# Neural Network Toolbox™ Reference

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# MATLAB®



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Neural Network Toolbox<sup>™</sup> Reference

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#### ${\bf Functions-Alphabetical\ List}$

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# 1

# Functions — Alphabetical List

#### adapt

Purpose	Adapt neural network to data as it is simulated		
Syntax	[net,Y,E,Pf,Af,tr] = adapt(net,P,T,Pi,Ai)		
To Get Help	Type help network/adapt.		
Description	This function calculates network outputs and errors after each presentation of an input. [net,Y,E,Pf,Af,tr] = adapt(net,P,T,Pi,Ai) takes		
	net	Network	
	Р	Network inputs	
	Т	Network targets (default = zeros)	
	Pi	Initial input delay conditions (default = zeros)	
	Ai	Initial layer delay conditions (default = zeros)	

and returns the following after applying the adapt function net.adaptFcn with the adaption parameters net.adaptParam:

net	Updated network
Y	Network outputs
E	Network errors
Pf	Final input delay conditions
Af	Final layer delay conditions
tr	Training record (epoch and perf)

Note that T is optional and is only needed for networks that require targets. Pi and Pf are also optional and only need to be used for networks that have input or layer delays.

adapt's signal arguments can have two formats: cell array or matrix.

The cell array format is easiest to describe. It is most convenient for networks with multiple inputs and outputs, and allows sequences of inputs to be presented,

Р	Ni-by-TS cell array	Each element P{i,ts} is an Ri-by-Q matrix.
т	Nt-by-TS cell array	Each element T{i,ts} is a Vi-by-Q matrix.
Pi	Ni-by-ID cell array	Each element Pi{i,k} is an Ri-by-Q matrix.
Ai	N1-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.
Y	No-by-TS cell array	Each element Y{i,ts} is a Ui-by-Q matrix.
Е	No-by-TS cell array	Each element E{i,ts} is a Ui-by-Q matrix.
Pf	Ni-by-ID cell array	Each element Pf{i,k} is an Ri-by-Q matrix.
Af	N1-by-LD cell array	Each element Af{i,k} is an Si-by-Q matrix.

#### where

Ni	=	net.numInputs
Nl	=	net.numLayers
No	=	net.numOutputs
ID	=	net.numInputDelays
LD	=	net.numLayerDelays
TS	=	Number of time steps

Q	=	Batch size
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Ui	=	net.outputs{i}.size

The columns of Pi, Pf, Ai, and Af are ordered from oldest delay condition to most recent:

Pi{i,k}	=	Input i at time $ts = k - ID$
Pf{i,k}	=	Input i at time ts = TS + $k$ - ID
Ai{i,k}	=	Layer output i at time $ts = k - LD$
Af{i,k}	=	Layer output i at time $ts = TS + k - LD$

The matrix format can be used if only one time step is to be simulated (TS = 1). It is convenient for networks with only one input and output, but can be used with networks that have more.

Each matrix argument is found by storing the elements of the corresponding cell array argument in a single matrix:

Р	(sum	of	Ri)-by-Q matrix
Т	(sum	of	Vi)-by-Q matrix
Pi	(sum	of	Ri)-by-(ID*Q) matrix
Ai	(sum	of	Si)-by-(LD*Q) matrix
Υ	(sum	of	Ui)-by-Q matrix
E	(sum	of	Ui)-by-Q matrix
Pf	(sum	of	Ri)-by-(ID*Q) matrix
Af	(sum	of	Si)-by-(LD*Q) matrix

## **Examples** Here two sequences of 12 steps (where T1 is known to depend on P1) are used to define the operation of a filter.

 $p1 = \{-1 \quad 0 \quad 1 \quad 0 \quad 1 \quad 1 \quad -1 \quad 0 \quad -1 \quad 1 \quad 0 \quad 1\};$  $t1 = \{-1 \quad -1 \quad 1 \quad 1 \quad 1 \quad 2 \quad 0 \quad -1 \quad -1 \quad 0 \quad 1 \quad 1\};$ 

Here linearlayer is used to create a layer with an input range of [-1 1], one neuron, input delays of 0 and 1, and a learning rate of 0.5. The linear layer is then simulated.

net = linearlayer([0 1],0.5);

Here the network adapts for one pass through the sequence.

The network's mean squared error is displayed. (Because this is the first call to adapt, the default Pi is used.)

```
[net,y,e,pf] = adapt(net,p1,t1);
mse(e)
```

Note that the errors are quite large. Here the network adapts to another 12 time steps (using the previous Pf as the new initial delay conditions).

```
p2 = {1 -1 -1 1 1 -1 0 0 0 1 -1 -1};
t2 = {2 0 -2 0 2 0 -1 0 0 1 0 -1};
[net,y,e,pf] = adapt(net,p2,t2,pf);
mse(e)
```

Here the network adapts for 100 passes through the entire sequence.

```
p3 = [p1 p2];
t3 = [t1 t2];
net.adaptParam.passes = 100;
[net,y,e] = adapt(net,p3,t3);
mse(e)
```

#### adapt

AlgorithmsThe error after 100 passes through the sequence is very small. The<br/>network has adapted to the relationship between the input and target<br/>signals.Algorithmsadapt calls the function indicated by net.adaptFcn, using the adaption<br/>parameter values indicated by net.adaptParam.<br/>Given an input sequence with TS steps, the network is updated as<br/>follows: Each step in the sequence of inputs is presented to the network<br/>one at a time. The network's weight and bias values are updated after<br/>each step, before the next step in the sequence is presented. Thus the<br/>network is updated TS times.See Alsosim | init | train | revert

Purpose	Adapt network with weight and bias learning rules		
Syntax	[net,ar,Ac] = adapt(net,Pd,T,Ai)		
Description	This function is normally not called directly, but instead called indirectly through the function adapt after setting a network's adaption function (net.adaptFcn) to this function.		
	[net,ar,Ac] = ada	apt(net,Pd,T,Ai) takes these arguments,	
	net	Neural network	
	Pd	Delayed processed input states and inputs	
	т	Targets	
	Ai	Initial layer delay states	
	and returns net Neural network after adaption ar Adaption record		
Examples	Ac Linear layers use th	Combined initial layer states and layer outputs nis adaption function. Here a linear layer with input	
-Xumpios	<pre>http://delays.org/construction/interform/</pre>		

#### adaptwb

See Also adapt

Purpose	Add delay to neural network response		
Syntax	<pre>net = adddelay(net,n)</pre>		
Description	<pre>net = adddelay(net,n) takes these arguments,</pre>		
	net Neural network n Number of delays		
	and returns the network with input delay connections increased, and output feedback delays decreased, by the specified number of delays n. The result is a network which behaves identically, except that outputs are produced n timesteps later.		
	If the number of delays n is not specified, a default of one delay is used.		
Examples	Here a time delay network is created, trained and simulated in its original form on an input time series X and target series T. It is then simulated with a delay removed and then added back. These first and third outputs will be identical, while the second will be shifted by one timestep.		
	<pre>[X,T] = simpleseries_dataset; net = timedelaynet(1:2,20); [Xs,Xi,Ai,Ts] = preparets(net,X,T); net = train(net,Xs,Ts,Xi); y1 = net(Xs) net2 = removedelay(net); [Xs,Xi,Ai,Ts] = preparets(net2,X,T); y2 = net2(Xs,Xi) net3 = adddelay(net2) [Xs,Xi,Ai,Ts] = preparets(net3,X,T); y3 = net3(Xs,Xi)</pre>		
See Also	closeloop   openloop   removedelay		

#### boxdist

Purpose	Distance between two position vectors		
Syntax	<pre>d = boxdist(pos)</pre>		
Description	boxdist is a layer distance function that is used to find the distances between the layer's neurons, given their positions.		
	d = boxdist(pos) takes one argument,		
	pos N-by-S matrix of neuron po	ositions	
	and returns the S-by-S matrix of distances.		
	boxdist is most commonly used with layers whose topology function is gridtop.		
Examples	Here you define a random matrix of positions for 10 neurons arranged in three-dimensional space and then find their distances.		
	<pre>pos = rand(3,10); d = boxdist(pos)</pre>		
Network Use	To change a network so that a layer's topolo net.layers{i}.distanceFcn to 'boxdist'		
	In either case, call <b>sim</b> to simulate the network	ork with boxdist.	
Algorithms	The box distance D between two position vec S vectors is	tors Pi and Pj from a set of	
	Dij = max(abs(Pi-Pj))		
See Also	dist   linkdist   mandist   sim		

Purpose	Backpropagation through time derivative function		
Syntax		dwb',net,X,T,Xi,Ai,EW) ',net,X,T,Xi,Ai,EW)	
Description	This function calculates derivatives using the chain rule from a network's performance back through the network, and in the case of dynamic networks, back through time.		
	bttderiv('dperf_	dwb',net,X,T,Xi,Ai,EW) takes these arguments,	
	net	Neural network	
	Х	Inputs, an RxQ matrix (or NxTS cell array of RixQ matrices)	
	T Targets, an SxQ matrix (or MxTS cell array of SixQ matrices)		
	Xi	Initial input delay states (optional)	
	Ai	Initial layer delay states (optional)	
	EW Error weights (optional)		
	and returns the gradient of performance with respect to the network's weights and biases, where R and S are the number of input and output elements and Q is the number of samples (and N and M are the number of input and output signals, Ri and Si are the number of each input and outputs elements, and TS is the number of timesteps). bttderiv('de_dwb',net,X,T,Xi,Ai,EW) returns the Jacobian of errors with respect to the network's weights and biases.		
Examples	Here a feedforward network is trained and both the gradient and Jacobian are calculated.		
	[x,t] = simplefi net = feedforwar		

```
net = train(net,x,t);
y = net(x);
perf = perform(net,t,y);
gwb = bttderiv('dperf_dwb',net,x,t)
jwb = bttderiv('de_dwb',net,x,t)
```

See Also defaultderiv | fpderiv | num2deriv | num5deriv | staticderiv

Purpose	Cascade-forward neural network		
Syntax	cascadeforwardnet(	hiddenSizes,trainFcn)	
Description	Cascade-forward networks are similar to feed-forward networks, but include a connection from the input and every previous layer to following layers.		
	As with feed-forward networks, a two-or more layer cascade-network can learn any finite input-output relationship arbitrarily well given enough hidden neurons.		
	cascadeforwardnet(	hiddenSizes,trainFcn) takes these arguments,	
	hiddenSizes	Row vector of one or more hidden layer sizes (default = 10)	
	trainFcn	Training function (default = 'trainlm')	
	and returns a new cascade-forward neural network.		
Examples	Here a cascade network is created and trained on a simple fitting problem.		
	<pre>[x,t] = simplefit_dataset; net = cascadeforwardnet(10); net = train(net,x,t); view(net) y = net(x) perf = perform(net,y,t)</pre>		
See Also	feedforwardnet		

#### catelements

Purpose	Concatenate neural network data elements		
Syntax	catelements(x1,x2,,xn) [x1; x2; xn]		
Description	catelements(x1,x2,,xn) takes any number of neural network data values, and merges them along the element dimension (i.e., the matrix row dimension).		
	If all arguments are matrices, this operation is the same as [x1; x2; xn].		
	If any argument is a cell array, then all non-cell array arguments are enclosed in cell arrays, and then the matrices in the same positions in each argument are concatenated.		
Examples	This code concatenates the elements of two matrix data values.		
	x1 = [1 2 3; 4 7 4] x2 = [5 8 2; 4 7 6; 2 9 1] y = catelements(x1,x2)		
	This code concatenates the elements of two cell array data values.		
	x1 = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} x2 = {[2 1 3] [4 5 6]; [2 5 4] [9 7 5]} y = catelements(x1,x2)		
See Also	nndata   numelements   getelements   setelements   catsignals   catsamples   cattimesteps		

Purpose	Concatenate neural network data samples		
Syntax	catsamples(x1,x2,,xn) [x1 x2 xn] catsamples(x1,x2,,xn,'pad',v)		
Description	catsamples(x1,x2,,xn) takes any number of neural network data values, and merges them along the samples dimension (i.e., the matrix column dimension).		
	If all arguments are matrices, this operation is the same as $[x1 x2 \dots xn]$ .		
	If any argument is a cell array, then all non-cell array arguments are enclosed in cell arrays, and then the matrices in the same positions in each argument are concatenated.		
	catsamples(x1,x2,,xn,'pad',v) allows samples with varying numbers of timesteps (columns of cell arrays) to be concatenated by padding the shorter time series with the value v, until they are the same length as the longest series. If v is not specified, then the value NaN is used, which is often used to represent unknown or don't-care inputs or targets.		
Examples	This code concatenates the samples of two matrix data values.		
	x1 = [1 2 3; 4 7 4] x2 = [5 8 2; 4 7 6] y = catsamples(x1,x2)		
	This code concatenates the samples of two cell array data values.		
	x1 = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} x2 = {[2 1 3; 5 4 1] [4 5 6; 9 4 8]; [2 5 4] [9 7 5]} y = catsamples(x1,x2)		
	Here the samples of two cell array data values, with unequal numbers of timesteps, are concatenated.		

```
x1 = {1 2 3 4 5};
x2 = {10 11 12};
y = catsamples(x1,x2,'pad')
```

# See Also nndata | numsamples | getsamples | setsamples | catelements | catsignals | cattimesteps

Purpose	Concatenate neural network data signals		
Syntax	catsignals(x1,x2,,xn) {x1; x2;; xn}		
Description	catsignals(x1,x2,,xn) takes any number of neural network data values, and merges them along the element dimension (i.e., the cell row dimension).		
	If all arguments are matrices, this operation is the same as {x1; x2;; xn}.		
	If any argument is a cell array, then all non-cell array arguments are enclosed in cell arrays, and the cell arrays are concatenated as [x1; x2;; xn].		
Examples	This code concatenates the signals of two matrix data values.		
	x1 = [1 2 3; 4 7 4] x2 = [5 8 2; 4 7 6] y = catsignals(x1,x2)		
	This code concatenates the signals of two cell array data values.		
	x1 = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} x2 = {[2 1 3; 5 4 1] [4 5 6; 9 4 8]; [2 5 4] [9 7 5]} y = catsignals(x1,x2)		
See Also	nndata   numsignals   getsignals   setsignals   catelements   catsamples   cattimesteps		

#### cattimesteps

Purpose	Concatenate neural network data timesteps		
Syntax	cattimesteps(x1,x2,,xn) {x1 x2 xn}		
Description	cattimesteps(x1, x2,, xn) takes any number of neural network data values, and merges them along the element dimension (i.e., the cell column dimension).		
	If all arguments are matrices, this operation is the same as $\{x1 \ x2 \ \dots \ xn\}$ .		
	If any argument is a cell array, all non-cell array arguments are enclosed in cell arrays, and the cell arrays are concatenated as $[x1 x2 \dots xn]$ .		
Examples	This code concatenates the elements of two matrix data values.		
	x1 = [1 2 3; 4 7 4] x2 = [5 8 2; 4 7 6] y = cattimesteps(x1,x2)		
	This code concatenates the elements of two cell array data values.		
	x1 = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} x2 = {[2 1 3; 5 4 1] [4 5 6; 9 4 8]; [2 5 4] [9 7 5]} y = cattimesteps(x1,x2)		
See Also	nndata   numtimesteps   gettimesteps   settimesteps   catelements   catsignals   catsamples		

Purpose	Create cell array of matrices		
Syntax	<pre>cellmat(A,B,C,D,v)</pre>		
Description	cellmat(A,B,C,D,v) takes four integer values and one scalar value v, and returns an A-by-B cell array of C-by-D matrices of value v. If the value v is not specified, zero is used.		
Examples	Here two cell arrays of matrices are created.		
	cm1 = cellmat(2,3,5,4) cm2 = cellmat(3,4,2,2,pi)		
See Also	nndata		

### closeloop

Purpose	Convert neural network open-loop feedback to closed loop		
Syntax	<pre>net = closeloop(net)</pre>		
Description	<pre>net = closeloop(net) takes a neural network and closes any open-loop feedback. For each feedback output i whose property net.outputs{i}.feedbackMode is 'open', it replaces its associated feedback input and their input weights with layer weight connections coming from the output. The net.outputs{i}.feedbackMode property is set to 'closed', and the net.outputs{i}.feedbackInput property is set to an empty matrix. Finally, the value of net.outputs{i}.feedbackDelays is added to the delays of the feedback layer weights (i.e., to the delays values of the replaced input weights).</pre>		
Examples	<pre>Here a NARX network is designed in open-loop form and then converted to closed-loop form. [X,T] = simplenarx_dataset; net = narxnet(1:2,1:2,10); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts,Xi,Ai); view(net) Yopen = net(Xs,Xi,Ai) net = closeloop(net) view(net) [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); Ycloesed = net(Xs,Xi,Ai);</pre>		
See Also	noloop   openloop		

Purpose	Create all combinations of vectors		
Syntax	combvec(A1,A2)		
Description	combvec(A1,A2) takes any number of inputs,		
	A1	Matrix of N1 (column) vectors	
	A2	Matrix of N2 (column) vectors	
	and returns a matrix of $(N1*N2*)$ column vectors, where the columns consist of all possibilities of A2 vectors, appended to A1 vectors, etc.		
Examples	a1 = [1 2 3; 4 5 6]; a2 = [7 8; 9 10]; a3 = combvec(a1,a2)		

#### compet

Purpose	Competitive transfer function		
Graph and Symbol	Input n 2 1 4 3 <i>a = comp</i> Compet Transfe		
Syntax	A = compet(N, info = compet		
Description	<pre>compet is a neural transfer function. Transfer functions calculate a layer's output from its net input. A = compet(N,FP) takes N and optional function parameters,</pre>		
	Ν	S-by-Q matrix of net input (column) vectors	
	FP	Struct of function parameters (ignored)	
	and returns the S-by-Q matrix A with a 1 in each column where the sa column of N has its maximum value, and O elsewhere.		
	<pre>info = compet('code') returns information according to the code string specified:</pre>		
	compet('name') returns the name of this function.		
	compet('outpu	t',FP) returns the [min max] output range.	
	compet('active',FP) returns the [min max] active input range.		
	compet('fullderiv') returns 1 or 0, depending on whether dA_d S-by-S-by-Q or S-by-Q.		
	compet('fpnam	es') returns the names of the function parameters.	
	compet('fpdef	aults') returns the default function parameters.	

