error distribution, and a log link.

Parameters

- \textit{terms} (expression specifying terms to model, optional.) – By default a univariate spline term will be allocated for each feature.

For example:

```
>>> GAM(s(0) + l(1) + f(2) + te(3, 4))
```

will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

- \textit{callbacks} (list of str or list of CallBack objects, optional) – Names of callback objects to call during the optimization loop.

- \textit{fit_intercept} (bool, optional) – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.
• **max_iter** (*int, optional*) – Maximum number of iterations allowed for the solver to converge.

• **tol** (*float, optional*) – Tolerance for stopping criteria.

• **verbose** (*bool, optional*) – Whether to show pyGAM warnings.

**coef_**

Coefficient of the features in the decision function. If fit_intercept is True, then self.coef_[0] will contain the bias.

Type array, shape (n_classes, m_features)

**statistics_**

Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

Type dict

**logs_**

Dictionary containing the outputs of any callbacks at each optimization loop.

The logs are structured as {callback: [...]}

Type dict

**References**


**confidence_intervals** *(X, width=0.95, quantiles=None)*

Estimate confidence intervals for the model.

**Parameters**

• **X** (*array-like of shape (n_samples, m_features]*) – Input data matrix

• **width** (*float on [0,1], optional*) –

• **quantiles** (*array-like of floats in (0, 1), optional*) – Instead of specifying the prediction width, one can specify the quantiles. So width=.95 is equivalent to quantiles=[.025, .975]

**Returns** intervals

Return type np.array of shape (n_samples, 2 or len(quantiles))

**Notes**

Wood 2006, section 4.9 Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.
deviance_residuals($X$, $y$, weights=None, scaled=False)
method to compute the deviance residuals of the model
these are analogous to the residuals of an OLS.

Parameters

- $X$ (array-like) – Input data array of shape (n_samples, m_features)
- $y$ (array-like) – Output data vector of shape (n_samples,)
- weights (array-like shape (n_samples,) or None, optional) – Sample weights. if None, defaults to array of ones
- scaled (bool, optional) – whether to scale the deviance by the (estimated) distribution scale

Returns deviance_residuals – with shape (n_samples,)
Return type np.array

fit($X$, $y$, exposure=None, weights=None)
Fit the generalized additive model.

Parameters

- $X$ (array-like, shape (n_samples, m_features)) – Training vectors, where n_samples is the number of samples and m_features is the number of features.
- $y$ (array-like, shape (n_samples,)) – Target values (integers in classification, real numbers in regression) For classification, labels must correspond to classes.
- exposure (array-like shape (n_samples,) or None, default: None) – containing exposures if None, defaults to array of ones
- weights (array-like shape (n_samples,) or None, default: None) – containing sample weights if None, defaults to array of ones

Returns self – Returns fitted GAM object
Return type object

generate_X_grid(term, n=100, meshgrid=False)
create a nice grid of X data
array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong
if term is >= 0, we generate n samples per feature, which results in n^deg samples, where deg is the degree of the interaction of the term

Parameters

- term (int,) – Which term to process.
- n (int, optional) – number of data points to create
- meshgrid (bool, optional) – Whether to return a meshgrid (useful for 3d plotting) or a feature matrix (useful for inference like partial predictions)

Returns

- if meshgrid is False – np.array of shape (n, n_features) where m is the number of (sub)terms in the requested (tensor)term.
- else – tuple of len m, where m is the number of (sub)terms in the requested (tensor)term. each element in the tuple contains a np.ndarray of size (n)^m
**pyGAM Documentation**

**Raises**  
*ValueError:* If the term requested is an intercept since it does not make sense to process the intercept term.

**get_params** *(deep=False)*  
returns a dict of all of the object’s user-facing parameters

**Parameters**  
- **deep** *(boolean, default: False)* – when True, also gets non-user-facing parameters

**Returns**

**Return type**  
dict

**gridsearch** *(X, y, exposure=None, weights=None, return_scores=False, keep_best=True, objective='auto', **param_grids)*  
performs a grid search over a space of parameters for a given objective  
NOTE: gridsearch method is lazy and will not remove useless combinations from the search space, eg.

```python
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```

will result in 10 loops, of which 5 are equivalent because even though fit_splines==False  
it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

**Parameters**

- **X** *(array)* – input data of shape *(n_samples, m_features)*
- **y** *(array)* – label data of shape *(n_samples,)*
- **exposure** *(array-like shape *(n_samples,)* or None, default: None)* – containing exposures if None, defaults to array of ones
- **weights** *(array-like shape *(n_samples,)* or None, default: None)* – containing sample weights if None, defaults to array of ones
- **return_scores** *(boolean, default False)* – whether to return the hyperparameters and score for each element in the grid
- **keep_best** *(boolean)* – whether to keep the best GAM as self. default: True
- **objective** *(string, default: 'auto')* – metric to optimize. must be in ['AIC', 'AICc', 'GCV', 'UBRE', 'auto'] if ‘auto’, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.
- **,**kwargs *(dict, default {'lam': np.logspace(-3, 3, 11)})* – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.

if iterable of iterables of floats, the outer iterable must have length m_features.

the method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

**Returns**

- **if return_values == True** –

  **model_scores** *(dict)* Contains each fitted model as keys and corresponding objective scores as values

- **else** – self, ie possibly the newly fitted model

**loglikelihood** *(X, y, exposure=None, weights=None)*  
compute the log-likelihood of the dataset using the current model
Parameters

- **X** (array-like of shape \((n_{\text{samples}}, m_{\text{features}})\)) – containing the input dataset
- **y** (array-like of shape \((n,)\)) – containing target values
- **exposure** (array-like shape \((n_{\text{samples}},)\) or None, default: None) – containing exposures if None, defaults to array of ones
- **weights** (array-like of shape \((n,)\)) – containing sample weights

Returns **log-likelihood** – containing log-likelihood scores

Return type **np.array of shape \((n,)\)**

**partial_dependence** *(term, X=None, width=None, quantiles=None, meshgrid=False)*

Computes the term functions for the GAM and possibly their confidence intervals.

if both width=None and quantiles=None, then no confidence intervals are computed

Parameters

- **term** (int, optional) – Term for which to compute the partial dependence functions.
- **X** (array-like with input data, optional) – if meshgrid=False, then \(X\) should be an array-like of shape \((n_{\text{samples}}, m_{\text{features}})\).
  
  if meshgrid=True, then \(X\) should be a tuple containing an array for each feature in the term.

  if None, an equally spaced grid of points is generated.
- **width** (float on \((0, 1)\), optional) – Width of the confidence interval.
- **quantiles** (array-like of floats on \((0, 1)\), optional) – instead of specifying the prediction width, one can specify the quantiles. so width=.95 is equivalent to quantiles=[.025, .975]. if None, defaults to width.
- **meshgrid** (bool, whether to return and accept meshgrids.) – Useful for creating outputs that are suitable for 3D plotting.

Note, for simple terms with no interactions, the output of this function will be the same for meshgrid=True and meshgrid=False, but the inputs will need to be different.

Returns

- **pdeps** (np.array of shape \((n_{\text{samples}},)\))
- **conf_intervals** (list of length \(len(term)\)) – containing np.arrays of shape \((n_{\text{samples}}, 2\) or \(len(quantiles)\))

Raises **ValueError** : – If the term requested is an intercept since it does not make sense to process the intercept term.

See also:

**generate_X_grid()** for help creating meshgrids.

**predict** *(X, exposure=None)*

predict expected value of target given model and input X often this is done via expected value of GAM given input X

Parameters
• **X** (array-like of shape (n_samples, m_features), default: None) – containing the input dataset

• **exposure** (array-like shape (n_samples,) or None, default: None) – containing exposures if None, defaults to array of ones

Returns **y** – containing predicted values under the model

Return type np.array of shape (n_samples,)

**predict_mu** (**X**)  
Predict expected value of target given model and input **X**

Parameters **X** (array-like of shape (n_samples, m_features),) – containing the input dataset

Returns **y** – containing expected values under the model

Return type np.array of shape (n_samples,)

**sample** (**X**, **y**, **quantity**='y', **sample_at_X**=None, **weights**=None, **n_draws**=100, **n_bootstraps**=5, **objective**='auto')  
Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with gridsearch (with the parameter keep_best=True) before calling sample, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. **n_bootstraps** many “bootstrap samples” of the response (**y**) are simulated by drawing random samples from the model’s distribution evaluated at the expected values (**mu**) for each sample in **X**.

2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter **lam** given the response data **y**.

3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.

**Notes**

A gridsearch is done **n_bootstraps** many times, so keep **n_bootstraps** small. Make **n_bootstraps** < **n_draws** to take advantage of the expensive bootstrap samples of the smoothing parameters.

Parameters

• **X** (array of shape (n_samples, m_features)) – empirical input data

• **y** (array of shape (n_samples,)) – empirical response vector

• **quantity** ((['y', 'coef', 'mu'], default: 'y') – What quantity to return pseudorandom samples of. If **sample_at_X** is not None and **quantity** is either 'y' or 'mu', then samples are drawn at the values of **X** specified in **sample_at_X**.

• **sample_at_X** (array of shape (n_samples_to_simulate, m_features) or) –
• **optional** (*None*) – Input data at which to draw new samples.

Only applies for `quantity` equal to ‘y’ or to ‘mu’. If *None*, then `sample_at_X` is replaced by `X`.

• **weights** ([`np.array` of shape `(n_samples,)`]) – sample weights

• **n_draws** ([`positive int`, `optional` (default=100)]) – The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters

• **n_bootstraps** ([`positive int`, `optional` (default=5)]) – The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If `n_bootstraps` is 1, then only the already fitted model’s smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.

• **objective** ([`string`, `optional` (default='auto')] – metric to optimize in grid search. must be in ['AIC', 'AICc', 'GCV', 'UBRE', 'auto'] if ‘auto’, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

**Returns**

`draws` – Simulations of the given `quantity` using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

If `quantity == 'coef'`, then the number of columns of `draws` is the number of coefficients (`len(self.coef_)`).

Otherwise, the number of columns of `draws` is the number of rows of `sample_at_X` if `sample_at_X` is not `None` or else the number of rows of `X`.

**Return type** 2D array of length `n_draws`

**References**


**score** (*X*, *y*, `weights=None`)  
method to compute the explained deviance for a trained model for a given X data and y labels

**Parameters**

• **X** ([`array-like`]) – Input data array of shape (n_samples, m_features)

• **y** ([`array-like`]) – Output data vector of shape (n_samples,)

• **weights** ([`array-like shape (n_samples,) or None, optional`]) – Sample weights. if *None*, defaults to array of ones

**Returns** explained deviance score

**Return type** np.array() (n_samples,)

**set_params** (**deep=False, force=False, **parameters**)  
sets an object’s parameters

**Parameters**

• **deep** (boolean, default: False) – when True, also sets non-user-facing parameters
• **force** *(boolean, default: False)* – when True, also sets parameters that the object does not already have

• **parameters** *(parameters to set)* –

Returns

Return type: self

**summary()**

produce a summary of the model statistics

Parameters: None –

Returns

Return type: None

### ExpectileGAM

