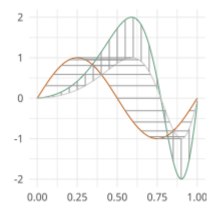
# **fdasrsf Documentation**

Release 2.5.11

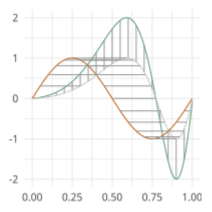
J. Derek Tucker

# **CONTENTS**

1.1 Elastic Functional Alignment 1.2 Elastic Functional Principal Component Analysis 1.3 Multivariate Functional Example 1.4 Elastic Curve Alignment  2 API Reference 2.1 Functional Alignment 2.2 Functional Principal Component Analysis 2.3 Elastic Functional Boxplots 2.4 Functional Principal Least Squares 2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering 2.11 Elastic Image Warping	
1.3 Multivariate Functional Example 1.4 Elastic Curve Alignment  2 API Reference 2.1 Functional Alignment 2.2 Functional Principal Component Analysis 2.3 Elastic Functional Boxplots 2.4 Functional Principal Least Squares 2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering	
1.4 Elastic Curve Alignment  2 API Reference 2.1 Functional Alignment 2.2 Functional Principal Component Analysis 2.3 Elastic Functional Boxplots 2.4 Functional Principal Least Squares 2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering	
2 API Reference 2.1 Functional Alignment 2.2 Functional Principal Component Analysis 2.3 Elastic Functional Boxplots 2.4 Functional Principal Least Squares 2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering	1
2.1 Functional Alignment 2.2 Functional Principal Component Analysis 2.3 Elastic Functional Boxplots 2.4 Functional Principal Least Squares 2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering	1
<ul> <li>2.2 Functional Principal Component Analysis</li> <li>2.3 Elastic Functional Boxplots</li> <li>2.4 Functional Principal Least Squares</li> <li>2.5 Elastic Regression</li> <li>2.6 Elastic Principal Component Regression</li> <li>2.7 Elastic Functional Changepoint</li> <li>2.8 Elastic GLM Regression</li> <li>2.9 Elastic Functional Tolerance Bounds</li> <li>2.10 Elastic Functional Clustering</li> </ul>	2
<ul> <li>2.2 Functional Principal Component Analysis</li> <li>2.3 Elastic Functional Boxplots</li> <li>2.4 Functional Principal Least Squares</li> <li>2.5 Elastic Regression</li> <li>2.6 Elastic Principal Component Regression</li> <li>2.7 Elastic Functional Changepoint</li> <li>2.8 Elastic GLM Regression</li> <li>2.9 Elastic Functional Tolerance Bounds</li> <li>2.10 Elastic Functional Clustering</li> </ul>	2
2.3 Elastic Functional Boxplots 2.4 Functional Principal Least Squares 2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering	
2.4       Functional Principal Least Squares         2.5       Elastic Regression         2.6       Elastic Principal Component Regression         2.7       Elastic Functional Changepoint         2.8       Elastic GLM Regression         2.9       Elastic Functional Tolerance Bounds         2.10       Elastic Functional Clustering	
2.5 Elastic Regression 2.6 Elastic Principal Component Regression 2.7 Elastic Functional Changepoint 2.8 Elastic GLM Regression 2.9 Elastic Functional Tolerance Bounds 2.10 Elastic Functional Clustering	
2.6       Elastic Principal Component Regression         2.7       Elastic Functional Changepoint         2.8       Elastic GLM Regression         2.9       Elastic Functional Tolerance Bounds         2.10       Elastic Functional Clustering	
2.8 Elastic GLM Regression	
2.8       Elastic GLM Regression         2.9       Elastic Functional Tolerance Bounds         2.10       Elastic Functional Clustering	4
<ul><li>2.9 Elastic Functional Tolerance Bounds</li><li>2.10 Elastic Functional Clustering</li></ul>	
2.10 Elastic Functional Clustering	
· · · · · · · · · · · · · · · · · · ·	
2.12 Curve Registration	
2.13 SRVF Geodesic Computation	
2.14 Utility Functions	
2.15 Curve Functions	
2.16 UMAP EFDA Metrics	
3 Installation	8
4 How do I start?	8
5 Contributions	8
6 License	8
7 References	9
8 Indices and tables	9.
Python Module Index	9
Index	,



A python package for functional data analysis using the square root slope framework and curves using the square root velocity framework which performs pair-wise and group-wise alignment as well as modeling using functional component analysis and regression.



CONTENTS 1

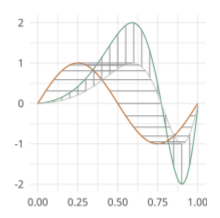
2 CONTENTS

**CHAPTER** 

**ONE** 

# **USER GUIDE**

Contents:



# 1.1 Elastic Functional Alignment

Otherwise known as time warping in the literature is at the center of elastic functional data analysis. Here our goal is to separate out the horizontal and vertical variability of the functional data

```
[1]: import fdasrsf as fs import numpy as np
```

Load in our example data

```
[2]: data = np.load('../../bin/simu_data.npz')
  time = data['arr_1']
  f = data['arr_0']
```

We will then construct the fdawarp object

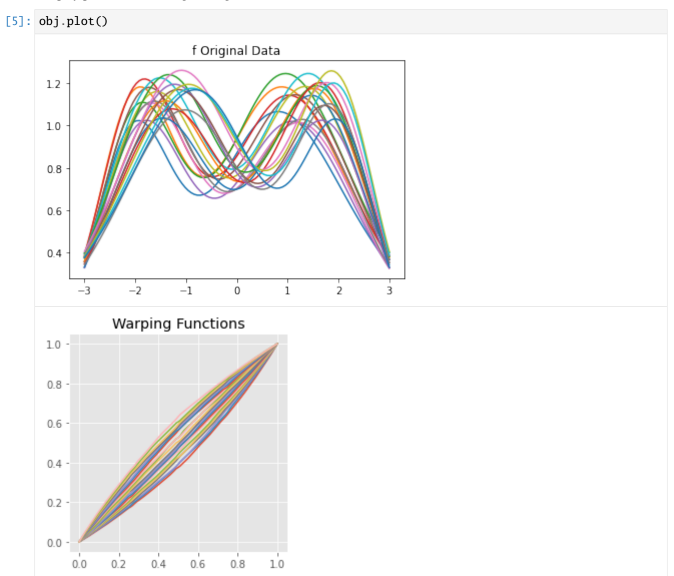
```
[3]: obj = fs.fdawarp(f,time)
```

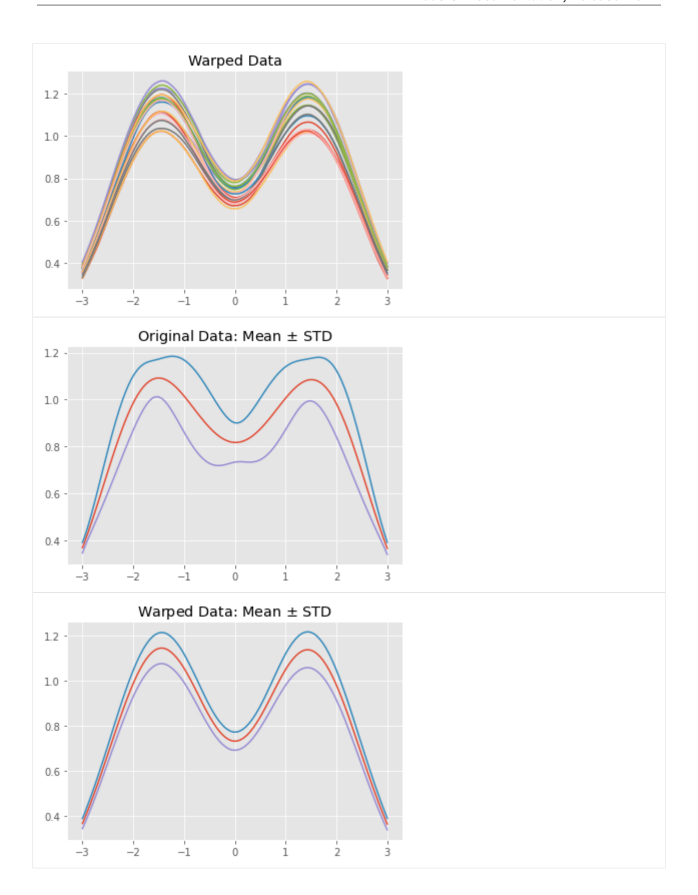
Next we will align the functions using the elastic framework

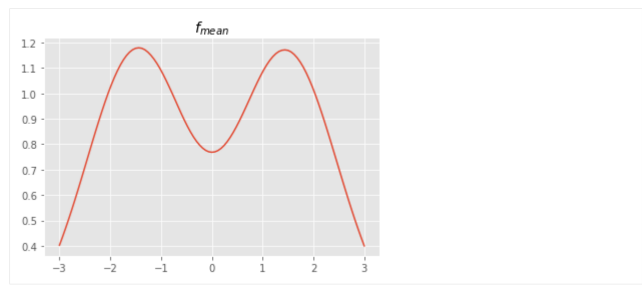
```
[4]: obj.srsf_align(parallel=True)

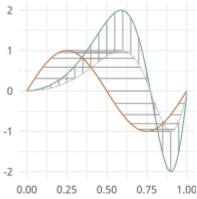
Initializing...
Compute Karcher Mean of 21 function in SRSF space...
updating step: r=1
updating step: r=2
```

Display plots demonstrating the alignment









# 1.2 Elastic Functional Principal Component Analysis

After we have aligned our data we can compute functional principal component analysis (fPCA) on the aligned data, warping functions, and jointly

```
[1]: import fdasrsf as fs import numpy as np
```

We will load in our example data again and compute the alignment

```
[2]: data = np.load('../../bin/simu_data.npz')
   time = data['arr_1']
   f = data['arr_0']
   obj = fs.fdawarp(f,time)
   obj.srsf_align(parallel=True)

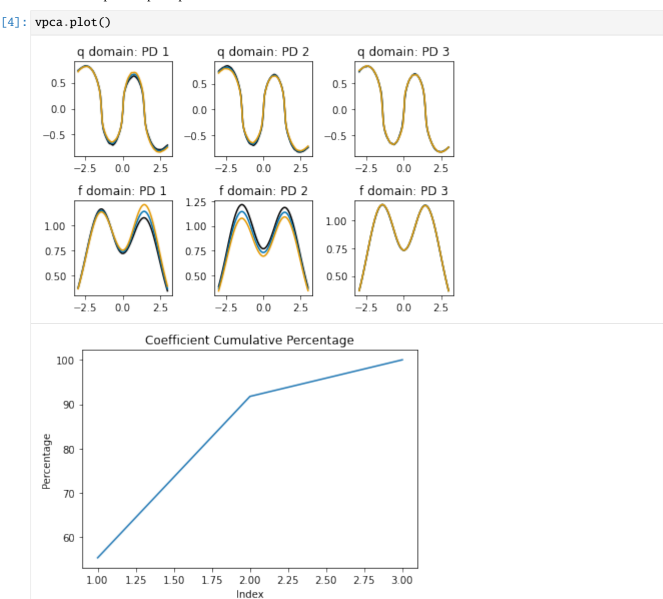
Initializing...
Compute Karcher Mean of 21 function in SRSF space...
   updating step: r=1
   updating step: r=2
```

# 1.2.1 Vertical fPCA

We will first compute fPCA on the aligned functions, by constructing the object and computing the PCA for the number of components, default=3)

[3]: vpca = fs.fdavpca(obj)
vpca.calc\_fpca(no=3)

We then can plot the principal directions

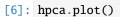


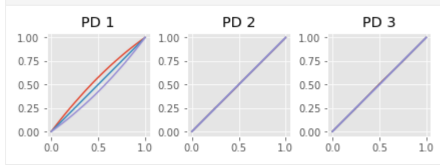
# 1.2.2 Horizontal fPCA

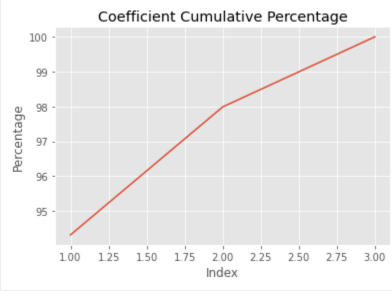
We can then compute PCA on the set of warping functions

[5]: hpca = fs.fdahpca(obj)
hpca.calc\_fpca(no=3)

We then can plot the principal directions







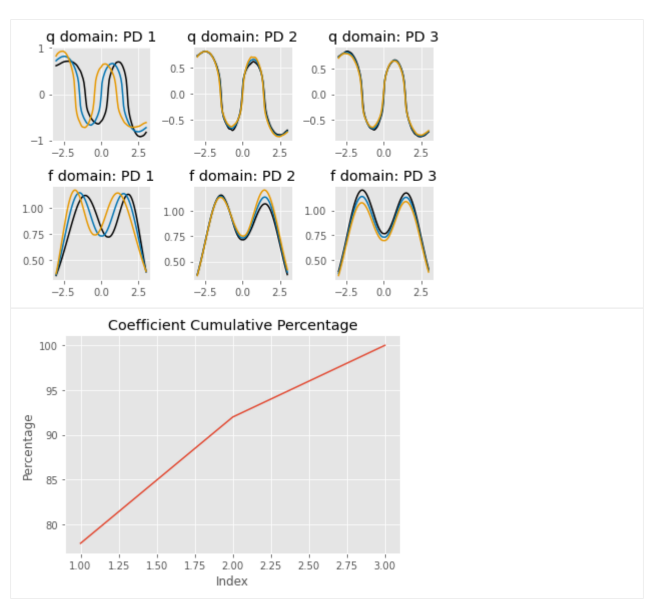
# 1.2.3 Joint fPCA

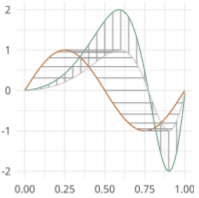
We can also compute the fPCA on jointly on the phase/amplitude space if we feel there is correlation between the variabilities

[7]: jpca = fs.fdajpca(obj) jpca.calc\_fpca(no=3)

We then can plot the principal directions

[8]: jpca.plot()





# 1.3 Multivariate Functional Example

This notebook will show how to use the fdasrsf package to align and statistically analyze a set of multivariate functions using the SRVF framework

# 1.3.1 Load Packages

We will load the required packages and the example data set (MPEG7)

```
[1]: import fdasrsf as fs
  import matplotlib.pyplot as plt
  import numpy as np
  data = np.load('.../bin/gait_data.npz',allow_pickle=True)
  f = data['f']
  g = data['g']
  time = data['time']
```

Now we will construct a 2-D array of a set of 1-D functions from the gait data

```
[2]: M,K = f.shape

beta = np.zeros((2,M,K))
beta[0,:,:] = f
beta[1,:,:] = g
```