Examples Here you define a net input vector N, calculate the output, and plot both with bar graphs. n = [0; 1; -0.5; 0.5]; a = compet(n); subplot(2,1,1), bar(n), ylabel('n') subplot(2,1,2), bar(a), ylabel('a') Assign this transfer function to layer i of a network. net.layers{i}.transferFcn = 'compet'; See Also sim | softmax

## competlayer

Purpose	Competitive layer		
Syntax	competlayer(numClasses,kohonenLR,conscienceLR)		
Description	Competitive layers learn to classify input vectors into a given number of classes, according to similarity between vectors, with a preference for equal numbers of vectors per class.		
	<pre>competlayer(numClasses,kohonenLR,conscienceLR) takes these arguments,</pre>		
	numClasses	Number of classes to classify inputs (default = 5)	
	kohonenLR	Learning rate for Kohonen weights (default = 0.01)	
	conscienceLR	Learning rate for conscience bias (default = 0.001)	
	and returns a competi	tive layer with numClasses neurons.	
Examples	<pre>Here a competitive layer is trained to classify 150 iris flowers into 6 classes. inputs = iris_dataset; net = competlayer(6); net = train(net,inputs); view(net) outputs = net(inputs); classes = vec2ind(outputs);</pre>		
See Also	selforgmap   patter	nnet   lvqnet	

Purpose	Convert concurrent vectors to sequential vectors			
•		Convert concurrent vectors to sequential vectors		
Syntax	• •	S = con2seq(b) S = con2seq(b,TS)		
Description	Neural Network Toolbox <sup>™</sup> software arranges concurrent vectors with a matrix, and sequential vectors with a cell array (where the second index is the time step).			
		q2con allow concurrent vectors to be converted to ors, and back again.		
	S = con2seq(b	) takes one input,		
	b	R-by-TS matrix		
	and returns one	e output,		
	S	1-by-TS cell array of R-by-1 vectors		
	<ul> <li>S = con2seq(b,TS) can also convert multiple batches,</li> <li>b N-by-1 cell array of matrices with M*TS columns</li> </ul>			
	TS	Time steps		
	and returns			
	S	N-by-TS cell array of matrices with ${\tt M}$ columns		
Examples	Here a batch of	three values is converted to a sequence.		
	p1 = [1 4 2] p2 = con2seq(p1)			

Here, two batches of vectors are converted to two sequences with two time steps.

p1 = {[1 3 4 5; 1 1 7 4]; [7 3 4 4; 6 9 4 1]} p2 = con2seq(p1,2)

See Also seq2con | concur

Purpose	Create concurrent bias vectors		
Syntax	concur(B,Q)		
Description	concur(B,Q)		
	B Q	S-by-1 bias vector (or an N1-by-1 cell array of vectors) Concurrent size	
	and returns an of matrices).	S-by-B matrix of copies of B (or an Nl-by-1 cell array	
Examples	Here concur cre	eates three copies of a bias vector.	
	b = [1; 3; 2; -1]; concur(b,3)		
Network Use	To calculate a layer's net input, the layer's weighted inputs must be combined with its biases. The following expression calculates the net input for a layer with the netsum net input function, two input weights, and a bias:		
	n = netsum(z1,z2,b)		
	The above expression works if Z1, Z2, and B are all S-by-1 vectors. However, if the network is being simulated by sim (or adapt or train) in response to Q concurrent vectors, then Z1 and Z2 will be S-by-Q matrices. Before B can be combined with Z1 and Z2, you must make Q copies of it.		
	n = netsum(z1	,z2,concur(b,q))	
See Also	con2seq   netprod   netsum   seq2con   sim		

## configure

Purpose	Configure network inputs and outputs to best match input and target data		
Syntax	<pre>net = configure(net,x,t) net = configure(net,x) net = configure(net,'inputs',x,i) net = configure(net,'outputs',t,i)</pre>		
Description	Configuration is the process of setting network input and output sizes and ranges, input preprocessing settings and output postprocessing settings, and weight initialization settings to match input and target data.		
	Configuration must happen before a network's weights and biases can be initialized. Unconfigured networks are automatically configured and initialized the first time train is called. Alternately, a network can be configured manually either by calling this function or by setting a network's input and output sizes, ranges, processing settings, and initialization settings properties manually.		
	<pre>net = configure(net,x,t) takes input data x and target data t, and configures the network's inputs and outputs to match.</pre>		
	<pre>net = configure(net,x) configures only inputs.</pre>		
	<pre>net = configure(net, 'inputs',x,i) configures the inputs specified with the index vector i. If i is not specified all inputs are configured.</pre>		
	<pre>net = configure(net,'outputs',t,i) configures the outputs specified with the index vector i. If i is not specified all targets are configured.</pre>		
Examples	Here a feedforward network is created and manually configured for a simple fitting problem (as opposed to allowing train to configure it).		
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(20); view(net) net = configure(net,x,t); view(net)</pre>		

See Also isconfigured | unconfigure | init | train

#### confusion

Purpose	Classification confusion matrix	
Syntax	<pre>[c,cm,ind,per] = confusion(targets,outputs)</pre>	
Description	<pre>[c,cm,ind,per] = confusion(targets,outputs) takes these values</pre>	
	targets	S-by-Q matrix, where each column vector contains a single 1 value, with all other elements 0. The index of the 1 indicates which of S categories that vector represents.
	outputs	S-by-Q matrix, where each column contains values in the range [0,1]. The index of the largest element in the column indicates which of S categories that vector represents.
	and returns these values:	
	С	Confusion value = fraction of samples misclassified
	CM	S-by-S confusion matrix, where cm(i,j) is the number of samples whose target is the ith class that was classified as j
	ind	S-by-S cell array, where ind{i,j} contains the indices of samples with the ith target class, but jth output class
	per	S-by-4 matrix, where each row summarizes four percentages associated with the ith class:
		<pre>per(i,1) false negative rate</pre>

	<pre>[c,cm,ind,per] = confusion(TARGETS,OUTPUTS) takes these values:</pre>		
	targets outputs	<ul> <li>1-by-Q vector of 1/0 values representing membership</li> <li>S-by-Q matrix, of value in [0,1] interval, where values greater than or equal to 0.5 indicate class membership</li> </ul>	
	and returns	these values:	
	С	Confusion value = fraction of samples misclassified	
	cm	2-by-2 confusion matrix	
	ind	2-by-2 cell array, where ind{i,j} contains the indices of samples whose target is 1 versus 0, and whose output was greater than or equal to 0.5 versus less than 0.5	
	per	2-by-4 matrix where each ith row represents the percentage of false negatives, false positives, true positives, and true negatives for the class and out-of-class	
Examples	<pre>load simpleclass_dataset net = newpr(simpleclassInputs,simpleclassTargets,20); net = train(net,simpleclassInputs,simpleclassTargets); simpleclassOutputs = sim(net,simpleclassInputs); [c,cm,ind,per] = confusion(simpleclassTargets,simpleclassOutputs)</pre>		

See Also plotconfusion | roc

#### convwf

Purpose	Convolution weight function		
Syntax	Z = convwf(W,P) dim = convwf('size dw = convwf('dw',W info = convwf('cod	, P, Z, FP)	
Description	Weight functions apply weights to an input to get weighted inputs.		
	Z = convwf(W,P) ret an input P.	urns the convolution of a weight matrix ${\tt W}$ and	
	dim = convwf('size',S,R,FP) takes the layer dimension S dimension R, and function parameters, and returns the weigh		
	<pre>dw = convwf('dw',W,P,Z,FP) returns the derivative of Z with response to W. info = convwf('code') returns information about this function. The following codes are defined:     'deriv' Name of derivative function</pre>		
	'fullderiv'	Reduced derivative = 2, full derivative = 1, linear derivative = 0	
=		Input: reduced derivative = 2, full derivative = 1, linear derivative = 0	
	'wfullderiv'	Weight: reduced derivative = 2, full derivative = 1, linear derivative = 0	
	'name'	Full name	
	'fpnames'	Returns names of function parameters	
	'fpdefaults'	Returns default function parameters	

# **Examples** Here you define a random weight matrix W and input vector P and calculate the corresponding weighted input Z.

	<pre>W = rand(4,1); P = rand(8,1); Z = convwf(W,P)</pre>
Network Use	To change a network so an input weight uses convwf, set net.inputWeight{i,j}.weightFcn to 'convwf'. For a layer weight, set net.layerWeight{i,j}.weightFcn to 'convwf'.
	In either ease, call sim to simulate the network with convert

In either case, call sim to simulate the network with convwf.

# defaultderiv

Purpose	Default derivative function		
Syntax	defaultderiv('dperf_dwb',net,X,T,Xi,Ai,EW) defaultderiv('de_dwb',net,X,T,Xi,Ai,EW)		
Description	This function chooses the recommended derivative algorithm for the type of network whose derivatives are being calculated. For static networks, defaultderiv calls staticderiv; for dynamic networks it calls bttderiv to calculate the gradient and fpderiv to calculate the Jacobian.		
	defaultderiv('dperf_dwb',net,X,T,Xi,Ai,EW) takes these arguments,		
	net Neural network X Inputs, an R-by-Q matrix (or N-by-TS cell array of Ri-by-Q matrices)		
	T Targets, an S-by-Q matrix (or M-by-TS cell array Si-by-Q matrices)		
	Xi	Initial input delay states (optional)	
	Ai	Initial layer delay states (optional)	
	EW Error weights (optional)		
	and returns the gradient of performance with respect to the network weights and biases, where R and S are the number of input and our elements and Q is the number of samples (or N and M are the numb input and output signals, Ri and Si are the number of each input outputs elements, and TS is the number of timesteps). defaultderiv('de_dwb',net,X,T,Xi,Ai,EW) returns the Jacobi errors with respect to the network's weights and biases.		
Examples	Here a feedforward network is trained and both the gradient and Jacobian are calculated.		

```
[x,t] = simplefit_dataset;
net = feedforwardnet(10);
net = train(net,x,t);
y = net(x);
perf = perform(net,t,y);
dwb = defaultderiv('dperf_dwb',net,x,t)
See Also bttderiv | fpderiv | num2deriv | num5deriv | staticderiv
```

# disp

Purpose	Neural network properties		
Syntax	disp(net)		
To Get Help	Type help network/disp.		
Description	disp(net) displays a network's properties.		
Examples	Here a perceptron is created and displayed.		
	net = newp([-1 1; 0 2],3); disp(net)		
See Also	display   sim   init   train   adapt		

Purpose	Name and properties of neural network variables	
Syntax	display(net)	
To Get Help	Type help network/display.	
Description	display(net) displays a network variable's name and properties.	
Examples	Here a perceptron variable is defined and displayed.	
	net = newp([-1 1; 0 2],3); display(net)	
	display is automatically called as follows:	
	net	
See Also	disp   sim   init   train   adapt	

Purpose	Euclidean distance weight function		
Syntax	<pre>Z = dist(W,P,FP) dim = dist('size',S,R,FP) dw = dist('dw',W,P,Z,FP) D = dist(pos) info = dist('code')</pre>		
Description	Weight functions apply weights to an input to get weighted inputs.		
	Z = dist(W,P,F	P) takes these inputs,	
	W	S-by-R weight matrix	
	Р	R-by-Q matrix of Q input (column) vectors	
	FP	Struct of function parameters (optional, ignored)	
	and returns the	S-by-Q matrix of vector distances.	
	dim = dist('size',S,R,FP) takes the layer dimension S, input dimension R, and function parameters, and returns the weight size [S-by-R].		
	<pre>dw = dist('dw',W,P,Z,FP) returns the derivative of Z with respec to W.</pre>		
	<pre>dist is also a layer distance function which can be used to find the distances between neurons in a layer. D = dist(pos) takes one argument,</pre>		
	pos	N-by-S matrix of neuron positions	
	and returns the S-by-S matrix of distances. info = dist('code') returns information about this funct following codes are supported:		

	'deriv'	Name of derivative function
	'fullderiv'	Full derivative = 1, linear derivative = 0
	'pfullderiv'	Input: reduced derivative = 2, full derivative = 1, linear derivative = 0
	'name'	Full name
	'fpnames'	Returns names of function parameters
	'fpdefaults'	Returns default function parameters
Examples	-	ndom weight matrix W and input vector P and onding weighted input Z.
	W = rand(4,3); P = rand(3,1); Z = dist(W,P)	
	-	dom matrix of positions for 10 neurons arranged space and find their distances.
	pos = rand(3,10); D = dist(pos)	
Network Use	You can create a stan or newgrnn.	dard network that uses dist by calling newpnn
	net.inputWeight{i,	so an input weight uses dist, set j}.weightFcn to 'dist'. For a layer weight, set j}.weightFcn to 'dist'.
	To change a network net.layers{i}.dist	so that a layer's topology uses dist, set anceFcn to 'dist'.
	In either case, call <b>s</b> i	m to simulate the network with dist.
	See newpnn or newgrn	n for simulation examples.

Algorithms	The Euclidean distance $\boldsymbol{d}$ between two vectors $\boldsymbol{X}$ and $\boldsymbol{Y}$ is		
	d = sum((x-y).^2).^0.5		
See Also	sim   dotprod   negdist   normprod   mandist   linkdist		

Purpose	Distributed delay network		
Syntax	distdelaynet(delays,hiddenSizes,trainFcn)		
Description	Distributed delay networks are similar to feedforward networks, except that each input and layer weights has a tap delay line associated with it. This allows the network to have a finite dynamic response to time series input data. This network is also similar to the time delay neural network (timedelaynet), which only has delays on the input weight.		
	<pre>distdelaynet(delays,hiddenSizes,trainFcn) takes these arguments, delays</pre>		
	hiddenSizes	Row vector of one or more hidden layer sizes (default = 10)	
	trainFcn	Training function (default = 'trainlm')	
	and returns a distribu	ited delay neural network.	
Examples	<pre>Here a distributed delay neural network is used to solve a simple time series problem. [X,T] = simpleseries_dataset; net = distdelaynet({1:2,1:2},10) [Xs,Xi,Ai,Ts] = preparets(net,X,T) net = train(net,Xs,Ts,Xi,Ai); view(net) Y = net(Xs,Xi,Ai); perf = perform(net,Y,Ts)</pre>		
See Also	preparets   removedelay   timedelaynet   narnet   narxnet		

# divideblock

Purpose	Divide targets into three sets using blocks of indices		
Syntax	[trainInd,valInd,testInd] = divideblock(Q,trainRatio,valRatio, testRatio)		
Description	<pre>[trainInd,valInd,testInd] = divideblock(Q,trainRatio,valRatio,testRatio) separates targets into three sets: training, validation, and testing. It takes the following inputs:</pre>		
	Q	Number of targets to divide up.	
	trainRatio	Ratio of targets for training. Default = $0.7$ .	
	valRatioRatio of targets for validation. Default = 0.testRatioRatio of targets for testing. Default = 0.15.		
	and returns trainInd valInd testInd	Training indices Validation indices Test indices	
Examples Network Use	<pre>[trainInd,valInd,testInd] = divideblock(3000,0.6,0.2,0.2); Here are the network properties that define which data division function to use, what its parameters are, and what aspects of targets are divided up, when train is called. net.divideFcn</pre>		
	net.divideParam net.divideMode		
See Also	divideind   divideint   dividerand   dividetrain		

Purpose	Divide targets into three sets using specified indices	
Syntax	[trainInd,valInd,t testInd)	estInd] = divideind(Q,trainInd,valInd,
Description	ption[trainInd,valInd,testInd] = divideind(Q,trainInd,valInd,testInd) separates targets into three sets: training, validation, and testing, according to indices provided. It actually returns the same indices it receives as arguments; its purpose 	
	trainInd	Training indices
	valInd	Validation indices
	testInd	Test indices
	and returns	
	trainIndTraining indices (unchanged)valIndValidation indices (unchanged)	
testInd		Test indices (unchanged)
Examples	[trainInd,valInd,testInd] = divideind(3000,1:2000,2001:2500,2501:3000);	
Network Use	Here are the network properties that define which data division function to use, what its parameters are, and what aspects of targets are divided up, when train is called.	

# divideind

net.divideFcn net.divideParam net.divideMode

See Also divideblock | divideint | dividerand | dividetrain

## divideint

Purpose	Divide targets into th	Divide targets into three sets using interleaved indices	
Syntax	[trainInd,valInd,testInd] = divideint(Q,trainRatio,valRatio, testRatio)		
Description	<pre>[trainInd,valInd,testInd] = divideint(Q,trainRatio,valRatio,testRatio) separates targets into three sets: training, validation, and testing. It takes the following inputs,</pre>		
	Q	Number of targets to divide up.	
	trainRatio	Ratio of vectors for training. Default = $0.7$ .	
	valRatio	Ratio of vectors for validation. Default = $0.15$ .	
	testRatio Ratio of vectors for testing. Default = 0.15.		
	and returns trainInd Training indices		
	valInd	Validation indices	
	testInd	Test indices	
Examples Network Use	<pre>[trainInd,valInd,testInd] = divideint(3000,0.6,0.2,0.2); Here are the network properties that define which data division function to use, what its parameters are, and what aspects of targets</pre>		
	<pre>are divided up, when train is called. net.divideFcn net.divideParam net.divideMode</pre>		
See Also	divideblock   divideind   dividerand   dividetrain		

# dividerand

Purpose	Divide targets into three sets using random indices		
Syntax	[trainInd,valInd,testInd] = dividerand(Q,trainRatio,valRatio, testRatio)		
Description	<pre>[trainInd,valInd,testInd] = dividerand(Q,trainRatio,valRatio,testRatio) separates targets into three sets: training, validation, and testing. It takes the following inputs,</pre>		
	Q	Number of targets to divide up.	
	trainRatio	Ratio of vectors for training. Default = $0.7$ .	
	valRatio	Ratio of vectors for validation. Default = $0.15$ .	
	testRatio Ratio of vectors for testing. Default = 0.15		
	and returns trainInd valInd testInd	Training indices Validation indices Test indices	
Examples Network	<pre>[trainInd,valInd,testInd] = dividerand(3000,0.6,0.2,0.2); Here are the network properties that define which data division</pre>		
Use	function to use, what its parameters are, and what aspects of targets are divided up, when train is called.		
	net.divideFcn net.divideParam net.divideMode		
See Also	divideblock   divideind   divideint   dividetrain		

Purpose	Assign all targets to training set		
Syntax	[trainInd,valInd,t testRatio)	[trainInd,valInd,testInd] = dividetrain(Q,trainRatio,valRatio, testRatio)	
Description	<pre>[trainInd,valInd,testInd] = dividetrain(Q,trainRatio,valRatio,testRatio) assigns all targets to the training set and no targets to either the validation or test sets. It takes the following inputs,</pre>		
	Q	Number of targets to divide up.	
	and returns		
	trainInd	Training indices equal to 1:Q	
	valInd	Empty validation indices, []	
	testInd	Empty test indices, []	
Examples	<pre>[trainInd,valInd,testInd] = dividetrain(3000);</pre>		
Network Use	Here are the network properties that define which data division function to use, what its parameters are, and what aspects of targets are divided up, when train is called. net.divideFcn net.divideParam net.divideMode		
See Also	divideblock   divideind   divideint   dividerand		

# dotprod

Purpose	Dot product weight function	
Syntax	<pre>Z = dotprod(W,P,FP) dim = dotprod('size',S,R,FP) dw = dotprod('dw',W,P,Z,FP) info = dotprod('code')</pre>	
Description	Weight functions apply weights to an input to get weighted inputs.	
	Z = dotprod(W,P,FP)	) takes these inputs,
	W	S-by-R weight matrix
	Р	R-by-Q matrix of Q input (column) vectors
	FP	Struct of function parameters (optional, ignored)
	<pre>and returns the S-by-Q dot product of W and P. dim = dotprod('size',S,R,FP) takes the layer dimension S, input dimension R, and function parameters, and returns the weight size [S-by-R]. dw = dotprod('dw',W,P,Z,FP) returns the derivative of Z with respect to W. info = dotprod('code') returns information about this function. The following codes are defined:</pre>	
	'deriv'	Name of derivative function
	'pfullderiv'	Input: reduced derivative = 2, full derivative = 1, linear derivative = 0
-		Weight: reduced derivative = 2, full derivative = 1, linear derivative = 0

	'name'	Full name
	'fpnames' 'fpdefaults'	Returns names of function parameters Returns default function parameters
Examples	Here you define a random weight matrix W and input vector P and calculate the corresponding weighted input Z.	
	W = rand(4,3); P = rand(3,1); Z = dotprod(W,P)	
Network Use	You can create a stan feedforwardnet.	dard network that uses dotprod by calling
	To change a network so an input weight uses dotprod, set net.inputWeight{i,j}.weightFcn to 'dotprod'. For a layer weight, set net.layerWeight{i,j}.weightFcn to 'dotprod'.	
	In either case, call <b>s</b> i	m to simulate the network with dotprod.
See Also	sim   dist   feedfor	wardnet   negdist   normprod