```python
from pygam import ExpectileGAM
from pygam.datasets import mcycle

X, y = mcycle(return_X_y=True)

# lets fit the mean model first by CV
gam50 = ExpectileGAM(expectile=0.5).gridsearch(X, y)

# and copy the smoothing to the other models
lam = gam50.lam

# now fit a few more models
gam95 = ExpectileGAM(expectile=0.95, lam=lam).fit(X, y)
gam75 = ExpectileGAM(expectile=0.75, lam=lam).fit(X, y)
gam25 = ExpectileGAM(expectile=0.25, lam=lam).fit(X, y)
gam05 = ExpectileGAM(expectile=0.05, lam=lam).fit(X, y)

from matplotlib import pyplot as plt

XX = gam50.generate_X_grid(term=0, n=500)

plt.scatter(X, y, c='k', alpha=0.2)
plt.plot(XX, gam95.predict(XX), label='0.95')
plt.plot(XX, gam75.predict(XX), label='0.75')
plt.plot(XX, gam50.predict(XX), label='0.50')
plt.plot(XX, gam25.predict(XX), label='0.25')
plt.plot(XX, gam05.predict(XX), label='0.05')
plt.legend()
```

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class pygam.pygam.ExpectileGAM(terms='auto', max_iter=100, tol=0.0001, scale=None, callbacks=['deviance', 'diffs'], fit_intercept=True, expectile=0.5, verbose=False, **kwargs)

Bases: pygam.pygam.GAM

Expectile GAM

This is a GAM with a Normal distribution and an Identity Link, but minimizing the Least Asymmetrically Weighted Squares

Parameters

- **terms** *(expression specifying terms to model, optional.)* – By default a univariate spline term will be allocated for each feature.

  For example:

  >>> GAM(s(0) + l(1) + f(2) + te(3, 4))

  will fit a spline term on feature 0, a linear term on feature 1, a factor term on feature 2, and a tensor term on features 3 and 4.

- **callbacks** *(list of str or list of CallBack objects, optional)* – Names of callback objects to call during the optimization loop.

- **fit_intercept** *(bool, optional)* – Specifies if a constant (a.k.a. bias or intercept) should be added to the decision function. Note: the intercept receives no smoothing penalty.

- **max_iter** *(int, optional)* – Maximum number of iterations allowed for the solver to converge.

- **tol** *(float, optional)* – Tolerance for stopping criteria.

- **verbose** *(bool, optional)* – whether to show pyGAM warnings.

**coef_**

Coefficient of the features in the decision function. If fit_intercept is True, then self.coef_[0] will contain the bias.

  Type array, shape (n_classes, m_features)

**statistics_**

Dictionary containing model statistics like GCV/UBRE scores, AIC/c, parameter covariances, estimated degrees of freedom, etc.

  Type dict

**logs_**

Dictionary containing the outputs of any callbacks at each optimization loop.

  The logs are structured as {callback: [...]}

  Type dict

References


confidence_intervals \((X, width=0.95, quantiles=None)\)
estimate confidence intervals for the model.

Parameters

- \(X\) (array-like of shape \((n\_samples, m\_features)\)) – Input data matrix
- \(width\) (float on \([0,1]\), optional) –
- \(quantiles\) (array-like of floats in \((0, 1)\), optional) – Instead of specifying the prediction width, one can specify the quantiles. So \(width=.95\) is equivalent to \(quantiles=[.025, .975]\)

Returns intervals

Return type np.array of shape \((n\_samples, 2 \text{ or } \text{len}(quantiles))\)

Notes

Wood 2006, section 4.9 Confidence intervals based on section 4.8 rely on large sample results to deal with non-Gaussian distributions, and treat the smoothing parameters as fixed, when in reality they are estimated from the data.

deviance_residuals \((X, y, weights=None, scaled=False)\)
method to compute the deviance residuals of the model
these are analogous to the residuals of an OLS.

Parameters

- \(X\) (array-like) – Input data array of shape \((n\_samples, m\_features)\)
- \(y\) (array-like) – Output data vector of shape \((n\_samples,)\)
- \(weights\) (array-like shape \((n\_samples,)\) or None, optional) – Sample weights. If None, defaults to array of ones
- \(scaled\) (bool, optional) – Whether to scale the deviance by the (estimated) distribution scale

Returns deviance_residuals – with shape \((n\_samples,)\)

Return type np.array

fit \((X, y, weights=None)\)
Fit the generalized additive model.

Parameters

- \(X\) (array-like, shape \((n\_samples, m\_features)\)) – Training vectors.
- \(y\) (array-like, shape \((n\_samples,)\)) – Target values, ie integers in classification, real numbers in regression
- \(weights\) (array-like shape \((n\_samples,)\) or None, optional) – Sample weights. If None, defaults to array of ones

Returns self – Returns fitted GAM object

Return type object

fit_quantile \((X, y, quantile, max_iter=20, tol=0.01, weights=None)\)
fit ExpectileGAM to a desired quantile via binary search

Parameters
• \textbf{X} (array-like, shape (n\_samples, m\_features)) – Training vectors, where \( n\_\text{samples} \) is the number of samples and \( m\_\text{features} \) is the number of features.
• \textbf{y} (array-like, shape (n\_samples,)) – Target values (integers in classification, real numbers in regression) For classification, labels must correspond to classes.
• \textbf{quantile} (float on (0, 1)) – desired quantile to fit.
• \textbf{max\_iter} (int, default: 20) – maximum number of binary search iterations to perform
• \textbf{tol} (float > 0, default: 0.01) – maximum distance between desired quantile and fitted quantile

\begin{itemize}
\item \textbf{weights} (array-like shape (n\_samples,) or None, default: None) – containing sample weights if None, defaults to array of ones
\end{itemize}

\textbf{Returns} \hspace{1em} self

\textbf{Return type} \hspace{1em} fitted GAM object

\textbf{generate\_X\_grid} (term, n=100, meshgrid=False)
create a nice grid of X data

array is sorted by feature and uniformly spaced, so the marginal and joint distributions are likely wrong
if \( \text{term} \) is \( \geq 0 \), we generate \( n \) samples per feature, which results in \( n^\text{deg} \) samples, where \( \text{deg} \) is the degree of the interaction of the term

\textbf{Parameters}

\begin{itemize}
\item \textbf{term} (int,) – Which term to process.
\item \textbf{n} (int, optional) – number of data points to create
\item \textbf{meshgrid} (bool, optional) – Whether to return a meshgrid (useful for 3d plotting) or a feature matrix (useful for inference like partial predictions)
\end{itemize}

\textbf{Returns}

\begin{itemize}
\item if meshgrid is False – np.array of shape \( (n, n\_\text{features}) \) where \( m \) is the number of (sub)terms in the requested (tensor)term.
\item else – tuple of len \( m \), where \( m \) is the number of (sub)terms in the requested (tensor)term. each element in the tuple contains a np.ndarray of size \( (n)^m \)
\end{itemize}

\textbf{Raises} \hspace{1em} \textbf{ValueError} – If the term requested is an intercept since it does not make sense to process the intercept term.

\textbf{get\_params} (deep=False)
returns a dict of all of the object’s user-facing parameters

\textbf{Parameters} \hspace{1em} \textbf{deep} (boolean, default: False) – when True, also gets non-user-facing parameters

\textbf{Returns} \hspace{1em} \textbf{dict}

\textbf{gridsearch} (X, y, weights=None, return\_scores=False, keep\_best=True, objective='auto', progress=True, **param\_grids)
Performs a grid search over a space of parameters for a given objective
**pyGAM Documentation**

**Warning:** `gridsearch` is lazy and will not remove useless combinations from the search space, eg.

