
fdasrsf Documentation

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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.

CHAPTER 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 fn 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 qfn: 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 wf: srsf principal weight functions :return wqf: 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

- **f** – (M,N): matrix defining N functions of M samples
- **time** – time vector of length M
- **fn** – aligned functions
- **qn** – aligned srvfs
- **q0** – initial srvfs
- **fmean** – function 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, 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)` `obj.multiple_align_functions(lambda)`

`obj.multiple_align_functions(lambda, ...)`

Parameters

- **mu** – vector of function to align to
- **omethod** – optimization method (DP, DP2, RBFGS) (default = DP)
- **smoothdata** (*bool*) – Smooth the data using a box filter (default = F)
- **parallel** – run in parallel (default = F)
- **lam** (*double*) – controls the elasticity (default = 0)
- **cores** – number of cores for parallel (default = -1 (all))
- **grid_dim** – size of the grid, for the DP2 method only (default = 7)

plot()

plot plot functional alignment results

Usage: `obj.plot()`

srsf_align (*method*=’mean’, *omethod*=’DP2’, *smoothdata*=*False*, *MaxItr*=20, *parallel*=*False*, *lam*=0.0, *cores*=-1, *grid_dim*=7)

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) (default = DP2)
- **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)
- **cores** – number of cores for parallel (default = -1 (all))
- **grid_dim** – size of the grid, for the DP2 method only (default = 7)

Examples >>> import tables >>> fun=tables.open_file(“..//Data/simu_data.h5”) >>> f = fun.root.f[:] >>> f = f.transpose() >>> time = fun.root.time[:] >>> obj = fs.fda_warp(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 independently², 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 distribution².

Note: New code should use the `normal` method of a `default_rng()` instance instead; please see the random-quick-start.

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.

out [ndarray or scalar] Drawn samples from the parameterized normal distribution.

scipy.stats.norm [probability density function, distribution or] cumulative density function, etc.

`Generator.normal`: which should be used for new code.

The probability density for the Gaussian distribution is

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

² P. R. Peebles Jr., “Central Limit Theorem” in “Probability, Random Variables and Random Signal Principles”, 4th ed., 2001, pp. 51, 51, 125.

where μ is the mean and σ the standard deviation. The square of the standard deviation, σ^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.

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:

```
>>> abs(mu - np.mean(s))
0.0 # may vary
```

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

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

```
>>> import matplotlib.pyplot as plt
>>> count, bins, ignored = plt.hist(s, 30, density=True)
>>> plt.plot(bins, 1/(sigma * np.sqrt(2 * np.pi)) *
...             np.exp(- (bins - mu)**2 / (2 * sigma**2)),
...             linewidth=2, color='r')
>>> plt.show()
```

Two-by-four array of samples from $N(3, 6.25)$:

```
>>> 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(f1i, f2i, time, mcmc_opts=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)` `out = pairwise_align_bayes(f1i, f2i, time, mcmc_opts)`

Parameters

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

default mcmc options: `tmp = {“betas”:np.array([0.5,0.5,0.005,0.0001]),“probs”:np.array([0.1,0.1,0.7,0.1])}`
`mcmc_opts = {“iter”:2*(10**4),“burnin”:np.minimum(5*(10**3),2*(10**4)//2),`

```

"alpha0":0.1, "beta0":0.1,"zpcn":tmp,"propvar":1, "initcoef":np.repeat(0,20), "npoints":200, "ex-
trainfo":True}

: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 poste-
rior :return ydist: amplitude distance posterior)

time_warping.pairwise_align_bayes_infHMC (y1i, y2i, time, mcmc_opts=None)
This function aligns two functions using Bayesian framework. It uses a hierarchical Bayesian framework assum-
ing measurement 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 infy-HMC algorithm is used
to explore both local and global structure in the posterior distribution.

Usage: out = pairwise_align_bayes_infHMC(f1i, f2i, time) out = pairwise_align_bayes_infHMC(f1i, f2i,
time, mcmc_opts)

```

Parameters

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

default mcmc options: mcmc_opts = {"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, grid_dim=7)
```

This function aligns f2 to f1 using the elastic square-root slope (srdf) framework.