# elliotsig

Purpose	Elliot symmetric sigmoid transfer function		
Syntax	A = elliotsig(N)		
Description	Transfer functions convert a neural network layer's net input into its net output.		
	A = elliotsig(N) takes an S-by-Q matrix of S N-element net input column vectors and returns an S-by-Q matrix A of output vectors, where each element of N is squashed from the interval [-inf inf] to the interval [-1 1] with an "S-shaped" function.		
	The advantage of this transfer function over other sigmoids is that it is fast to calculate on simple computing hardware as it does not require any exponential or trigonometric functions. Its disadvantage is that it only flattens out for large inputs, so its effect is not as local as other sigmoid functions. This might result in more training iterations, or require more neurons to achieve the same accuracy.		
Examples	Calculate a layer output from a single net input vector:		
	n = [0; 1; -0.5; 0.5]; a = elliotsig(n);		
	<pre>a = elliotsig(n);</pre>		
	<pre>a = elliotsig(n); Plot the transfer function: n = -5:0.01:5; plot(n, elliotsig(n))</pre>		
	<pre>a = elliotsig(n); Plot the transfer function: n = -5:0.01:5; plot(n, elliotsig(n)) set(gca,'dataaspectratio',[1 1 1],'xgrid','on','ygrid','on') For a network you have already defined, change the transfer function</pre>		

Purpose	Elliot 2 symmetric sigmoid transfer function	
Syntax	A = elliot2sig(N)	
Description	Transfer functions convert a neural network layer's net input into its net output. This function is a variation on the original Elliot sigmoid function. It has a steeper slope, closer to tansig, but is not as smooth at the center.	
	A = elliot2sig(N) takes an S-by-Q matrix of S N-element net input column vectors and returns an S-by-Q matrix A of output vectors, where each element of N is squashed from the interval [-inf inf] to the interval [-1 1] with an "S-shaped" function.	
	The advantage of this transfer function over other sigmoids is that it is fast to calculate on simple computing hardware as it does not require any exponential or trigonometric functions. Its disadvantage is that it departs from the classic sigmoid shape around zero.	
Examples	Calculate a layer output from a single net input vector:	
	n = [0; 1; -0.5; 0.5]; a = elliot2sig(n);	
	Plot the transfer function:	
	n = -5:0.01:5; plot(n, elliot2sig(n)) set(gca,'dataaspectratio',[1 1 1],'xgrid','on','ygrid','on')	
	For a network you have already defined, change the transfer function for layer i:	
	<pre>net.layers{i}.transferFcn = 'elliot2sig';</pre>	
See Also	elliotsig   logsig   tansig	

#### elmannet

Purpose	Elman neural network		
Syntax	elmannet(layerdelays,hiddenSizes,trainFcn)		
Description	Elman networks are feedforward networks (feedforwardnet) with the addition of layer recurrent connections with tap delays.		
	With the availability of full dynamic derivative calculations (fpderiv and bttderiv), the Elman network is no longer recommended except for historical and research purposes. For more accurate learning try time delay (timedelaynet), layer recurrent (layrecnet), NARX (narxnet), and NAR (narnet) neural networks.		
	Elman networks with one or more hidden layers can learn any dynamic input-output relationship arbitrarily well, given enough neurons in the hidden layers. However, Elman networks use simplified derivative calculations (using staticderiv, which ignores delayed connections) at the expense of less reliable learning.		
	elmannet(layerdelays,hiddenSizes,trainFcn) takes these arguments,		
	layerdelaysRow vector of increasing 0 or positive delays (default = 1:2)		
	hiddenSizes	Row vector of one or more hidden layer sizes (default = 10)	
	trainFcn	Training function (default = 'trainlm')	
	and returns an Elman neural network.		
Examples	<pre>Here an Elman neural network is used to solve a simple time series problem. [X,T] = simpleseries_dataset; net = elmannet(1:2,10); [Xs,Xi,Ai,Ts] = preparets(net,X,T);</pre>		

```
net = train(net,Xs,Ts,Xi,Ai);
view(net)
Y = net(Xs,Xi,Ai);
perf = perform(net,Ts,Y)
See Also preparets | removedelay | timedelaynet | layrecnet | narnet
| narxnet
```

## errsurf

Purpose	Error surface of single-input neuron	
Syntax	errsurf(P,T,WV,BV,F)	
Description	errsurf(P,T,WV,BV,F) takes these arguments,	
	Р	1-by-Q matrix of input vectors
	Т	1-by-Q matrix of target vectors
	WV	Row vector of values of W
	BV	Row vector of values of B
	F	Transfer function (string)
	and returns a m	atrix of error values over WV and BV.
Examples	<pre>p = [-6.0 -6.1 -4.1 -4.0 +4.0 +4.1 +6.0 +6.1]; t = [+0.0 +0.0 +.97 +.99 +.01 +.03 +1.0 +1.0]; wv = -1:.1:1; bv = -2.5:.25:2.5; es = errsurf(p,t,wv,bv,'logsig'); plotes(wv,bv,es,[60 30])</pre>	
See Also	plotes	

Purpose	Extend time series data to given number of timesteps		
Syntax	<pre>extendts(x,ts,v)</pre>		
Description	extendts(x,ts	<pre>extendts(x,ts,v) takes these values,</pre>	
	x	Neural network time series data	
	ts	Number of timesteps	
	v	Value	
Examples	<pre>v value and returns the time series data either extended or truncated to match the specified number of timesteps. If the value v is specified, then extended series are filled in with that value, otherwise they are extended with random values. Here, a 20-timestep series is created and then extended to 25 timesteps with the value zero. x = nndata(5,4,20); y = extendts(x,25,0)</pre>		
See Also	nndata   catsamples   preparets		

# feedforwardnet

Purpose	Feedforward neural network			
Syntax	feedforwardnet(hiddenSizes,trainFcn)			
Description	Feedforward networks consist of a series of layers. The first layer has a connection from the network input. Each subsequent layer has a connection from the previous layer. The final layer produces the network's output.			
	Feedforward networks can be used for any kind of input to output mapping. A feedforward network with one hidden layer and enough neurons in the hidden layers, can fit any finite input-output mapping problem.			
	Specialized versions of the feedforward network include fitting (fitnet) and pattern recognition (patternnet) networks. A variation on the feedforward network is the cascade forward network (cascadeforwardnet) which has additional connections from the input to every layer, and from each layer to all following layers.			
	feedforwardnet(hiddenSizes,trainFcn) takes these arguments,			
	hiddenSizes Row vector of one or more hidden layer sizes (default = 10)			
	<pre>trainFcn Training function (default = 'trainlm')</pre>			
	and returns a feedforward neural network.			
Examples	Here a feedforward neural network is used to solve a simple problem.			
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(10) net = train(net,x,t); view(net) y = net(x); perf = perform(net,y,t)</pre>			

**See Also** fitnet | patternnet | cascadeforwardnet

## fitnet

Purpose	Function fitting neural network	
Syntax	fitnet(hiddenSize	es,trainFcn)
Description	Fitting networks are feedforward neural networks (feedforwardnet) used to fit an input-output relationship.	
	fitnet(hiddenSize	es,trainFcn) takes these arguments,
	hiddenSizes	Row vector of one or more hidden layer sizes (default = 10)
	trainFcn	Training function (default = 'trainlm')
Examples	and returns a fitting neural network. Here a fitting neural network is used to solve a simple problem.	
-	<pre>[x,t] = simplefit_dataset; net = fitnet(10) net = train(net,x,t); view(net) y = net(x); perf = perform(net,y,t)</pre>	
See Also	feedforwardnet   nftool	

Purpose	Process data by marking rows with unknown values		
Syntax	<pre>[y,ps] = fixunknowns(X) [y,ps] = fixunknowns(X,FP) Y = fixunknowns('apply',X,PS) X = fixunknowns('reverse',Y,PS) name = fixunknowns('name') fp = fixunknowns('pdefaults') pd = fixunknowns('pdesc') fixunknowns('pcheck',fp)</pre>		
Description	fixunknowns processes matrixes by replacing each row containing unknown values (represented by NaN) with two rows of information.		
	The first row contains the original row, with NaN values replaced by t row's mean. The second row contains 1 and 0 values, indicating whic values in the first row were known or unknown, respectively.		
	[y,ps] = fixunknowns(X) takes these inputs,		
	X	Single N-by-Q matrix or a 1-by-TS row cell array of N-by-Q matrices	
	and returns		
	Y	Each M-by-Q matrix with M - N rows added (optional)	
	PS	Process settings that allow consistent processing of values	
	[y,ps] = fixun	knowns(X,FP) takes an empty struct FP of parameters.	
	Y = fixunknown	s('apply',X,PS) returns Y, given X and settings PS.	
	X = fixunknowns('reverse',Y,PS) returns X, given Y and setting		

	<pre>name = fixunknowns('name') returns the name of this process method.</pre>		
	<pre>fp = fixunknowns('pdefaults') returns the default process parameter structure.</pre>		
	<pre>pd = fixunknowns('pdesc') returns the process parameter descriptions.</pre>		
	fixunknowns('pcheck',fp) throws an error if any parameter is illegal.		
Examples	Here is how to format a matrix with a mixture of known and unknown values in its second row:		
	x1 = [1 2 3 4; 4 NaN 6 5; NaN 2 3 NaN] [y1,ps] = fixunknowns(x1)		
	Next, apply the same processing settings to new values:		
	x2 = [4 5 3 2; NaN 9 NaN 2; 4 9 5 2] y2 = fixunknowns('apply',x2,ps)		
	Reverse the processing of y1 to get x1 again.		
	x1_again = fixunknowns('reverse',y1,ps)		
Definitions	If you have input data with unknown values, you can represent them with NaN values. For example, here are five 2-element vectors with unknown values in the first element of two of the vectors:		
	p1 = [1 NaN 3 2 NaN; 3 1 -1 2 4];		
	The network will not be able to process the NaN values properly. Use the function fixunknowns to transform each row with NaN values (in this case only the first row) into two rows that encode that same information numerically.		
	<pre>[p2,ps] = fixunknowns(p1);</pre>		

Here is how the first row of values was recoded as two rows.

p2 = 1 2 3 2 2 1 0 1 1 0 3 1 -1 2 4

The first new row is the original first row, but with the mean value for that row (in this case 2) replacing all NaN values. The elements of the second new row are now either 1, indicating the original element was a known value, or 0 indicating that it was unknown. The original second row is now the new third row. In this way both known and unknown values are encoded numerically in a way that lets the network be trained and simulated.

Whenever supplying new data to the network, you should transform the inputs in the same way, using the settings ps returned by fixunknowns when it was used to transform the training input data.

p2new = fixunknowns('apply',p1new,ps);

The function fixunkowns is only recommended for input processing. Unknown targets represented by NaN values can be handled directly by the toolbox learning algorithms. For instance, performance functions used by backpropagation algorithms recognize NaN values as unknown or unimportant values.

See Also mapminmax | mapstd | processpca

## formwb

Purpose	Form bias and weights into single vector		
Syntax	<pre>formwb(net,b,IW,LW)</pre>		
Description	formwb(net,b,IW,LW) takes a neural network and bias b, input weight IW, and layer weight LW values, and combines the values into a single vector.		
Examples	Here a network is created, configured, and its weights and biases formed into a vector.		
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(10); net = configure(net,x,t); wb = formwb(net,net.b,net.IW,net.LW)</pre>		
See Also	getwb   setwb   separatewb		

Purpose	Forward propagation derivative function		
Syntax	fpderiv('dperf_dwb',net,X,T,Xi,Ai,EW) fpderiv('de_dwb',net,X,T,Xi,Ai,EW)		
Description	This function calculates derivatives using the chain rule from inputs to outputs, and in the case of dynamic networks, forward through time. fpderiv('dperf_dwb',net,X,T,Xi,Ai,EW) takes these arguments,		
	net	Neural network	
	х	Inputs, an R-by-Q matrix (or N-by-TS cell array of Ri-by-Q matrices)	
	т	Targets, an S-by-Q matrix (or M-by-TS cell array of Si-by-Q matrices)	
	Xi	Initial input delay states (optional)	
	Ai	Initial layer delay states (optional)	
	EW	Error weights (optional)	
	and returns the gradient of performance with respect to the network weights and biases, where R and S are the number of input and out elements and Q is the number of samples (or N and M are the number input and output signals, Ri and Si are the number of each input a outputs elements, and TS is the number of timesteps).		
	fpderiv('de_dwb',net,X,T,Xi,Ai,EW) returns the Jacobian of error with respect to the network's weights and biases.		
Examples	<pre>Here a feedforward network is trained and both the gradient and Jacobian are calculated. [x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t);</pre>		

```
y = net(x);
perf = perform(net,t,y);
gwb = fpderiv('dperf_dwb',net,x,t)
jwb = fpderiv('de_dwb',net,x,t)
See Also bttderiv | defaultderiv | num2deriv | num5deriv | staticderiv
```

# fromnndata

Purpose	Convert data from standard neural network cell array form			
Syntax	<pre>fromnndata(x,toMatrix,columnSample,cellTime)</pre>			
Description	<pre>fromnndata(x,toMatrix,columnSample,cellTime) takes these arguments,</pre>			
	net	Neural network		
	toMatrix	True if result is to be in matrix form		
	columnSample	True if samples are to be represented as columns, false if rows		
	cellTime	True if time series are to be represented as a cell array, false if represented with a matrix		
	and returns the original data reformatted accordingly.			
Examples	Here time-series data is converted from a matrix representation standard cell array representation, and back. The original data co of a 5-by-6 matrix representing one time-series sample consisting a 5-element vector over 6 timesteps arranged in a matrix with th samples as columns.			
	<pre>x = rands(5,6) columnSamples = true; % samples are by columns. cellTime = false; % time-steps represented by a matrix, not cell. [y,wasMatrix] = tonndata(x,columnSamples,cellTime) x2 = fromnndata(y,wasMatrix,columnSamples,cellTime)</pre>			
	Here data is defined in standard neural network data cell form. Converting this data does not change it. The data consists of three time series samples of 2-element signals over 3 timesteps.			
	<pre>x = {rands(2,3); rands(2,3); rands(2,3)} columnSamples = true; cellTime = true; [y,wasMatrix] = tonndata(x)</pre>			

# fromnndata

x2 = fromnndata(y,wasMatrix,columnSamples)

See Also tonndata

Purpose	Generalized addition
Syntax	gadd(a,b)
Description	This function generalizes matrix addition to the addition of cell arrays of matrices combined in an element-wise fashion.
	gadd(a,b) takes two matrices or cell arrays, and adds them in an element-wise manner.
Examples	Here matrix and cell array values are added.
	gadd([1 2 3; 4 5 6],[10;20]) gadd({1 2; 3 4},{1 3; 5 2}) gadd({1 2 3 4},{10;20;30})
See Also	gsubtract   gmultiply   gdivide   gnegate   gsqrt

# <u>gdivide</u>

Purpose	Generalized division
Syntax	gdivide(a,b)
Description	This function generalizes matrix element-wise division to the division of cell arrays of matrices combined in an element-wise fashion.
	gdivide(a,b) takes two matrices or cell arrays, and divides them in an element-wise manner.
Examples	Here matrix and cell array values are added.
	gdivide([1 2 3; 4 5 6],[10;20]) gdivide({1 2; 3 4},{1 3; 5 2}) gdivide({1 2 3 4},{10;20;30})
See Also	gadd   gsubtract   gmultiply   gnegate   gsqrt

Purpose	Generate Simulir	nk block for neural network simulation
Syntax	gensim(net,st)	
To Get Help	Type help network/gensim.	
Description	gensim(net,st) creates a Simulink <sup>®</sup> system containing a block that simulates neural network net.	
	gensim(net,st) takes these inputs:	
	net	Neural network
	st	Sample time (default = 1)
	and creates a Simulink system containing a block that simulates ne network net with a sampling time of st. If net has no input or layer delays (net.numInputDelays and net.numLayerDelays are both 0), you can use -1 for st to get a net that samples continuously.	
Examples	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(10); net = train(net,x,t) gensim(net)</pre>	

# <u>ge</u>telements

Purpose	Get neural network data elements
Syntax	<pre>getelements(x,ind)</pre>
Description	<pre>getelements(x,ind) returns the elements of neural network data x indicated by the indices ind. The neural network data may be in matrix or cell array form.</pre>
	If $x$ is a matrix, the result is the ind rows of $x$ .
	If x is a cell array, the result is a cell array with as many columns as x, whose elements (1,i) are matrices containing the ind rows of [x{:,i}].
Examples	This code gets elements 1 and 3 from matrix data:
	x = [1 2 3; 4 7 4] y = getelements(x,[1 3])
	This code gets elements 1 and 3 from cell array data:
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} y = getelements(x,[1 3])</pre>
See Also	nndata   numelements   setelements   catelements   getsamples   gettimesteps   getsignals

Purpose	Get neural network data samples
Syntax	getsamples(x,ind)
Description	getsamples(x,ind) returns the samples of neural network data x indicated by the indices ind. The neural network data may be in matrix or cell array form.
	If x is a matrix, the result is the ind columns of x.
	If $x$ is a cell array, the result is a cell array the same size as $x$ , whose elements are the ind columns of the matrices in $x$ .
Examples	This code gets samples 1 and 3 from matrix data:
	x = [1 2 3; 4 7 4] y = getsamples(x,[1 3])
	This code gets elements 1 and 3 from cell array data:
	x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} y = getsamples(x,[1 3])
See Also	nndata   numsamples   setsamples   catsamples   getelements   gettimesteps   getsignals

# getsignals

Purpose	Get neural network data signals
Syntax	getsignals(x,ind)
Description	getsignals(x,ind) returns the signals of neural network data x indicated by the indices ind. The neural network data may be in matrix or cell array form.
	If x is a matrix, ind may only be 1, which will return x, or [] which will return an empty matrix.
	If $x$ is a cell array, the result is the ind rows of $x$ .
Examples	This code gets signal 2 from cell array data:
	$x = \{[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]\}$ y = getsignals(x,2)
See Also	nndata   numsignals   setsignals   catsignals   getelements   getsamples   gettimesteps

Purpose	Get Simulink neura	l network block initial input and layer delays states
Syntax	[xi,ai] = getsiminit(sysName,netName,net)	
Description	<pre>[xi,ai] = getsiminit(sysName,netName,net) takes these arguments,</pre>	
	sysName	The name of the Simulink system containing the neural network block
	netName	The name of the Simulink neural network block
	net	The original neural network
	and returns,	
	xi	Initial input delay states
	ai	Initial layer delay states
Examples	<pre>Here a NARX network is designed. The NARX network has a stand input and an open-loop feedback output to an associated feedback in [x,t] = simplenarx_dataset; net = narxnet(1:2,1:2,20);</pre>	
<pre>view(net) [xs,xi,ai,ts] = preparets(net,x,{},t); net = train(net,xs,ts,xi,ai); y = net(xs,xi,ai);</pre>		<pre>s] = preparets(net,x,{},t); net,xs,ts,xi,ai);</pre>
	Now the network is converted to closed-loop, and the data is reformatted to simulate the network's closed-loop response.	
	net = closeloop(n view(net) [xs,xi,ai,ts] = p	uet); preparets(net,x,{},t);

y = net(xs,xi,ai);

Here the network is converted to a Simulink system with workspace input and output ports. Its delay states are initialized, inputs X1 defined in the workspace, and it is ready to be simulated in Simulink.

```
[sysName,netName] = gensim(net,'InputMode','Workspace',...
'OutputMode','WorkSpace','SolverMode','Discrete');
setsiminit(sysName,netName,net,xi,ai,1);
x1 = nndata2sim(x,1,1);
```

Finally the initial input and layer delays are obtained from the Simulink model. (They will be identical to the values set with setsiminit.)