# 1.3.2 Analyze

We now will construct a fdacurve object

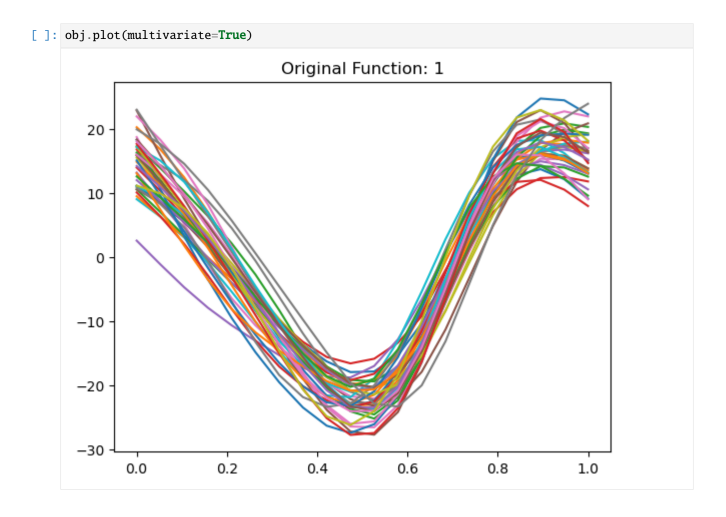
```
[]: obj = fs.fdacurve(beta,N=M)
```

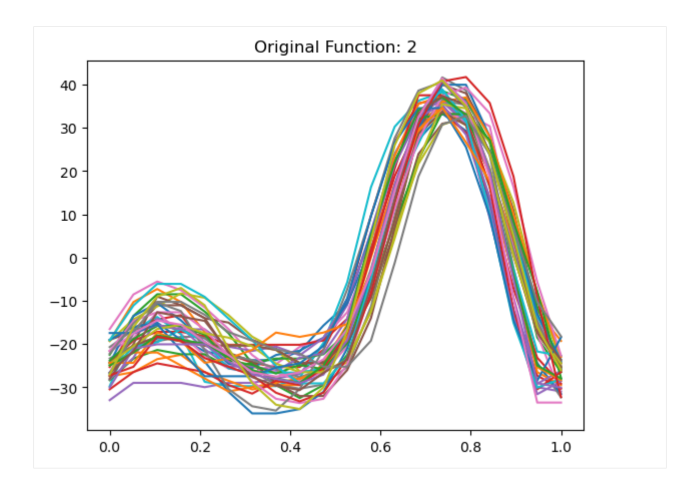
Next, find the Karcher mean and align the curves to the mean

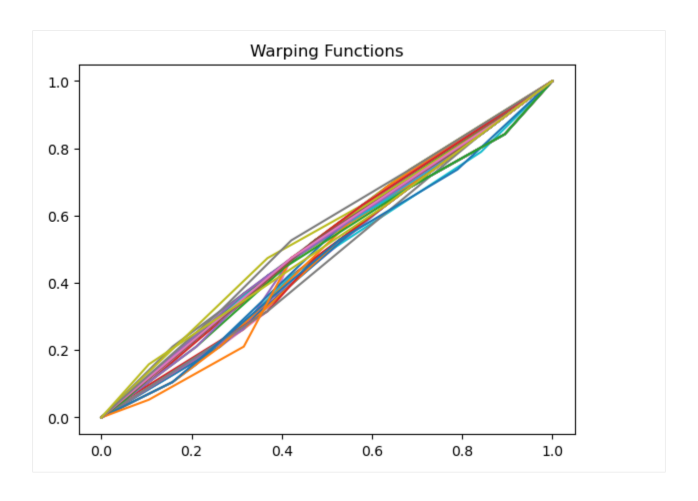
```
[ ]: obj.karcher_mean(rotation=False)
    obj.srvf_align(rotation=False)

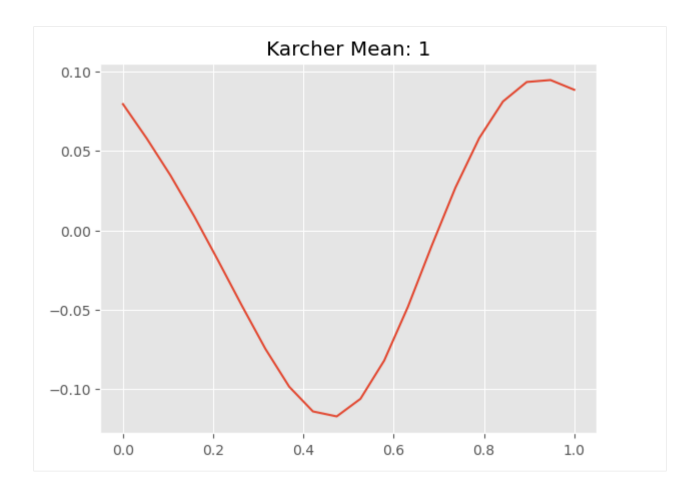
Computing Karcher Mean of 39 curves in SRVF space with lam=0
    updating step: 1
    updating step: 2
    updating step: 3
    updating step: 4
    updating step: 5
    updating step: 6
    updating step: 7
    updating step: 8
    updating step: 9
    updating step: 10
    updating step: 10
```

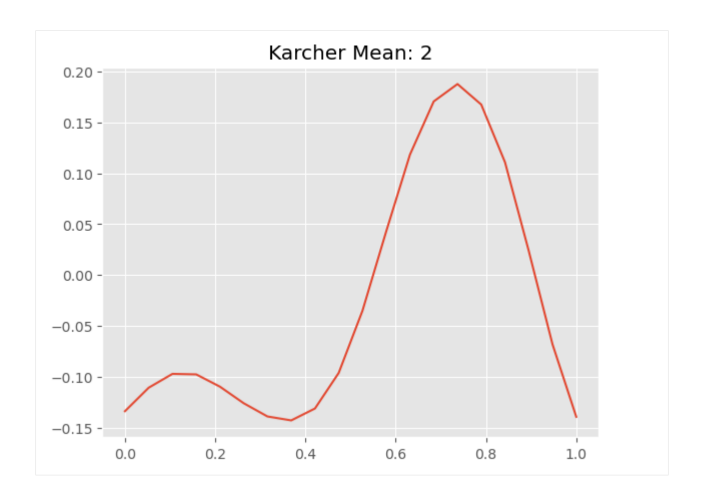
We will now plot the results

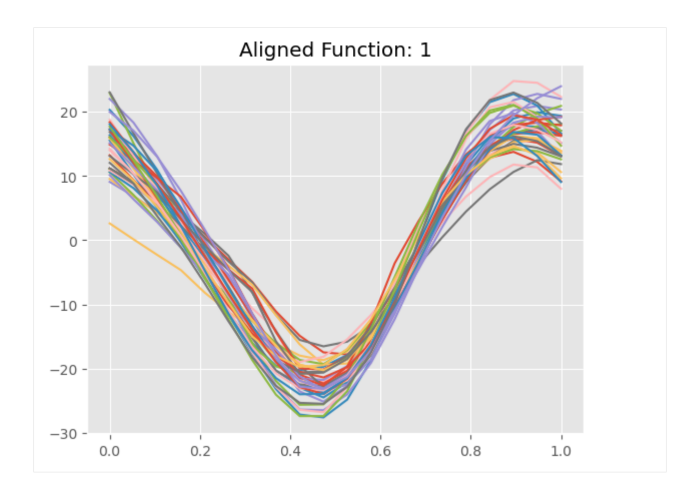


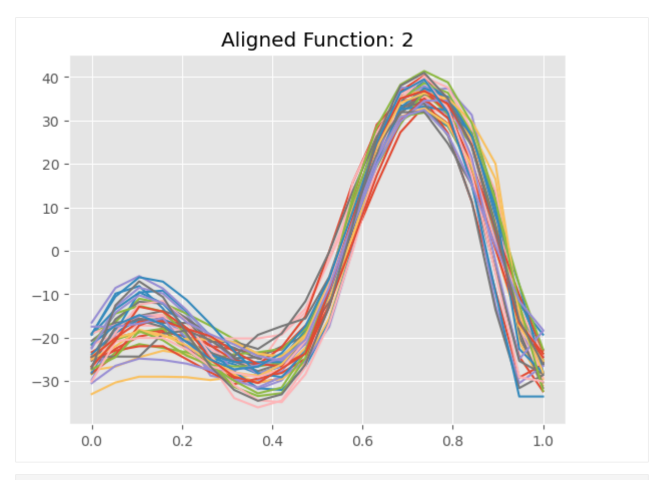




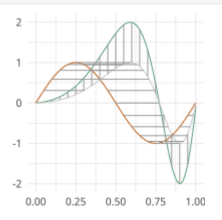








[]:



# 1.4 Elastic Curve Alignment

Otherwise known as time warping in the literature is at the center of elastic functional data analysis. Here our goal is to separate out the horizontal and vertical variability of the open/closed curves

```
[1]: import fdasrsf as fs import numpy as np
```

Load in our example data

```
[2]: data = np.load('../../bin/MPEG7.npz',allow_pickle=True)
    Xdata = data['Xdata']
    curve = Xdata[0,1]
    n,M = curve.shape
    K = Xdata.shape[1]

beta = np.zeros((n,M,K))
    for i in range(0,K):
        beta[:,:,i] = Xdata[0,i]
```

We will then construct the fdacurve object

```
[3]: obj = fs.fdacurve(beta, N=M)
```

We then will compute karcher mean of the curves

```
[4]: obj.karcher_mean()

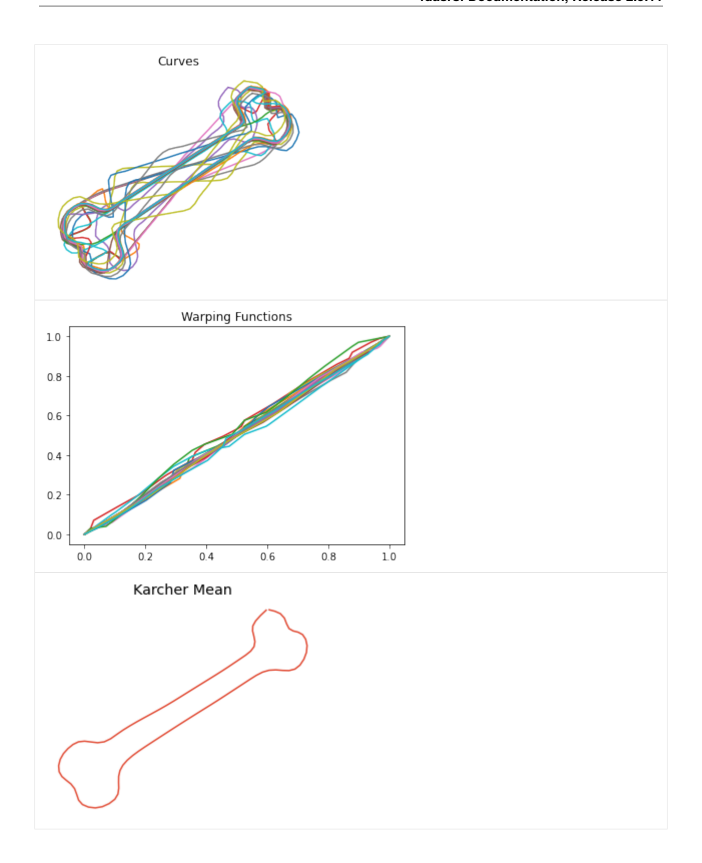
Computing Karcher Mean of 20 curves in SRVF space..
    updating step: 1
    updating step: 2
    updating step: 3
    updating step: 4
    updating step: 5
    updating step: 6
    updating step: 7
```

We then can align the curves to the karcher mean

```
[5]: obj.srvf_align(rotation=False)
```

Plot the results

```
[6]: obj.plot()
```

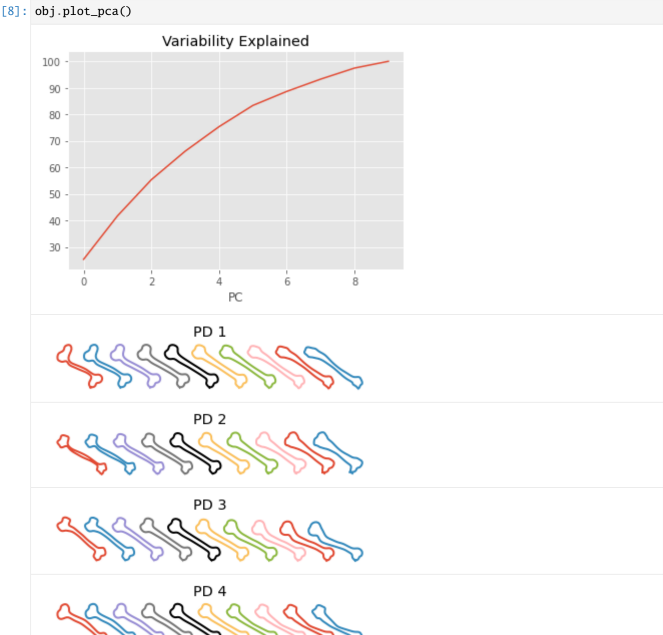


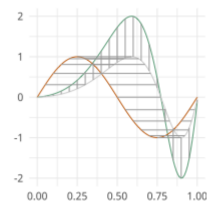
# 1.4.1 Shape PCA

We then can compute the Karcher covariance and compute the shape pca

[7]: obj.karcher\_cov() obj.shape\_pca()

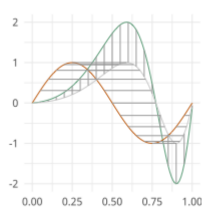
Plot the principal directions





# **TWO**

# **API REFERENCE**



# 2.1 Functional Alignment

Group-wise function alignment using SRSF framework and Dynamic Programming

moduleauthor:: J. Derek Tucker < jdtuck@sandia.gov>

time\_warping.align\_fPCA(f, time, num\_comp=3, showplot=True, smoothdata=False, cores=-1)

aligns a collection of functions while extracting principal components. The functions are aligned to the principal components

#### **Parameters**

- **f** (*np.ndarray*) numpy ndarray of shape (M,N) of N functions with M samples
- time (np.ndarray) vector of size M describing the sample points
- **num\_comp** number of fPCA components
- **showplot** Shows plots of results using matplotlib (default = T)
- **smooth\_data** (*bool*) Smooth the data using a box filter (default = F)
- **cores** number of cores for parallel (default = -1 (all))

#### Return type

tuple of numpy array

#### Return fr

aligned functions - numpy ndarray of shape (M,N) of N functions with M samples

### Return qn

aligned srvfs - similar structure to fn

#### Return q0

original srvf - similar structure to fn

# Return mqn

srvf mean or median - vector of length M

#### Return gam

warping functions - similar structure to fn

#### Return q pca

srsf principal directions

### Return f\_pca

functional principal directions

#### **Return latent**

latent values

#### Return coef

coefficients

### Return U

eigenvectors

# Return orig\_var

Original Variance of Functions

#### Return amp\_var

Amplitude Variance

#### Return phase var

Phase Variance

time\_warping.align\_fPLS(f, g, time, comps=3, showplot=True, smoothdata=False, delta=0.01, max\_itr=100)

This function aligns a collection of functions while performing principal least squares

#### **Parameters**

- **f** (np.ndarray) numpy ndarray of shape (M,N) of N functions with M samples
- g (np.ndarray) numpy ndarray of shape (M,N) of N functions with M samples
- time (np.ndarray) vector of size M describing the sample points
- **comps** number of fPLS components
- **showplot** Shows plots of results using matplotlib (default = T)
- **smooth\_data** (*bool*) Smooth the data using a box filter (default = F)
- **delta** gradient step size
- max\_itr maximum number of iterations