```python
>>> n_splines=np.arange(5,10), fit_splines=[True, False]
```

will result in 10 loops, of which 5 are equivalent because `fit_splines = False`

Also, it is not recommended to search over a grid that alternates between known scales and unknown scales, as the scores of the candidate models will not be comparable.

**Parameters**

- **X** *(array-like)* – input data of shape (n_samples, m_features)
- **y** *(array-like)* – label data of shape (n_samples,)
- **weights** *(array-like shape (n_samples,), optional)* – sample weights
- **return_scores** *(boolean, optional)* – whether to return the hyperparameters and score for each element in the grid
- **keep_best** *(boolean, optional)* – whether to keep the best GAM as self.
- **objective** *({'auto', 'AIC', 'AICc', 'GCV', 'UBRE'}, optional)* – Metric to optimize. If *auto*, then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.
- **progress** *(bool, optional)* – whether to display a progress bar
- ****kwargs** – pairs of parameters and iterables of floats, or parameters and iterables of iterables of floats.

If no parameter are specified, `lam=np.logspace(-3, 3, 11)` is used. This results in a 11 points, placed diagonally across lam space.

If grid is iterable of iterables of floats, the outer iterable must have length `m_features`. the cartesian product of the subgrids in the grid will be tested.

If grid is a 2d numpy array, each row of the array will be tested.

The method will make a grid of all the combinations of the parameters and fit a GAM to each combination.

**Returns**

- if `return_scores=True` – `model_scores`: dict containing each fitted model as keys and corresponding objective scores as values
- `else` – self: ie possibly the newly fitted model

**Examples**

For a model with 4 terms, and where we expect 4 lam values, our search space for lam must have 4 dimensions.

We can search the space in 3 ways:

1. via cartesian product by specifying the grid as a list. our grid search will consider `11 ** 4` points:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = [lam] * 4
>>> gam.gridsearch(X, y, lam=lams)
```
2. directly by specifying the grid as a np.ndarray. This is useful for when the dimensionality of the search space is very large, and we would prefer to execute a randomized search:

```python
>>> lams = np.exp(np.random.random(50, 4) * 6 - 3)
>>> gam.gridsearch(X, y, lam=lams)
```

3. copying grids for parameters with multiple dimensions. If we specify a 1D np.ndarray for lam, we are implicitly testing the space where all points have the same value

```python
>>> gam.gridsearch(lam=np.logspace(-3, 3, 11))
```

is equivalent to:

```python
>>> lam = np.logspace(-3, 3, 11)
>>> lams = np.array([lam] * 4)
>>> gam.gridsearch(X, y, lam=lams)
```

### loglikelihood

**loglikelihood** *(X, y, weights=None)*

Computes the log-likelihood of the dataset using the current model.

**Parameters**

- **X** *(array-like of shape (n_samples, m_features))* — containing the input dataset.
- **y** *(array-like of shape (n,))* — containing target values.
- **weights** *(array-like of shape (n,), optional)* — containing sample weights.

**Returns**

- **log-likelihood** — containing log-likelihood scores.

**Return type**

np.array of shape (n,)

### partial_dependence

**partial_dependence** *(term, X=None, width=None, quantiles=None, meshgrid=False)*

Computes the term functions for the GAM and possibly their confidence intervals.

If both width=None and quantiles=None, then no confidence intervals are computed.

**Parameters**

- **term** *(int, optional)* — Term for which to compute the partial dependence functions.
- **X** *(array-like with input data, optional)* — if meshgrid=False, then X should be an array-like of shape (n_samples, m_features).

If meshgrid=True, then X should be a tuple containing an array for each feature in the term.

If None, an equally spaced grid of points is generated.
- **width** *(float on (0, 1), optional)* — Width of the confidence interval.
- **quantiles** *(array-like of floats on (0, 1), optional)* — instead of specifying the prediction width, one can specify the quantiles. So width=.95 is equivalent to quantiles=[.025, .975]. If None, defaults to width.
- **meshgrid** *(bool, whether to return and accept meshgrids.)* — Useful for creating outputs that are suitable for 3D plotting.

Note, for simple terms with no interactions, the output of this function will be the same for meshgrid=True and meshgrid=False, but the inputs will need to be different.
Returns

- `pdeps` *(np.array of shape (n_samples,))*
- `conf_intervals` *(list of length len(term)) – containing np.arrays of shape (n_samples, 2 or len(quantiles))*

Raises `ValueError` – If the term requested is an intercept since it does not make sense to process the intercept term.

See also:

`generate_X_grid()` for help creating meshgrids.

**predict** (*X*)

Predict expected value of target given model and input X often this is done via expected value of GAM given input X

**Parameters**

- `X` *(array-like of shape (n_samples, m_features)) – containing the input dataset*

**Returns**

- `y` – containing predicted values under the model

**Return type**

- np.array of shape (n_samples,)

**predict_mu** (*X*)

Predict expected value of target given model and input X

**Parameters**

- `X` *(array-like of shape (n_samples, m_features)) – containing the input dataset*

**Returns**

- `y` – containing expected values under the model

**Return type**

- np.array of shape (n_samples,)

**sample** (*X, y*, *quantity='y', sample_at_X=None, weights=None, n_draws=100, n_bootstraps=5, objective='auto'* )

Simulate from the posterior of the coefficients and smoothing params.

Samples are drawn from the posterior of the coefficients and smoothing parameters given the response in an approximate way. The GAM must already be fitted before calling this method; if the model has not been fitted, then an exception is raised. Moreover, it is recommended that the model and its hyperparameters be chosen with gridsearch (with the parameter `keep_best=True`) before calling sample, so that the result of that gridsearch can be used to generate useful response data and so that the model’s coefficients (and their covariance matrix) can be used as the first bootstrap sample.

These samples are drawn as follows. Details are in the reference below.

1. `n_bootstraps` many “bootstrap samples” of the response (*y*) are simulated by drawing random samples from the model’s distribution evaluated at the expected values (*mu*) for each sample in *X*.

2. A copy of the model is fitted to each of those bootstrap samples of the response. The result is an approximation of the distribution over the smoothing parameter *lam* given the response data *y*.

3. Samples of the coefficients are simulated from a multivariate normal using the bootstrap samples of the coefficients and their covariance matrices.