[xi,ai] = getsiminit(sysName,netName,net);

See Also gensim | setsiminit | nndata2sim | sim2nndata

Purpose	Get neural network data timesteps	
Syntax	gettimesteps(x,ind)	
Description	gettimesteps(x,ind) returns the timesteps of neural network data x indicated by the indices ind. The neural network data may be in matrix or cell array form.	
	If x is a matrix, ind can only be 1, which will return x; or [], which will return an empty matrix.	
	If $x$ is a cell array the result is the ind columns of $x$ .	
Examples	This code gets timestep 2 from cell array data:	
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} y = gettimesteps(x,2)</pre>	
See Also	nndata   numtimesteps   settimesteps   cattimesteps   getelements   getsamples   getsignals	

# getwb

Purpose	Get network weight and bias values as single vector
Syntax	getwb(net)
Description	getwb(net) returns a neural network's weight and bias values as a single vector.
Examples	Here a feedforward network is trained to fit some data, then its bias and weight values are formed into a vector.
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); wb = getwb(net)</pre>
See Also	setwb   formwb   separatewb

# gmultiply

Purpose	Generalized multiplication	
Syntax	gmultiply(a,b)	
Description	This function generalizes matrix multiplication to the multiplication of cell arrays of matrices combined in an element-wise fashion.	
	gmultiply(a,b) takes two matrices or cell arrays, and multiplies them in an element-wise manner.	
Examples	Here matrix and cell array values are added.	
	gmultiply([1 2 3; 4 5 6],[10;20]) gmultiply({1 2; 3 4},{1 3; 5 2}) gmultiply({1 2 3 4},{10;20;30})	
See Also	gadd   gsubtract   gdivide   gnegate   gsqrt	

### gnegate

Purpose	Generalized negation
Syntax	gnegate(x)
Description	This function generalizes matrix negation to the negation of cell arrays of matrices combined in an element-wise fashion.
	gnegate(x) takes a matrix or cell array of matrices, and negates the matrices.
Examples	Here is an example of negating a cell array:
	<pre>x = {[1 2; 3 4],[1 3; 5 2]}; y = gnegate(x); y{1}, y{2}</pre>
See Also	gadd   gsubtract   gdivide   gmultiply   gsqrt

Purpose	Reformat neural data back from GPU
Syntax	X = gpu2nndata(Y,Q) X = gpu2nndata(Y) X = gpu2nndata(Y,Q,N,TS)
Description	Training and simulation of neural networks require that matrices be transposed. But they do not require (although they are more efficient with) padding of column length so that each column is memory aligned. This function copies data back from the current GPU and reverses this transform. It can be used on data formatted with nndata2gpu or on the results of network simulation.
	X = gpu2nndata(Y,Q) copies the QQ-by-N gpuArray Y into RAM, takes the first Q rows and transposes the result to get an N-by-Q matrix representing Q N-element vectors.
	X = gpu2nndata(Y) calculates Q as the index of the last row in Y that is not all NaN values (those rows were added to pad Y for efficient GPU computation by nndata2gpu). Y is then transformed as before.
	X = gpu2nndata(Y,Q,N,TS) takes a QQ-by-(N*TS) gpuArray where N is a vector of signal sizes, Q is the number of samples (less than or equal to the number of rows after alignment padding QQ), and TS is the number of time steps.
	The gpuArray Y is copied back into RAM, the first Q rows are taken, and then it is partitioned and transposed into an M-by-TS cell array, where M is the number of elements in N. Each Y{i,ts} is an N(i)-by-Q matrix.
Examples	Copy a matrix to the GPU and back:
	x = rand(5,6) [y,q] = nndata2gpu(x) x2 = gpu2nndata(y,q)
	Copy from the GPU a neural network cell array data representing four time series, each consisting of five time steps of 2-element and

3-element signals.

x = nndata([2;3],4,5)
[y,q,n,ts] = nndata2gpu(x)
x2 = gpu2nndata(y,q,n,ts)

#### See Also

nndata2gpu

Purpose	Grid layer topology function	
Syntax	gridtop(dim1,dim2,,dimN)	
Description	<pre>pos = gridtop calculates neuron positions for layers whose neurons are arranged in an N-dimensional grid. gridtop(dim1,dim2,,dimN) takes N arguments,</pre>	
	dimi Length of layer in dimension i	
	and returns an N-by-S matrix of N coordinate vectors where S is the product of dim1*dim2**dimN.	
Examples	This code uses gridtop to directly create a two-dimensional layer with 40 neurons arranged in an 8-by-5 grid; then uses the function as an input to selforgmap to create weight positions of neurons for a self-organizing map and plots the neuron topology.	
	<pre>pos = gridtop(8,5); net = selforgmap([8 5],'topologyFcn','gridtop'); plotsomtop(net)</pre>	
See Also	hextop   randtop   tritop	

## gsqrt

Purpose	Generalized square root	
Syntax	gsqrt(x)	
Description	This function generalizes matrix element-wise square root to the square root of cell arrays of matrices combined in an element-wise fashion.	
	gsqrt(x) takes a matrix or cell array of matrices, and takes the element-wise square root of the matrices.	
Examples	Here is an example of taking the element-wise square root of a cell array:	
	gsqrt({1 2; 3 4},{1 3; 5 2})	
See Also	gadd   gsubtract   gdivide   gmultiply   gnegate	

# gsubtract

Purpose	Generalized subtraction	
Syntax	gsubtract(a,b)	
Description	This function generalizes matrix subtraction to the subtraction of cell arrays of matrices combined in an element-wise fashion.	
	gsubtract(a,b) takes two matrices or cell arrays, and subtracts them in an element-wise manner.	
Examples	Here matrix and cell array values are added.	
	gsubtract([1 2 3; 4 5 6],[10;20]) gsubtract({1 2; 3 4},{1 3; 5 2}) gsubtract({1 2 3 4},{10;20;30})	
See Also	gadd   gmultiply   gdivide   gnegate   gsqrt	

#### hardlim

Purpose	Hard-limit tran	sfer function
Graph and Symbol	a h+1 0 -1 a = hardlim(n) Hard-Limit Trans	- ▶ n ⊥⊥ fer Function
Syntax	A = hardlim(N	,FP)
Description	hardlim is a neural transfer function. Transfer functions calculate a layer's output from its net input.	
	A = hardlim(N	,FP) takes N and optional function parameters,
	Ν	S-by-Q matrix of net input (column) vectors
	FP	Struct of function parameters (ignored)
	and returns A, t	he S-by-Q Boolean matrix with 1s where $N \ge 0$ .
	<pre>info = hardlim('code') returns information according to the code string specified:</pre>	
	hardlim('name') returns the name of this function.	
	hardlim('output', FP) returns the [min max] output range.	
	hardlim('acti	ve', FP) returns the [min max] active input range.
	hardlim('full S-by-S-by-Q or S	deriv') returns 1 or 0, depending on whether dA_dN is -by-Q.
	hardlim('fpna	mes') returns the names of the function parameters.
	hardlim('fpde	faults') returns the default function parameters.

Examples	Here is how to create a plot of the hardlim transfer funct		
	n = -5:0.1:5; a = hardlim(n); plot(n,a)		
	Assign this transfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'hardlim';</pre>		
Algorithms	$\begin{aligned} \texttt{hardlim}(\texttt{n}) &= 1 \text{ if } \texttt{n} \geq 0 \\ 0 \text{ otherwise} \end{aligned}$		
See Also	sim   hardlims		

#### hardlims

Purpose	Symmetric hard-limit transfer function		
Graph and Symbol	$a \rightarrow n$ $a \rightarrow n$ $a = hardlims(n)$ Symmetric Hard-Limit Transfer Function		
Syntax	A = hardlims(N,FP)		
Description	hardlims is a neural transfer function. Transfer functions calculate a layer's output from its net input.		
	A = hardlims(N,FP) takes N and optional function parameters,		
	N 0 h 0 h for a first in such (as larger) most and		
	N S-by-Q matrix of net input (column) vectors		
	NS-by-Q matrix of net input (column) vectorsFPStruct of function parameters (ignored)		
	· · · · · · · · · · · · · · · · · · ·		
	FP Struct of function parameters (ignored)		
	FPStruct of function parameters (ignored)and returns A, the S-by-Q +1/-1 matrix with +1s where $N \ge 0$ .info = hardlims('code') returns information according to the code		
	FPStruct of function parameters (ignored)and returns A, the S-by-Q +1/-1 matrix with +1s where $N \ge 0$ .info = hardlims('code') returns information according to the code string specified:		
	<pre>FP Struct of function parameters (ignored) and returns A, the S-by-Q +1/−1 matrix with +1s where N ≥ 0. info = hardlims('code') returns information according to the code string specified: hardlims('name') returns the name of this function.</pre>		
	<pre>FP Struct of function parameters (ignored) and returns A, the S-by-Q +1/-1 matrix with +1s where N ≥ 0. info = hardlims('code') returns information according to the code string specified: hardlims('name') returns the name of this function. hardlims('output',FP) returns the [min max] output range.</pre>		
	<pre>FP Struct of function parameters (ignored) and returns A, the S-by-Q +1/-1 matrix with +1s where N ≥ 0. info = hardlims('code') returns information according to the code string specified: hardlims('name') returns the name of this function. hardlims('output',FP) returns the [min max] output range. hardlims('active',FP) returns the [min max] active input range. hardlims('fullderiv') returns 1 or 0, depending on whether dA_dN is</pre>		

#### hardlims

Examples	Here is how to create a plot of the hardlims transfer function		
	<pre>n = -5:0.1:5; a = hardlims(n); plot(n,a)</pre>		
	Assign this transfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'hardlims';</pre>		
Algorithms	hardlims(n) = 1 if $n \ge 0, -1$ otherwise.		
See Also	sim   hardlim		

### hextop

Purpose	Hexagonal layer topology function	
Syntax	hextop(dim1,dim2,,dimN)	
Description	hextop calculates the neuron positions for layers whose neurons are arranged in an N-dimensional hexagonal pattern.	
	hextop(dim1,dim2,,dimN) takes N arguments,	
	dimi Length of layer in dimension i	
	and returns an N-by-S matrix of N coordinate vectors where S is the product of dim1*dim2**dimN.	
Examples	This code creates and displays a two-dimensional layer with 40 neurons arranged in an 8-by-5 hexagonal pattern.	
	<pre>pos = hextop(8,5); net = selforgmap([8 5],'topologyFcn','hextop'); plotsomtop(net)</pre>	
See Also	gridtop   randtop   tritop	

Purpose	Convert indices to vectors	
Syntax	ind2vec(ind)	
Description	ind2vec and vec2ind allow indices to be represented either by themselves, or as vectors containing a 1 in the row of the index they represent.	
	<pre>ind2vec(ind) takes one argument,</pre>	
	ind Row vector of indices	
	and returns a sparse matrix of vectors, with one 1 in each column, as indicated by ind.	
Examples	Here four indices are defined and converted to vector representation.	
	ind = [1 3 2 3] vec = ind2vec(ind)	
See Also	vec2ind	

#### init

Purpose	Initialize neural network
Syntax	<pre>net = init(net)</pre>
To Get Help	Type help network/init.
Description	<pre>net = init(net) returns neural network net with weight and bias values updated according to the network initialization function, indicated by net.initFcn, and the parameter values, indicated by net.initParam.</pre>
Examples	<pre>Here a perceptron is created, and then configured so that its input, output, weight, and bias dimensions match the input and target data. x = [0 1 0 1; 0 0 1 1]; t = [0 0 0 1]; net = perceptron; net = configure(net,x,t); net.iw{1,1} net.b{1} Training the perceptron alters its weight and bias values. net = train(net,x,t); net.iw{1,1} net.b{1} init reinitializes those weight and bias values. net = init(net); net.iw{1,1} net.b{1} The weights and biases are zeros again, which are the initial values</pre>
	The weights and biases are zeros again, which are the initial values used by perceptron networks.

# Algorithms init calls net.initFcn to initialize the weight and bias values according to the parameter values net.initParam.

Typically, net.initFcn is set to 'initlay', which initializes each layer's weights and biases according to its net.layers{i}.initFcn.

Backpropagation networks have net.layers{i}.initFcn set to 'initnw', which calculates the weight and bias values for layer i using the Nguyen-Widrow initialization method.

Other networks have net.layers{i}.initFcn set to 'initwb', which initializes each weight and bias with its own initialization function. The most common weight and bias initialization function is rands, which generates random values between -1 and 1.

See Also sim | adapt | train | initlay | initnw | initwb | rands | revert

#### initcon

Purpose	Conscience bias	initialization function	
Syntax	initcon (S,PR)		
Description	initcon is a bias initialization function that initializes biases for learning with the learncon learning function.		
	initcon (S,PR	) takes two arguments,	
	S	Number of rows (neurons)	
	PR	R-by-2 matrix of R = [Pmin Pmax] (default = [1 1])	
	and returns an S-by-1 bias vector.		
		ases, R is always 1. initcon could also be used to ts, but it is not recommended for that purpose.	
Examples	Here initial bias	s values are calculated for a five-neuron layer.	
	b = initcon(5	)	
Network Use		a standard network that uses initcon to initialize ng competlayer.	
	To prepare the bias of layer i of a custom network to initialize with initcon,		
1 Set net.initFcn to 'initlay'. (net.initParam automatic becomes initlay's default parameters.)			
	<b>2</b> Set net.laye	ers{i}.initFcn to 'initwb'.	
	<b>3</b> Set net.bias	ses{i}.initFcn to 'initcon'.	
To initialize the network, call init.		e network, call init.	

Algorithms	learncon updates biases so that each bias value $b(i)$ is a function of the average output $c(i)$ of the neuron $i$ associated with the bias.		
	initcon gets initial bias values by assuming that each neuron has responded to equal numbers of vectors in the past.		
See Also	competlayer   init   initlay   initwb   learncon		

# initlay

Purpose	Layer-by-layer network initialization function		
Syntax	net = initlay(net) info = initlay(' <i>code</i> ')		
Description	<pre>initlay is a network initialization function that initializes each layer i according to its own initialization function net.layers{i}.initFcn. net = initlay(net) takes</pre>		
	net	Neural network	
	and returns the network with each layer updated.		
	<pre>info = initlay('code') returns useful information for each supported code string:</pre>		
	'pnames'	Names of initialization parameters	
	'pdefaults'	Default initialization parameters	
	initlay does not hav	e any initialization parameters.	
Network Use	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
		Set net.initFcn to 'initlay'. This sets net.initParam to the empty matrix [], because initlay has no initialization parameters.	
	2 Set each net.layers{i}.initFcn to a layer initialization function. (Examples of such functions are initwb and initnw.)		
	To initialize the network, call init.		

# Algorithms The weights and biases of each layer i are initialized according to net.layers{i}.initFcn.

#### See Also cascadeforwardnet | feedforwardnet | init | initnw | initwb

### initlvq

Purpose	LVQ weight initialization function
Syntax	<pre>initlvq('configure',x) initlvq('configure',net,'IW',i,j,settings) initlvq('configure',net,'LW',i,j,settings) initlvq('configure',net,'b',i,)</pre>
Description	<pre>initlvq('configure',x) takes input data x and returns initialization settings for an LVQ weights associated with that input.</pre>
	<pre>initlvq('configure',net,'IW',i,j,settings) takes a network, and indices indicating an input weight to layer i from input j, and that weights settings, and returns new weight values.</pre>
	<pre>initlvq('configure',net,'LW',i,j,settings) takes a network, and indices indicating a layer weight to layer i from layer j, and that weights settings, and returns new weight values.</pre>
	initlvq('configure',net,'b',i,) takes a network, and an index indicating a bias for layer i, and returns new bias values.
See Also	lvqnet   init

Purpose	Nguyen-Widrow layer in	itialization function
Syntax	net = initnw(net,i)	
Description	initnw is a layer initialization function that initializes a layer's weights and biases according to the Nguyen-Widrow initialization algorithm. This algorithm chooses values in order to distribute the active region of each neuron in the layer approximately evenly across the layer's input space. The values contain a degree of randomness, so they are not the same each time this function is called.	
	initnw requires that the layer it initializes have a transfer function with a finite active input range. This includes transfer functions such as tansig and satlin, but not purelin, whose active input range is the infinite interval [-inf, inf]. Transfer functions, such as tansig, will return their active input range as follows:	
	activeInputRange = tansig('active') activeInputRange = -2 2	
	net = initnw(net,i) takes two arguments,	
	net ]	Neural network
	i ]	Index of a layer
	and returns the network with layer i's weights and biases upda	
	default random generate	ent to Nguyen-Widrow initialization. Unless the or is set to the same seed before each call to different weight and bias values each time.
Network Use	You can create a standard network that uses initnw by calling feedforwardnet or cascadeforwardnet. To prepare a custom network to be initialized with initnw,	
	- · propure a custom not	

#### initnw

	Set net.initFcn to 'initlay'. This sets net.initParam to the empty matrix [], because initlay has no initialization parameters.
	<b>2</b> Set net.layers{i}.initFcn to 'initnw'.
	To initialize the network, call init.
Algorithms	The Nguyen-Widrow method generates initial weight and bias values for a layer so that the active regions of the layer's neurons are distributed approximately evenly over the input space.
	Advantages over purely random weights and biases are
	• Few neurons are wasted (because all the neurons are in the input space).
	• Training works faster (because each area of the input space has neurons). The Nguyen-Widrow method can only be applied to layers
	• With a bias
	<ul> <li>With weights whose weightFcn is dotprod</li> </ul>
	<ul> <li>With netInputFcn set to netsum</li> </ul>
	<ul> <li>With transferFcn whose active region is finite</li> </ul>
	If these conditions are not met, then initnw uses rands to initialize the layer's weights and biases.