#### Return type

tuple of numpy array

#### Return fn

aligned functions - numpy ndarray of shape (M,N) of N functions with M samples

#### Return gn

aligned functions - numpy ndarray of shape (M,N) of N functions with M samples

#### Return afr

aligned srvfs - similar structure to fn

#### Return qgn

aligned srvfs - similar structure to fn

# Return qf0

original srvf - similar structure to fn

#### Return qg0

original srvf - similar structure to fn

#### Return gam

warping functions - similar structure to fn

### Return wqf

srsf principal weight functions

### Return wqg

srsf principal weight functions

#### Return wf

srsf principal weight functions

#### Return wg

srsf principal weight functions

#### Return cost

cost function value

### class time\_warping.fdawarp(f, time)

This class provides alignment methods for functional data using the SRVF framework

Usage: obj = fdawarp(f,t)

# **Parameters**

- $\mathbf{f} (M,N)$ : matrix defining N functions of M samples
- $\bullet \ \, \text{time} \text{time vector of length} \,\, M$
- **fn** aligned functions
- qn aligned srvfs
- **q0** initial srvfs
- **fmean** Karcher mean
- mqn mean srvf
- gam warping functions
- **psi** srvf of warping functions
- **stats** alignment statistics
- qun cost function
- lambda lambda
- method optimization method
- gamI inverse warping function
- rsamps random samples
- **fs** random aligned functions
- gams random warping functions

- **ft** random warped functions
- qs random aligned srvfs
- **type** alignment type
- mcmc mcmc output if bayesian

Author: J. D. Tucker (JDT) < jdtuck AT sandia.gov > Date: 15-Mar-2018

```
gauss_model(n=1, sort_samples=False)
```

This function models the functional data using a Gaussian model extracted from the principal components of the srvfs

#### **Parameters**

- n (integer) number of random samples
- **sort\_samples** (*bool*) sort samples (default = T)

```
joint_gauss_model(n=1, no=3)
```

This function models the functional data using a joint Gaussian model extracted from the principal components of the srsfs

#### **Parameters**

- n (integer) number of random samples
- **no** (*integer*) number of principal components (default = 3)

```
multiple_align_functions(mu, omethod='DP2', smoothdata=False, parallel=False, lam=0.0, pen='roughness', cores=-1, grid_dim=7)
```

This function aligns a collection of functions using the elastic square-root slope (srsf) framework.

Usage: obj.multiple\_align\_functions(mu)

#### **Parameters**

- **mu** vector of function to align to
- omethod optimization method (DP, DP2, RBFGS, cRBFGS) (default = DP2)
- **smoothdata** (*bool*) Smooth the data using a box filter (default = F)
- parallel run in parallel (default = F)
- lam(double) controls the elasticity (default = 0)
- **penalty** penalty type (default="roughness") options are "roughness", "l2gam", "l2psi", "geodesic". Only roughness implemented in all methods. To use others method needs to be "RBFGS" or "cRBFGS"
- **cores** number of cores for parallel (default = -1 (all))
- **grid\_dim** size of the grid, for the DP2 method only (default = 7)

# plot()

plot functional alignment results

Usage: obj.plot()

srsf\_align(method='mean', omethod='DP2', center=True, smoothdata=False, MaxItr=20, parallel=False,
lam=0.0, pen='roughness', cores=-1, grid\_dim=7, verbose=True)

This function aligns a collection of functions using the elastic square-root slope (srsf) framework.

#### **Parameters**

- **method** (string) warp calculate Karcher Mean or Median (options = "mean" or "median") (default="mean")
- **omethod** optimization method (DP, DP2, RBFGS, cRBFGS) (default = DP2)
- **center** center warping functions (default = T)
- **smoothdata** (bool) Smooth the data using a box filter (default = F)
- MaxItr Maximum number of iterations (default = 20)
- parallel run in parallel (default = F)
- lam (double) controls the elasticity (default = 0)
- **penalty** penalty type (default="roughness") options are "roughness", "l2gam", "l2psi", "geodesic". Only roughness implemented in all methods. To use others method needs to be "RBFGS" or "cRBFGS"
- **cores** number of cores for parallel (default = -1 (all))
- **grid\_dim** size of the grid, for the DP2 method only (default = 7)
- **verbose** print status output (default = T)

Examples >>> import tables >>> fun=tables.open\_file("../Data/simu\_data.h5") >>> f = fun.root.f[:] >>> f = f.transpose() >>> time = fun.root.time[:] >>> obj = fs.fdawarp(f,time) >>> obj.srsf\_align()

#### time\_warping.normal(loc=0.0, scale=1.0, size=None)

Draw random samples from a normal (Gaussian) distribution.

The probability density function of the normal distribution, first derived by De Moivre and 200 years later by both Gauss and Laplace independently<sup>2</sup>, is often called the bell curve because of its characteristic shape (see the example below).

The normal distributions occurs often in nature. For example, it describes the commonly occurring distribution of samples influenced by a large number of tiny, random disturbances, each with its own unique distribution<sup>2</sup>.

**Note:** New code should use the *~numpy.random.Generator.normal* method of a *~numpy.random.Generator* instance instead; please see the random-quick-start.

#### **Parameters**

- loc (float or array\_like of floats) Mean ("centre") of the distribution.
- scale (float or array\_like of floats) Standard deviation (spread or "width") of the distribution. Must be non-negative.
- **size** (*int or tuple of ints, optional*) Output shape. If the given shape is, e.g., (m, n, k), then m \* n \* k samples are drawn. If size is None (default), a single value is returned if loc and scale are both scalars. Otherwise, np.broadcast(loc, scale). size samples are drawn.

#### Returns

out – Drawn samples from the parameterized normal distribution.

#### **Return type**

ndarray or scalar

#### See also:

<sup>&</sup>lt;sup>2</sup> P. R. Peebles Jr., "Central Limit Theorem" in "Probability, Random Variables and Random Signal Principles", 4th ed., 2001, pp. 51, 51, 125.

#### scipy.stats.norm

probability density function, distribution or cumulative density function, etc.

#### random.Generator.normal

which should be used for new code.

#### **Notes**

The probability density for the Gaussian distribution is

$$p(x) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{(x-\mu)^2}{2\sigma^2}},$$

where  $\mu$  is the mean and  $\sigma$  the standard deviation. The square of the standard deviation,  $\sigma^2$ , is called the variance.

The function has its peak at the mean, and its "spread" increases with the standard deviation (the function reaches 0.607 times its maximum at  $x + \sigma$  and  $x - \sigma^2$ ). This implies that normal is more likely to return samples lying close to the mean, rather than those far away.

#### References

# **Examples**

Draw samples from the distribution:

```
>>> mu, sigma = 0, 0.1 # mean and standard deviation
>>> s = np.random.normal(mu, sigma, 1000)
```

Verify the mean and the variance:

0.1 # may vary

```
>>> abs(mu - np.mean(s))
0.0 # may vary
>>> abs(sigma - np.std(s, ddof=1))
```

Display the histogram of the samples, along with the probability density function:

Two-by-four array of samples from the normal distribution with mean 3 and standard deviation 2.5:

```
>>> np.random.normal(3, 2.5, size=(2, 4))
array([[-4.49401501, 4.00950034, -1.81814867, 7.29718677], # random
[ 0.39924804, 4.68456316, 4.99394529, 4.84057254]]) # random
```

time\_warping.pairwise\_align\_bayes(fli, f2i, time, mcmcopts=None)

This function aligns two functions using Bayesian framework. It will align f2 to f1. It is based on mapping warping functions to a hypersphere, and a subsequent exponential mapping to a tangent space. In the tangent space, the Z-mixture pCN algorithm is used to explore both local and global structure in the posterior distribution.

The Z-mixture pCN algorithm uses a mixture distribution for the proposal distribution, controlled by input parameter zpcn. The zpcn\$betas must be between 0 and 1, and are the coefficients of the mixture components, with larger coefficients corresponding to larger shifts in parameter space. The zpcn["probs"] give the probability of each shift size.

Usage: out = pairwise\_align\_bayes(f1i, f2i, time)

#### **Parameters**

- **f1i** vector defining M samples of function 1
- **f2i** vector defining M samples of function 2
- time time vector of length M
- mcmopts dict of mcmc parameters

 $\begin{array}{lll} \mbox{default mcmc options:} & tmp = \{\mbox{"ibetas":np.array}([0.5,0.5,0.005,0.0001]),\mbox{"probs":np.array}([0.1,0.1,0.7,0.1])\} \\ \mbox{mcmcopts} & = \{\mbox{"iter":}2*(10**4) \mbox{,"burnin":np.minimum}(5*(10**3),2*(10**4)//2), \mbox{"alpha0":}0.1,\mbox{"beta0":}0.1,\mbox{"zpcn":tmp,"propvar":}1,\mbox{"initcoef":np.repeat}(0,20),\mbox{"npoints":}200,\mbox{"extrainfo":True}\} \\ \end{array}$ 

:rtype collection containing :return f2\_warped: aligned f2 :return gamma: warping function :return g\_coef: final g\_coef :return psi: final psi :return sigma1: final sigma

if extrainfo :return accept: accept of psi samples :return betas\_ind :return logl: log likelihood :return gamma\_mat: posterior gammas :return gamma\_stats: posterior gamma stats :return xdist: phase distance posterior :return ydist: amplitude distance posterior)

#### time\_warping.pairwise\_align\_bayes\_infHMC(y1i, y2i, time, mcmcopts=None)

This function aligns two functions using Bayesian framework. It uses a hierarchical Bayesian framework assuming mearsurement error error It will align f2 to f1. It is based on mapping warping functions to a hypersphere, and a subsequent exponential mapping to a tangent space. In the tangent space, the infty-HMC algorithm is used to explore both local and global structure in the posterior distribution.

Usage: out = pairwise\_align\_bayes\_infHMC(f1i, f2i, time)

#### **Parameters**

- y1i vector defining M samples of function 1
- y2i vector defining M samples of function 2
- time time vector of length M
- mcmopts dict of mcmc parameters

default mcmc options: mcmcopts = {"iter":1\*(10\*\*4), "nchains":4, "vpriorvar":1, "burnin":np.minimum(5\*(10\*\*3),2\*(10\*\*4)//2), "alpha0":0.1, "beta0":0.1, "alpha":1, "beta":1, "h":0.01, "L":4, "f1propvar":0.0001, "f2propvar":0.0001, "L1propvar":0.3, "L2propvar":0.3, "npoints":200, "thin":1, "sampfreq":1, "initcoef":np.repeat(0,20), "nbasis":10, "basis":'fourier', "extrainfo":True}

Basis can be 'fourier' or 'legendre'

:rtype collection containing :return f2\_warped: aligned f2 :return gamma: warping function :return v\_coef: final v\_coef :return psi: final psi :return sigma1: final sigma

if extrainfo :return theta\_accept: accept of psi samples :return f2\_accept: accept of f2 samples :return SSE: SSE :return gamma\_mat: posterior gammas :return gamma\_stats: posterior gamma stats :return xdist: phase distance posterior :return ydist: amplitude distance posterior)

J. D. Tucker, L. Shand, and K. Chowdhary. "Multimodal Bayesian Registration of Noisy Functions using Hamiltonian Monte Carlo", Computational Statistics and Data Analysis, accepted, 2021.

```
time_warping.pairwise_align_functions(f1, f2, time, omethod='DP2', lam=0, pen='roughness', grid dim=7)
```

# This function aligns f2 to f1 using the elastic square-root

slope (srsf) framework.

Usage: out = pairwise\_align\_functions(f1, f2, time)

#### **Parameters**

- **f1** vector defining M samples of function 1
- **f2** vector defining M samples of function 2
- **time** time vector of length M
- omethod optimization method (DP, DP2, RBFGS, cRBFGS) (default = DP)
- lam controls the elasticity (default = 0)
- **penalty** penalty type (default="roughness") options are "roughness", "l2gam", "l2psi", "geodesic". Only roughness implemented in all methods. To use others method needs to be "RBFGS" or "cRBFGS"
- **grid\_dim** size of the grid, for the DP2 method only (default = 7)

:rtype list containing :return f2n: aligned f2 :return gam: warping function :return q2n: aligned q2 (srsf)

Random values in a given shape.

time\_warping.rand(d0, d1, ..., dn)

**Note:** This is a convenience function for users porting code from Matlab, and wraps *random\_sample*. That function takes a tuple to specify the size of the output, which is consistent with other NumPy functions like *numpy.zeros* and *numpy.ones*.

Create an array of the given shape and populate it with random samples from a uniform distribution over [0, 1).

# **Parameters**

- **d0** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- **d1** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- ... (int, optional) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- **dn** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.

### Returns

out – Random values.

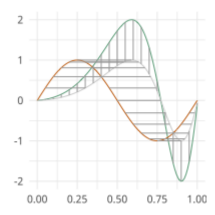
#### **Return type**

ndarray, shape (d0, d1, ..., dn)

#### See also:

random

# **Examples**



# 2.2 Functional Principal Component Analysis

Vertical and Horizontal Functional Principal Component Analysis using SRSF

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

# class fPCA.fdahpca(fdawarp)

This class provides horizontal fPCA using the SRVF framework

Usage: obj = fdahpca(warp\_data)

# **Parameters**

- warp\_data fdawarp class with alignment data
- gam\_pca warping functions principal directions
- psi\_pca srvf principal directions
- latent latent values
- **U** eigenvectors
- coef coefficients
- **vec** shooting vectors
- mu Karcher Mean
- tau principal directions

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 15-Mar-2018

```
calc_fpca(no=3, var_exp=None, stds=array([-1, 0, 1]))
```

This function calculates horizontal functional principal component analysis on aligned data

#### **Parameters**

- **no** (*int*) number of components to extract (default = 3)
- var\_exp compute no based on value percent variance explained (example: 0.95)
- stds number of standard deviations along geodesic to compute (default = -1,0,1)

# Return type

fdahpca object of numpy ndarray

# Return q\_pca

srsf principal directions

#### Return f\_pca

functional principal directions

#### **Return latent**

latent values

#### Return coef

coefficients

#### Return U

eigenvectors

# plot()

plot plot elastic horizontal fPCA results

Usage: obj.plot()

# project(f)

project new data onto fPCA basis

Usage: obj.project(f)

#### **Parameters**

**f** – numpy array (MxN) of N functions on M time

# class fPCA.fdajpca(fdawarp)

This class provides joint fPCA using the SRVF framework

Usage: obj = fdajpca(warp\_data)