**Notes**

A gridsearch is done `n_bootstraps` many times, so keep `n_bootstraps` small. Make `n_bootstraps` < `n_draws` to take advantage of the expensive bootstrap samples of the smoothing parameters.
Parameters

- **X** *(array of shape (n_samples, m_features))* – empirical input data
- **y** *(array of shape (n_samples,))* – empirical response vector
- **quantity** *(\{'y', 'coef', 'mu\}', default: 'y')* – What quantity to return pseudorandom samples of. If **sample_at_X** is not None and **quantity** is either ‘y’ or ‘mu’, then samples are drawn at the values of X specified in **sample_at_X**.
- **sample_at_X** *(array of shape (n_samples_to_simulate, m_features) or None)* – Input data at which to draw new samples. Only applies for **quantity** equal to ‘y’ or to ‘mu’. If None, then **sample_at_X** is replaced by X.
- **weights** *(np.array of shape (n_samples,))* – sample weights
- **n_draws** *(positive int, optional (default=100))* – The number of samples to draw from the posterior distribution of the coefficients and smoothing parameters
- **n_bootstraps** *(positive int, optional (default=5))* – The number of bootstrap samples to draw from simulations of the response (from the already fitted model) to estimate the distribution of the smoothing parameters given the response data. If **n_bootstraps** is 1, then only the already fitted model’s smoothing parameter is used, and the distribution over the smoothing parameters is not estimated using bootstrap sampling.
- **objective** *(string, optional (default='auto'))* – metric to optimize in grid search. must be in ['AIC', 'AICc', 'GCV', 'UBRE', 'auto'] if 'auto', then grid search will optimize GCV for models with unknown scale and UBRE for models with known scale.

Returns

- **draws** – Simulations of the given **quantity** using samples from the posterior distribution of the coefficients and smoothing parameter given the response data. Each row is a pseudorandom sample.

  If **quantity** == ‘coef’, then the number of columns of **draws** is the number of coefficients (len(self.coef_)).

  Otherwise, the number of columns of **draws** is the number of columns of **sample_at_X** if **sample_at_X** is not None or else the number of rows of X.

Return type 2D array of length n_draws

References


**score** *(X, y, weights=None)*
method to compute the explained deviance for a trained model for a given X data and y labels

Parameters

- **X** *(array-like)* – Input data array of shape (n_samples, m_features)
- **y** *(array-like)* – Output data vector of shape (n_samples,)
• **weights** *(array-like shape (n_samples,) or None, optional)* – Sample weights. If None, defaults to an array of ones.

**Returns** explained deviance score

**Return type** np.array() (n_samples,)

**set_params** *(deep=False, force=False, **parameters)*

Sets an object’s parameters

**Parameters**

• **deep** *(boolean, default: False)* – When True, also sets non-user-facing parameters.

• **force** *(boolean, default: False)* – When True, also sets parameters that the object does not already have.

• **parameters** *(parameters to set)*

**Returns**

**Return type** self

**summary** ()

Produce a summary of the model statistics

**Parameters** None –

**Returns**

**Return type** None

### 6.3.2 Terms

#### Linear Term

`pygam.terms.L(feature, lam=0.6, penalties='auto', verbose=False)`

Creates an instance of a LinearTerm

**feature** *(int)* Index of the feature to use for the feature function.

**lam** *(float or iterable of floats)* Strength of smoothing penalty. Must be a positive float. Larger values enforce stronger smoothing.

If single value is passed, it will be repeated for every penalty.

If iterable is passed, the length of `lam` must be equal to the length of `penalties`.

**penalties** *[{'auto', 'derivative', 'l2', None} or callable or iterable]* Type of smoothing penalty to apply to the term.

If an iterable is used, multiple penalties are applied to the term. The length of the iterable must match the length of `lam`.

If ‘auto’, then 2nd derivative smoothing for ‘numerical’ dtypes, and L2/ridge smoothing for ‘categorical’ dtypes.

Custom penalties can be passed as a callable.

**n_coefs** *(int)* Number of coefficients contributed by the term to the model.

**istensor** *(bool)* Whether the term is a tensor product of sub-terms.
isintercept  [bool] whether the term is an intercept

hasconstraint  [bool] whether the term has any constraints

info  [dict] contains dict with the sufficient information to duplicate the term

See also:

LinearTerm()  for developer details

Spline Term

pygam.terms.s(feature, n_splines=20, spline_order=3, lam=0.6, penalties='auto', constraints=None, dtype='numerical', basis='ps', by=None, edge_knots=None, verbose=False)

creates an instance of a SplineTerm

feature  [int] Index of the feature to use for the feature function.

n_splines  [int] Number of splines to use for the feature function. Must be non-negative.

spline_order  [int] Order of spline to use for the feature function. Must be non-negative.

lam  [float or iterable of floats] Strength of smoothing penalty. Must be a positive float. Larger values enforce stronger smoothing.

If single value is passed, it will be repeated for every penalty.

If iterable is passed, the length of lam must be equal to the length of penalties

penalties  [{‘auto’, ‘derivative’, ‘l2’, None} or callable or iterable] Type of smoothing penalty to apply to the term.

If an iterable is used, multiple penalties are applied to the term. The length of the iterable must match the length of lam.

If ‘auto’, then 2nd derivative smoothing for ‘numerical’ dtypes, and L2/ridge smoothing for ‘categorical’ dtypes.

Custom penalties can be passed as a callable.

constraints  [{None, ‘convex’, ‘concave’, ‘monotonic_inc’, ‘monotonic_dec’}] or callable or iterable

Type of constraint to apply to the term.

If an iterable is used, multiple penalties are applied to the term.

dtype  [{‘numerical’, ‘categorical’}] String describing the data-type of the feature.

basis  [{‘ps’, ‘cp’}] Type of basis function to use in the term.

‘ps’ : p-spline basis

‘cp’ [cyclic p-spline basis, useful for building periodic functions.] by default, the maximum and minimum of the feature values are used to determine the function’s period.

edge_knots : optional, array-like of floats of length 2

to specify a custom period use argument edge_knots

these values specify minimum and maximum domain of the spline function.
in the case that spline_basics="cp", edge_knots determines the period of the cyclic function.
when edge_knots=none these values are inferred from the data.
default: None

**by** [int, optional] Feature to use as a by-variable in the term.

For example, if `feature = 2 by = 0`, then the term will produce: x0 * f(x2)

**n_coefs** [int] Number of coefficients contributed by the term to the model

**istensor** [bool] whether the term is a tensor product of sub-terms

**isintercept** [bool] whether the term is an intercept

**hasconstraint** [bool] whether the term has any constraints

**info** [dict] contains dict with the sufficient information to duplicate the term

See also:

*SplineTerm() for developer details*

**Factor Term**

`pygam.terms.f(feature, lam=0.6, penalties='auto', coding='one-hot', verbose=False)`

creates an instance of a FactorTerm

**feature** [int] Index of the feature to use for the feature function.

**lam** [float or iterable of floats] Strength of smoothing penalty. Must be a positive float. Larger values enforce stronger smoothing.

If single value is passed, it will be repeated for every penalty.

If iterable is passed, the length of `lam` must be equal to the length of `penalties`

**penalties** [('auto', 'derivative', 'l2', None) or callable or iterable] Type of smoothing penalty to apply to the term.

If an iterable is used, multiple penalties are applied to the term. The length of the iterable must match the length of `lam`.

If ‘auto’, then 2nd derivative smoothing for ‘numerical’ dtypes, and L2/ridge smoothing for ‘categorical’ dtypes.

Custom penalties can be passed as a callable.

**coding** ['one-hot'] type of contrast encoding to use. currently, only ‘one-hot’ encoding has been developed. this means that we fit one coefficient per category.

**n_coefs** [int] Number of coefficients contributed by the term to the model

**istensor** [bool] whether the term is a tensor product of sub-terms

**isintercept** [bool] whether the term is an intercept

**hasconstraint** [bool] whether the term has any constraints

**info** [dict] contains dict with the sufficient information to duplicate the term

See also:

*FactorTerm() for developer details*
**Tensor Term**

`pygam.terms.te(*args, **kwargs)` creates an instance of a TensorTerm

This is useful for creating interactions between features, or other terms.

*args : marginal Terms to combine into a tensor product

**feature** [list of integers] Indices of the features to use for the marginal terms.

**n_splines** [list of integers] Number of splines to use for each marginal term. Must be of same length as feature.

**spline_order** [list of integers] Order of spline to use for the feature function. Must be of same length as feature.

**lam** [float or iterable of floats] Strength of smoothing penalty. Must be a positive float. Larger values enforce stronger smoothing.

If single value is passed, it will be repeated for every penalty.

If iterable is passed, the length of lam must be equal to the length of penalties

**penalties** [{'auto', 'derivative', 'l2', None} or callable or iterable] Type of smoothing penalty to apply to the term.

If an iterable is used, multiple penalties are applied to the term. The length of the iterable must match the length of lam.

If ‘auto’, then 2nd derivative smoothing for ‘numerical’ dtypes, and L2/ridge smoothing for ‘categorical’ dtypes.

Custom penalties can be passed as a callable.

**constraints** [{None, ‘convex’, ‘concave’, ‘monotonic_inc’, ‘monotonic_dec’}] or callable or iterable

Type of constraint to apply to the term.

If an iterable is used, multiple penalties are applied to the term.