See Also cascadeforwardnet | feedforwardnet | init | initlay | initwb

Purpose	Initialize SOM weights	with principal components	
Syntax	<pre>weights = initsom(inputs,dimensions,positions) weights = initsom(inputs,dimensions,topologyFcn)</pre>		
Description	initsompc initializes the weights of an N-dimensional self-organizing map so that the initial weights are distributed across the space spanned by the most significant N principal components of the inputs. Distributing the weight significantly speeds up SOM learning, as the map starts out with a reasonable ordering of the input space.		
	<pre>weights = initsom(inputs,dimensions,positions) takes these arguments:</pre>		
	inputs	$R\text{-}\mathrm{by-}Q$ matrix of $Q$ R-element input vectors	
	dimensions	D-by-1 vector of positive integer SOM dimensions	
	positions	D-by-S matrix of $S$ $D-dimension$ neuron positions	
	and returns the following:		
	weights	S-by-R matrix of weights	
	alternative specifying	reights = initsom(inputs,dimensions,topologyFcn) is an lternative specifying the name of a layer topology function instead f positions. topologyFcn is called with dimensions to obtain ositions.	
Examples	<pre>inputs = rand(2,100)+[2;3]*ones(1,100); dimensions = [3 4]; positions = gridtop(dimensions); weights = initsompc(inputs,dimensions,positions);</pre>		
See Also	gridtop   hextop   r	andtop	

#### initwb

Purpose	By weight and bias layer initialization function	
Syntax	initwb(net,i)	
Description	initwb is a layer initialization function that initializes a layer's weights and biases according to their own initialization functions.	
	<pre>initwb(net,i) takes two arguments,</pre>	
	net Neural network	
	i Index of a layer	
	and returns the network with layer i's weights and biases updated.	
Network Use	You can create a standard network that uses initwb by calling perceptron or linearlayer.	
	To prepare a custom network to be initialized with initwb,	
	Set net.initFcn to 'initlay'. This sets net.initParam to the empty matrix [], because initlay has no initialization parameters	
	<b>2</b> Set net.layers{i}.initFcn to 'initwb'.	
	3 Set each net.inputWeights{i,j}.initFcn to a weight initialization function. Set each net.layerWeights{i,j}.initFcn to a weight initialization function. Set each net.biases{i}.initFcn to a bias initialization function. (Examples of such functions are rands and midpoint.)	
	To initialize the network, call init.	
Algorithms	Each weight (bias) in layer i is set to new values calculated according to its weight (bias) initialization function.	
See Also	init   initlay   initnw   linearlayer   perceptron	

Purpose	Zero weight and bias initialization function	
Syntax	W = initzero(S,PR) b = initzero(S,[1 1])	
Description	W = initzero(S,PR) takes two arguments,	
	S Number of rows (neurons)	
	PR R-by-2 matrix of input value ranges = [Pmin Pmax]	
	and returns an S-by-R weight matrix of zeros.	
	<pre>b = initzero(S,[1 1]) returns an S-by-1 bias vector of zeros.</pre>	
Examples	Here initial weights and biases are calculated for a layer with two inputs ranging over [0 1] and [-2 2] and four neurons.	
	W = initzero(5,[0 1; -2 2]) b = initzero(5,[1 1])	
Network Use	You can create a standard network that uses initzero to initialize its weights by calling newp or newlin.	
	To prepare the weights and the bias of layer i of a custom network to be initialized with midpoint,	
	Set net.initFcn to 'initlay'. (net.initParam automatically becomes initlay's default parameters.)	
	<b>2</b> Set net.layers{i}.initFcn to 'initwb'.	
	<b>3</b> Set each net.inputWeights{i,j}.initFcn to 'initzero'.	
	<b>4</b> Set each net.layerWeights{i,j}.initFcn to 'initzero'.	
	<b>5</b> Set each net.biases{i}.initFcn to 'initzero'.	

#### initzero

To initialize the network, call init.

See help newp and help newlin for initialization examples.

See Also initwb | initlay | init

## isconfigured

Purpose	Indicate if network inputs and outputs are configured		
Syntax	[flag,inputflags,outputflags] = isconfigured(net)		
Description	[flag,inputflags,outputflags] = isconfigured(net) takes a neural network and returns three values,		
	flag True if all network inputs and outputs are configured (have non-zero sizes)		
	inputflags Vector of true/false values for each configured/unconfigured input		
	outputflags Vector of true/false values for each configured/unconfigured output		
Examples	Here are the flags returned for a new network before and after being configured:		
	<pre>net = feedforwardnet; [flag,inputFlags,outputFlags] = isconfigured(net) [x,t] = simplefit_dataset; net = configure(net,x,t); [flag,inputFlags,outputFlags] = isconfigured(net)</pre>		
See Also	configure   unconfigure		

## layrecnet

Purpose	Layer recurrent neural network		
Syntax	layrecnet(layerDelays,hiddenSizes,trainFcn)		
Description	Layer recurrent neural networks are similar to feedforward networks, except that each layer has a recurrent connection with a tap delay associated with it. This allows the network to have an infinite dynamic response to time series input data. This network is similar to the time delay (timedelaynet) and distributed delay (distdelaynet) neural networks, which have finite input responses.		
	layrecnet(layerDelays,hiddenSizes,trainFcn) takes these arguments,		
	layerDelaysRow vector of increasing 0 or positive delays (default = 1:2)		
	hiddenSizesRow vector of one or more hidden layer sizes (default = 10)trainFcnTraining function (default = 'trainlm')		
	and returns a layer recurrent neural network.		
Examples	Use a layer recurrent neural network to solve a simple time series problem:		
	<pre>[X,T] = simpleseries_dataset; net = layrecnet(1:2,10) [Xs,Xi,Ai,Ts] = preparets(net,X,T) net = train(net,Xs,Ts,Xi,Ai); view(net) Y = net(Xs,Xi,Ai); perf = perform(net,Y,Ts)</pre>		
See Also	preparets   removedelay   distdelaynet   timedelaynet   narnet   narxnet		

Purpose	Conscience bias learning function		
Syntax	<pre>[dB,LS] = learncon(B,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learncon('code')</pre>		
Description	<b>learncon</b> is the conscience bias learning function used to increase the net input to neurons that have the lowest average output until each neuron responds approximately an equal percentage of the time.		
	<pre>[dB,LS] = learncon(B,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>		
	В	S-by-1 bias vector	
	P1-by-Q ones vectorZS-by-Q weighted input vectorsNS-by-Q net input vectorsAS-by-Q output vectors		
	T S-by-Q layer target vectors		
	ES-by-Q layer error vectorsgWS-by-R gradient with respect to performancegAS-by-Q output gradient with respect to performance		
	D	S-by-S neuron distances	
	LP	Learning parameters, none, LP = []	
	LS Learning state, initially should be = []		

dB	S-by-1 weight (or bias) change matrix
LS	New learning state

### learncon

	Learning occurs according to <b>learncon</b> 's learning parameter, shown here with its default value.	
	LP.lr - 0.001	Learning rate
	<pre>info = learncon('code') returns useful information for each supported code string:</pre>	
	'pnames' Names of learning parameters	
	'pdefaults'	Default learning parameters
	'needg'	Returns 1 if this function uses gW or gA $% \left( {{{\left[ {{{K_{{\rm{B}}}} \right]}} \right]_{\rm{A}}}} \right)$
Examples	<pre>equals 1 minus the bi Network Toolbox 2.0 Here you define a ran three neurons. You a a = rand(3,1); b = rand(3,1); lp.lr = 0.5;</pre>	ndom output A and bias vector W for a layer with lso define the learning rate LR.
	Because learncon only needs these values to calculate a bias change (see "Algorithm" below), use them to do so.	
	dW = learncon(b,[]	,[],[],a,[],[],[],[],1p,[])
Network Use	To prepare the bias o learncon,	f layer i of a custom network to learn with
		to 'trainr'. (net.trainParam automatically default parameters.)

	2 Set net.adaptFcn to 'trains'. (net.adaptParam automatically becomes trains's default parameters.)		
	<b>3</b> Set net.inputWeights{i}.learnFcn to 'learncon'		
	4 Set each net.layerWeights{i,j}.learnFcn to 'learncon'. .(Each weight learning parameter property is automatically set to learncon's default parameters.)		
	To train the network (or enable it to adapt),		
	<b>1</b> Set net.trainParam (or net.adaptParam) properties as desired.		
	<b>2</b> Call train (or adapt).		
Algorithms	<b>Learncon</b> calculates the bias change <b>db</b> for a given neuron by first updating each neuron's <i>conscience</i> , i.e., the running average of its output:		
	c = (1-lr)*c + lr*a		
	The conscience is then used to compute a bias for the neuron that is greatest for smaller conscience values.		
	b = exp(1-log(c)) - b		
	(learncon recovers C from the bias values each time it is called.)		
See Also	learnk   learnos   adapt   train		

## learngd

Purpose	Gradient descent weight and bias learning function		
Syntax	<pre>[dW,LS] = learngd(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learngd('code')</pre>		
Description	learngd is the gradient descent weight and bias learning function.		
	[dW,LS] = learngd(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs:		
	W	S-by-R weight matrix (or S-by-1 bias vector)	
	Р	R-by-Q input vectors (or ones(1,Q))	
	ZS-by-Q output gradient with respect to performance x Q weighted input vectorsNS-by-Q net input vectorsAS-by-Q output vectorsTS-by-Q layer target vectorsES-by-Q layer error vectorsgWS-by-R gradient with respect to performancegAS-by-Q output gradient with respect to performance		
	D	S-by-S neuron distances	
	LP Learning parameters, none, LP = []		
	LS	Learning state, initially should be []	

and returns

dW	$\ensuremath{S}\xspace$ -by-R weight (or bias) change matrix
LS	New learning state

	Learning occurs according to learngd's learning parameter, shown here with its default value.		
	LP.lr - 0.01	Learning rate	
	<pre>info = learngd('code') returns useful information for each supported code string:</pre>		
	'pnames' Names of learning parameters		
	'pdefaults'	Default learning parameters	
	'needg'	Returns 1 if this function uses $gW$ or $gA$	
Examples	<pre>Here you define a random gradient gW for a weight going to a laye with three neurons from an input with two elements. Also define learning rate of 0.5. gW = rand(3,2); lp.lr = 0.5;</pre>		
	Because learngd only needs these values to calculate a weight cha (see "Algorithm" below), use them to do so.		
	dW = learngd([]	,[],[],[],[],[],[],gW,[],[],lp,[])	
Network Use	You can create a standard network that uses learngd with newff, newcf, or newelm. To prepare the weights and the bias of layer i of a custom network to adapt with learngd,		
1 Set net.adaptFcn to 'trains'. net.adaptParan becomes trains's default parameters.			
	2 Set each net.inputWeights{i,j}.learnFcn to 'learngd'. S each net.layerWeights{i,j}.learnFcn to 'learngd'. Set net.biases{i}.learnFcn to 'learngd'. Each weight and bia		

#### learngd

learning parameter property is automatically set to learngd's default parameters.
 To allow the network to adapt,
 1 Set net.adaptParam properties to desired values.
 2 Call adapt with the network.
 See help newff or help newcf for examples.
 Algorithms
 learngd calculates the weight change dW for a given neuron from the neuron's input P and error E, and the weight (or bias) learning rate LR, according to the gradient descent dw = lr\*gW.

See Also adapt | learngdm | train

Purpose	Gradient descent with momentum weight and bias learning function	
Syntax	[dW,LS] = learngdm(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learngdm(' <i>code</i> ')	
Description	learngdm is the gradient descent with momentum weight and bias learning function.	
	<pre>[dW,LS] = learngdm(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>	
	W S-by-R weight matrix (or S-by-1 bias vector)	
	P R-by-Q input vectors (or ones(1,Q))	
	Z S-by-Q weighted input vectors	
	N S-by-Q net input vectors	
	A S-by-Q output vectors	
	TS-by-Q layer target vectorsES-by-Q layer error vectorsgWS-by-R gradient with respect to performance	
	gA	$\ensuremath{S}\xspace$ output gradient with respect to performance
	D S-by-S neuron distances	
	LP	Learning parameters, none, LP = []
	LS Learning state, initially should be = []	

dW	S-by-R weight (or bias) change matrix
LS	New learning state

Learning occurs according to  ${\tt learngdm}$ 's learning parameters, shown here with their default values.

LP.lr - 0.01	Learning rate
LP.mc - 0.9	Momentum constant

info = learngdm('code') returns useful information for each code
string:

'pnames'	Names of learning parameters
'pdefaults'	Default learning parameters
'needg'	Returns 1 if this function uses gW or gA

# **Examples** Here you define a random gradient G for a weight going to a layer with three neurons from an input with two elements. Also define a learning rate of 0.5 and momentum constant of 0.8:

gW = rand(3,2); lp.lr = 0.5; lp.mc = 0.8;

Because learngdm only needs these values to calculate a weight change (see "Algorithm" below), use them to do so. Use the default initial learning state.

```
ls = [];
[dW,ls] = learngdm([],[],[],[],[],[],gW,[],[],lp,ls)
```

learngdm returns the weight change and a new learning state.

Network Use	You can create a standard network that uses learngdm with newff, newcf, or newelm.
	To prepare the weights and the bias of layer i of a custom network to adapt with learngdm,
	1 Set net.adaptFcn to 'trains'. net.adaptParam automatically becomes trains's default parameters.
	2 Set each net.inputWeights{i,j}.learnFcn to 'learngdm'. Set each net.layerWeights{i,j}.learnFcn to 'learngdm'. Set net.biases{i}.learnFcn to 'learngdm'. Each weight and bias learning parameter property is automatically set to learngdm's default parameters.
	To allow the network to adapt,
	1 Set net.adaptParam properties to desired values.
	2 Call adapt with the network.
	See help newff or help newcf for examples.
Algorithms	learngdm calculates the weight change dW for a given neuron from the neuron's input P and error E, the weight (or bias) W, learning rate LR, and momentum constant MC, according to gradient descent with momentum:
	dW = mc*dWprev + (1-mc)*lr*gW
	The previous weight change dWprev is stored and read from the learning state LS.
See Also	adapt   learngd   train

#### learnh

Purpose	Hebb weight learning rule	
Syntax	[dW,LS] = learnh(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnh(' <i>code</i> ')	
Description	learnh is the Hebb weight learning function.	
	[dW,LS] = lear inputs,	nh(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	Z	S-by-Q weighted input vectors
	Ν	S-by-Q net input vectors
	А	S-by-Q output vectors
	Т	S-by-Q layer target vectors
	E	S-by-Q layer error vectors
	gW	S-by-R gradient with respect to performance
	gA	S-by-Q output gradient with respect to performance
	D	S-by-S neuron distances
	LP	Learning parameters, none, LP = []
	LS	Learning state, initially should be = []
	and returns	
	dW	S-by-R weight (or bias) change matrix

Learning occurs according to  ${\tt learnh}$  's learning parameter, shown here with its default value.

	LP.lr - 0.01 Le	arning rate
	<pre>info = learnh('code') returns useful information for each code string:</pre>	
	'pnames'	Names of learning parameters
	'pdefaults'	Default learning parameters
	'needg'	Returns 1 if this function uses $gW  \mathrm{or}  gA$
Examples	Here you define a random input P and output A for a layer with a two-element input and three neurons. Also define the learning rate LR.	
	p = rand(2,1); a = rand(3,1); lp.lr = 0.5;	
	Because learnh only (see "Algorithm" below	needs these values to calculate a weight change v), use them to do so.
	dW = learnh([],p,[]	],[],a,[],[],[],[],[],lp,[])
Network Use	To prepare the weights and the bias of layer i of a custom network to learn with learnh,	
	1 Set net.trainFcn t becomes trainr's d	to 'trainr'. (net.trainParam automatically efault parameters.)
	2 Set net.adaptFcn t becomes trains's d	to 'trains'. (net.adaptParam automatically efault parameters.)
	<b>3</b> Set each net.input	Weights{i,j}.learnFcn to 'learnh'.

## learnh

	<pre>4 Set each net.layerWeights{i,j}.learnFcn to 'learnh'. (Each weight learning parameter property is automatically set to learnh's default parameters.)</pre>
	To train the network (or enable it to adapt),
	1 Set net.trainParam (or net.adaptParam) properties to desired values.
	<b>2</b> Call train (adapt).
Algorithms	learnh calculates the weight change dW for a given neuron from the neuron's input P, output A, and learning rate LR according to the Hebb learning rule:
	dw = lr*a*p'
References	Hebb, D.O., The Organization of Behavior, New York, Wiley, 1949
See Also	learnhd   adapt   train

Purpose	Hebb with decay weight learning rule	
Syntax	[dW,LS] = learnhd(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnhd(' <i>code</i> ')	
Description	learnhd is the Hebb weight learning function.	
	[dW,LS] = learnhd(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,	
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	Z	S-by-Q weighted input vectors
	Ν	S-by-Q net input vectors
	А	S-by-Q output vectors
	т	S-by-Q layer target vectors
	E	S-by-Q layer error vectors
	gW	S-by-R gradient with respect to performance
	gA	$\ensuremath{S}\xspace$ output gradient with respect to performance
	D	S-by-S neuron distances
	LP	Learning parameters, none, LP = []
	LS	Learning state, initially should be = []

dW	S-by-R weight (or bias) change matrix
LS	New learning state

Learning occurs according to  ${\tt learnhd}{}^{\rm s}$  learning parameters, shown here with default values.

## learnhd

LP.dr -	0.01	Decay rate
LP.lr -	0.1	Learning rate

info = learnhd('code') returns useful information for each code
string:

'pnames'	Names of learning parameters
'pdefaults'	Default learning parameters
'needg'	Returns 1 if this function uses ${\tt gW}{\tt or}{\tt gA}$

Examples	Here you define a random input $P$ , output $A$ , and weights $W$ for a layer with a two-element input and three neurons. Also define the decay and learning rates.		
	<pre>p = rand(2,1); a = rand(3,1); w = rand(3,2); lp.dr = 0.05; lp.lr = 0.5;</pre>		
	Because learnhd only needs these values to calculate a weight change (see "Algorithm" below), use them to do so.		
	dW = learnhd(w,p,[],[],a,[],[],[],[],[],lp,[])		
Network Use	To prepare the weights and the bias of layer i of a custom network to learn with learnhd,		
	1 Set net.trainFcn to 'trainr'. (net.trainParam automatically becomes trainr's default parameters.)		
	2 Set net.adaptFcn to 'trains'. (net.adaptParam automatically becomes trains's default parameters.)		

	<b>3</b> Set each net.inputWeights{i,j}.learnFcn to 'learnhd'.
	<b>4</b> Set each net.layerWeights{i,j}.learnFcn to 'learnhd'. (Each weight learning parameter property is automatically set to learnhd's default parameters.)
	To train the network (or enable it to adapt),
	1 Set net.trainParam (or net.adaptParam) properties to desired values.
	<b>2</b> Call train (adapt).
Algorithms	learnhd calculates the weight change dW for a given neuron from the neuron's input P, output A, decay rate DR, and learning rate LR according to the Hebb with decay learning rule:
	dw = lr*a*p' - dr*w
See Also	learnh   adapt   train

#### learnis

Purpose	Instar weight learning function	
Syntax	<pre>[dW,LS] = learnis(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnis('code')</pre>	
Description	<pre>learnis is the instar weight learning function. [dW,LS] = learnis(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>	
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	Z	S-by-Q weighted input vectors
	NS-by-Q net input vectorsAS-by-Q output vectorsTS-by-Q layer target vectorsES-by-Q layer error vectorsgWS-by-R gradient with respect to performance	
	gA	S-by-Q output gradient with respect to performance
	DS-by-S neuron distancesLPLearning parameters, none, LP = []	
	LS	Learning state, initially should be = []
	and returns	
	dW	S-by-R weight (or bias) change matrix

LS	New learning state
----	--------------------

Learning occurs according to learnis's learning parameter, shown here with its default value.