#### **Parameters**

- warp\_data fdawarp class with alignment data
- q\_pca srvf principal directions
- **f\_pca** f principal directions
- latent latent values
- coef principal coefficients
- id point used for f(0)
- mqn mean srvf
- **U** eigenvectors
- mu\_psi mean psi

```
• mu_g – mean g
                 • C – scaling value
                 • stds – geodesic directions
      Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 18-Mar-2018
      calc_fpca(no=3, var exp=None, stds=array([-1., 0., 1.]), id=None, parallel=False, cores=-1)
           This function calculates joint functional principal component analysis on aligned data
               Parameters
                    • no (int) – number of components to extract (default = 3)
                    • var_exp – compute no based on value percent variance explained (example: 0.95)
                    • id (int) – point to use for f(0) (default = midpoint)
                    • stds – number of standard deviations along gedoesic to compute (default = -1,0,1)
                    • parallel (bool) – run in parallel (default = F)
                    • cores (int) – number of cores for parallel (default = -1 (all))
               Return type
                    fdajpca object of numpy ndarray
               Return q_pca
                    srsf principal directions
               Return f_pca
                    functional principal directions
               Return latent
                    latent values
               Return coef
                    coefficients
               Return U
                    eigenvectors
      plot()
           plot plot elastic joint fPCA result
           Usage: obj.plot()
      project(f)
           project new data onto fPCA basis
           Usage: obj.project(f)
               Parameters
                    \mathbf{f} – numpy array (MxN) of N functions on M time
class fPCA.fdajpcah(fdawarp)
      This class provides joint fPCA using the SRVF framework using MFPCA
      Usage: obj = fdajpcah(warp_data)
```

2.2. Functional Principal Component Analysis

• q\_pca - srvf principal directions

• warp\_data - fdawarp class with alignment data

- **f\_pca** f principal directions
- latent latent values
- **coef** principal coefficients
- id point used for f(0)
- mqn mean srvf
- **U\_q** eigenvectors for q
- **U\_h** eigenvectors for gam
- C scaling value
- **stds** geodesic directions

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 06-April-2024

calc\_fpca(var\_exp=0.99, stds=array([-1., 0., 1.]), id=None, parallel=False, cores=-1, srsf=True)

This function calculates joint functional principal component analysis on aligned data

#### **Parameters**

- var\_exp compute no based on value percent variance explained (default: None)
- **id** (*int*) point to use for f(0) (default = midpoint)
- stds number of standard deviations along gedoesic to compute (default = -1,0,1)
- **parallel** (*bool*) run in parallel (default = F)
- **cores** (*int*) number of cores for parallel (default = -1 (all))
- **U\_q** eigenvectors for q
- **U\_h** eigenvectors for gam

# **Return type**

fdajpcah object of numpy ndarray

#### Return q\_pca

srsf principal directions

# Return f\_pca

functional principal directions

## **Return latent**

latent values

#### Return coef

coefficients

# plot()

plot plot elastic joint fPCA result

Usage: obj.plot()

## project(f)

project new data onto fPCA basis

Usage: obj.project(f)

## **Parameters**

 $\mathbf{f}$  – numpy array (MxN) of N functions on M time

# class fPCA.fdavpca(fdawarp)

This class provides vertical fPCA using the SRVF framework

Usage: obj = fdavpca(warp\_data)

#### **Parameters**

- warp\_data fdawarp class with alignment data
- **q\_pca** srvf principal directions
- **f\_pca** f principal directions
- latent latent values
- coef principal coefficients
- id point used for f(0)
- mqn mean srvf
- **U** eigenvectors
- stds geodesic directions
- new\_coef principal coefficients of new data

Author: J. D. Tucker (JDT) < jdtuck AT sandia.gov > Date: 15-Mar-2018

```
calc_fpca(no=3, var_exp=None, id=None, stds=array([-1, 0, 1]))
```

This function calculates vertical functional principal component analysis on aligned data

#### **Parameters**

- **no** (*int*) number of components to extract (default = 3)
- var\_exp compute no based on value percent variance explained (example: 0.95)
- **id** (*int*) point to use for f(0) (default = midpoint)
- stds number of standard deviations along geodesic to compute (default = -1,0,1)

## Return type

fdavpca object containing

# Return q\_pca

srsf principal directions

# Return f\_pca

functional principal directions

#### **Return latent**

latent values

#### Return coef

coefficients

## Return U

eigenvectors

#### plot()

plot plot elastic vertical fPCA result Usage: obj.plot()

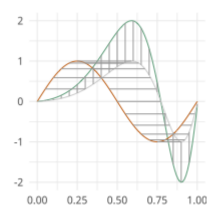
# project(f)

project new data onto fPCA basis

Usage: obj.project(f)

#### **Parameters**

 $\mathbf{f}$  – numpy array (MxN) of N functions on M time



# 2.3 Elastic Functional Boxplots

Elastic Functional Boxplots

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

class boxplots.ampbox(fdawarp)

This class provides amplitude boxplot for functional data using the SRVF framework

Usage: obj = ampbox(warp\_data)

#### **Parameters**

- warp\_data (fdawarp) fdawarp class with alignment data
- Q1 First quartile
- **Q3** Second quartile
- Q1a First quantile based on alpha
- Q3a Second quantile based on alpha
- minn minimum extreme function
- maxx maximum extreme function
- **outlier\_index** indexes of outlier functions
- **f\_median** median function
- q\_median median srvf
- plt surface plot mesh

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 15-Mar-2018

# $construct\_boxplot(alpha=0.05, k\_a=1)$

This function constructs the amplitude boxplot using the elastic square-root slope (srsf) framework.

- **alpha** quantile value (e.g.,=.05, i.e., 95%)
- **k\_a** scalar for outlier cutoff (e.g.,=1)

# plot()

plot box plot and surface plot

Usage: obj.plot()

# class boxplots.phbox(fdawarp)

This class provides phase boxplot for functional data using the SRVF framework

Usage: obj = phbox(warp\_data)

## **Parameters**

- warp\_data (fdawarp) fdawarp class with alignment data
- Q1 First quartile
- **Q3** Second quartile
- Q1a First quantile based on alpha
- Q3a Second quantile based on alpha
- minn minimum extreme function
- maxx maximum extreme function
- **outlier\_index** indexes of outlier functions
- median\_x median warping function
- **psi\_median** median srvf of warping function
- plt surface plot mesh

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 15-Mar-2018

# construct\_boxplot(alpha=0.05, k\_a=1)

This function constructs phase boxplot for functional data using the elastic square-root slope (srsf) framework.

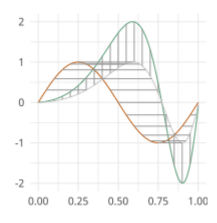
#### **Parameters**

- **alpha** quantile value (e.g.,=.05, i.e., 95%)
- **k\_a** scalar for outlier cutoff (e.g.,=1)

# plot()

plot box plot and surface plot

Usage: obj.plot()



# 2.4 Functional Principal Least Squares

Partial Least Squares using SVD

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

fPLS.pls\_svd(time, qf, qg, no, alpha=0.0)

This function computes the partial least squares using SVD

#### **Parameters**

- **time** vector describing time samples
- $\mathbf{qf}$  numpy ndarray of shape (M,N) of N functions with M samples
- qg numpy ndarray of shape (M,N) of N functions with M samples
- **no** number of components
- **alpha** amount of smoothing (Default = 0.0 i.e., none)

# Return type

numpy ndarray

#### Return wqf

f weight function

## Return wqg

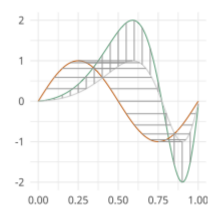
g weight function

# Return alpha

smoothing value

## **Return values**

singular values



# 2.5 Elastic Regression

Warping Invariant Regression using SRSF

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

class regression.elastic\_logistic(f, y, time)

This class provides elastic logistic regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic\_logistic(f,y,time)

#### **Parameters**

- **f** (*np.ndarray*) numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy array of N responses
- time (np.ndarray) vector of size M describing the sample points
- **B** optional matrix describing Basis elements
- alpha alpha parameter of model
- **beta** beta(t) of model
- fn aligned functions numpy ndarray of shape (M,N) of M functions with N samples
- qn aligned srvfs similar structure to fn
- gamma calculated warping functions
- **q** original training SRSFs
- **b** basis coefficients
- Loss logistic loss

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 29-Oct-2021

calc\_model(B=None, lam=0, df=20, max\_itr=20, cores=-1, smooth=False)

This function identifies a regression model with phase-variability using elastic pca

#### **Parameters**

- **B** optional matrix describing Basis elements
- **lam** regularization parameter (default 0)
- **df** number of degrees of freedom B-spline (default 20)
- max\_itr maximum number of iterations (default 20)
- cores number of cores for parallel processing (default all)

# predict(newdata=None)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

#### **Parameters**

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- **time** vector of time points
- **y** truth if available
- smooth smooth data if needed
- **sparam** number of times to run filter

# class regression.elastic\_mlogistic(f, y, time)

This class provides elastic multinomial logistic regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic\_mlogistic(f,y,time)

#### **Parameters**

- **f** (np.ndarray) numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy array of N responses
- time (np.ndarray) vector of size M describing the sample points
- B optional matrix describing Basis elements
- alpha alpha parameter of model
- **beta** beta(t) of model
- fn aligned functions numpy ndarray of shape (M,N) of N functions with M samples
- qn aligned srvfs similar structure to fn
- gamma calculated warping functions
- **q** original training SRSFs
- **b** basis coefficients
- Loss logistic loss

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 29-Oct-2021

calc\_model(B=None, lam=0, df=20, max\_itr=20, delta=0.01, cores=-1, smooth=False)

This function identifies a regression model with phase-variability using elastic pca

#### **Parameters**

- **B** optional matrix describing Basis elements
- **lam** regularization parameter (default 0)
- **df** number of degrees of freedom B-spline (default 20)
- max\_itr maximum number of iterations (default 20)
- cores number of cores for parallel processing (default all)

#### predict(newdata=None)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

#### **Parameters**

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- time vector of time points
- y truth if available
- smooth smooth data if needed
- sparam number of times to run filter

## class regression.elastic\_regression(f, y, time)

This class provides elastic regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic\_regression(f,y,time)

- $\mathbf{f}$  numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy array of N responses
- time vector of size M describing the sample points
- B optional matrix describing Basis elements
- alpha alpha parameter of model
- **beta** beta(t) of model
- fn aligned functions numpy ndarray of shape (M,N) of M functions with N samples
- qn aligned srvfs similar structure to fn
- gamma calculated warping functions
- **q** original training SRSFs
- **b** basis coefficients
- SSE sum of squared error

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 29-Oct-2021

calc\_model(B=None, lam=0, df=20, max\_itr=20, cores=-1, smooth=False)

This function identifies a regression model with phase-variability using elastic pca

#### **Parameters**

- B optional matrix describing Basis elements
- lam regularization parameter (default 0)
- **df** number of degrees of freedom B-spline (default 20)
- max\_itr maximum number of iterations (default 20)
- cores number of cores for parallel processing (default all)

#### predict(newdata=None)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

#### **Parameters**

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- **time** vector of time points
- **y** truth if available
- **smooth** smooth data if needed
- **sparam** number of times to run filter

# $\verb|regression.logistic_warp|(beta, time, q, y)|$

calculates optimal warping for function logistic regression

- beta numpy ndarray of shape (M,N) of N functions with M samples
- $\mbox{time}$  vector of size N describing the sample points

- **q** numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy ndarray of shape (1,N) responses

# Return type

numpy array

#### Return gamma

warping function

## regression.logit\_gradient(b, X, y)

calculates gradient of the logistic loss

#### **Parameters**

- **b** numpy ndarray of shape (M,N) of N functions with M samples
- X numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy ndarray of shape (1,N) responses

## **Return type**

numpy array

#### Return grad

gradient of logistic loss

# regression.logit\_hessian(s, b, X, y)

calculates hessian of the logistic loss

#### **Parameters**

- s numpy ndarray of shape (M,N) of N functions with M samples
- **b** numpy ndarray of shape (M,N) of N functions with M samples
- X numpy ndarray of shape (M,N) of N functions with M samples
- y numpy ndarray of shape (1,N) responses

## Return type

numpy array

# Return out

hessian of logistic loss

# regression. $logit_loss(b, X, y)$

logistic loss function, returns Sum{-log(phi(t))}

#### **Parameters**

- **b** numpy ndarray of shape (M,N) of N functions with M samples
- X numpy ndarray of shape (M,N) of N functions with M samples
- y numpy ndarray of shape (1,N) of N responses

## **Return type**

numpy array

# Return out

loss value

# regression.mlogit\_gradient(b, X, Y)

calculates gradient of the multinomial logistic loss

#### **Parameters**

- **b** numpy ndarray of shape (M,N) of N functions with M samples
- X numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy ndarray of shape (1,N) responses

## **Return type**

numpy array

# Return grad

gradient

## regression.mlogit\_loss(b, X, Y)

calculates multinomial logistic loss (negative log-likelihood)

#### **Parameters**

- **b** numpy ndarray of shape (M,N) of N functions with M samples
- X numpy ndarray of shape (M,N) of N functions with M samples
- y numpy ndarray of shape (1,N) responses

## **Return type**

numpy array

#### Return nll

negative log-likelihood

regression.mlogit\_warp\_grad(alpha, beta, time, q, y, max\_itr=8000, tol=1e-10, delta=0.008, display=0) calculates optimal warping for functional multinomial logistic regression

#### **Parameters**

- alpha scalar
- beta numpy ndarray of shape (M,N) of N functions with M samples
- time vector of size M describing the sample points
- **q** numpy ndarray of shape (M,N) of N functions with M samples
- y numpy ndarray of shape (1,N) responses
- max\_itr maximum number of iterations (Default=8000)
- **tol** stopping tolerance (Default=1e-10)
- **delta** gradient step size (Default=0.008)
- **display** display iterations (Default=0)

#### Return type

tuple of numpy array

## Return gam\_old

warping function

# regression.phi(t)

calculates logistic function, returns  $1/(1 + \exp(-t))$ 

#### **Parameters**

t – scalar

## Return type

numpy array

#### Return out

return value

regression.regression\_warp(beta, time, q, y, alpha)

calculates optimal warping for function linear regression

#### **Parameters**

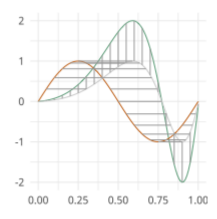
- beta numpy ndarray of shape (M,N) of M functions with N samples
- time vector of size N describing the sample points
- $\mathbf{q}$  numpy ndarray of shape (M,N) of M functions with N samples
- y numpy ndarray of shape (1,N) of M functions with N samples responses
- alpha numpy scalar

# Return type

numpy array

## Return gamma\_new

warping function



# 2.6 Elastic Principal Component Regression

Warping Invariant PCR Regression using SRSF

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

class pcr\_regression.elastic\_lpcr\_regression(f, y, time)

This class provides elastic logistic per regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic\_lpcr\_regression(f,y,time)

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- y response vector of length N (-1/1)

- warp\_data fdawarp object of alignment
- pca class dependent on fPCA method used object of fPCA
- information -
- alpha intercept
- **b** coefficient vector
- Loss logistic loss
- PC probability of classification
- ylabels predicted labels

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 18-Mar-2018

calc\_model(pca\_method='combined', no=5, smooth\_data=False, sparam=25, parallel=False)

This function identifies a logistic regression model with phase-variability using elastic pca

#### **Parameters**

- **pca\_method** string specifying pca method (options = "combined", "vert", or "horiz", default = "combined")
- **no** scalar specify number of principal components (default=5)
- **smooth\_data** smooth data using box filter (default = F)
- **sparam** number of times to apply box filter (default = 25)
- **parallel** calculate in parallel (default = F)