**dtype** [list of {'numerical', 'categorical'}] String describing the data-type of the feature.

Must be of same length as feature.

**basis** [list of {'ps'}] Type of basis function to use in the term.

`'ps'` : p-spline basis

NotImplemented: `cp` : cyclic p-spline basis

Must be of same length as feature.

**by** [int, optional] Feature to use as a by-variable in the term.

For example, if `feature = [1, 2] by = 0`, then the term will produce: \(x0 \times \text{te}(x1, x2)\)

**n_coefs** [int] Number of coefficients contributed by the term to the model

**istensor** [bool] whether the term is a tensor product of sub-terms

**isintercept** [bool] whether the term is an intercept

**hasconstaint** [bool] whether the term has any constraints

**info** [dict] contains dict with the sufficient information to duplicate the term
See also:

*TensorTerm()* for developer details

### 6.4 Developer API

#### 6.4.1 Terms

```python
class pygam.terms.Term(feature, lam=0.6, dtype='numerical', fit_linear=False, fit_splines=True, penalties='auto', constraints=None, verbose=False)
```

**Bases:** pygam.core.Core

**build_columns** (*X*, *verbose=False*)
construct the model matrix columns for the term

**Parameters**
- *X* (*array-like*) – Input dataset with n rows
- *verbose* (*bool*) – whether to show warnings

**Returns**
- **Return type** scipy sparse array with n rows

**build_constraints** (*coef*, *constraint_lam*, *constraint_l2*)
builds the GAM block-diagonal constraint matrix in quadratic form out of constraint matrices specified for each feature.

behaves like a penalty, but with a very large lambda value, ie 1e6.

**Parameters**
- *coef* (*array-like containing the coefficients of a term*) –
- *constraint_lam* (*float*) – penalty to impose on the constraint.
  typically this is a very large number.
- *constraint_l2* (*float*) – loading to improve the numerical conditioning of the constraint matrix.
  typically this is a very small number.

**Returns**
- **C**
  - **Return type** sparse CSC matrix containing the model constraints in quadratic form

**Class method** **build_from_info**(info)
build a Term instance from a dict

**Parameters**
- *cls* (*class*) –
- *info* (*dict*) – contains all information needed to build the term

**Returns**
- **Return type** Term instance
build_penalties(verbos=False)
builds the GAM block-diagonal penalty matrix in quadratic form out of penalty matrices specified for each feature.

   each feature penalty matrix is multiplied by a lambda for that feature.
   so for m features: P = block_diag[lam0 * P0, lam1 * P1, lam2 * P2, . . . , lamm * Pm]

Parameters None –
Returns P
Return type sparse CSC matrix containing the model penalties in quadratic form

compile(X, verbos=False)
method to validate and prepare data-dependent parameters

Parameters
   • X(array-like) – Input dataset
   • verbose(bool) – whether to show warnings

Returns

Return type None

hasconstraint
bool, whether the term has any constraints

info
get information about this term

Returns

Return type dict containing information to duplicate this term

isintercept

istensor

n_coefs
Number of coefficients contributed by the term to the model

class pygam.terms.LinearTerm(feature, lam=0.6, penalties='auto', verbose=False)
Bases: pygam.terms.Term

build_columns(X, verbos=False)
construct the model matrix columns for the term

Parameters
   • X(array-like) – Input dataset with n rows
   • verbose(bool) – whether to show warnings

Returns

Return type scipy sparse array with n rows

build_constraints(coef, constraint_lam, constraint_l2)
builds the GAM block-diagonal constraint matrix in quadratic form out of constraint matrices specified for each feature.

behaves like a penalty, but with a very large lambda value, ie 1e6.

Parameters
• **coefs** *(array-like containing the coefficients of a term)* –

• **constraint_lam** *(float)* – penalty to impose on the constraint. Typically this is a very large number.

• **constraint_l2** *(float)* – loading to improve the numerical conditioning of the constraint matrix. Typically this is a very small number.

**Return** C

**Return type** sparse CSC matrix containing the model constraints in quadratic form

**classmethod build_from_info**(info)

build a Term instance from a dict

**Parameters**

• **cls** *(class)* –

• **info** *(dict)* – contains all information needed to build the term

**Returns**

**Return type** Term instance

**build_penalties**(verbose=False)

builds the GAM block-diagonal penalty matrix in quadratic form out of penalty matrices specified for each feature. Each feature penalty matrix is multiplied by a lambda for that feature. So for m features: P = block_diag[lam0 * P0, lam1 * P1, lam2 * P2, …, lam_m * Pm]

**Parameters** None –

**Returns** P

**Return type** sparse CSC matrix containing the model penalties in quadratic form

**compile**(X, verbose=False)

method to validate and prepare data-dependent parameters

**Parameters**

• **X** *(array-like)* – Input dataset

• **verbose** *(bool)* – whether to show warnings

**Returns**

**Return type** None

**get_params**(deep=False)

returns a dict of all of the object’s user-facing parameters

**Parameters** **deep** *(boolean, default: False)* – when True, also gets non-user-facing parameters

**Returns**

**Return type** dict

**hasconstraint**

bool, whether the term has any constraints
**pyGAM Documentation**

**info**
get information about this term

Returns

**Return type** dict containing information to duplicate this term

**isintercept**

**istensor**

**n_coefs**
Number of coefficients contributed by the term to the model

**set_params** *(deep=False, force=False, **parameters)*
sets an object’s parameters

**Parameters**

- **deep** *(boolean, default: False)* – when True, also sets non-user-facing parameters

- **force** *(boolean, default: False)* – when True, also sets parameters that the object does not already have

- ****parameters**(parameters to set)**

Returns

**Return type** self

**class** *pygam.terms.SplineTerm*(feature, n_splines=20, spline_order=3, lam=0.6, penalties='auto', constraints=None, dtype='numerical', basis='ps', by=None, edge_knots=None, verbose=False)

**Bases:** *pygam.terms.Term*

**build_columns** *(X, verbose=False)*
construct the model matrix columns for the term

**Parameters**

- **X** *(array-like)* – Input dataset with n rows

- **verbose** *(bool)* – whether to show warnings

Returns

**Return type** scipy sparse array with n rows

**build_constraints** *(coef, constraint_lam, constraint_l2)*
builds the GAM block-diagonal constraint matrix in quadratic form out of constraint matrices specified for each feature.

behaves like a penalty, but with a very large lambda value, ie 1e6.

**Parameters**

- **coef** *(array-like containing the coefficients of a term)* –

- **constraint_lam**(float, ) – penalty to impose on the constraint.

  typically this is a very large number.

- **constraint_l2**(float, ) – loading to improve the numerical conditioning of the constraint matrix.

  typically this is a very small number.
Returns C

Return type sparse CSC matrix containing the model constraints in quadratic form

classmethod build_from_info(info)

build a Term instance from a dict

Parameters

• cls (class) –

• info (dict) – contains all information needed to build the term

Returns

Return type Term instance

build_penalties(verbosel=sFalse)

builds the GAM block-diagonal penalty matrix in quadratic form out of penalty matrices specified for each feature.

each feature penalty matrix is multiplied by a lambda for that feature.

so for m features: P = block_diag[lam0 * P0, lam1 * P1, lam2 * P2,. . . , lamn * Pn]

Parameters None –

Returns P

Return type sparse CSC matrix containing the model penalties in quadratic form

compile(X, verbosel=sFalse)

method to validate and prepare data-dependent parameters

Parameters

• X (array-like) – Input dataset

• verbosel (bool) – whether to show warnings

Returns

Return type None

get_params(deep=sFalse)

returns a dict of all of the object’s user-facing parameters

Parameters deep (boolean, default: False) – when True, also gets non-user-facing parameters

Returns

Return type dict

hasconstraint

bool, whether the term has any constraints

info

get information about this term

Returns

Return type dict containing information to duplicate this term

isintercept

istensor
n_coefs
Number of coefficients contributed by the term to the model

set_params (deep=False, force=False, **parameters)
sets an object’s parameters

Parameters
• deep (boolean, default: False) – when True, also sets non-user-facing parameters
• force (boolean, default: False) – when True, also sets parameters that the object does not already have
• **parameters (parameters to set) –

Returns
Return type self

class pygam.terms.FactorTerm (feature, lam=0.6, penalties='auto', coding='one-hot', verbose=False)
Bases: pygam.terms.SplineTerm

build_columns (X, verbose=False)
construct the model matrix columns for the term

Parameters
• X (array-like) – Input dataset with n rows
• verbose (bool) – whether to show warnings

Returns
Return type scipy sparse array with n rows

build_constraints (coef, constraint_lam, constraint_l2)
builds the GAM block-diagonal constraint matrix in quadratic form out of constraint matrices specified for each feature.

behaves like a penalty, but with a very large lambda value, ie 1e6.