Usage: out = pairwise_align_functions(f1, f2, time) out = pairwise_align_functions(f1, f2, time, omethod,
lam, grid_dim)

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) (default = DP)

- **lam** – controls the elasticity (default = 0)
- **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 (srdf)
time_warping.**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).

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

out [ndarray, shape (d0, d1, ..., dn)] Random values.

random

```
>>> np.random.rand(3,2)
array([[ 0.14022471,  0.96360618], #random
       [ 0.37601032,  0.25528411], #random
       [ 0.49313049,  0.94909878]]) #random
```


CHAPTER 2

Functional Principal Component Analysis

Vertical and Horizontal Functional Principal Component Analysis using SRSF

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

class fPCA.fdahtpca(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@sandia.gov> Date : 15-Mar-2018

calc_fpca (no=3, 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)
- **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()

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** – srsvf principal directions
- **f_pca** – f principal directions
- **latent** – latent values
- **coef** – principal coefficients
- **id** – point used for f(0)
- **mgn** – mean srsvf
- **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, 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)
- **id** (*int*) – point to use for f(0) (default = midpoint)
- **stds** – number of standard deviations along geodesic 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 vertical fPCA result
    Usage: obj.plot()
```

```
class fPCA.fdapca(fdawarp)
```

This class provides vertical fPCA using the SRVF framework

Usage: obj = fdapca(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)
- **mgn** – mean srvf
- **U** – eigenvectors
- **stds** – geodesic directions

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

```
calc_fPCA (no=3, 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)
- **id** (*int*) – point to use for f(0) (default = midpoint)
- **stds** – number of standard deviations along gedoesic to compute (default = -1,0,1)

Return type fdapca 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()
```


CHAPTER 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@ 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.

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()

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()

CHAPTER 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
- **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

CHAPTER 5

Elastic Regression

Warping Invariant Regression using SRSF

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

`regression.elastic_logistic(f, y, time, B=None, df=20, max_itr=20, cores=-1, smooth=False)`

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

Parameters

- `f` (`np.ndarray`) – numpy ndarray of shape (M,N) of N functions with M samples
- `y` – numpy array of labels (1/-1)
- `time` (`np.ndarray`) – vector of size M describing the sample points
- `B` – optional matrix describing Basis elements
- `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)

Return type tuple of numpy array

Return alpha alpha parameter of model

Return beta beta(t) of model

Return fn aligned functions - numpy ndarray of shape (M,N) of M

functions with N samples :return qn: aligned srvfs - similar structure to fn :return gamma: calculated warping functions :return q: original training SRSFs :return B: basis matrix :return b: basis coefficients :return Loss: logistic loss

`regression.elastic_mlogistic(f, y, time, B=None, df=20, max_itr=20, cores=-1, delta=0.01, parallel=True, smooth=False)`

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

Parameters

- **f** (*np.ndarray*) – numpy ndarray of shape (M,N) of N functions with M samples
- **y** – numpy array of labels {1,2,...,m} for m classes
- **time** (*np.ndarray*) – vector of size M describing the sample points
- **B** – optional matrix describing Basis elements
- **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)

Return type tuple of numpy array

Return alpha alpha parameter of model

Return beta beta(t) of model

Return fn aligned functions - numpy ndarray of shape (M,N) of N

functions with M samples :return qn: aligned svrfs - similar structure to fn :return gamma: calculated warping functions :return q: original training SRSFs :return B: basis matrix :return b: basis coefficients :return Loss: logistic loss

`regression.elastic_prediction(f, time, model, y=None, smooth=False)`

This function performs prediction from an elastic regression model with phase-variability

Parameters

- **f** – numpy ndarray of shape (M,N) of N functions with M samples
- **time** – vector of size M describing the sample points
- **model** – identified model from elastic_regression
- **y** – truth, optional used to calculate SSE

Return type tuple of numpy array

Return alpha alpha parameter of model

Return beta beta(t) of model

Return fn aligned functions - numpy ndarray of shape (M,N) of N

functions with M samples :return qn: aligned svrfs - similar structure to fn :return gamma: calculated warping functions :return q: original training SRSFs :return B: basis matrix :return b: basis coefficients :return SSE: sum of squared error

`regression.elastic_regression(f, y, time, B=None, lam=0, df=20, max_itr=20, cores=-1, smooth=False)`

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

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
- **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)

Return type tuple of numpy array

Return alpha alpha parameter of model

Return beta beta(t) of model

Return fn aligned functions - numpy ndarray of shape (M,N) of M

functions with N samples :return qn: aligned srvfs - similar structure to fn :return gamma: calculated warping functions :return q: original training SRSFs :return B: basis matrix :return b: basis coefficients :return SSE: sum of squared error

regression.**logistic_warp**(beta, time, q, y)

calculates optimal warping for function logistic regression

Parameters

- **beta** – numpy ndarray of shape (M,N) of N functions with M samples
- **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
- **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 :param alpha: numpy scalar

Return type numpy array

Return gamma_new warping function

CHAPTER 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 pcr regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic_lpcr_regression(f,y,time)

Parameters

- **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

:param information :param alpha: intercept :param b: coefficient vector :param Loss: logistic loss :param PC: probability of classification :param ylabels: predicted labels

Author : J. D. Tucker (JDT) <jdtuck@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()