#### predict(newdata=None)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

#### **Parameters**

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- **time** vector of time points
- **y** truth if available
- smooth smooth data if needed
- sparam number of times to run filter

## class pcr\_regression.elastic\_mlpcr\_regression(f, y, time)

This class provides elastic multinomial logistic per regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic\_mlpcr\_regression(f,y,time)

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- y response vector of length N
- **Y** coded label matrix
- warp\_data fdawarp object of alignment

- pca class dependent on fPCA method used object of fPCA
- information -
- alpha intercept
- **b** coefficient vector
- Loss logistic loss
- **PC** probability of classification
- ylabels predicted labels

Author: J. D. Tucker (JDT) < jdtuck AT sandia.gov> Date: 18-Mar-2018

calc\_model(pca\_method='combined', no=5, smooth\_data=False, sparam=25, parallel=False)

This function identifies a logistic regression model with phase-variability using elastic pca

#### **Parameters**

- **f** (*np.ndarray*) numpy ndarray of shape (M,N) of N functions with M samples
- **y** numpy array of N responses
- time (np.ndarray) vector of size M describing the sample points
- **pca\_method** string specifying pca method (options = "combined", "vert", or "horiz", default = "combined")
- **no** scalar specify number of principal components (default=5)
- **smooth\_data** smooth data using box filter (default = F)
- **sparam** number of times to apply box filter (default = 25)
- **parallel** run model in parallel (default = F)

# predict(newdata=None)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

#### **Parameters**

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- **time** vector of time points
- y truth if available
- smooth smooth data if needed
- sparam number of times to run filter

# class pcr\_regression.elastic\_pcr\_regression(f, y, time)

This class provides elastic pcr regression for functional data using the SRVF framework accounting for warping Usage: obj = elastic\_pcr\_regression(f,y,time)

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- y response vector of length N
- warp\_data fdawarp object of alignment

- pca class dependent on fPCA method used object of fPCA
- alpha intercept
- **b** coefficient vector
- SSE sum of squared errors

Author: J. D. Tucker (JDT) < jdtuck AT sandia.gov > Date: 18-Mar-2018

**calc\_model**(pca\_method='combined', no=5, var\_exp=None, smooth\_data=False, sparam=25, parallel=False, C=None)

This function identifies a regression model with phase-variability using elastic pca

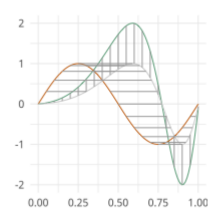
#### **Parameters**

- **pca\_method** string specifying pca method (options = "combined", "vert", or "horiz", default = "combined")
- **no** scalar specify number of principal components (default=5)
- var\_exp compute no based on value percent variance explained (example: 0.95)
- **smooth\_data** smooth data using box filter (default = F)
- **sparam** number of times to apply box filter (default = 25)
- **parallel** run in parallel (default = F)
- **C** scale balance parameter for combined method (default = None)

predict(newdata=None, alpha=0.05)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- **time** vector of time points
- **y** truth if available
- smooth smooth data if needed
- **sparam** number of times to run filter



# 2.7 Elastic Functional Changepoint

Elastic functional change point detection

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

"This class provides elastic changepoint using elastic FDA. It is fully-functional and an extension of the methodology of Aue et al.

Usage: obj = elastic\_amp\_change\_ff(f,time)

#### **Parameters**

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- time time vector of length M
- warp\_data aligned data (default: None)
- **Sn** test statistic values
- Tn max of test statistic
- **p** p-value
- **k\_star** change point
- values values of computed Brownian Bridges
- dat\_a data before changepoint
- dat\_b data after changepoint
- warp\_a warping functions before changepoint
- warp\_b warping functions after changepoint
- mean\_a mean function before changepoint
- mean\_b mean function after changepoint
- warp\_mean\_a mean warping function before changepoint
- warp\_mean\_b mean warping function after changepoint

Author: J. Derek Tucker <jdtuck AT sandia.gov> and Drew Yarger <anyarge AT sandia.gov> Date: 24-Aug-2022

```
compute(d=1000, h=0, M_approx=365, compute_epidemic=False)
```

Compute elastic change detection :param d: number of monte carlo iterations to compute p-value :param h: index of window type to compute long run covariance :param  $M_approx$ : number of time points to compute p-value :param compute\_epidemic: compute epidemic changepoint model (default: False)

#### plot()

plot elastic changepoint results

Usage: obj.plot()

"This class provides elastic changepoint using elastic fpca

Usage: obj = elastic\_change(f,time)

#### **Parameters**

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- time time vector of length M
- **BBridges** precomputed Brownian Bridges (default: None)
- use\_BBridges use precomputed Brownian Bridges (default: False)
- warp\_data aligned data (default: None)
- Sn test statistic values
- Tn max of test statistic
- **p** p-value
- **k\_star** change point
- values values of computed Brownian Bridges
- dat\_a data before changepoint
- dat\_b data after changepoint
- warp\_a warping functions before changepoint
- warp\_b warping functions after changepoint
- mean\_a mean function before changepoint
- **mean\_b** mean function after changepoint
- warp\_mean\_a mean warping function before changepoint
- warp\_mean\_b mean warping function after changepoin

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 27-Apr-2022

**compute**(pca\_method='vert', pc=0.95, d=1000, compute\_epidemic=False, n\_pcs=5, preset\_pcs=False)

Compute elastic change detection

#### **Parameters**

- **pca\_method** string specifying pca method (options = "combined","vert", or "horiz", default = "combined")
- pc percentage of cumulative variance to use (default: 0.95)
- compute\_epidemic compute epidemic changepoint model (default: False)
- **n\_pcs** scalar specify number of principal components (default: 5)
- preset\_pcs use all PCs (default: False)

## plot()

plot elastic changepoint results

Usage: obj.plot()

"This class provides elastic changepoint using elastic FDA on warping functions. It is fully-functional and an extension of the methodology of Aue et al.

Usage: obj = elastic\_ph\_change\_ff(f,time)

#### **Parameters**

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- time time vector of length M
- warp\_data aligned data (default: None)
- **Sn** test statistic values
- Tn max of test statistic
- **p** p-value
- **k\_star** change point
- values values of computed Brownian Bridges
- dat\_a data before changepoint
- dat\_b data after changepoint
- warp\_a warping functions before changepoint
- warp\_b warping functions after changepoint
- mean\_a mean function before changepoint
- **mean\_b** mean function after changepoint
- warp\_mean\_a mean warping function before changepoint
- warp\_mean\_b mean warping function after changepoint

Author: J. Derek Tucker < jdtuck AT sandia.gov > Date: 17-Nov-2022

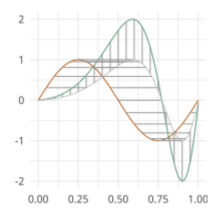
**compute**(*d*=1000, *h*=0, *M*\_approx=365)

Compute elastic change detection :param d: number of monte carlo iterations to compute p-value :param h: index of window type to compute long run covariance :param  $M_approx$ : number of time points to compute p-value

#### plot()

plot elastic changepoint results

Usage: obj.plot()



# 2.8 Elastic GLM Regression

Warping Invariant GML Regression using SRSF

moduleauthor:: Derek Tucker <jdtuck@sandia.gov>

class elastic\_glm\_regression.elastic\_glm\_regression(f, y, time)

This class provides elastic glm regression for functional data using the SRVF framework accounting for warping Usage: obj = elastic\_glm\_regression(f,y,time)

#### **Parameters**

- $\mathbf{f} (M,N)$  % matrix defining N functions of M samples
- y response vector of length N
- time time vector of length M
- alpha intercept
- **b** coefficient vector
- **B** basis matrix
- lambda regularization parameter
- SSE sum of squared errors

Author: J. D. Tucker (JDT) < jdtuck AT sandia.gov > Date: 18-Mar-2018

This function identifies a regression model with phase-variability using elastic pca

- link string of link function ('linear', 'quadratic', 'cubic')
- **B** optional matrix describing Basis elements
- **lam** regularization parameter (default 0)
- **df** number of degrees of freedom B-spline (default 20)
- max\_itr maximum number of iterations (default 20)
- **smooth\_data** smooth data using box filter (default = F)

- **sparam** number of times to apply box filter (default = 25)
- parallel run in parallel (default = F)

# predict(newdata=None, parallel=True)

This function performs prediction on regression model on new data if available or current stored data in object Usage: obj.predict(newdata)

#### **Parameters**

- **newdata** (*dict*) dict containing new data for prediction (needs the keys below, if None predicts on training data)
- $\mathbf{f} (M,N)$  matrix of functions
- **time** vector of time points
- **y** truth if available
- smooth smooth data if needed
- **sparam** number of times to run filter

```
elastic_glm_regression.rand(d0, d1, ..., dn)
```

Random values in a given shape.

**Note:** This is a convenience function for users porting code from Matlab, and wraps *random\_sample*. That function takes a tuple to specify the size of the output, which is consistent with other NumPy functions like *numpy.zeros* and *numpy.ones*.

Create an array of the given shape and populate it with random samples from a uniform distribution over [0, 1).

#### **Parameters**

- **d0** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- **d1** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- ... (int, optional) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- **dn** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.

#### **Returns**

out - Random values.

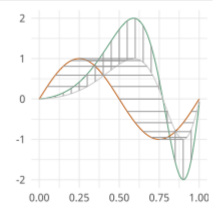
# Return type

ndarray, shape (d0, d1, ..., dn)

#### See also:

random

# **Examples**



# 2.9 Elastic Functional Tolerance Bounds

Functional Tolerance Bounds using SRSF

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

tolerance.bootTB(f, time, a=0.05, p=0.99, B=500, no=5, parallel=True)

This function computes tolerance bounds for functional data containing phase and amplitude variation using bootstrap sampling

#### **Parameters**

- f (np.ndarray) numpy ndarray of shape (M,N) of N functions with M samples
- **time** (*np.ndarray*) vector of size M describing the sample points
- $\mathbf{a}$  confidence level of tolerance bound (default = 0.05)
- $\mathbf{p}$  coverage level of tolerance bound (default = 0.99)
- $\mathbf{B}$  number of bootstrap samples (default = 500)
- **no** number of principal components (default = 5)
- **parallel** enable parallel processing (default = T)

# Return type

tuple of boxplot objects

#### Return amp

amplitude tolerance bounds

## Rtype out\_med

ampbox object

# Return ph

phase tolerance bounds

# Rtype out\_med

phbox object

#### Return out med

alignment results

# Rtype out\_med

fdawarp object

## tolerance.mvtol\_region(x, alpha, P, B)

Computes tolerance factor for multivariate normal

Krishnamoorthy, K. and Mondal, S. (2006), Improved Tolerance Factors for Multivariate Normal Distributions, Communications in Statistics - Simulation and Computation, 35, 461–478.

#### **Parameters**

- $\mathbf{x} (M,N)$  matrix defining N variables of M samples
- alpha confidence level
- P coverage level
- **B** number of bootstrap samples

#### Return type

double

#### Return tol

tolerance factor

# tolerance.pcaTB(f, time, a=0.5, p=0.99, no=5, parallel=True)

This function computes tolerance bounds for functional data containing phase and amplitude variation using fPCA

#### **Parameters**

- **f** (np.ndarray) numpy ndarray of shape (M,N) of N functions with M samples
- time (np.ndarray) vector of size M describing the sample points
- $\mathbf{a}$  confidence level of tolerance bound (default = 0.05)
- $\mathbf{p}$  coverage level of tolerance bound (default = 0.99)
- **no** number of principal components (default = 5)
- parallel enable parallel processing (default = T)

#### Return type

tuple of boxplot objects

# Return warp

alignment data from time\_warping

# Return pca

functional pca from jointFPCA

#### Return tol

tolerance factor

# tolerance.rwishart(df, p)

Computes a random wishart matrix

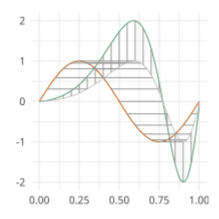
- **df** degree of freedom
- **p** number of dimensions

# Return type

double

#### Return R

matrix



# 2.10 Elastic Functional Clustering

Elastic Functional Clustering

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

kmeans.kmeans\_align(f, time, K, seeds=None, lam=0, showplot=True, smooth\_data=False, parallel=False, alignment=True, omethod='DP2', MaxItr=50, thresh=0.01)

This function clusters functions and aligns using the elastic square-root slope (srsf) framework.

# **Parameters**

- **f** (*np.ndarray*) numpy ndarray of shape (M,N) of N functions with M samples
- time (np.ndarray) vector of size M describing the sample points
- **K** number of clusters
- **seeds** indexes of cluster center functions (default = None)
- lam controls the elasticity (default = 0)
- **showplot** shows plots of functions (default = T)
- **smooth\_data** smooth data using box filter (default = F)
- **parallel** enable parallel mode using code{link{joblib}} and code{doParallel} package (default=F)
- **alignment** whether to perform alignment (default = T)
- **omethod** optimization method (DP,DP2,RBFGS)
- MaxItr maximum number of iterations
- thresh cost function threshold

# Return type

dictionary

## Return fn

aligned functions - matrix (N x M) of M functions with N samples which is a list for each cluster

#### Return qn

aligned SRSFs - similar structure to fn

#### Return q0

original SRSFs

#### **Return labels**

cluster labels

## **Return templates**

cluster center functions

# Return templates\_q

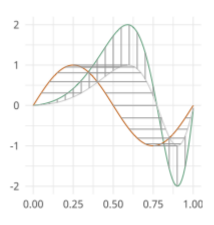
cluster center SRSFs

# Return gam

warping functions - similar structure to fn

#### Return qun

Cost Function



# 2.11 Elastic Image Warping

image warping using SRVF framework

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

image.reparam\_image(It, Im, gam=None, b=None, stepsize=0.0001, itermax=20)

This function warps an image to another using SRVF framework

# **Parameters**

- Im numpy ndarray of shape (N,N) representing a NxN image
- Im numpy ndarray of shape (N,N) representing a NxN image
- gam numpy ndarray of shape (N,N) representing an initial warping function
- **b** numpy ndarray representing basis matrix

#### Return type

numpy ndarray

Return gamnew

diffeomorphism

**Return Inew** 

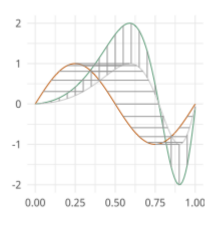
warped image

Return H

energy

**Return stepsize** 

final stepsize



# 2.12 Curve Registration

statistic calculation for SRVF (curves) open and closed using Karcher Mean and Variance

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

**class** curve\_stats.**fdacurve**(beta, mode='O', N=200, scale=False)

This class provides alignment methods for open and closed curves using the SRVF framework

Usage: obj = fdacurve(beta, mode, N, scale) :param beta: numpy ndarray of shape (n, M, N) describing N curves in R^M :param mode: Open ('O') or closed curve ('C') (default 'O') :param N: resample curve to N points :param scale: scale curve to length 1 (true/false) :param q: (n,T,K) matrix defining n dimensional srvf on T samples with K srvfs :param betan: aligned curves :param qn: aligned srvfs :param basis: calculated basis :param beta\_mean: karcher mean curve :param q\_mean: karcher mean srvf :param gams: warping functions :param v: shooting vectors :param C: karcher covariance :param s: pca singular values :param U: pca singular vectors :param coef: pca coefficients :param pca principal directions :param qun: cost function :param lambda: lambda :param samples: random samples :param gamr: random warping functions :param cent: center :param scale: scale :param len: length of curve :param len\_q: length of srvf :param mean\_scale mean length :param mean\_scale q mean length srvf :param E: energy

Author: J. D. Tucker (JDT) <jdtuck AT sandia.gov> Date: 26-Aug-2020

# karcher\_cov()

This calculates the mean of a set of curves

**karcher\_mean**(rotation=True, parallel=False, lam=0.0, cores=-1, method='DP')

This calculates the mean of a set of curves :param rotation: compute optimal rotation (default = T) :param parallel: run in parallel (default = F) :param lam: controls the elasticity (default = 0) :param cores: number of cores for parallel (default = -1 (all)) :param method: method to apply optimization (default="DP") options are "DP" or "RBFGS"

#### plot(multivariate=False)

plot curve mean results

#### **Parameters**

**multivariate** – plot as multivariate functions instead of curves (default=False)

```
sample_shapes(no=3, numSamp=10)
```

Computes sample shapes from mean and covariance

#### **Parameters**

- **no** number of direction (default 3)
- **numSamp** number of samples (default 10)

#### $shape_pca(no=10)$

Computes principal direction of variation specified by no. N is Number of shapes away from mean. Creates 2\*N+1 shape sequence

## **Parameters**

**no** – number of direction (default 3)

```
srvf_align(rotation=True, lam=0.0, parallel=False, cores=- 1, method='DP')
```

This aligns a set of curves to the mean and computes mean if not computed :param rotation: compute optimal rotation (default = T) :param lam: controls the elasticity (default = T) :param parallel: run in parallel (default = T) :param cores: number of cores for parallel (default = T) :param method: method to apply optimization (default="T) options are "T0P" or "RBFGS"

```
curve_stats.randn(d0, d1, ..., dn)
```

Return a sample (or samples) from the "standard normal" distribution.