Parameters
• coefs (array-like containing the coefficients of a term) –
• constraint_lam (float,) – penalty to impose on the constraint.
  typically this is a very large number.
• constraint_l2 (float,) – loading to improve the numerical conditioning of the constraint matrix.
  typically this is a very small number.

Returns C
Return type sparse CSC matrix containing the model constraints in quadratic form

classmethod build_from_info (info)
build a Term instance from a dict

Parameters
• cls (class) –
• info (dict) – contains all information needed to build the term
Returns

Return type Term instance

build_penalties (verbose=False)
builds the GAM block-diagonal penalty matrix in quadratic form out of penalty matrices specified for each feature.
each feature penalty matrix is multiplied by a lambda for that feature.
so for m features: P = block_diag[lam0 * P0, lam1 * P1, lam2 * P2, …, lam_m * Pm]

Parameters None –

Returns P

Return type sparse CSC matrix containing the model penalties in quadratic form

compile (X, verbose=False)
method to validate and prepare data-dependent parameters

Parameters

• X (array-like) – Input dataset

• verbose (bool) – whether to show warnings

Returns

Return type None

get_params (deep=False)
returns a dict of all of the object’s user-facing parameters

Parameters deep (boolean, default: False) – when True, also gets non-user-facing parameters

Returns

Return type dict

hasconstraint
bool, whether the term has any constraints

info
get information about this term

Returns

Return type dict containing information to duplicate this term

isintercept

istensor

n_coefs
Number of coefficients contributed by the term to the model

set_params (deep=False, force=False, **parameters)
sets an object’s parameters

Parameters

• deep (boolean, default: False) – when True, also sets non-user-facing parameters

• force (boolean, default: False) – when True, also sets parameters that the object does not already have
**parameters** (parameters to set)

Returns

Return type  self

class  pygam.terms.TensorTerm  (*args, **kwargs)

Bases:  pygam.terms.SplineTerm, pygam.terms.MetaTermMixin

build_columns  (X, verbose=False)

construct the model matrix columns for the term

Parameters

•  X  (array-like) – Input dataset with n rows

•  verbose  (bool) – whether to show warnings

Returns

Return type  scipy sparse array with n rows

build_constraints  (coef, constraint_lam, constraint_l2)

builds the GAM block-diagonal constraint matrix in quadratic form out of constraint matrices specified for each feature.

Parameters

•  coefs  (array-like containing the coefficients of a term) –

•  constraint_lam  (float,) – penalty to impose on the constraint.

  typically this is a very large number.

•  constraint_l2  (float,) – loading to improve the numerical conditioning of the constraint matrix.

  typically this is a very small number.

Returns  C

Return type  sparse CSC matrix containing the model constraints in quadratic form

classmethod  build_from_info  (info)

build a TensorTerm instance from a dict

Parameters

•  cls  (class) –

•  info  (dict) – contains all information needed to build the term

Returns

Return type  TensorTerm instance

build_penalties()

builds the GAM block-diagonal penalty matrix in quadratic form out of penalty matrices specified for each feature.

each feature penalty matrix is multiplied by a lambda for that feature.

so for m features:  P = block_diag[lam0 * P0, lam1 * P1, lam2 * P2, … , lamm * Pm]

Parameters  None –

Returns  P

Return type  sparse CSC matrix containing the model penalties in quadratic form
**compile** \((X, \text{verbose}=\text{False})\)
method to validate and prepare data-dependent parameters

**Parameters**

- **X** *(array-like)* – Input dataset
- **verbose** *(bool)* – whether to show warnings

**Returns**

**Return type** *None*

**get_params** \((\text{deep}=\text{False})\)
returns a dict of all of the object’s user-facing parameters

**Parameters**

- **deep** *(boolean, default: False)* – when True, also gets non-user-facing parameters

**Returns**

**Return type** *dict*

**hasconstraint**
bool, whether the term has any constraints

**info**
get information about this term

**Returns**

**Return type** *dict containing information to duplicate this term*

**isintercept**

**istensor**

**n_coefs**
Number of coefficients contributed by the term to the model

**set_params** \((\text{deep}=\text{False}, \text{force}=\text{False}, **\text{parameters})\)
sets an object’s parameters

**Parameters**

- **deep** *(boolean, default: False)* – when True, also sets non-user-facing parameters
- **force** *(boolean, default: False)* – when True, also sets parameters that the object does not already have
- ****parameters**(\text{parameters to set})**

**Returns**

**Return type** *self*

**class** *pygam.terms.TermList\(*terms, \text{**kwargs}\)*

**Bases:** *pygam.core.Core, pygam.terms.MetaTermMixint*

**build_columns** \((X, \text{term}=-1, \text{verbose}=\text{False})\)
construct the model matrix columns for the term

**Parameters**

- **X** *(array-like)* – Input dataset with \(n\) rows
- **verbose** *(bool)* – whether to show warnings
Returns

Return type scipy sparse array with n rows

build_constraints (coefs, constraint_lam, constraint_l2)

builds the GAM block-diagonal constraint matrix in quadratic form out of constraint matrices specified for each feature.

behaves like a penalty, but with a very large lambda value, ie 1e6.

Parameters

• coefs (array-like containing the coefficients of a term) –

• constraint_lam (float) – penalty to impose on the constraint.

typically this is a very large number.

• constraint_l2 (float) – loading to improve the numerical conditioning of the constraint matrix.

typically this is a very small number.

Returns C

Return type sparse CSC matrix containing the model constraints in quadratic form

classmethod build_from_info (info)

build a TermList instance from a dict

Parameters

• cls (class) –

• info (dict) – contains all information needed to build the term

Returns

Return type TermList instance

build_penalties ()

builds the GAM block-diagonal penalty matrix in quadratic form out of penalty matrices specified for each feature.

each feature penalty matrix is multiplied by a lambda for that feature.

so for m features: P = block_diag[lam0 * P0, lam1 * P1, lam2 * P2, … , lamm * Pm]

Parameters None –

Returns P

Return type sparse CSC matrix containing the model penalties in quadratic form

compile (X, verbose=False)

method to validate and prepare data-dependent parameters

Parameters

• X (array-like) – Input dataset

• verbose (bool) – whether to show warnings

Returns

Return type None

get_coef_indices (i=-1)

get the indices for the coefficients of a term in the term list
Parameters  `i (int)` – by default `int=-1`, meaning that coefficient indices are returned for all terms in the term list

Returns

Return type  list of integers

**get_params** *(deep=False)*
returns a dict of all of the object’s user-facing parameters

Parameters  `deep (boolean, default: False)` – when True, also gets non-user-facing parameters

Returns

Return type  dict

**hasconstraint**
bool, whether the term has any constraints

**info**
get information about the terms in the term list

Returns

Return type  dict containing information to duplicate the term list

**n_coefs**
Total number of coefficients contributed by the terms in the model

**pop** *(i=None)*
remove the *i*th term from the term list

Parameters  `i (int, optional)` – term to remove from term list
by default the last term is popped.

Returns  *term*

Return type  *Term*

**set_params** *(deep=False, force=False, **parameters)*
sets an object’s parameters

Parameters

•  `deep (boolean, default: False)` – when True, also sets non-user-facing parameters

•  `force (boolean, default: False)` – when True, also sets parameters that the object does not already have

•  `**parameters (parameters to set)–`

Returns

Return type  self

6.4.2 Distributions

Distributions

**class**  *pygam.distributions.BinomialDist*( `levels=1`)

Bases: *pygam.distributions.Distribution*

Binomial Distribution
V (mu)
glm Variance function
computes the variance of the distribution

Parameters

- mu (array-like of length n) – expected values

Returns

- variance

Return type

np.array of length n

deviance (y, mu, scaled=True)
model deviance
for a bernoulli logistic model, this is equal to the twice the negative loglikelihood.

Parameters

- y (array-like of length n) – target values
- mu (array-like of length n) – expected values
- scaled (boolean, default: True) – whether to divide the deviance by the distribution scaled

Returns

- deviances

Return type

np.array of length n

log_pdf (y, mu, weights=None)
computes the log of the pdf or pmf of the values under the current distribution

Parameters

- y (array-like of length n) – target values
- mu (array-like of length n) – expected values
- weights (array-like shape (n,) or None, default: None) – sample weights if None, defaults to array of ones

Returns

- pdf/pmf

Return type

np.array of length n

sample (mu)
Return random samples from this Normal distribution.