	LP.lr - 0.01	Learning rate	
	info = learnis(' string:	code') returns useful information for each code	
	'pnames'	Names of learning parameters	
	'pdefaults'	Default learning parameters	
	'needg'	Returns 1 if this function uses $gW$ or $gA$	
Examples	Here you define a random input P, output A, and weight matrix W for a layer with a two-element input and three neurons. Also define the learning rate LR.		
	<pre>p = rand(2,1); a = rand(3,1); w = rand(3,2); lp.lr = 0.5;</pre>		
		nly needs these values to calculate a weight change low), use them to do so.	
	dW = learnis(w,p	,[],[],a,[],[],[],[],[],1p,[])	
Network Use	To prepare the weig so that it can learn	ghts and the bias of layer i of a custom network with learnis,	
		en to 'trainr'. (net.trainParam automatically s default parameters.)	
		en to 'trains'. (net.adaptParam automatically s default parameters.)	
	<b>3</b> Set each net.in	<pre>outWeights{i,j}.learnFcn to 'learnis'.</pre>	

### learnis

	<b>4</b> Set each net.layerWeights{i,j}.learnFcn to 'learnis'. (Each weight learning parameter property is automatically set to learnis's default parameters.)
	To train the network (or enable it to adapt),
	1 Set net.trainParam (net.adaptParam) properties to desired values.
	<b>2</b> Call train (adapt).
Algorithms	learnis calculates the weight change dW for a given neuron from the neuron's input P, output A, and learning rate LR according to the instar learning rule:
	dw = lr*a*(p'-w)
References	Grossberg, S., <i>Studies of the Mind and Brain</i> , Drodrecht, Holland, Reidel Press, 1982
See Also	learnk   learnos   adapt   train

Purpose	Kohonen weight learning function	
Syntax	[dW,LS] = learnk(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnk(' <i>code</i> ')	
Description	<pre>learnk is the Kohonen weight learning function. [dW,LS] = learnk(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>	
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	ZS-by-Q weighted input vectorsNS-by-Q net input vectorsAS-by-Q output vectors	
	т	S-by-Q layer target vectors
	E	S-by-Q layer error vectors
	gW	S-by-R gradient with respect to performance
	gA	$\ensuremath{S}\xspace$ output gradient with respect to performance
	D	S-by-S neuron distances
	LPLearning parameters, none, LP = []LSLearning state, initially should be = []	

dW	S-by-R weight (or bias) change matrix
LS	New learning state

Learning occurs according to  ${\tt learnk}$  's learning parameter, shown here with its default value.

## learnk

	LP.lr - 0.01	Learning rate	
	<pre>info = learnk('code string:</pre>	') returns useful information for each <i>code</i>	
	'pnames'	Names of learning parameters	
	'pdefaults'	Default learning parameters	
	'needg'	Returns 1 if this function uses $gW$ or $gA$	
Examples	Here you define a random input P, output A, and weight matrix W for a layer with a two-element input and three neurons. Also define the learning rate LR.		
	<pre>p = rand(2,1); a = rand(3,1); w = rand(3,2); lp.lr = 0.5;</pre>		
	Because learnk only n (see "Algorithm" below	eeds these values to calculate a weight change ), use them to do so.	
	dW = learnk(w,p,[],	[],a,[],[],[],[],[],lp,[])	
Network Use	To prepare the weights learnk,	of layer i of a custom network to learn with	
	Set net.trainFcn to 'trainr'. (net.trainParam automatically becomes trainr's default parameters.)		
	2 Set net.adaptFcn to becomes trains's de	o ' <b>trains</b> '. ( <b>net.adaptParam</b> automatically fault parameters.)	
	<b>3</b> Set each net.inputV	Veights{i,j}.learnFcn to 'learnk'.	

	4 Set each net.layerWeights{i,j}.learnFcn to 'learnk'. (Each weight learning parameter property is automatically set to learnk's default parameters.)		
	To train the network (or enable it to adapt),		
	1 Set net.trainParam (or net.adaptParam) properties as desired.		
	<b>2</b> Call train (or adapt).		
Algorithms	learnk calculates the weight change dW for a given neuron from the neuron's input P, output A, and learning rate LR according to the Kohonen learning rule:		
	dw = $lr^{(p'-w)}$ , if a ~= 0; = 0, otherwise		
References	Kohonen, T., Self-Organizing and Associative Memory, New York, Springer-Verlag, 1984		
See Also	learnis   learnos   adapt   train		

#### learnlv1

Purpose	LVQ1 weight learning function	
Syntax	[dW,LS] = learnlv1(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnlv1(' <i>code</i> ')	
Description	<pre>learnlv1 is the LVQ1 weight learning function. [dW,LS] = learnlv1(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>	
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	Z	S-by-Q weighted input vectors
	NS-by-Q net input vectorsAS-by-Q output vectorsTS-by-Q layer target vectorsES-by-Q layer error vectorsgWS-by-R gradient with respect to performancegAS-by-Q output gradient with respect to performanceDS-by-S neuron distancesLPLearning parameters, none, LP = []	
	LS	Learning state, initially should be = []
	and returns	
	dW	S-by-R weight (or bias) change matrix

LS	New learning state
L3	New learning state

Learning occurs according to learn1v1's learning parameter, shown here with its default value.

	LP.lr - 0.01	Learning rate
	info = learnlv1(' string:	code') returns useful information for each code
	'pnames'	Names of learning parameters
	'pdefaults'	Default learning parameters
	'needg'	Returns 1 if this function uses gW or $gA$
Examples	output gradient gA f	andom input P, output A, weight matrix W, and for a layer with a two-element input and three e the learning rate LR.
	<pre>p = rand(2,1); w = rand(3,2); a = compet(negdis gA = [-1;1; 1]; lp.lr = 0.5;</pre>	<pre>st(w,p));</pre>
		nly needs these values to calculate a weight change ow), use them to do so.
	dW = learnlv1(w,p	o,[],[],a,[],[],[],gA,[],lp,[])
Network Use		undard network that uses learnlv1 with lvqnet. hts of layer i of a custom network to learn with
		n to 'trainr'. (net.trainParam automatically default parameters.)
	-	n to 'trains'. (net.adaptParam automatically default parameters.)

## learnlv1

	<b>3</b> Set each net.inputWeights{i,j}.learnFcn to 'learnlv1'.		
	4 Set each net.layerWeights{i,j}.learnFcn to 'learnlv1'. (Each weight learning parameter property is automatically set to learnlv1's default parameters.)		
	To train the network (or enable it to adapt),		
	1 Set net.trainParam (or net.adaptParam) properties as desired.		
	<b>2</b> Call train (or adapt).		
Algorithms	<code>learnlv1</code> calculates the weight change dW for a given neuron from the neuron's input P, output A, output gradient gA, and learning rate LR, according to the LVQ1 rule, given i, the index of the neuron whose output $a(i)$ is 1:		
	dw(i,:) = +lr*(p-w(i,:)) if gA(i) = 0;= -lr*(p-w(i,:)) if gA(i) = -1		
See Also	learnlv2   adapt   train		

Purpose	LVQ2.1 weight learning function	
Syntax	[dW,LS] = learnlv2(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnlv2(' <i>code</i> ')	
Description	<pre>learnlv2 is the LVQ2 weight learning function. [dW,LS] = learnlv2(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>	
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	ZS-by-Q weighted input vectorsNS-by-Q net input vectors	
	А	S-by-Q output vectors
	т	S-by-Q layer target vectors
	E	S-by-Q layer error vectors
	gW	S-by-R weight gradient with respect to performance
	gA	S-by-Q output gradient with respect to performance
	D S-by-S neuron distances	
	LP	Learning parameters, none, LP = []
	LS	Learning state, initially should be = []

dW	S-by-R weight (or bias) change matrix
LS	New learning state

Learning occurs according to  $\verb"learnlv2"$ 's learning parameter, shown here with its default value.

	LP.lr - 0.01	Learning rate
	LP.window - 0.25	Window size (0 to 1, typically 0.2 to 0.3)
	<pre>info = learnlv2('co string:</pre>	ode') returns useful information for each code
	'pnames'	Names of learning parameters
	'pdefaults'	Default learning parameters
	'needg'	Returns 1 if this function uses $gW$ or $gA$
Examples	-	ple input P, output A, weight matrix W, and output r with a two-element input and three neurons. ng rate LR.
	<pre>p = rand(2,1); w = rand(3,2); n = negdist(w,p); a = compet(n); gA = [-1;1; 1]; lp.lr = 0.5;</pre>	
	Because learnlv2 onl (see "Algorithm" belov	y needs these values to calculate a weight change v), use them to do so.
	dW = learnlv2(w,p,	[],n,a,[],[],[],gA,[],lp,[])
Network	You can create a stand	dard network that uses learnlv2 with lvqnet.
Use	To prepare the weight learn1v2,	ts of layer i of a custom network to learn with
	1 Set net.trainFcn t becomes trainr's d	to 'trainr'. (net.trainParam automatically efault parameters.)

	<b>2</b> Set net.adaptFcn to 'trains'. (net.adaptParam automatically becomes trains's default parameters.)		
	<b>3</b> Set each net.inputWeights{i,j}.learnFcn to 'learnlv2'.		
	4 Set each net.layerWeights{i,j}.learnFcn to 'learnlv2'. (Each weight learning parameter property is automatically set to learnlv2's default parameters.)		
	To train the network (or enable it to adapt),		
	1 Set net.trainParam (or net.adaptParam) properties as desired.		
	<b>2</b> Call train (or adapt).		
Algorithms	learnlv2 implements Learning Vector Quantization 2.1, which works as follows:		
	For each presentation, if the winning neuron i should not have won, and the runnerup j should have, and the distance di between the winning neuron and the input p is roughly equal to the distance dj from the runnerup neuron to the input p according to the given window,		
	<pre>min(di/dj, dj/di) &gt; (1-window)/(1+window)</pre>		
	then move the winning neuron $i$ weights away from the input vector, and move the runnerup neuron $j$ weights toward the input according to		
	<pre>dw(i,:) = - lp.lr*(p'-w(i,:)) dw(j,:) = + lp.lr*(p'-w(j,:))</pre>		
See Also	learnlv1   adapt   train		

#### learnos

Purpose	Outstar weight learning function	
Syntax	[dW,LS] = learnos(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnos(' <i>code</i> ')	
Description	learnos is the c	outstar weight learning function.
	[dW,LS] = learnos(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,	
	W	S-by-R weight matrix (or S-by-1 bias vector)
	Р	R-by-Q input vectors (or ones(1,Q))
	Z	S-by-Q weighted input vectors
	Ν	S-by-Q net input vectors
	А	S-by-Q output vectors
	т	S-by-Q layer target vectors
	E	S-by-Q layer error vectors
	gW	S-by-R weight gradient with respect to performance
	gA	S-by-Q output gradient with respect to performance
	D	S-by-S neuron distances
	LP	Learning parameters, none, LP = []
	LS	Learning state, initially should be = []
	and returns	
	dW	S-by-R weight (or bias) change matrix

LS New learning state

Learning occurs according to learnos's learning parameter, shown here with its default value.

	LP.lr - 0.01	Learning rate
	info = learnos('c string:	code') returns useful information for each code
	'pnames'	Names of learning parameters
	'pdefaults'	Default learning parameters
	'needg'	Returns 1 if this function uses $gW$ or $gA$
Examples	-	andom input P, output A, and weight matrix W for lement input and three neurons. Also define the
	p = rand(2,1); a = rand(3,1); w = rand(3,2); lp.lr = 0.5;	
		ly needs these values to calculate a weight change low), use them to do so.
	dW = learnos(w,p,	[],[],a,[],[],[],[],1p,[])
Network Use	To prepare the weig learn with learnos,	ths and the bias of layer i of a custom network to
		n to 'trainr'. (net.trainParam automatically s default parameters.)
		n to 'trains'. (net.adaptParam automatically s default parameters.)
	<b>3</b> Set each net.inp	outWeights{i,j}.learnFcn to 'learnos'.

#### learnos

	<b>4</b> Set each net.layerWeights{i,j}.learnFcn to 'learnos'. (Each weight learning parameter property is automatically set to learnos's default parameters.)		
	To train the network (or enable it to adapt),		
	1 Set net.trainParam (or net.adaptParam) properties to desired values.		
	<b>2</b> Call train (adapt).		
Algorithms	learnos calculates the weight change dW for a given neuron from the neuron's input P, output A, and learning rate LR according to the outstar learning rule:		
	dw = lr*(a-w)*p'		
References	Grossberg, S., <i>Studies of the Mind and Brain</i> , Drodrecht, Holland, Reidel Press, 1982		
See Also	learnis   learnk   adapt   train		

Purpose	Perceptron weight and bias learning function		
	reception weight and blas learning function		
Syntax	[dW,LS] = lea info = learnp	rnp(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) ('code')	
Description	learnp is the perceptron weight/bias learning function.		
	[dW,LS] = learnp(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,		
	W	S-by-R weight matrix (or b, and S-by-1 bias vector)	
	Р	R-by-Q input vectors (or ones(1,Q))	
	Z	S-by-Q weighted input vectors	
	Ν	S-by-Q net input vectors	
	А	S-by-Q output vectors	
	Т	S-by-Q layer target vectors	
	E	S-by-Q layer error vectors	
	gW	S-by-R weight gradient with respect to performance	
	gA	S-by-Q output gradient with respect to performance	
	D	S-by-S neuron distances	
	LP	Learning parameters, none, LP = []	
	LS	Learning state, initially should be = []	
	and returns		
	dW	S-by-R weight (or bias) change matrix	
	LS	New learning state	
	info = learno	('code') returns useful information for each code	

info = learnp('code') returns useful information for each code
string:

	'pnames'	Names of learning parameters	
	'pdefaults'	Default learning parameters	
	'needg'	Returns 1 if this function uses $gW  \mathrm{or}  gA$	
Examples	Here you define a ran two-element input an	dom input P and error E for a layer with a d three neurons.	
	p = rand(2,1); e = rand(3,1);		
	Because learnp only : (see "Algorithm" below	needs these values to calculate a weight change v), use them to do so.	
	dW = learnp([],p,[	],[],[],[],e,[],[],[],[])	
Network	You can create a stan	dard network that uses learnp with newp.	
Use	To prepare the weights and the bias of layer i of a custom network to learn with learnp,		
	1 Set net.trainFcn to 'trainb'. (net.trainParam automatically becomes trainb's default parameters.)		
	2 Set net.adaptFcn to 'trains'. (net.adaptParam automatically becomes trains's default parameters.)		
	<b>3</b> Set each net.input	Weights{i,j}.learnFcn to 'learnp'.	
	<b>4</b> Set each net.layerWeights{i,j}.learnFcn to 'learnp'.		
	5 Set net.biases{i}.learnFcn to 'learnp'. (Each weight and bias learning parameter property automatically becomes the empty matrix, because learnp has no learning parameters.)		
	To train the network	(or enable it to adapt),	

	1 Set net.trainParam (or net.adaptParam) properties to desired values.
	<b>2</b> Call train (adapt).
	See help newp for adaption and training examples.
Algorithms	learnp calculates the weight change dW for a given neuron from the neuron's input P and error E according to the perceptron learning rule:
	dw = 0, if e = 0 = p', if e = 1 = -p', if e = -1
	This can be summarized as
	dw = e*p'
References	Rosenblatt, F., <i>Principles of Neurodynamics</i> , Washington, D.C., Spartan Press, 1961
See Also	adapt   learnpn   train

## learnpn

Purpose	Normalized perceptron weight and bias learning function		
Syntax	[dW,LS] = learnpn(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnpn(' <i>code</i> ')		
Description	<pre>learnpn is a weight and bias learning function. It can result in faster learning than learnp when input vectors have widely varying magnitudes. [dW,LS] = learnpn(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>		
	W	W S-by-R weight matrix (or S-by-1 bias vector)	
	Р	R-by-Q input vectors (or ones(1,Q))	
	ZS-by-Q weighted input vectorsNS-by-Q net input vectors		
	А	S-by-Q output vectors	
	T S-by-Q layer target vectors		
	E	S-by-Q layer error vectors	
	gW	$\ensuremath{S}\xspace$ by $\ensuremath{R}\xspace$ weight gradient with respect to performance	
	gAS-by-Q output gradient with respect to performanceDS-by-S neuron distances		
	LP	Learning parameters, none, LP = []	
	LS Learning state, initially should be = []		

and returns

dW	S-by-R weight (or bias) change matrix
LS	New learning state

	<pre>info = learnpn('code') returns useful information for each code string:</pre>			
	'pnames'	Names of learning parameters		
	'pdefaults'	Default learning parameters		
	'needg'	Returns 1 if this function uses $gW$ or $gA$		
Examples	Here you define a random input P and error E for a layer with a two-element input and three neurons.			
	<pre>p = rand(2,1); e = rand(3,1);</pre>			
	Because <b>learnpn</b> only needs these values to calculate a weight c (see "Algorithm" below), use them to do so.			
	dW = learnpn([],p,	[],[],[],[],e,[],[],[],[],[])		
Network	You can create a standard network that uses learnpn with newp.			
Use	To prepare the weights and the bias of layer i of a custom network to learn with learnpn,			
	Set net.trainFcn to 'trainb'. (net.trainParam automatically becomes trainb's default parameters.)			
	<b>2</b> Set net.adaptFcn to 'trains'. (net.adaptParam automatically becomes trains's default parameters.)			
	<b>3</b> Set each net.input	Weights{i,j}.learnFcn to 'learnpn'.		
	<b>4</b> Set each net.layer	Weights{i,j}.learnFcn to 'learnpn'.		
	bias learning paran	.learnFcn to 'learnpn'. (Each weight and neter property automatically becomes the empty arnpn has no learning parameters.)		

## learnpn

	To train the network (or enable it to adapt),
	1 Set net.trainParam (or net.adaptParam) properties to desired values.
	<b>2</b> Call train (adapt).
	See help newp for adaption and training examples.
Algorithms	learnpn calculates the weight change dW for a given neuron from the neuron's input P and error E according to the normalized perceptron learning rule:
	<pre>pn = p / sqrt(1 + p(1)^2 + p(2)^2) + + p(R)^2) dw = 0, if e = 0</pre>
	The expression for dW can be summarized as
	dw = e*pn'
Limitations	Perceptrons do have one real limitation. The set of input vectors must be linearly separable if a solution is to be found. That is, if the input vectors with targets of 1 cannot be separated by a line or hyperplane from the input vectors associated with values of 0, the perceptron will never be able to classify them correctly.
See Also	adapt   learnp   train

#### learnsom

Purpose	Self-organizing map weight learning function		
Syntax	<pre>[dW,LS] = learnsom(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnsom('code')</pre>		
Description	learnsom is the self-organizing map weight learning function.		
	<pre>[dW,LS] = learnsom(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>		
	W	W S-by-R weight matrix (or S-by-1 bias vector)	
	P R-by-Q input vectors (or ones(1,Q))		
	Z S-by-Q weighted input vectors		
	N S-by-Q net input vectors		
	A S-by-Q output vectors		
	T S-by-Q layer target vectors		
	E S-by-Q layer error vectors		
	gW S-by-R weight gradient with respect to performance		
	gA S-by-Q output gradient with respect to performance		
	D	S-by-S neuron distances	
	LP Learning parameters, none, LP = []		
	LS	Learning state, initially should be = []	

and returns

dW	S-by-R weight (or bias) change matrix
LS	New learning state

Learning occurs according to  $\verb"learnsom"$ 's learning parameters, shown here with their default values.