```
obj.predict(newdata)
```

Parameters

- **newdata** (*dict*) – dict containing new data for prediction (needs the keys below, if None predicts on training data)
- **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 pcr regression for functional data using the SRVF framework accounting for warping

Usage: obj = elastic_mlpcr_regression(f,y,time)

Parameters

- **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

:param information :param alpha: intercept :param b: coefficient vector :param Loss: logistic loss :param PC: probability of classification :param ylabels: predicted labels :param

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()

```
obj.predict(newdata)
```

Parameters

- **newdata** (*dict*) – dict containing new data for prediction (needs the keys below, if None predicts on training data)
- **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)

Parameters

- **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*, *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)
- **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*)

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

```
obj.predict(newdata)
```

Parameters

- **newdata** (*dict*) – dict containing new data for prediction (needs the keys below, if None predicts on training data)
- **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

CHAPTER 7

Elastic Functional Tolerance Bounds

Functional Tolerance Bounds using SRSF

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

`tolerance.bootTB(f, time, a=0.5, 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
- `a` – confidence level of tolerance bound (default = 0.05)
- `p` – coverage level of tolerance bound (default = 0.99)
- `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

- **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
- **a** – confidence level of tolerance bound (default = 0.05)
- **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

Parameters

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

Return type double

Return R matrix

CHAPTER 8

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 qun: cost function :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@sandia.gov> Date : 26-Aug-2020

karcher_cov()

This calculates the mean of a set of curves

karcher_mean(parallel=False, cores=-1, method='DP')

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

plot()

plot curve mean results

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 (*parallel=False, cores=-1, method='DP'*)

This aligns a set of curves to the mean and computes mean if not computed :param parallel: run in parallel (default = F) :param cores: number of cores for parallel (default = -1 (all)) :param method: method to apply optimization (default="DP") options are "DP" 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 *standard_normal* method of a *default_rng()* 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, d1, ..., dn [int, optional] The dimensions of the returned array, must be non-negative. If no argument is given a single Python float is returned.

Z [ndarray or float] 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.

standard_normal : Similar, but takes a tuple as its argument. *normal* : Also accepts mu and sigma arguments. *Generator.standard_normal*: which should be used for new code.

For random samples from $N(\mu, \sigma^2)$, use:

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

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

Two-by-four array of samples from $N(3, 6.25)$:

```
>>> 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
```

CHAPTER 9

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)
- **k** – number of samples along path (Default = 5)

Return type numpy ndarray

Return utilde translated vector

`geodesic.calc_alpha_dot(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)
- **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)
- **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)
- **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)
- **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)
- **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

- **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 beta1 and beta2 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
- **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 O rotation matrix

`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)
- **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 \mathcal{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)
- **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*, *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 = NULL, only needed for init “rand”)
- **init** – initialize path geodesic or random (Default = “rand”)
- **T** – Number of samples of curve (Default = 100)
- **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.**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 parameterer
- **T** – Number of samples of curve (Default = 100)
- **k** – number of samples along path (Default = 5)

Return type numpy scalar

Return alpha updated path of svrfs

Return beta updated path of curves

CHAPTER 10

Utility Functions

Utility functions for SRSF Manipulations

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

`utility_functions.SqrtMean(gam, parallel=False, cores=-1)`

calculates the srsf of warping functions with corresponding shooting vectors

Parameters

- **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))

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 M warping functions with N 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"
- **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)`
" 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"
- **lam** – controls the elasticity (default = 0.0)

Return type scalar

Return Dy amplitude distance

Return Dx phase 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
- **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 descrbing 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, 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"
- `lam` – controls the amount of elasticity (default = 0.0)
- `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
- `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()`

- **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)`
Generates random warping functions

Parameters

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

Returns gam: numpy ndarray of warping functions

`utility_functions.smooth_data(f, sparam)`
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.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

`utility_functions.zero_crossing(Y, q, bt, time, y_max, y_min, gmax, gmin)`

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

Parameters

- `Y` – response
- `q` – predictive function
- `bt` – basis function
- `time` – time samples
- `y_max` – maximum response for warping function `gmax`
- `y_min` – minimum response for warping function `gmin`
- `gmax` – max warping function
- `gmin` – min warping function

Return type numpy array

Return gamma optimal warping function

CHAPTER 11

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'*)

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')

Return type numpy ndarray**Return q** srvf of curve**Return lenb** length of curve**Return lenq** length of srvf`curve_functions.curve_zero_crossing(Y, beta, bt, y_max, y_min, gmax, gmin)`finds zero-crossing of optimal gamma, $\text{gam} = s*\text{gmax} + (1-s)*\text{gmin}$ from elastic curve regression model**Parameters**

- **Y** – response
- **beta** – predictive function
- **bt** – basis function
- **y_max** – maximum response for warping function gmax
- **y_min** – minimum response for warping function gmin
- **gmax** – max warping function
- **gmin** – min warping function

Return type numpy array**Return gamma** optimal warping function**Return O_hat** rotation matrix`curve_functions.elastic_distance_curve(beta1, beta2, closed=0, scale=False, method='DP')`

Calculates the two elastic distances between two curves :param beta1: numpy ndarray of shape (2,M) of M samples :param beta2: numpy ndarray of shape (2,M) of M samples :param closed: open (0) or closed (1) curve (default=0) :param scale: include scale (default=False) :param method: method to apply optimization (default="DP") options are "DP" or "RBFGS"

Return type scalar**Return dist** shape distance`curve_functions.elastic_shooting(q1, v)`

Calculates shooting vector from v to q1

Parameters

- **q1** – vector of srvf
- **v** – shooting vector

:rtype numpy ndarray :return q2n: vector of 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 svrfs 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 rotated beta2 to beta1

Return O rotation matrix

Return tau seed

```
curve_functions.find_rotation_and_seed_q(q1, q2)
```

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

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 beta2new optimal rotated beta2 to beta1

Return O rotation matrix

Return tau seed

```
curve_functions.find_rotation_and_seed_unique(q1, q2, closed=0, 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)
- **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 Orthogonalization 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 orthogonalized basis

curve_functions.**group_action_by_gamma** (*q, gamma*)

This function reparameterized srvf q 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 qn reparameterized srvf

curve_functions.**group_action_by_gamma_coord** (*f, gamma*)

This function reparameterized 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 reparameterized 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)
- **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 preprocessed 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 wproj 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

- **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')

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

- **f** – numpy ndarray of shape (2,M) of M samples
- **tau** – scalar

Return type numpy ndarray

Return fn shifted curve

CHAPTER 12

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`

” 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
- `q2` – vector of size N

Return type

scalar

Return dist amplitude distance

`umap_metric.efda_distance_curve`

” 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 13

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