Parameters

- mu (array-like of shape n_samples or shape (n_simulations, n_samples)) – expected values

Returns

- random_samples

Return type

np.array of same shape as mu

class pygam.distributions.Distribution (name=None, scale=None)
Bases: pygam.core.Core

phi (y, mu, edof, weights)
GLM scale parameter. for Binomial and Poisson families this is unity for Normal family this is variance

Parameters

- y (array-like of length n) – target values
- mu (array-like of length n) – expected values
- edof (float) – estimated degrees of freedom
• **weights** *(array-like shape (n,) or None, default: None)* – sample weights if None, defaults to array of ones

Returns *scale*

Return type *estimated model scale*

**sample**(mu)

Return random samples from this distribution.

Parameters **mu** *(array-like of shape n_samples or shape (n_simulations, n_samples)) – expected values*

Returns *random_samples*

Return type *np.array of same shape as mu*

class pygam.distributions.GammaDist(scale=None)

Bases: pygam.distributions.Distribution

Gamma Distribution

V(mu)

glm Variance function

computes the variance of the distribution

Parameters **mu** *(array-like of length n) – expected values*

Returns *variance*

Return type *np.array of length n*

deviance**(y, mu, scaled=True)**

model deviance

for a bernoulli logistic model, this is equal to the twice the negative loglikelihood.

Parameters

• **y** *(array-like of length n) – target values*

• **mu** *(array-like of length n) – expected values*

• **scaled** *(boolean, default: True) – whether to divide the deviance by the dis-


tribution scaled*

Returns *deviances*

Return type *np.array of length n*

log_pdf**(y, mu, weights=None)**

computes the log of the pdf or pmf of the values under the current distribution

Parameters

• **y** *(array-like of length n) – target values*

• **mu** *(array-like of length n) – expected values*

• **weights** *(array-like shape (n,) or None, default: None) – contain-


ing sample weights if None, defaults to array of ones*

Returns *pdf/pmf*

Return type *np.array of length n*
**sample**(*mu*)

Return random samples from this Gamma distribution.

Parameters

* mu (array-like of shape n_samples or shape (n_simulations, n_samples)) – expected values

Returns random_samples

Return type np.array of same shape as mu

class **pygam.distributions.InvGaussDist**(scale=None)

Bases: **pygam.distributions.Distribution**

Inverse Gaussian (Wald) Distribution

V(*mu*)

glm Variance function

computes the variance of the distribution

Parameters

* mu (array-like of length n) – expected values

Returns variance

Return type np.array of length n

deviceance(y, mu, scaled=True)

model deviance

for a bernoulli logistic model, this is equal to the twice the negative loglikelihood.

Parameters

* y (array-like of length n) – target values
* mu (array-like of length n) – expected values
* scaled (boolean, default: True) – whether to divide the deviance by the distribution scaled

Returns deviances

Return type np.array of length n

log_pdf(y, mu, weights=None)

computes the log of the pdf or pmf of the values under the current distribution

Parameters

* y (array-like of length n) – target values
* mu (array-like of length n) – expected values
* weights (array-like shape (n,) or None, default: None) – containing sample weights if None, defaults to array of ones

Returns pdf/pmf

Return type np.array of length n

sample(*mu*)

Return random samples from this Inverse Gaussian (Wald) distribution.

Parameters

* mu (array-like of shape n_samples or shape (n_simulations, n_samples)) – expected values

Returns random_samples
**Return type**
np.array of same shape as mu

class pygam.distributions.NormalDist(scale=None)
    Bases: pygam.distributions.Distribution
    Normal Distribution
    
    V(mu)
    glm Variance function.
    
    if Y ~ ExpFam(theta, scale=phi)
    such that E[Y] = mu = b'(theta)
    and Var[Y] = b''(theta) * phi / w
    
    then we seek V(mu) such that we can represent Var[y] as a fn of mu: Var[Y] = V(mu) * phi
    ie V(mu) = b''(theta) / w

    Parameters
    mu (array-like of length n) – expected values
    
    Returns V(mu)
    
    Return type
    np.array of length n

    deviance(y, mu, scaled=True)
        model deviance
        for a gaussian linear model, this is equal to the SSE
        
        Parameters
        • y (array-like of length n) – target values
        • mu (array-like of length n) – expected values
        • scaled (boolean, default: True) – whether to divide the deviance by the distribution scaled
        
        Returns 
        deviances
        
        Return type
        np.array of length n

    log_pdf(y, mu, weights=None)
        computes the log of the pdf or pmf of the values under the current distribution
        
        Parameters
        • y (array-like of length n) – target values
        • mu (array-like of length n) – expected values
        • weights (array-like shape (n,) or None, default: None) – sample weights if None, defaults to array of ones
        
        Returns 
        pdf/pmf
        
        Return type
        np.array of length n

    sample(mu)
        Return random samples from this Normal distribution.
        Samples are drawn independently from univariate normal distributions with means given by the values in mu and with standard deviations equal to the scale attribute if it exists otherwise 1.0.
Parameters mu

(array-like of shape n_samples or shape (n_simulations, n_samples))—expected values

Returns random_samples

Return type np.array of same shape as mu

class pygam.distributions.PoissonDist

Bases: pygam.distributions.Distribution

Poisson Distribution

V(mu)

glm Variance function

computes the variance of the distribution

Parameters mu (array-like of length n)—expected values

Returns variance

Return type np.array of length n

deviance(y, mu, scaled=True)

model deviance

for a bernoulli logistic model, this is equal to the twice the negative loglikelihood.

Parameters

• y (array-like of length n)—target values
• mu (array-like of length n)—expected values
• scaled (boolean, default: True)—whether to divide the deviance by the distribution scaled

Returns deviances

Return type np.array of length n

log_pdf(y, mu, weights=None)

computes the log of the pdf or pmf of the values under the current distribution

Parameters

• y (array-like of length n)—target values
• mu (array-like of length n)—expected values
• weights (array-like shape (n,) or None, default: None)—containing sample weights if None, defaults to array of ones

Returns pdf/pmf

Return type np.array of length n

sample(mu)

Return random samples from this Poisson distribution.

Parameters mu

(array-like of shape n_samples or shape (n_simulations, n_samples))—expected values

Returns random_samples

Return type np.array of same shape as mu

pygam.distributions.divide_weights(V)
pygam.distributions.\texttt{multiply_weights}(\textit{deviance})

### 6.4.3 Links

\begin{verbatim}
class pygam.links.Link(
    name=None
):
    bases: pygam.core.Core
class pygam.links.IdentityLink(
    bases: pygam.links.Link
    
    \texttt{gradient}(\textit{mu}, \textit{dist})
    
    derivative of the link function wrt \textit{mu}

    \begin{verbatim}
    Parameters
    \begin{itemize}
    \item \textit{mu} (array-like of length n) –
    \item \textit{dist} (Distribution instance) –
    \end{itemize}
    \end{verbatim}
    
    Returns \textit{grad}

    \begin{verbatim}
    Return type np.array of length n
    \end{verbatim}

    \texttt{link}(\textit{mu}, \textit{dist})
    
    glm link function this is useful for going from \textit{mu} to the linear prediction

    \begin{verbatim}
    Parameters
    \begin{itemize}
    \item \textit{mu} (array-like of length n) –
    \item \textit{dist} (Distribution instance) –
    \end{itemize}
    \end{verbatim}
    
    \begin{verbatim}
    Returns \textit{lp}
    \end{verbatim}

    \begin{verbatim}
    Return type np.array of length n
    \end{verbatim}

    \texttt{mu}(\textit{lp}, \textit{dist})
    
    glm mean function, ie inverse of link function this is useful for going from the linear prediction to \textit{mu}

    \begin{verbatim}
    Parameters
    \begin{itemize}
    \item \textit{lp} (array-like of length n) –
    \item \textit{dist} (Distribution instance) –
    \end{itemize}
    \end{verbatim}
    
    \begin{verbatim}
    Returns \textit{mu}
    \end{verbatim}

    \begin{verbatim}
    Return type np.array of length n
    \end{verbatim}

    class pygam.links.InvSquaredLink(
    bases: pygam.links.Link
    
    \texttt{gradient}(\textit{mu}, \textit{dist})
    
    derivative of the link function wrt \textit{mu}

    \begin{verbatim}
    Parameters
    \begin{itemize}
    \item \textit{mu} (array-like of length n) –
    \item \textit{dist} (Distribution instance) –
    \end{itemize}
    \end{verbatim}
    
    \begin{verbatim}
    Returns \textit{grad}
    \end{verbatim}

    \begin{verbatim}
    Return type np.array of length n
    \end{verbatim}

    \texttt{link}(\textit{mu}, \textit{dist})
    
    glm link function this is useful for going from \textit{mu} to the linear prediction

    \begin{verbatim}
    \end{verbatim}
\end{verbatim}

6.4. Developer API
Parameters

- $\mu$ (array-like of length n)
- $\text{dist}$ (Distribution instance)