LP.order_lr	0.9	Ordering phase learning rate
LP.order_steps	1000	Ordering phase steps
LP.tune_lr	0.02	Tuning phase learning rate
LP.tune_nd	1	Tuning phase neighborhood distance

info = learnsom('code') returns useful information for each code
string:

'pnames'	Names of learning parameters
'pdefaults'	Default learning parameters
'needg'	Returns 1 if this function uses gW or gA

#### **Examples**

Here you define a random input P, output A, and weight matrix W for a layer with a two-element input and six neurons. You also calculate positions and distances for the neurons, which are arranged in a 2-by-3 hexagonal pattern. Then you define the four learning parameters.

```
p = rand(2,1);
a = rand(6,1);
w = rand(6,2);
pos = hextop(2,3);
d = linkdist(pos);
lp.order_lr = 0.9;
lp.order_steps = 1000;
lp.tune_lr = 0.02;
lp.tune_nd = 1;
```

Because learnsom only needs these values to calculate a weight change (see "Algorithm" below), use them to do so.

ls = [];

	[dW,ls] = learnsom(w,p,[],[],a,[],[],[],[],d,lp,ls)			
Network	You can create a standard network that uses learnsom with newsom.			
Use	1 Set net.trainFcn to 'trainr'. (net.trainParam automatically becomes trainr's default parameters.)			
	<b>2</b> Set net.adaptFcn to 'trains'. (net.adaptParam automatically becomes trains's default parameters.)			
	<b>3</b> Set each net.inputWeights{i,j}.learnFcn to 'learnsom'.			
	<b>4</b> Set each net.layerWeights{i,j}.learnFcn to 'learnsom'.			
	<b>5</b> Set net.biases{i}.learnFcn to 'learnsom'. (Each weight learning parameter property is automatically set to learnsom's default parameters.)			
	To train the network (or enable it to adapt):			
	Set net.trainParam (or net.adaptParam) properties to desired values.			
	<b>2</b> Call train (adapt).			
Algorithms	learnsom calculates the weight change dW for a given neuron from the neuron's input P, activation A2, and learning rate LR:			
	dw = lr*a2*(p'-w)			
	where the activation $A2$ is found from the layer output A, neuron distances D, and the current neighborhood size ND:			
	a2(i,q) = 1, if a(i,q) = 1 = 0.5, if a(j,q) = 1 and D(i,j) <= nd = 0, otherwise			

The learning rate LR and neighborhood size NS are altered through two phases: an ordering phase and a tuning phase.

The ordering phases lasts as many steps as LP.order\_steps. During this phase LR is adjusted from LP.order\_lr down to LP.tune\_lr, and ND is adjusted from the maximum neuron distance down to 1. It is during this phase that neuron weights are expected to order themselves in the input space consistent with the associated neuron positions.

During the tuning phase LR decreases slowly from LP.tune\_lr, and ND is always set to LP.tune\_nd. During this phase the weights are expected to spread out relatively evenly over the input space while retaining their topological order, determined during the ordering phase.

See Also adapt | train

### learnsomb

Purpose	Batch self-organizing map weight learning function		
Syntax		[dW,LS] = learnsomb(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnsomb(' <i>code</i> ')	
Description	learnsomb is the batch self-organizing map weight learning function.		
	[dW,LS] = inputs:	[dW,LS] = learnsomb(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs:	
	W	S-by-R weight matrix (or S-by-1 bias vector)	
	Р	R-by-Q input vectors (or ones(1,Q))	
	Z	S-by-Q weighted input vectors	
	Ν	S-by-Q net input vectors	
	А	S-by-Q output vectors	
	T S-by-Q layer target vectors		
	E S-by-Q layer error vectors		
	gW	gW S-by-R gradient with respect to performance	
	gA	gA S-by-Q output gradient with respect to performance	
	D	DS-by-S neuron distancesLPLearning parameters, none, LP = []	
	LP		
	LS	Learning state, initially should be = []	
	and returns the following:		
	dW	dW S-by-B weight (or bias) change matrix	

dW	S-by-R weight (or bias) change matrix
LS	New learning state

Learning occurs according to learnsomb's learning parameter, shown here with its default value:

LP.init_neighborhood	3	Initial neighborhood size
LP.steps	100	Ordering phase steps

info = learnsomb('code') returns useful information for each code
string:

'pnames'	Returns names of learning parameters.
'pdefaults'	Returns default learning parameters.
'needg'	Returns 1 if this function uses $gW\ {\rm or}\ gA.$

**Examples** This example defines a random input P, output A, and weight matrix W for a layer with a 2-element input and 6 neurons. This example also calculates the positions and distances for the neurons, which appear in a 2-by-3 hexagonal pattern.

```
p = rand(2,1);
a = rand(6,1);
w = rand(6,2);
pos = hextop(2,3);
d = linkdist(pos);
lp = learnsomb('pdefaults');
```

Because learnsom only needs these values to calculate a weight change (see Algorithm).

```
ls = [];
[dW,ls] = learnsomb(w,p,[],[],a,[],[],[],[],d,lp,ls)
```

NetworkYou can create a standard network that uses learnsomb withUseselforgmap. To prepare the weights of layer i of a custom network<br/>to learn with learnsomb:

	Set NET.trainFcn to 'trainr'. (NET.trainParam automatically becomes trainr's default parameters.)
	<b>2</b> Set NET.adaptFcn to 'trains'. (NET.adaptParam automatically becomes trains's default parameters.)
	<b>3</b> Set each NET.inputWeights{i,j}.learnFcn to 'learnsomb'.
	4 Set each NET.layerWeights{i,j}.learnFcn to 'learnsomb'. (Each weight learning parameter property is automatically set to learnsomb's default parameters.)
	To train the network (or enable it to adapt):
	1 Set NET.trainParam (or NET.adaptParam) properties as desired.
	<b>2</b> Call train (or adapt).
Algorithms learnsomb calculates the weight changes so that each new weight vector is the weighted average of the input vector neuron and neurons in its neighborhood responded to with of 1.	
	The ordering phase lasts as many steps as LP.steps.
	During this phase, the neighborhood is gradually reduced from a maximum size of LP.init_neighborhood down to 1, where it remains from then on.
See Also	adapt   selforgmap   train

### learnwh

Purpose	Widrow-Hoff weight/bias learning function		
Syntax	[dW,LS] = learnwh(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) info = learnwh(' <i>code</i> ')		
Description	learnwh is the Widrow-Hoff weight/bias learning function, and is also known as the delta or least mean squared (LMS) rule.		
	<pre>[dW,LS] = learnwh(W,P,Z,N,A,T,E,gW,gA,D,LP,LS) takes several inputs,</pre>		
	W S-by-R weight matrix (or b, and S-by-1 bias vector)		
	P R-by-Q input vectors (or ones(1,Q))		
	Z	S-by-Q weighted input vectors	
	Ν	S-by-Q net input vectors	
	А	S-by-Q output vectors	
	Т	S-by-Q layer target vectors	
	E	S-by-Q layer error vectors	
	gW	S-by-R weight gradient with respect to performance	
	gA	S-by-Q output gradient with respect to performance	
	D	S-by-S neuron distances	
	LP	Learning parameters, none, LP = []	
	LS	Learning state, initially should be = []	

and returns

dW	S-by-R weight (or bias) change matrix
LS	New learning state

	Learning occurs according to <b>learn</b> wh's learning parameter, shown here with its default value.	
	LP.lr Le 0.01	earning rate
	<pre>info = learnwh('code') returns useful information for each code string:</pre>	
	'pnames'	Names of learning parameters
	'pdefaults'	Default learning parameters
	'needg'	Returns 1 if this function uses $gW$ or $gA$
Examples	Here you define a random input P and error E for a layer with a two-element input and three neurons. You also define the learning rate LR learning parameter.	
	p = rand(2,1) e = rand(3,1) lp.lr = 0.5;	
		/h only needs these values to calculate a weight change " below), use them to do so.
	dW = learnwh(	[],p,[],[],[],[],e,[],[],[],lp,[])
Network Use	You can create a standard network that uses learnwh with linearlayer.	
	To prepare the learn with lear	weights and the bias of layer i of a custom network to nwh,
		InFcn to 'trainb'. net.trainParam automatically inb's default parameters.

### learnwh

	<b>2</b> Set net.adaptFcn to 'trains'. net.adaptParam automatically becomes trains's default parameters.
	<b>3</b> Set each net.inputWeights{i,j}.learnFcn to 'learnwh'.
	<b>4</b> Set each net.layerWeights{i,j}.learnFcn to 'learnwh'.
	<b>5</b> Set net.biases{i}.learnFcn to 'learnwh'. Each weight and bias learning parameter property is automatically set to learnwh's default parameters.
	To train the network (or enable it to adapt),
	1 Set net.trainParam (net.adaptParam) properties to desired values.
	<b>2</b> Call train (adapt).
Algorithms	learnwh calculates the weight change dW for a given neuron from the neuron's input P and error E, and the weight (or bias) learning rate LR, according to the Widrow-Hoff learning rule:
	dw = lr*e*pn'
References	Widrow, B., and M.E. Hoff, "Adaptive switching circuits," <i>1960 IRE WESCON Convention Record</i> , New York IRE, pp. 96–104, 1960
	Widrow, B., and S.D. Sterns, <i>Adaptive Signal Processing</i> , New York, Prentice-Hall, 1985
See Also	adapt   linearlayer   train

## linearlayer

Purpose	Linear layer			
Syntax	linearlayer(inpu	tDelays,widrowHoffLR)		
Description	Linear layers are single layers of linear neurons. They may be static, with input delays of 0, or dynamic, with input delays greater than 0. They can be trained on simple linear time series problems, but often are used adaptively to continue learning while deployed so they can adjust to changes in the relationship between inputs and outputs while being used.			
	If a network is needed to solve a nonlinear time series relationship, then better networks to try include timedelaynet, narxnet, and narnet.			
	linearlayer(inpu	tDelays,widrowHoffLR) takes these arguments,		
	<pre>inputDelays Row vector of increasing 0 or positive delays     (default = 1:2)</pre>			
	widrowHoffLR	Widrow-Hoff learning rate (default = $0.01$ )		
	and returns a linear layer. If the learning rate is too small, learning will happen very slowly. However, a greater danger is that it may be too large and learning will become unstable resulting in large changes to weight vectors and errors increasing instead of decreasing. If a data set is available which characterizes the relationship the layer is to learn, the maximum stable learning rate can be calculated with maxlinlr.			
Examples	Here a linear layer	is trained on a simple time series problem.		
	<pre>x = {0 -1 1 1 0 -1 1 0 0 1}; t = {0 -1 0 2 1 -1 0 1 0 1} net = linearlayer(1:2,0.01) [Xs,Xi,Ai,Ts] = preparets(net,X,T) net = train(net,Xs,Ts,Xi,Ai); view(net)</pre>			

Y = net(Xs,Xi); perf = perform(net,Ts,Y)

#### See Also preparets | removedelay | timedelaynet | narnet | narxnet

## linkdist

Purpose	Link distance function		
Syntax	d = linkdist(pos)		
Description	linkdist is a layer distance function used to find the distances between the layer's neurons given their positions.		
	<pre>d = linkdist(pos) takes one argument,</pre>		
	pos N-by-S matrix of neuron positions		
	and returns the S-by-S matrix of distances.		
Examples	Here you define a random matrix of positions for 10 neurons arranged in three-dimensional space and find their distances.		
	<pre>pos = rand(3,10); D = linkdist(pos)</pre>		
Network Use	You can create a standard network that uses linkdist as a distance function by calling selforgmap.		
	To change a network so that a layer's topology uses linkdist, set net.layers{i}.distanceFcn to 'linkdist'.		
	In either case, call sim to simulate the network with dist.		
Algorithms	The link distance $D$ between two position vectors $\ensuremath{\texttt{Pi}}$ and $\ensuremath{\texttt{Pj}}$ from a set of $S$ vectors is		
	<pre>Dij = 0, if i == j = 1, if (sum((Pi-Pj).^2)).^0.5 is &lt;= 1 = 2, if k exists, Dik = Dkj = 1 = 3, if k1, k2 exist, Dik1 = Dk1k2 = Dk2j = 1 = N, if k1kN exist, Dik1 = Dk1k2 == DkNj = 1 = S, if none of the above conditions apply</pre>		

## linkdist

See Also dist | mandist | selforgmap | sim

Purpose L	og-sigmoid transfer function
-----------	------------------------------

Graph and Symbol	a 0 a = logs Log-Sigmoid	•
Syntax		g(N,FP) ogsig('dn',N,A,FP) gsig(' <i>code</i> ')
Description	<pre>logsig is a transfer function. Transfer functions calculate a layer's output from its net input. A = logsig(N,FP) takes N and optional function parameters,</pre>	
	Ν	S-by-Q matrix of net input (column) vectors
	FP	Struct of function parameters (ignored)
	and returns A, the S-by-Q matrix of N's elements squashed into $[0, 1]$ .	
	$dA_dN = logsig('dn', N, A, FP)$ returns the S-by-Q derivative of A with respect to N. If A or FP is not supplied or is set to [], FP reverts to the default parameters, and A is calculated from N.	
	info = log string:	gsig('code') returns useful information for each code
	logsig('na	ame') returns the name of this function.

logsig('output',FP) returns the [min max] output range.

logsig('active',FP) returns the [min max] active input range.

	logsig('fullderiv') returns 1 or 0, depending on whether dA_dN is S-by-S-by-Q or S-by-Q.		
	logsig('fpnames') returns the names of the function parameters.		
	logsig('fpdefaults') returns the default function parameters.		
Examples	Here is the code to create a plot of the logsig transfer function.		
	n = -5:0.1:5; a = logsig(n); plot(n,a)		
	Assign this transfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'logsig';</pre>		
Algorithms	logsig(n) = 1 / (1 + exp(-n))		
See Also	sim   tansig		

Purpose	Learning vector quantization neural network		
Syntax	lvqnet(hiddenSize,lvqLR,lvqLF)		
Description	LVQ (learning vector quantization) neural networks consist of two layers. The first layer maps input vectors into clusters that are found by the network during training. The second layer maps merges groups of first layer clusters into the classes defined by the target data.		
	The total number of first layer clusters is determined by the number of hidden neurons. The larger the hidden layer the more clusters the first layer can learn, and the more complex mapping of input to target classes can be made. The relative number of first layer clusters assigned to each target class are determined according to the distribution of target classes at the time of network initialization. This occurs when the network is automatically configured the first time train is called, or manually configured with the function configure, or manually initialized with the function init is called.		
	<pre>lvqnet(hiddenSize,lvqLR,lvqLF) takes these arguments,</pre>		
	hiddenSize Size of hidden layer (default = 10)		
	lvqLR LVQ learning rate (default = 0.01)		
	lvqLF	LVQ learning function (default = 'learnlv1')	
	and returns an LVQ neural network. The other option for the lvq learning function is learnlv2.		
Examples	Here, an LVQ network is trained to classify iris flowers.		
	<pre>[x,t] = iris_dataset; net = lvqnet(10) net = train(net,x,t); view(net) y = net(x);</pre>		

#### lvqnet

```
perf = perform(net,y,t)
classes = vec2ind(y)
See Also
preparets | removedelay | timedelaynet | narnet | narxnet
```

Purpose	LVQ outputs processing function
Syntax	<pre>[X,settings] = lvqoutputs(X) X = lvqoutputs('apply',X,PS) X = lvqoutputs('reverse',X,PS) dx_dy = lvqoutputs('dx_dy',X,X,PS)</pre>
Description	[X,settings] = lvqoutputs(X) returns its argument unchanged, but stores the ratio of target classes in the settings for use by initlvq to initialize weights.
	<pre>X = lvqoutputs('apply',X,PS) returns X.</pre>
	<pre>X = lvqoutputs('reverse',X,PS) returns X.</pre>
	<pre>dx_dy = lvqoutputs('dx_dy',X,X,PS) returns the identity derivative.</pre>
See Also	lvqnet   initlvq

#### mae

Purpose	Mean absolute error performance function		
Syntax	<pre>perf = mae(E,Y,X,FP)</pre>		
Description	mae is a network performance function. It measures network performance as the mean of absolute errors.		
	perf = mae	e(E,Y,X,FP) takes E and optional function parameters,	
	E	Matrix or cell array of error vectors	
	Y	Matrix or cell array of output vectors (ignored)	
	Х	Vector of all weight and bias values (ignored)	
	FP	Function parameters (ignored)	
	and returns the mean absolute error. dPerf_dx = mae('dx',E,Y,X,perf,FP) returns the derivative of perf with respect to X.		
	<pre>info = mae('code') returns useful information for each code string:</pre>		
	mae('name') returns the name of this function.		
	mae('pnames') returns the names of the training parameters.		
	mae('pdefaults') returns the default function parameters.		
Examples	Create and configure a perceptron to have one input and one neuron:		
	<pre>net = perceptron; net = configure(net,0,0);</pre>		
	The network is given a batch of inputs P. The error is calculated by subtracting the output A from target T. Then the mean absolute error is calculated.		
		p = [-10 -5 0 5 10]; t = [0 0 1 1 1];	

	y = net(p) e = t-y perf = mae(e)
	Note that mae can be called with only one argument because the other arguments are ignored. mae supports those arguments to conform to the standard performance function argument list.
Network Use	You can create a standard network that uses mae with perceptron. To prepare a custom network to be trained with mae, set net.performFcn to 'mae'. This automatically sets net.performParam to the empty matrix [], because mae has no performance parameters.
	In either case, calling train or adapt, results in mae being used to calculate performance.
See Also	mse   perceptron

## mandist

Purpose	Manhattan	Manhattan distance weight function	
Syntax	Z = mandist(W,P) D = mandist(pos)		
Description	mandist is the Manhattan distance weight function. Weight functions apply weights to an input to get weighted inputs.		
	Z = mandis	t(W,P) takes these inputs,	
	W	S-by-R weight matrix	
	Р	R-by-Q matrix of Q input (column) vectors	
	and returns	the S-by-Q matrix of vector distances.	
mandist is also a layer distance fun distances between neurons in a lay		also a layer distance function, which can be used to find the etween neurons in a layer.	
	<pre>D = mandist(pos) takes one argument,</pre>		
	pos	S row matrix of neuron positions	
	and returns the S-by-S matrix of distances.		
<b>Examples</b> Here you define a random weight matrix W and input ve calculate the corresponding weighted input Z.			
	W = rand(4,3); P = rand(3,1); Z = mandist(W,P)		
	Here you define a random matrix of positions for 10 neurons arranged in three-dimensional space and then find their distances.		
	pos = rand(3,10); D = mandist(pos)		

Network Use	To change a network so an input weight uses mandist, set net.inputWeight{i,j}.weightFcn to 'mandist'. For a layer weight, set net.layerWeight{i,j}.weightFcn to 'mandist'.		
	To change a network so a layer's topology uses mandist, set net.layers{i}.distanceFcn to 'mandist'.		
	In either case, call sim to simulate the network with dist. See newpnn or newgrnn for simulation examples.		
Algorithms	The Manhattan distance D between two vectors X and Y is		
	D = sum(abs(x-y))		
See Also	dist   linkdist   sim		

# mapminmax

Process matrices by mapping row minimum and maximum values to [-1 1]		
<pre>[Y,PS] = mapminmax(X,YMIN,YMAX) [Y,PS] = mapminmax(X,FP) Y = mapminmax('apply',X,PS) X = mapminmax('reverse',Y,PS) dx_dy = mapminmax('dx_dy',X,Y,PS)</pre>		
mapminmax processes matrices by normalizing the minimum and maximum values of each row to [YMIN, YMAX].		
[Y,PS] = ma	apminmax(X,YMIN,YMAX) takes X and optional parameters	
Х	N-by-Q matrix or a 1-by-TS row cell array of N-by-Q matrices	
YMIN	Minimum value for each row of Y (default is $-1$ )	
YMAX	Maximum value for each row of Y (default is +1)	
and returns		
Y	Each $M$ -by- $Q$ matrix (where $M == N$ ) (optional)	
PS	Process settings that allow consistent processing of values	
[Y,PS] = mapminmax(X,FP) takes parameters as a struct: FP.ymin, FP.ymax.		
Y = mapmin	max('apply',X,PS) returns Y, given X and settings PS.	
<pre>X = mapminmax('reverse',Y,PS) returns X, given Y and settings PS. dx_dy = mapminmax('dx_dy',X,Y,PS) returns the reverse derivative.</pre>		

Examples	Here is how to format a matrix so that the minimum and maximum values of each row are mapped to default interval [-1,+1].
	x1 = [1 2 4; 1 1 1; 3 2 2; 0 0 0] [y1,PS] = mapminmax(x1)
	Next, apply the same processing settings to new values.
	x2 = [5 2 3; 1 1 1; 6 7 3; 0 0 0] y2 = mapminmax('apply',x2,PS)
	Reverse the processing of y1 to get x1 again.
	x1_again = mapminmax('reverse',y1,PS)
Algorithms	It is assumed that X has only finite real values, and that the elements of each row are not all equal. (If xmax=xmin or if either xmax or xmin are non-finite, then y=x and no change occurs.)
	y = (ymax-ymin)*(x-xmin)/(xmax-xmin) + ymin;
Definitions	Before training, it is often useful to scale the inputs and targets so that they always fall within a specified range. The function mapminmax scales inputs and targets so that they fall in the range $[-1,1]$ . The following code illustrates how to use this function.
	<pre>[pn,ps] = mapminmax(p); [tn,ts] = mapminmax(t); net = train(net,pn,tn);</pre>
	The original network inputs and targets are given in the matrices $p$ and $t$ . The normalized inputs and targets $pn$ and $tn$ that are returned will all fall in the interval $[-1,1]$ . The structures $ps$ and $ts$ contain the settings, in this case the minimum and maximum values of the original inputs and targets. After the network has been trained, the $ps$ settings should be used to transform any future inputs that are applied to the network. They effectively become a part of the network, just like the network weights and biases.