**Note:** This is a convenience function for users porting code from Matlab, and wraps *standard\_normal*. That function takes a tuple to specify the size of the output, which is consistent with other NumPy functions like *numpy.zeros* and *numpy.ones*.

**Note:** New code should use the *~numpy.random.Generator.standard\_normal* method of a *~numpy.random.Generator* instance instead; please see the random-quick-start.

If positive int\_like arguments are provided, *randn* generates an array of shape (d0, d1, ..., dn), filled with random floats sampled from a univariate "normal" (Gaussian) distribution of mean 0 and variance 1. A single float randomly sampled from the distribution is returned if no argument is provided.

- **d0** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- **d1** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- ... (int, optional) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.
- **dn** (*int*, *optional*) The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.

#### Returns

Z - A (d0, d1, ..., dn)-shaped array of floating-point samples from the standard normal distribution, or a single such float if no parameters were supplied.

## **Return type**

ndarray or float

See also:

#### standard\_normal

Similar, but takes a tuple as its argument.

## normal

Also accepts mu and sigma arguments.

## random.Generator.standard\_normal

which should be used for new code.

#### **Notes**

For random samples from the normal distribution with mean mu and standard deviation sigma, use:

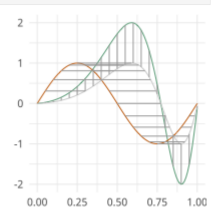
```
sigma * np.random.randn(...) + mu
```

# **Examples**

```
>>> np.random.randn()
2.1923875335537315 # random
```

Two-by-four array of samples from the normal distribution with mean 3 and standard deviation 2.5:

```
>>> 3 + 2.5 * np.random.randn(2, 4)
array([[-4.49401501, 4.00950034, -1.81814867, 7.29718677], # random
[ 0.39924804, 4.68456316, 4.99394529, 4.84057254]]) # random
```



# 2.13 SRVF Geodesic Computation

geodesic calculation for SRVF (curves) open and closed

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

geodesic.back\_parallel\_transport(u1, alpha, basis, T=100, k=5)

backwards parallel translates q1 and q2 along manifold

#### **Parameters**

- **u1** numpy ndarray of shape (2,M) of M samples
- alpha numpy ndarray of shape (2,M) of M samples
- basis list numpy ndarray of shape (2,M) of M samples
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

#### Return type

numpy ndarray

#### Return utilde

translated vector

geodesic.calc\_alphadot(alpha, basis, T=100, k=5)

calculates derivative along the path alpha

## **Parameters**

- alpha numpy ndarray of shape (2,M) of M samples
- basis list of numpy ndarray of shape (2,M) of M samples
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

# Return type

numpy ndarray

#### Return alphadot

derivative of alpha

 $geodesic.calculate\_energy(alphadot, T=100, k=5)$ 

calculates energy along path

## **Parameters**

- alphadot numpy ndarray of shape (2,M) of M samples
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

# **Return type**

numpy scalar

#### Return E

energy

# geodesic.calculate\_gradE(u, utilde, T=100, k=5)

calculates gradient of energy along path

#### **Parameters**

- **u** numpy ndarray of shape (2,M) of M samples
- utilde numpy ndarray of shape (2,M) of M samples
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

# Return type

numpy scalar

## Return gradE

gradient of energy

#### Return normgradE

norm of gradient of energy

# geodesic.cov\_integral(alpha, alphadot, basis, T=100, k=5)

Calculates covariance along path alpha

#### **Parameters**

- alpha numpy ndarray of shape (2,M) of M samples (first curve)
- alphadot numpy ndarray of shape (2,M) of M samples
- basis list numpy ndarray of shape (2,M) of M samples
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

# Return type

numpy ndarray

#### Return u

covariance

# geodesic.find\_basis\_normal\_path(alpha, k=5)

computes orthonormalized basis vectors to the normal space at each of the k points (q-functions) of the path alpha

#### **Parameters**

- **alpha** numpy ndarray of shape (2,M) of M samples (path)
- $\mathbf{k}$  number of samples along path (Default = 5)

# Return type

numpy ndarray

#### Return basis

basis vectors along the path

## geodesic.geod\_dist\_path\_strt(beta, k=5)

calculate geodisc distance for path straightening

# **Parameters**

• beta – numpy ndarray of shape (2,M) of M samples

•  $\mathbf{k}$  – number of samples along path (Default = 5)

#### **Return type**

numpy scalar

## Return dist

geodesic distance

geodesic.geod\_sphere(beta1, beta2, k=5, scale=False, rotation=True, center=True)

This function calculates the geodesics between open curves beta 1 and beta 2 with k steps along path

## **Parameters**

- **beta1** numpy ndarray of shape (2,M) of M samples
- beta2 numpy ndarray of shape (2,M) of M samples
- $\mathbf{k}$  number of samples along path (Default = 5)
- **scale** include length (Default = False)
- **rotation** include rotation (Default = True)
- **center** center curves at origin (Default = True)

# Return type

numpy ndarray

#### Return dist

geodesic distance

#### Return path

geodesic path

# Return PsiO

geodesic path in SRVF

```
geodesic.init_path_geod(beta1, beta2, T=100, k=5)
```

Initializes a path in  $\mathcal{C}$ . beta1, beta2 are already standardized curves. Creates a path from beta1 to beta2 in shape space, then projects to the closed shape manifold.

# **Parameters**

- **beta1** numpy ndarray of shape (2,M) of M samples (first curve)
- **beta2** numpy ndarray of shape (2,M) of M samples (end curve)
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

#### Return type

numpy ndarray

# Return alpha

a path between two q-functions

#### Return beta

a path between two curves

## Return O

rotation matrix

#### geodesic.init\_path\_rand(beta1, beta\_mid, beta2, T=100, k=5)

Initializes a path in C. beta1, beta\_mid beta2 are already standardized curves. Creates a path from beta1 to beta mid to beta2 in shape space, then projects to the closed shape manifold.

#### **Parameters**

- **beta1** numpy ndarray of shape (2,M) of M samples (first curve)
- **betamid** numpy ndarray of shape (2,M) of M samples (mid curve)
- **beta2** numpy ndarray of shape (2,M) of M samples (end curve)
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

# Return type

numpy ndarray

# Return alpha

a path between two q-functions

#### Return beta

a path between two curves

## **Return O**

rotation matrix

# geodesic.path\_straightening(beta1, beta2, betamid=None, init='rand', T=100, k=5)

Perform path straightening to find geodesic between two shapes in either the space of closed curves or the space of affine standardized curves. This algorithm follows the steps outlined in section 4.6 of the manuscript.

#### **Parameters**

- **beta1** numpy ndarray of shape (2,M) of M samples (first curve)
- **beta2** numpy ndarray of shape (2,M) of M samples (end curve)
- **betamid** numpy ndarray of shape (2,M) of M samples (mid curve Default = None, only needed for init "geod")
- init initialize path geodesic or random (Default = "rand")
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

#### Return type

numpy ndarray

#### Return dist

geodesic distance

# Return path

geodesic path

#### Return pathsqnc

geodesic path sequence

# Return E

energy

# geodesic.plot\_geod(path)

Plots the geodesic path as a sequence of curves

#### **Parameters**

path – numpy ndarray of shape (2,M,K) of M sample points of K samples along path

geodesic.update\_path(alpha, beta, gradE, delta, T=100, k=5)

Update the path along the direction -gradE

#### **Parameters**

- alpha numpy ndarray of shape (2,M) of M samples
- **beta** numpy ndarray of shape (2,M) of M samples
- gradE numpy ndarray of shape (2,M) of M samples
- **delta** gradient paramenter
- **T** Number of samples of curve (Default = 100)
- $\mathbf{k}$  number of samples along path (Default = 5)

# Return type

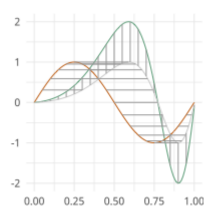
numpy scalar

# Return alpha

updated path of srvfs

#### Return beta

updated path of curves



# 2.14 Utility Functions

Utility functions for SRSF Manipulations

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

utility\_functions.SqrtMean(gam, parallel=False, cores=-1, smooth=True)

calculates the srsf of warping functions with corresponding shooting vectors

- gam numpy ndarray of shape (M,N) of M warping functions with N samples
- parallel run in parallel (default = F)
- **cores** number of cores for parallel (default = -1 (all))
- **smooth** smooth warping functions before gradient calculation

#### **Return type**

2 numpy ndarray and vector

#### Return mu

Karcher mean psi function

#### Return gam mu

vector of dim N which is the Karcher mean warping function

#### Return psi

numpy ndarray of shape (M,N) of M SRSF of the warping functions

#### Return vec

numpy ndarray of shape (M,N) of M shooting vectors

# utility\_functions.SqrtMeanInverse(gam)

finds the inverse of the mean of the set of the diffeomorphisms gamma

#### **Parameters**

gam – numpy ndarray of shape (M,N) of N warping functions with M samples

## **Return type**

vector

# Return gamI

inverse of gam

# utility\_functions.SqrtMedian(gam)

calculates the median srsf of warping functions with corresponding shooting vectors

#### **Parameters**

gam – numpy ndarray of shape (M,N) of M warping functions with N samples

# Return type

2 numpy ndarray and vector

#### Return gam median

Karcher median warping function

# Return psi\_meidan

vector of dim N which is the Karcher median srsf function

# Return psi

numpy ndarray of shape (M,N) of M SRSF of the warping functions

#### Return vec

numpy ndarray of shape (M,N) of M shooting vectors

# utility\_functions.cumtrapzmid(x, y, c, mid)

cumulative trapezoidal numerical integration taken from midpoint

#### **Parameters**

- **x** vector of size N describing the time samples
- **y** vector of size N describing the function
- **c** midpointtic
- **mid** midpiont location

# Return type

vector

#### Return fa

cumulative integration

utility\_functions.diffop(n, binsize=1)

Creates a second order differential operator

#### **Parameters**

- **n** dimension
- **binsize** dx (default = 1)

#### **Return type**

numpy ndarray

## Return m

matrix describing differential operator

utility\_functions.elastic\_depth(f, time, method='DP2', lam=0.0, parallel=True)

calculates the elastic depth between functions in matrix f

#### **Parameters**

- **f** matrix of size MxN (M time points for N functions)
- **time** vector of size M describing the sample points
- **method** method to apply optimization (default="DP2") options are "DP","DP2","RBFGS","cRBFGS"
- lam controls the elasticity (default = 0.0)

# Return type

scalar

# Return amp

amplitude depth

## Return phase

phase depth

utility\_functions.elastic\_distance( $f1, f2, time, method='DP2', lam=0.0, alpha=None, return_dt_only=True$ )

" calculates the distances between function, where f1 is aligned to f2. In other words calculates the elastic distances

## **Parameters**

- **f1** vector of size N
- **f2** vector of size N
- **time** vector of size N describing the sample points
- **method** method to apply optimization (default="DP2") options are "DP","DP2","RBFGS","cRBFGS"
- lam controls the elasticity (default = 0.0)
- alpha makes alpha \* dx + (1-alpha) \* dy
- return\_dt\_only returns only dt if alpha is set

# Return type

scalar

# **Return Dy**

amplitude distance

#### Return Dx

phase distance

#### **Return Dt**

combined distance

## utility\_functions.f\_K\_fold(Nobs, K=5)

generates sample indices for K-fold cross validation

:param Nobs number of observations :param K number of folds

# Return type

numpy ndarray

#### Return train

train indexes (Nobs\*(K-1)/K X K)

#### Return test

test indexes (Nobs\*(1/K) X K)

utility\_functions.f\_to\_srsf(f, time, smooth=False)

converts f to a square-root slope function (SRSF)

#### **Parameters**

- **f** vector of size N samples
- **time** vector of size N describing the sample points

#### **Return type**

vector

#### Return q

srsf of f

# utility\_functions.geigen(Amat, Bmat, Cmat)

generalized eigenvalue problem of the form

max tr L'AM / sqrt(tr L'BL tr M'CM) w.r.t. L and M

:param Amat numpy ndarray of shape (M,N) :param Bmat numpy ndarray of shape (M,N) :param Bmat numpy ndarray of shape (M,N)

#### Return type

numpy ndarray

# **Return values**

eigenvalues

# **Return Lmat**

left eigenvectors

#### **Return Mmat**

right eigenvectors

# utility\_functions.gradient\_spline(time, f, smooth=False)

This function takes the gradient of f using b-spline smoothing

## **Parameters**

• time – vector of size N describing the sample points

- $\mathbf{f}$  numpy ndarray of shape (M,N) of M functions with N samples
- **smooth** smooth data (default = F)

# Return type

tuple of numpy ndarray

#### Return f0

smoothed functions functions

#### Return g

first derivative of each function

#### Return g2

second derivative of each function

# utility\_functions.innerprod\_q(time, q1, q2)

calculates the innerproduct between two srsfs

:param time vector describing time samples :param q1 vector of srsf 1 :param q2 vector of srsf 2