Returns $lp$

Return type np.array of length n

$\text{mu} (lp, \text{dist})$

glm mean function, ie inverse of link function this is useful for going from the linear prediction to $\mu$

Parameters

- $lp$ (array-like of length n)
- $\text{dist}$ (Distribution instance)

Returns $\mu$

Return type np.array of length n

class pygam.links.LogitLink
Bases: pygam.links.Link

gradient ($\mu$, $\text{dist}$)

derivative of the link function wrt $\mu$

Parameters

- $\mu$ (array-like of length n)
- $\text{dist}$ (Distribution instance)

Returns grad

Return type np.array of length n

link ($\mu$, $\text{dist}$)

glm link function this is useful for going from $\mu$ to the linear prediction

Parameters

- $\mu$ (array-like of length n)
- $\text{dist}$ (Distribution instance)

Returns $lp$

Return type np.array of length n

$\text{mu} (lp, \text{dist})$

glm mean function, ie inverse of link function this is useful for going from the linear prediction to $\mu$

Parameters

- $lp$ (array-like of length n)
- $\text{dist}$ (Distribution instance)

Returns $\mu$

Return type np.array of length n

class pygam.links.LogLink
Bases: pygam.links.Link
**gradient** \((\mu, \text{dist})\)

Derivative of the link function wrt \(\mu\)

**Parameters**
- \(\mu\) (array-like of length \(n\))
- \(\text{dist}\) (Distribution instance)

**Returns** \(\text{grad}\)

**Return type** np.array of length \(n\)

**link** \((\mu, \text{dist})\)

GLM link function, this is useful for going from \(\mu\) to the linear prediction

**Parameters**
- \(\mu\) (array-like of length \(n\))
- \(\text{dist}\) (Distribution instance)

**Returns** \(\text{lp}\)

**Return type** np.array of length \(n\)

**mu** \((\text{lp, dist})\)

GLM mean function, i.e., inverse of link function, this is useful for going from the linear prediction to \(\mu\)

**Parameters**
- \(\text{lp}\) (array-like of length \(n\))
- \(\text{dist}\) (Distribution instance)

**Returns** \(\mu\)

**Return type** np.array of length \(n\)

### 6.4.4 Callbacks

**class** pygam.callbacks.CallBack (name=None)

**Bases:** pygam.core.Core

CallBack class

**class** pygam.callbacks.Accuracy

**Bases:** pygam.callbacks.CallBack

**on_loop_start** (\(y, \mu\))

Runs the method at start of each optimization loop

**Parameters**
- \(y\) (array-like of length \(n\)) – target data
- \(\mu\) (array-like of length \(n\)) – expected value data

**Returns** accuracy

**Return type** np.array of length \(n\)

**class** pygam.callbacks.Coef

**Bases:** pygam.callbacks.CallBack

**on_loop_start** (\(gam\))

Runs the method at start of each optimization loop
Parameters

gam (float) –
Returns coef_

Return type list of floats

class pygam.callbacks.Deviance
    Bases: pygam.callbacks.CallBack

Deviance CallBack class

on_loop_start (gam, y, mu)
    runs the method at loop start

Parameters
    • gam (GAM instance) –
    • y (array-like of length n) – target data
    • mu (array-like of length n) – expected value data

Returns deviance

Return type np.array of length n

class pygam.callbacks.Diffs
    Bases: pygam.callbacks.CallBack

on_loop_end (diff)
    runs the method at end of each optimization loop

Parameters diff (float) –

Returns diff

Return type float

pygam.callbacks.validate_callback (callback)
    validates a callback’s on_loop_start and on_loop_end methods

Parameters callback (Callback object) –

Returns

Return type validated callback

pygam.callbacks.validate_callback_data (method)
    wraps a callback’s method to pull the desired arguments from the vars dict also checks to ensure the method’s arguments are in the vars dict

Parameters method (callable) –

Returns

Return type validated callable

6.4.5 Penalties

Penalty matrix generators

pygam.penalties.concave (n, coef)
    Builds a penalty matrix for P-Splines with continuous features. Penalizes violation of a concave feature function.

Parameters
    • n (int) – number of splines
• `coef (array-like)` – coefficients of the feature function

**Returns** penalty matrix

**Return type** sparse csc matrix of shape (n,n)

```python
pygam.penalties.convex(n, coef)
```
Builds a penalty matrix for P-Splines with continuous features. Penalizes violation of a convex feature function.

**Parameters**

• `n (int)` – number of splines

• `coef (array-like)` – coefficients of the feature function

**Returns** penalty matrix

**Return type** sparse csc matrix of shape (n,n)

```python
pygam.penalties.convexity_(n, coef, convex=True)
```
Builds a penalty matrix for P-Splines with continuous features. Penalizes violation of convexity in the feature function.

**Parameters**

• `n (int)` – number of splines

• `coef (array-like)` – coefficients of the feature function

• `convex (bool, default: True)` – whether to enforce convex, or concave functions

**Returns** penalty matrix

**Return type** sparse csc matrix of shape (n,n)

```python
pygam.penalties.derivative(n, coef, derivative=2, periodic=False)
```
Builds a penalty matrix for P-Splines with continuous features. Penalizes the squared differences between basis coefficients.

**Parameters**

• `n (int)` – number of splines

• `coef (unused)` – for compatibility with constraints

• `derivative (int, default: 2)` – which derivative do we penalize. derivative is 1, we penalize 1st order derivatives, derivative is 2, we penalize 2nd order derivatives, etc

**Returns** penalty matrix

**Return type** sparse csc matrix of shape (n,n)

```python
pygam.penalties.l2(n, coef)
```
Builds a penalty matrix for P-Splines with categorical features. Penalizes the squared value of each basis coefficient.

**Parameters**

• `n (int)` – number of splines

• `coef (unused)` – for compatibility with constraints

**Returns** penalty matrix

**Return type** sparse csc matrix of shape (n,n)
pygam.penalties.monotonic_dec(n, coef)
Builds a penalty matrix for P-Splines with continuous features. Penalizes violation of a monotonic decreasing feature function.

Parameters
- n (int) – number of splines
- coef (array-like) – coefficients of the feature function

Returns penalty matrix
Return type sparse csc matrix of shape (n,n)

pygam.penalties.monotonic_inc(n, coef)
Builds a penalty matrix for P-Splines with continuous features. Penalizes violation of a monotonic increasing feature function.

Parameters
- n (int) – number of splines
- coef (array-like, coefficients of the feature function)

Returns penalty matrix
Return type sparse csc matrix of shape (n,n)

pygam.penalties.monotonicity_(n, coef, increasing=True)
Builds a penalty matrix for P-Splines with continuous features. Penalizes violation of monotonicity in the feature function.

Parameters
- n (int) – number of splines
- coef (array-like) – coefficients of the feature function
- increasing (bool, default: True) – whether to enforce monotic increasing, or decreasing functions

Returns penalty matrix
Return type sparse csc matrix of shape (n,n)

pygam.penalties.none(n, coef)
Build a matrix of zeros for features that should go unpenalized

Parameters
- n (int) – number of splines
- coef (unused) – for compatibility with constraints

Returns penalty matrix
Return type sparse csc matrix of shape (n,n)

pygam.penalties.periodic(n, coef, derivative=2, _penalty=<function derivative>)

pygam.penalties.sparse_diff(array, n=1, axis=-1)
A ported sparse version of np.diff. Uses recursion to compute higher order differences

Parameters
- array (sparse array)
- n (int, default: 1) – differencing order
• `axis (int, default: -1)` – axis along which differences are computed
  
  **Returns** diff_array – same shape as input array, but ‘axis’ dimension is smaller by ‘n’.

  **Return type** sparse array

**pygam.penalties.wrap_penalty** *(p, fit_linear, linear_penalty=0.0)*

tool to account for unity penalty on the linear term of any feature.

**Example**

```python
p = wrap_penalty(derivative, fit_linear=True)(n, coef)
```

**Parameters**

- **p** *(callable.)* – penalty-matrix-generating function.
- **fit_linear** *(boolean.)* – whether the current feature has a linear term or not.
- **linear_penalty** *(float, default: 0.)* – penalty on the linear term

**Returns** wrapped_p – modified penalty-matrix-generating function

**Return type** callable
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