#### Return type

scalar

#### Return val

inner product value

## utility\_functions.invertGamma(gam)

finds the inverse of the diffeomorphism gamma

#### **Parameters**

gam – vector describing the warping function

# Return type

vector

# Return gamI

inverse of gam

utility\_functions.**optimum\_reparam**(q1, time, q2, method='DP2', lam=0.0, penalty='roughness', grid\_dim=7)

calculates the warping to align srsf q2 to q1

# **Parameters**

- q1 vector of size N or array of NxM samples of first SRSF
- time vector of size N describing the sample points
- q2 vector of size N or array of NxM samples samples of second SRSF
- method method to apply optimization (default="DP2") options are "DP","DP2","RBFGS","cRBFGS"
- lam controls the amount of elasticity (default = 0.0)
- **penalty** penalty type (default="roughness") options are "roughness", "l2gam", "l2psi", "geodesic". Only roughness implemented in all methods. To use others method needs to be "RBFGS" or "cRBFGS"
- grid\_dim size of the grid, for the DP2 method only (default = 7)

# Return type

vector

#### Return gam

describing the warping function used to align q2 with q1

utility\_functions.optimum\_reparam\_pair(q, time, q1, q2, lam=0.0)

calculates the warping to align srsf pair q1 and q2 to q

## **Parameters**

- **q** vector of size N or array of NxM samples of first SRSF
- $\bullet$   $\,$  time vector of size N describing the sample points
- q1 vector of size N or array of NxM samples samples of second SRSF
- q2 vector of size N or array of NxM samples samples of second SRSF
- lam controls the amount of elasticity (default = 0.0)

# Return type

vector

# Return gam

describing the warping function used to align q2 with q1

utility\_functions.outlier\_detection(q, time, mq, k=1.5)

calculates outlier's using geodesic distances of the SRSFs from the median

#### **Parameters**

- **q** numpy ndarray of N x M of M SRS functions with N samples
- time vector of size N describing the sample points
- mq median calculated using time\_warping.srsf\_align()
- $\mathbf{k}$  cutoff threshold (default = 1.5)

#### Returns

q\_outlier: outlier functions

utility\_functions.randomGamma(gam, num)

generates random warping functions

#### **Parameters**

- gam numpy ndarray of N x M of M of warping functions
- **num** number of random functions

#### Returns

rgam: random warping functions

utility\_functions.resamplefunction(x, n)

resample function using n points

#### **Parameters**

- **x** functions
- **n** number of points

#### **Return type**

numpy array

#### Return xn

resampled function

utility\_functions.rgam(N, sigma, num, mu\_gam=None)

Generates random warping functions

#### **Parameters**

- N length of warping function
- **sigma** variance of warping functions
- **num** number of warping functions

:param mu\_gam mean warping function (default identity) :return: gam: numpy ndarray of warping functions

```
utility_functions.smooth_data(f, sparam=1)
```

This function smooths a collection of functions using a box filter

#### **Parameters**

- **f** numpy ndarray of shape (M,N) of M functions with N samples
- **sparam** Number of times to run box filter (default = 25)

#### Return type

numpy ndarray

#### Return f

smoothed functions functions

```
utility_functions.srsf_to_f(q, time, f0=0.0)
```

converts q (srsf) to a function

#### **Parameters**

- **q** vector of size N samples of srsf
- **time** vector of size N describing time sample points
- **f0** initial value

## **Return type**

vector

#### Return f

function

# utility\_functions.svd\_flip(u, v, u\_based\_decision=True)

Sign correction to ensure deterministic output from SVD.

Adjusts the columns of u and the rows of v such that the loadings in the columns in u that are largest in absolute value are always positive.

If u\_based\_decision is False, then the same sign correction is applied to so that the rows in v that are largest in absolute value are always positive.

#### **Parameters**

- $\mathbf{u}$  (*ndarray*) Parameters u and v are the output of *linalg.svd* or randomized\_svd(), with matching inner dimensions so one can compute np.dot(u \* s, v).
- $\mathbf{v}$  (ndarray) Parameters u and v are the output of linalg.svd or randomized\_svd(), with matching inner dimensions so one can compute np.dot(u \* s, v). The input v should really be called vt to be consistent with scipy's output.
- u\_based\_decision (bool, default=True) If True, use the columns of u as the basis for sign flipping. Otherwise, use the rows of v. The choice of which variable to base the decision on is generally algorithm dependent.

#### Returns

- u\_adjusted (ndarray) Array u with adjusted columns and the same dimensions as u.
- v\_adjusted (ndarray) Array v with adjusted rows and the same dimensions as v.

# utility\_functions.update\_progress(progress)

This function creates a progress bar

#### **Parameters**

progress - fraction of progress

utility\_functions.warp\_f\_gamma(time, f, gam)

warps a function f by gam

:param time vector describing time samples :param q vector describing srsf :param gam vector describing warping function

# Return type

numpy ndarray

#### Return f temp

warped srsf

utility\_functions.warp\_q\_gamma(time, q, gam)

warps a srsf q by gam

:param time vector describing time samples :param q vector describing srsf :param gam vector describing warping function

#### Return type

numpy ndarray

# Return q\_temp

warped srsf

 $\verb|utility_functions.zero_crossing|(Y, q, bt, time, y_max, y_min, gmax, gmin)|$ 

finds zero-crossing of optimal gamma, gam = s\*gmax + (1-s)\*gmin from elastic regression model

#### **Parameters**

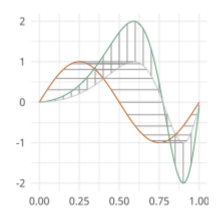
- Y response
- q predicitve function
- bt basis function
- **time** time samples
- **y\_max** maximum repsonse for warping function gmax
- **y\_min** minimum response for warping function gmin
- gmax max warping function
- gmin min warping fucntion

#### Return type

numpy array

# Return gamma

optimal warping function



# 2.15 Curve Functions

functions for SRVF curve manipulations

moduleauthor:: J. Derek Tucker <jdtuck@sandia.gov>

curve\_functions.Basis\_Normal\_A(q)

Find Normal Basis

#### **Parameters**

**q** – numpy ndarray (n,T) defining T points on n dimensional SRVF

:rtype list :return delg: basis

curve\_functions.calc\_j(basis)

Calculates Jacobian matrix from normal basis

#### **Parameters**

**basis** – list of numpy ndarray of shape (2,M) of M samples basis

# **Return type**

numpy ndarray

# Return j

Jacobian

# curve\_functions.calculate\_variance(beta)

This function calculates variance of curve beta

#### **Parameters**

**beta** – numpy ndarray of shape (2,M) of M samples

# Return type

numpy ndarray

# **Return variance**

variance

# curve\_functions.calculatecentroid(beta)

This function calculates centroid of a parameterized curve

# **Parameters**

beta – numpy ndarray of shape (2,M) of M samples

# **Return type**

numpy ndarray

#### Return centroid

center coordinates

curve\_functions.curve\_to\_q(beta, mode='O', scale=True)

This function converts curve beta to srvf q

#### **Parameters**

- **beta** numpy ndarray of shape (2,M) of M samples
- mode Open ('O') or closed curve ('C') (default 'O')
- **scale** scale curve to unit length (default = True)

# Return type

numpy ndarray

#### Return q

srvf of curve

#### Return lenb

length of curve

# Return leng

length of srvf

 $curve\_functions.curve\_zero\_crossing(Y, q, bt, y\_max, y\_min, gmax, gmin)$ 

finds zero-crossing of optimal gamma, gam = s\*gmax + (1-s)\*gmin from elastic curve regression model

#### **Parameters**

- Y response
- beta predicitve function
- **bt** basis function
- **y\_max** maximum repsonse for warping function gmax
- **y\_min** minimum response for warping function gmin
- gmax max warping function
- gmin min warping fucntion

# Return type

numpy array

## Return gamma

optimal warping function

## Return O\_hat

rotation matrix

curve\_functions.elastic\_distance\_curve(beta1, beta2, closed=0, rotation=True, scale=False, method='DP')

Calculates the two elastic distances between two curves in R^M :param beta1: numpy ndarray of shape (M,N) of N samples :param beta2: numpy ndarray of shape (M,N) of N samples :param closed: open (0) or closed (1) curve (default=0) :param rotation: compute optimal rotation (default=True) :param scale: include scale (default=False) :param method: method to apply optimization (default="DP") options are "DP" or "RBFGS"

#### Return type

tuple

2.15. Curve Functions 73

#### Return dist

shape distance

#### Return dx

phase distance

curve\_functions.elastic\_shooting(q1, v, mode=0)

Calculates shooting vector from v to q1

#### **Parameters**

- q1 vector of srvf
- **v** shooting vector
- mode closed or open (1/0)

:rtype numpy ndarray :return q2n: vector of srvf

curve\_functions.elastic\_shooting\_vector(q1, q2, mode=0)

Calculates shooting between two srvfs

#### **Parameters**

- q1 vector of srvf
- q2 vector of srvf
- mode closed or open (1/0)

:rtype numpy ndarray :return v: shooting vector :return d: distance :return q2n: aligned srvf

# curve\_functions.find\_basis\_normal(q)

Finds the basis normal to the srvf

## **Parameters**

**q1** – numpy ndarray of shape (2,M) of M samples

## Return type

list of numpy ndarray

#### Return basis

list containing basis vectors

curve\_functions.find\_best\_rotation(q1, q2, allow\_reflection=False, only\_xy=False)

This function calculates the best rotation between two srvfs using procustes rigid alignment

#### **Parameters**

- q1 numpy ndarray of shape (2,M) of M samples
- **q2** numpy ndarray of shape (2,M) of M samples
- **allow\_reflection** bool indicating if reflection is allowed (i.e. if the determinant of the optimal rotation can be -1)
- only\_xy bool indicating if rotation should only be allowed in the first two dimensions of the space

# Return type

numpy ndarray

# Return q2new

optimal rotated q2 to q1

#### Return R

rotation matrix

curve\_functions.find\_rotation\_and\_seed\_coord(beta1, beta2, closed=0, rotation=True, method='DP')

This function returns a candidate list of optimally oriented and registered (seed) shapes w.r.t. beta1

#### **Parameters**

- beta1 numpy ndarray of shape (2,M) of M samples
- beta2 numpy ndarray of shape (2,M) of M samples
- closed Open (0) or Closed (1)
- rotation find rotation (default=True)
- method method to apply optimization (default="DP") options are "DP" or "RBFGS"

#### Return type

numpy ndarray

#### Return beta2new

optimal aligned beta2 to beta1

#### Return q2best

optimal aligned q2 to q1

# **Return Rbest**

rotation matrix

#### Return gamIbest

warping function

curve\_functions.find\_rotation\_and\_seed\_q(q1, q2, closed=0, rotation=True, method='DP')

This function returns a candidate list of optimally oriented and registered (seed) srvs w.r.t. q1

#### **Parameters**

- q1 numpy ndarray of shape (2,M) of M samples
- **q2** numpy ndarray of shape (2,M) of M samples
- closed Open (0) or Closed (1)
- **rotation** find rotation (default=True)
- method method to apply optimization (default="DP") options are "DP" or "RBFGS"

## **Return type**

numpy ndarray

#### Return q2best

optimal aligned q2 to q1

#### **Return Rbest**

rotation matrix

#### Return gamIbest

warping function

curve\_functions.find\_rotation\_and\_seed\_unique(q1, q2, closed=0, lam=0.0, rotation=True, method='DP')

This function returns a candidate list of optimally oriented and registered (seed) shapes w.r.t. beta1

# **Parameters**

2.15. Curve Functions 75

- **beta1** numpy ndarray of shape (2,M) of M samples
- beta2 numpy ndarray of shape (2,M) of M samples
- closed Open (0) or Closed (1)
- **rotation** find rotation (default=True)
- lam controls the elasticity (default = 0)
- **method** method to apply optimization (default="DP") options are "DP" or "RBFGS"

# Return type

numpy ndarray

#### Return beta2new

optimal rotated beta2 to beta1

## **Return O**

rotation matrix

# Return tau

seed

# curve\_functions.gram\_schmidt(basis)

Performs Gram Schmidt Orthogonlization of a basis\_o

#### param basis

list of numpy ndarray of shape (2,M) of M samples

#### rtype

list of numpy ndarray

#### return basis\_o

orthogonlized basis

# $curve\_functions.group\_action\_by\_gamma(q, gamma)$

This function reparamerized srvf  $\boldsymbol{q}$  by gamma

#### **Parameters**

- $\mathbf{f}$  numpy ndarray of shape (2,M) of M samples
- gamma numpy ndarray of shape (2,M) of M samples

# Return type

numpy ndarray

## Return qn

reparatermized srvf

# curve\_functions.group\_action\_by\_gamma\_coord(f, gamma)

This function reparamerized curve f by gamma

#### **Parameters**

- **f** numpy ndarray of shape (2,M) of M samples
- gamma numpy ndarray of shape (2,M) of M samples

# Return type

numpy ndarray

#### Return fn

reparatermized curve

# curve\_functions.innerprod\_q2(q1, q2)

This function calculates the inner product in srvf space

#### **Parameters**

- q1 numpy ndarray of shape (2,M) of M samples
- **q2** numpy ndarray of shape (2,M) of M samples

## **Return type**

numpy ndarray

#### Return val

inner product

# curve\_functions.inverse\_exp(q1, q2, beta2)

Calculate the inverse exponential to obtain a shooting vector from q1 to q2 in shape space of open curves

#### **Parameters**

- **q1** numpy ndarray of shape (2,M) of M samples
- **q2** numpy ndarray of shape (2,M) of M samples
- **beta2** numpy ndarray of shape (2,M) of M samples

#### **Return type**

numpy ndarray

#### Return v

shooting vectors

## curve\_functions.inverse\_exp\_coord(beta1, beta2, closed=0, method='DP')

Calculate the inverse exponential to obtain a shooting vector from beta1 to beta2 in shape space of open curves

#### **Parameters**

- beta1 numpy ndarray of shape (2,M) of M samples
- beta2 numpy ndarray of shape (2,M) of M samples
- closed open (0) or closed (1) curve
- method method to apply optimization (default="DP") options are "DP" or "RBFGS"

# Return type

numpy ndarray

#### Return v

shooting vectors

# Return dist

distance

# curve\_functions.optimum\_reparam\_curve(q1, q2, lam=0.0, method='DP')

calculates the warping to align srsf q2 to q1

#### **Parameters**

- q1 matrix of size nxN or array of NxM samples of first SRVF
- time vector of size N describing the sample points
- q2 matrix of size nxN or array of NxM samples samples of second SRVF
- lam controls the amount of elasticity (default = 0.0)

2.15. Curve Functions 77

• method – method to apply optimization (default="DP") options are "DP" or "RBFGS"

# Return type

vector

# Return gam

describing the warping function used to align q2 with q1

curve\_functions.parallel\_translate(w, q1, q2, basis, mode=0)

parallel translates q1 and q2 along manifold

#### **Parameters**

- w numpy ndarray of shape (2,M) of M samples
- q1 numpy ndarray of shape (2,M) of M samples
- q2 numpy ndarray of shape (2,M) of M samples
- basis list of numpy ndarray of shape (2,M) of M samples
- mode open 0 or closed curves 1 (default 0)

# Return type

numpy ndarray

#### Return wbar

translated vector

curve\_functions.pre\_proc\_curve(beta, T=100)

This function prepcoessed a curve beta to set of closed curves

#### **Parameters**

- beta numpy ndarray of shape (2,M) of M samples
- T number of samples (default = 100)

#### Return type

numpy ndarray

# **Return betanew**

projected beta

# Return qnew

projected srvf

#### Return A

alignment matrix (not used currently)

curve\_functions.project\_curve(q)

This function projects srvf q to set of close curves

# **Parameters**

**q** – numpy ndarray of shape (2,M) of M samples

## **Return type**

numpy ndarray

# Return qproj

project srvf

curve\_functions.project\_tangent(w, q, basis)

projects srvf to tangent space w using basis

#### **Parameters**

- w numpy ndarray of shape (2,M) of M samples
- **q** numpy ndarray of shape (2,M) of M samples
- basis list of numpy ndarray of shape (2,M) of M samples

#### Return type

numpy ndarray

# Return wproj

projected q

curve\_functions.**psi**(x, a, q)

This function formats variance output

#### **Parameters**

- **x** numpy ndarray of shape (2,M) of M samples curve
- **a** numpy ndarray of shape (2,1) mean
- **q** numpy ndarray of shape (2,M) of M samples srvf

# Return type

numpy ndarray

## Return psi1

variance

#### Return psi2

cross variance

# Return psi3

curve end

# Return psi4

curve end

curve\_functions.q\_to\_curve(q, scale=1)

This function converts srvf to beta

#### **Parameters**

- **q** numpy ndarray of shape (n,M) of M samples
- scale scale of curve

#### **Return type**

numpy ndarray

# Return beta

parameterized curve

curve\_functions.resamplecurve(x, N=100, time=None, mode='O')

This function resamples a curve to have N samples

## **Parameters**

- $\mathbf{x}$  numpy ndarray of shape (2,M) of M samples
- N Number of samples for new curve (default = 100)
- time timing vector (Default=None)
- mode Open ('O') or closed curve ('C') (default 'O')

2.15. Curve Functions 79

# Return type

numpy ndarray

#### Return xn

resampled curve

# curve\_functions.scale\_curve(beta)

scales curve to length 1

#### **Parameters**

beta – numpy ndarray of shape (2,M) of M samples

## **Return type**

numpy ndarray

# Return beta\_scaled

scaled curve

#### Return scale

scale factor used

# curve\_functions. $shift_f(f, tau)$

shifts a curve f by tau

## **Parameters**

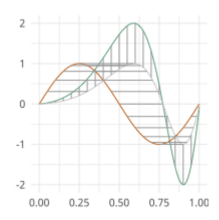
- $\mathbf{f}$  numpy ndarray of shape (2,M) of M samples
- tau scalar

# Return type

numpy ndarray

# Return fn

shifted curve



# 2.16 UMAP EFDA Metrics

Distance metrics for functions and curves in R^n for use with UMAP (https://github.com/lmcinnes/umap)

moduleauthor:: J. Derek Tucker < jdtuck@sandia.gov>

umap\_metric.efda\_distance(q1, q2, alpha=0)

" calculates the distances between two curves, where q2 is aligned to q1. In other words calculates the elastic distances/ This metric is set up for use with UMAP or t-sne from scikit-learn

## **Parameters**

- q1 vector of size N
- $\mathbf{q2}$  vector of size N
- **alpha** weight between phase and amplitude (default = 0, returns amplitude)

# Return type

scalar

## Return dist

amplitude distance

# umap\_metric.efda\_distance\_curve(beta1, beta2, closed)

" calculates the distances between two curves, where beta2 is aligned to beta1. In other words calculates the elastic distance. This metric is set up for use with UMAP or t-sne from scikit-learn

#### **Parameters**

- beta1 vector of size n\*M
- beta2 vector of size n\*M
- closed -
- (0) if open curves and (1) if closed curves

# Return type

scalar

# Return dist

shape distance

**CHAPTER** 

**THREE** 

# **INSTALLATION**

Currently, *fdasrsf* is available in Python versions above 3.8, regardless of the platform. The stable version can be installed via PyPI:

pip install fdasrsf

It is also available from conda-forge:

conda install -c conda-forge fdasrsf

It is possible to install the latest version of the package, available in the develop branch, by cloning this repository and doing a manual installation.

git clone https://github.com/jdtuck/fdasrsf\_python.git
pip install ./fdasrsf\_python

In this type of installation make sure that your default Python version is currently supported, or change the python and pip commands by specifying a version, such as python3.8.

# CHAPTER FOUR

# **HOW DO I START?**

If you want a quick overview of the package, we recommend you to look at the example notebooks in the *Users Guide* 

# **CHAPTER**

# **FIVE**

# **CONTRIBUTIONS**

All contributions are welcome. You can help this project grow in multiple ways, from creating an issue, reporting an improvement or a bug, to doing a repository fork and creating a pull request to the development branch.

# **CHAPTER**

SIX

# **LICENSE**

The package is licensed under the BSD 3-Clause License. A copy of the license can be found along with the code or in the project page.

90 Chapter 6. License

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# **CHAPTER**

# **EIGHT**

# **INDICES AND TABLES**

- genindex
- modindex
- search

# **PYTHON MODULE INDEX**

```
b
boxplots, 36
С
curve_functions, 72
curve_stats, 57
е
elastic_changepoint, 48
{\tt elastic\_glm\_regression}, {\tt 51}
fPCA, 31
fPLS, 38
geodesic, 60
image, 56
k
kmeans, 55
pcr_regression, 44
regression, 38
time_warping, 23
tolerance, 53
u
umap_metric, 80
utility_functions, 64
```

96 Python Module Index

# **INDEX**

A	<pre>construct_boxplot() (boxplots.ampbox method), 36</pre>
align_fPCA() (in module time_warping), 23 align_fPLS() (in module time_warping), 24 ampbox (class in boxplots), 36	<pre>construct_boxplot() (boxplots.phbox method), 37 cov_integral() (in module geodesic), 61 cumtrapzmid() (in module utility_functions), 65 curve_functions</pre>
В	module, 72
back_parallel_transport() (in module geodesic), 60 Basis_Normal_A() (in module curve_functions), 72 bootTB() (in module tolerance), 53 boxplots module, 36	<pre>curve_stats    module, 57 curve_to_q() (in module curve_functions), 73 curve_zero_crossing() (in module curve_functions),</pre>
C	D
calc_alphadot() (in module geodesic), 60	diffop() (in module utility_functions), 66
<pre>calc_fpca() (fPCA.fdahpca method), 31 calc_fpca() (fPCA.fdajpca method), 33 calc_fpca() (fPCA.fdajpcah method), 34 calc_fpca() (fPCA.fdavpca method), 35 calc_j() (in module curve_functions), 72 calc_model() (elastic_glm_regression.elastic_glm_regression_method), 51 calc_model() (pcr_regression.elastic_lpcr_regression_method), 45 calc_model() (pcr_regression.elastic_mlpcr_regression_method), 46 calc_model() (pcr_regression.elastic_pcr_regression_method), 47 calc_model() (regression.elastic_logistic method), 39</pre>	elastic_change (class in elastic_changepoint), 48 elastic_changepoint     module, 48 elastic_depth() (in module utility_functions), 66 elastic_distance() (in module utility_functions), 66 elastic_distance_curve() (in module
<pre>calc_model() (regression.elastic_mlogistic method), 40 calc_model() (regression.elastic_regression method),</pre>	module, 51 elastic_glm_regression (class in elastic_glm_regression), 51 elastic_logistic (class in regression), 38
<pre>calculate_energy() (in module geodesic), 60 calculate_gradE() (in module geodesic), 60 calculate_variance() (in module curve_functions),</pre>	elastic_lpcr_regression (class in pcr_regression), 44
72 calculatecentroid() (in module curve_functions), 72 compute() (elastic_changepoint.elastic_amp_change_ff	elastic_mlogistic (class in regression), 39 elastic_mlpcr_regression (class in pcr_regression), 45 elastic_pcr_regression (class in pcr_regression), 46 elastic_ph_change_ff (class in elastic_changepoint), 49 elastic_regression (class in regression), 40
<pre>compute() (elastic_changepoint.elastic_ph_change_ff</pre>	elastic_shooting() (in module curve_functions), 74

elastic_shooting_vector() (in module	J
curve_functions), 74	<pre>joint_gauss_model() (time_warping.fdawarp</pre>
F	method), 26
f_K_fold() (in module utility_functions), 67	K
f_to_srsf() (in module utility_functions), 67	karcher_cov() (curve_stats.fdacurve method), 57
fdacurve (class in curve_stats), 57	karcher_mean() (curve_stats.fdacurve method), 57
Edahpca (class in fPCA), 31	kmeans
Edajpca (class in fPCA), 32	module, 55
fdajpcah (class in fPCA), 33	kmeans_align() (in module kmeans), 55
Edavpca (class in fPCA), 34	killedits_allgit() (in module killedits), 33
Edawarp (class in time_warping), 25	1
find_basis_normal() (in module curve_functions), 74	
find_basis_normal_path() (in module geodesic), 61	logistic_warp() (in module regression), 41
<pre>find_best_rotation() (in module curve_functions),</pre>	<pre>logit_gradient() (in module regression), 42</pre>
74	<pre>logit_hessian() (in module regression), 42</pre>
find_rotation_and_seed_coord() (in module	logit_loss() (in module regression), 42
curve_functions), 75	1.4
find_rotation_and_seed_q() (in module	M
curve_functions), 75	<pre>mlogit_gradient() (in module regression), 42</pre>
find_rotation_and_seed_unique() (in module	mlogit_loss() (in module regression), 43
curve_functions), 75	<pre>mlogit_warp_grad() (in module regression), 43</pre>
	module
fPCA	boxplots, 36
module, 31	curve_functions, 72
FPLS	curve_stats, 57
module, 38	elastic_changepoint, 48
G	elastic_glm_regression, 51
	fPCA, 31
gauss_model() (time_warping.fdawarp method), 26	fPLS, 38
geigen() (in module utility_functions), 67	geodesic, 60
<pre>geod_dist_path_strt() (in module geodesic), 61</pre>	image, 56
geod_sphere() (in module geodesic), 62	kmeans, 55
geodesic	pcr_regression, 44
module, 60	regression, 38
<pre>gradient_spline() (in module utility_functions), 67</pre>	time_warping, 23
gram_schmidt() (in module curve_functions), 76	tolerance, 53
<pre>group_action_by_gamma()</pre>	
curve_functions), 76	<pre>umap_metric, 80 utility_functions, 64</pre>
<pre>group_action_by_gamma_coord() (in module</pre>	
curve_functions), 76	multiple_align_functions()
	(time_warping.fdawarp method), 26
	<pre>mvtol_region() (in module tolerance), 54</pre>
image	N
module, 56	
init_path_geod() (in module geodesic), 62	normal() (in module time_warping), 27
init_path_rand() (in module geodesic), 62	$\circ$
innerprod_q() (in module utility_functions), 68	O
innerprod_q2() (in module curve_functions), 76	<pre>optimum_reparam() (in module utility_functions), 68</pre>
inverse_exp() (in module curve_functions), 77	<pre>optimum_reparam_curve()</pre>
inverse_exp_coord() (in module curve_functions), 77	curve_functions), 77
invertGamma() (in module utility_functions), 68	optimum_reparam_pair() (in module util-
	ity_functions), 69
	<pre>outlier_detection() (in module utility_functions), 69</pre>

98 Index

P	R
<pre>pairwise_align_bayes() (in module time_warping),</pre>	rand() (in module elastic_glm_regression), 52 rand() (in module time_warping), 30
<pre>pairwise_align_bayes_infHMC() (in module</pre>	randn() (in module curve_stats), 58
time_warping), 29	randomGamma() (in module utility_functions), 69
<pre>pairwise_align_functions() (in module</pre>	regression module, 38
<pre>parallel_translate() (in module curve_functions),</pre>	regression_warp() (in module regression), 44
78	reparam_image() (in module image), 56
<pre>path_straightening() (in module geodesic), 63</pre>	resamplecurve() (in module curve_functions), 79
pcaTB() (in module tolerance), 54	resamplefunction() (in module utility_functions), 69
pcr_regression	rgam() (in module utility_functions), 69
module, 44	rwishart() (in module tolerance), 54
phbox (class in boxplots), 37	
phi() (in module regression), 43	S
plot() (boxplots.ampbox method), 36	<pre>sample_shapes() (curve_stats.fdacurve method), 58</pre>
plot() (boxplots.phbox method), 37	scale_curve() (in module curve_functions), 80
<pre>plot() (curve_stats.fdacurve method), 57</pre>	shape_pca() (curve_stats.fdacurve method), 58
<pre>plot() (elastic_changepoint.elastic_amp_change_ff</pre>	shift_f() (in module curve_functions), 80
method), 48	<pre>smooth_data() (in module utility_functions), 70</pre>
<pre>plot() (elastic_changepoint.elastic_change method), 49</pre>	<pre>SqrtMean() (in module utility_functions), 64</pre>
<pre>plot() (elastic_changepoint.elastic_ph_change_ff</pre>	<pre>SqrtMeanInverse() (in module utility_functions), 65</pre>
method), 50	<pre>SqrtMedian() (in module utility_functions), 65</pre>
plot() (fPCA.fdahpca method), 32	<pre>srsf_align() (time_warping.fdawarp method), 26</pre>
plot() (fPCA.fdajpca method), 33	<pre>srsf_to_f() (in module utility_functions), 70</pre>
plot() (fPCA.fdajpcah method), 34	<pre>srvf_align() (curve_stats.fdacurve method), 58</pre>
plot() (fPCA.fdavpca method), 35	<pre>svd_flip() (in module utility_functions), 70</pre>
plot() (time_warping.fdawarp method), 26	_
<pre>plot_geod() (in module geodesic), 63</pre>	T
pls_svd() (in module fPLS), 38	time_warping
pre_proc_curve() (in module curve_functions), 78	1 7 00
predict() (elastic_glm_regression.elastic_glm_regression	ntolerance
method), 52	module, 53
<pre>predict() (pcr_regression.elastic_lpcr_regression</pre>	
method), 45	U
<pre>predict() (pcr_regression.elastic_mlpcr_regression</pre>	umap_metric
method), 46	module, 80
<pre>predict() (pcr_regression.elastic_pcr_regression</pre>	update_path() (in module geodesic), 64
method), 47	update_progress() (in module utility_functions), 71
predict() (regression.elastic_logistic method), 39	utility_functions
predict() (regression.elastic_mlogistic method), 40	module, 64
predict() (regression.elastic_regression method), 41	
project() (fPCA.fdahpca method), 32	W
project() (fPCA.fdajpca method), 33	<pre>warp_f_gamma() (in module utility_functions), 71</pre>
project() (fPCA.fdajpcah method), 34	warp_q_gamma() (in module utility_functions), 71 warp_q_gamma() (in module utility_functions), 71
project() (fPCA.fdavpca method), 35	"" " " " " " " " " " " " " " " " " " "
project_curve() (in module curve_functions), 78	Z
project_tangent() (in module curve_functions), 78	
psi() (in module curve_functions), 79	zero_crossing() (in module utility_functions), 71
Q	
q_to_curve() (in module curve_functions), 79	

Index 99