If mapminmax is used to scale the targets, then the output of the network will be trained to produce outputs in the range [-1,1]. To convert these outputs back into the same units that were used for the original targets, use the settings ts. The following code simulates the network that was trained in the previous code, and then converts the network output back into the original units.

```
an = sim(net,pn);
a = mapminmax('reverse',an,ts);
```

The network output an corresponds to the normalized targets tn. The unnormalized network output a is in the same units as the original targets t.

If mapminmax is used to preprocess the training set data, then whenever the trained network is used with new inputs they should be preprocessed with the minimum and maximums that were computed for the training set stored in the settings ps. The following code applies a new set of inputs to the network already trained.

```
pnewn = mapminmax('apply',pnew,ps);
anewn = sim(net,pnewn);
anew = mapminmax('reverse',anewn,ts);
```

For most networks, including feedforwardnet, these steps are done automatically, so that you only need to use the sim command.

See Also fixunknowns | mapstd | processpca

Purpose	Process mat	Process matrices by mapping each row's means to 0 and deviations to 1		
Syntax	<pre>[Y,PS] = mapstd(X,ymean,ystd) [Y,PS] = mapstd(X,FP) Y = mapstd('apply',X,PS) X = mapstd('reverse',Y,PS) dx_dy = mapstd('dx_dy',X,Y,PS)</pre>			
Description	mapstd processes matrices by transforming the mean and standard deviation of each row to ymean and ystd.			
	[Y,PS] = m	napstd(X,ymean,ystd) takes X and optional parameters,		
	Х	N-by-Q matrix or a 1-by-TS row cell array of N-by-Q matrices		
	ymean	Mean value for each row of Y (default is 0)		
	ystd	Standard deviation for each row of Y (default is 1)		
	and returns Y Each M-by-Q matrix (where M == N) (optional)			
	PS	Process settings that allow consistent processing of values		
	[Y,PS] = m FP.ystd.	napstd(X,FP) takes parameters as a struct: FP.ymean,		
	Y = mapstd	('apply',X,PS) returns Y, given X and settings PS.		
	X = mapstd	X = mapstd('reverse',Y,PS) returns X, given Y and settings PS.		
	dx_dy = ma	apstd('dx_dy',X,Y,PS) returns the reverse derivative.		
Examples	Here you format a matrix so that the minimum and maximum values of each row are mapped to default mean and STD of 0 and 1.			

x1 = [1 2 4; 1 1 1; 3 2 2; 0 0 0][y1, PS] = mapstd(x1)Next, apply the same processing settings to new values.  $x^2 = [5 2 3; 1 1 1; 6 7 3; 0 0 0]$ y2 = mapstd('apply', x2, PS)Reverse the processing of y1 to get x1 again. x1 again = mapstd('reverse',v1,PS) **Algorithms** It is assumed that X has only finite real values, and that the elements of each row are not all equal. y = (x - xmean) \* (ystd/xstd) + ymean;**Definitions** Another approach for scaling network inputs and targets is to normalize the mean and standard deviation of the training set. The function mapstd normalizes the inputs and targets so that they will have zero mean and unity standard deviation. The following code illustrates the use of mapstd. [pn,ps] = mapstd(p); [tn,ts] = mapstd(t);The original network inputs and targets are given in the matrices p and t. The normalized inputs and targets pn and tn that are returned will have zero means and unity standard deviation. The settings structures ps and ts contain the means and standard deviations of the original inputs and original targets. After the network has been trained, you should use these settings to transform any future inputs that are applied to the network. They effectively become a part of the network, just like the network weights and biases. If mapstd is used to scale the targets, then the output of the network is trained to produce outputs with zero mean and unity standard

deviation. To convert these outputs back into the same units that were

used for the original targets, use ts. The following code simulates the network that was trained in the previous code, and then converts the network output back into the original units.

```
an = sim(net,pn);
a = mapstd('reverse',an,ts);
```

The network output an corresponds to the normalized targets tn. The unnormalized network output a is in the same units as the original targets t.

If mapstd is used to preprocess the training set data, then whenever the trained network is used with new inputs, you should preprocess them with the means and standard deviations that were computed for the training set using ps. The following commands apply a new set of inputs to the network already trained:

```
pnewn = mapstd('apply',pnew,ps);
anewn = sim(net,pnewn);
anew = mapstd('reverse',anewn,ts);
```

For most networks, including feedforwardnet, these steps are done automatically, so that you only need to use the sim command.

See Also fixunknowns | mapminmax | processpca

## maxlinlr

Purpose	Maximum learning rate for linear layer		
Syntax	<pre>lr = maxlinlr(P) lr = maxlinlr(P,'bias')</pre>		
Description	<pre>maxlinlr is used to calculate learning rates for linearlayer. lr = maxlinlr(P) takes one argument,</pre>		
	P R-by-Q matrix of input vectors		
	and returns the maximum learning rate for a linear layer without a bias that is to be trained only on the vectors in P.		
	<pre>lr = maxlinlr(P, 'bias') returns the maximum learning rate for a linear layer with a bias.</pre>		
Examples	Here you define a batch of four two-element input vectors and find th maximum learning rate for a linear layer with a bias.		
	P = [1 2 -4 7; 0.1 3 10 6]; lr = maxlinlr(P,'bias')		
See Also	learnwh   linearlayer		

### meanabs

Purpose	Mean of absolute elements of matrix or matrices	
Syntax	[m,n] = meanabs(x)	
Description	<pre>[m,n] = meanabs(x) takes a matrix or cell array of matrices and returns,</pre>	
	m Mean value of all absolute finite values	
	n Number of finite values	
	If x contains no finite values, the mean returned is 0.	
Examples	m = meanabs([1 2;3 4]) [m,n] = meanabs({[1 2; NaN 4], [4 5; 2 3]})	
See Also	meansqr   sumabs   sumsqr	

#### meansqr

Purpose	Mean of squared elements of matrix or matrices	
Syntax	[m,n] = meansqr(x)	
Description	[m,n] = meansqr(x) takes a matrix or cell array of matrices and returns,	
	m Mean value of all squared finite values	
	n Number of finite values	
	If <b>x</b> contains no finite values, the mean returned is 0.	
Examples	m = meansqr([1 2;3 4]) [m,n] = meansqr({[1 2; NaN 4], [4 5; 2 3]})	
See Also	meanabs   sumabs   sumsqr	

Purpose	Midpoint weight initialization function		
Syntax	<pre>W = midpoint(S,PR)</pre>		
Description	midpoint is a weight initialization function that sets weight (row) vectors to the center of the input ranges.		
	W = midpoint(	S,PR) takes two arguments,	
	S	Number of rows (neurons)	
	PR	R-by-Q matrix of input value ranges = [Pmin Pmax]	
	and returns an S-by-R matrix with rows set to (Pmin+Pmax)'/2.		
Examples	Here initial weight values are calculated for a five-neuron layer with input elements ranging over [0 1] and [-2 2].		
W = midpoint(5,[0 1; -2 2])		5,[0 1; -2 2])	
Network Use	You can create a standard network that uses midpoint to initial weights by calling newc.		
	To prepare the weights and the bias of layer i of a custom network to initialize with midpoint,		
	1 Set net.initFcn to 'initlay'. (net.initParam automatically becomes initlay's default parameters.)		
	<b>2</b> Set net.layers{i}.initFcn to 'initwb'.		
3 Set each net.inputWeights{i,j}.initFcn to 'midpoint net.layerWeights{i,j}.initFcn to 'midpoint'.			
	To initialize the network, call init.		
See Also	initwb   initl	ay   init	

## minmax

Purpose	Ranges of matrix	rows
Syntax	pr = minmax(P)	
Description	<pre>pr = minmax(P) takes one argument,</pre>	
	Р	R-by-Q matrix
	and returns the R-by-2 matrix PR of minimum and maximum values for each row of P. Alternatively, P can be an M-by-N cell array of matrices. Each matrix P{i,j} should have Ri rows and Q columns. In this case, minmax returns an M-by-1 cell array where the mth matrix is an Ri-by-2 matrix of the minimum and maximum values of elements for the matrix on the ith row of P.	
Examples	P = [0 1 2; -1 pr = minmax(P) P = {[0 1; -1 pr = minmax(P)	-2 -0.5] -2] [2 3 -2; 8 0 2]; [1 -2] [9 7 3]};

Purpose	Mean squared normalized error performance function	
Syntax	perf = mse(net,t,y,ew)	
Description	<pre>mse is a network performance function. It measures the network's performance according to the mean of squared errors. perf = mse(net,t,y,ew) takes these arguments:</pre>	
	netNeural networktMatrix or cell array of targets	
	У ew	Matrix or cell array of outputs Error weights (optional)

and returns the mean squared error.

This function has two optional parameters, which are associated with networks whose net.trainFcn is set to this function:

- 'regularization' can be set to any value between 0 and 1. The greater the regularization value, the more squared weights and biases are included in the performance calculation relative to errors. The default is 0, corresponding to no regularization.
- 'normalization' can be set to 'none' (the default); 'standard', which normalizes errors between -2 and 2, corresponding to normalizing outputs and targets between -1 and 1; and 'percent', which normalizes errors between -1 and 1. This feature is useful for networks with multi-element outputs. It ensures that the relative accuracy of output elements with differing target value ranges are treated as equally important, instead of prioritizing the relative accuracy of the output element with the largest target value range.

You can create a standard network that uses mse with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with mse, set net.performFcn to 'mse'. This automatically sets  $\verb|net.performParam|$  to a structure with the default optional parameter values.

# **Examples** Here a two-layer feedforward network is created and trained to predict median house prices using the mse performance function and a regularization value of 0.01, which is the default performance function for feedforwardnet.

```
[x,t] = house_dataset;
net = feedforwardnet(10);
net.performFcn = 'mse'; % Redundant, MSE is default
net.performParam.regularization = 0.01;
net = train(net,x,t);
y = net(x);
perf = perform(net,t,y);
```

Alternately, you can call this function directly.

perf = mse(net,x,t,'regularization',0.01);

See Also

mae

Purpose	Nonlinear autoregressive neural network			
Syntax	narnet(feedbackDelays,hiddenSizes,trainFcn)			
Description	NAR (nonlinear autoregressive) neural networks can be trained to predict a time series from that series past values.			
	narnet(feedbackDelays,hiddenSizes,trainFcn) takes these arguments,			
	feedbackDelays Row vector of increasing 0 or positive delays (default = 1:2)			
	hiddenSizes	Row vector of one or more hidden layer sizes (default = 10)		
	trainFcn	Training function (default = 'trainlm')		
	and returns a NAR neural network.			
Examples	<pre>Here a NAR network is used to solve a simple time series problem. T = simplenar_dataset; net = narnet(1:2,10) [Xs,Xi,Ai,Ts] = preparets(net,{},{},T) net = train(net,Xs,Ts,Xi,Ai); view(net) Y = net(Xs,Xi); perf = perform(net,Ts,Y)</pre>			
See Also	preparets   removedelay   timedelaynet   narnet   narxnet			

#### narxnet

Purpose	Nonlinear autoregressive neural network with external input		
Syntax	narxnet(inputDelays,feedbackDelays,hiddenSizes,trainFcn)		
Description	NARX (Nonlinear autoregressive with external input) networks can learn to predict one time series given past values of the same time series, the feedback input, and another time series, called the external or exogenous time series.		
	<pre>narxnet(inputDelays, takes these arguments,</pre>	feedbackDelays,hiddenSizes,trainFcn)	
	inputDelaysRow vector of increasing 0 or positive delays (default = 1:2)feedbackDelaysRow vector of increasing 0 or positive delays (default = 1:2)hiddenSizesRow vector of one or more hidden layer sizes (default = 10)trainFcnTraining function (default = 'trainlm')		
	and returns a NARX neural network.		
Examples	<pre>ples Here a NARX neural network is used to solve a simple time series problem. [X,T] = simpleseries_dataset; net = narxnet(1:2,1:2,10) [Xs,Xi,Ai,Ts] = preparets(net,X,{},T) net = train(net,Xs,Ts,Xi,Ai); view(net) Y = net(Xs,Xi,Ai); perf = perform(net,Ts,Y)</pre>		
	Here the NARX network is simulated in closed loop form.		

```
netc = closeloop(net);
view(netc)
[Xs,Xi,Ai,Ts] = preparets(netc,X,{},T);
y = netc(Xs,Xi,Ai)
```

Here the NARX network is used to predict the next output a timestep ahead of when it will actually appear.

```
netp = removedelay(net);
view(netp)
[Xs,Xi,Ai,Ts] = preparets(netp,X,{},T);
y = netp(Xs,Xi,Ai)
```

See Also closeloop | narnet | openloop | preparets | removedelay | timedelaynet

## nctool

Purpose	Neural network classification or clustering tool		
Syntax	nctool		
Description	nctool opens the neural network clustering GUI.		
Algorithms	nctool leads you through solving a clustering problem using a self-organizing map. The map forms a compressed representation of the inputs space, reflecting both the relative density of input vectors in that space, and a two-dimensional compressed representation of the input-space topology.		

Purpose	Negative distance weight function	
Syntax	Z = negdist(W,P) dim = negdist('size',S,R,FP) dw = negdist('dz_dw',W,P,Z,FP)	
Description	<b>negdist</b> is a weight function. Weight functions apply weights to an input to get weighted inputs.	
	Z = negdist(W,P)	takes these inputs,
	W	S-by-R weight matrix
	Р	R-by-Q matrix of Q input (column) vectors
	FP	Row cell array of function parameters (optional, ignored)
	and returns the S-by-Q matrix of negative vector distances.	
	dim = negdist('size',S,R,FP) takes the layer dimension S, input dimension R, and function parameters, and returns the weight size [S-by-R].	
	<pre>dw = negdist('dz_dw',W,P,Z,FP) returns the derivative of Z with respect to W.</pre>	
Examples	Here you define a random weight matrix W and input vector P and calculate the corresponding weighted input Z.	
	<pre>W = rand(4,3); P = rand(3,1); Z = negdist(W,P)</pre>	

## negdist

Network Use	You can create a standard network that uses negdist by calling competlayer or selforgmap.		
	To change a network so an input weight uses negdist, set net.inputWeight{i,j}.weightFcn to 'negdist'. For a layer weight, set net.layerWeight{i,j}.weightFcn to 'negdist'.		
	In either case, call sim to simulate the network with negdist.		
Algorithms	negdist returns the negative Euclidean distance:		
	$z = -sqrt(sum(w-p)^2)$		
See Also	competlayer   dist   dotprod   selforgmap   sim		

## netinv

Purpose	Inverse transfer function			
Syntax	A = netinv(N,	A = netinv(N,FP)		
Description	netinv is a transfer function. Transfer functions calculate a layer's output from its net input.			
	A = netinv(N,	FP) takes inputs		
	Ν	S-by-Q matrix of net input (column) vectors		
	FP	Struct of function parameters (ignored)		
	and returns 1/N	٨.		
	info = netinv following codes	('code') returns information about this function. The are supported:		
	netinv('name'	<pre>netinv('name') returns the name of this function.</pre>		
	<pre>netinv('output',FP) returns the [min max] output range. netinv('active',FP) returns the [min max] active input range. netinv('fullderiv') returns 1 or 0, depending on whether dA_dN is S-by-S-by-Q or S-by-Q. netinv('fpnames') returns the names of the function parameters. netinv('fpdefaults') returns the default function parameters.</pre>			
Examples	Here you define	e 10 five-element net input vectors N and calculate A.		
	n = rand(5,10 a = netinv(n)			
	Assign this trar	nsfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'netinv';</pre>			

## netinv

See Also tansig | logsig

Purpose	Product net input function		
Syntax	<pre>N = netprod({Z1,Z2,. info = netprod('code</pre>		
Description	netprod is a net input function. Net input functions calculate a layer's net input by combining its weighted inputs and biases.		
	N = netprod({Z1,Z2,.	,Zn}) takes	
	Zi	S-by-Q matrices in a row cell array	
	and returns an element-	wise product of Z1 to Zn.	
	<pre>info = netprod('code') returns information about this function. The following codes are supported:     'deriv' Name of derivative function</pre>		
	'fullderiv'	Full N-by-S-by-Q derivative = 1, element-wise S-by-Q derivative = 0	
	'name'	Full name	
	'fpnames'	Returns names of function parameters	
	'fpdefaults'	Returns default function parameters	
Examples	<pre>Here netprod combines two sets of weighted input vectors (user-defined). Z1 = [1 2 4;3 4 1]; Z2 = [-1 2 2; -5 -6 1]; Z = {Z1,Z2}; N = netprod({Z})</pre>		

	<pre>Here netprod combines the same weighted inputs with a bias vector. Because Z1 and Z2 each contain three concurrent vectors, three concurrent copies of B must be created with concur so that all sizes match. B = [0; -1]; Z = {Z1, Z2, concur(B,3)}; N = netprod(Z)</pre>	
Network Use	You can create a standard network that uses netprod by calling newpnn or newgrnn.	
	To change a network so that a layer uses netprod, set net.layers{i}.netInputFcn to 'netprod'.	
	In either case, call sim to simulate the network with netprod. See newpnn or newgrnn for simulation examples.	
See Also	sim   netsum   concur	

Purpose	Sum net input function	
Syntax	N = netsum({Z1,Z2,,Zn},FP) info = netsum(' <i>code</i> ')	
Description	netsum is a net input function. Net input functions calculate a layer's net input by combining its weighted inputs and biases.	
	N = netsum({Z1,Z2 parameters,	2,,Zn},FP) takes Z1 to Zn and optional function
	Zi	S-by-Q matrices in a row cell array
	FP	Row cell array of function parameters (ignored)
	and returns the elem	mentwise sum of Z1 to Zn.
	<pre>info = netsum('code') returns information about this function. The following codes are supported: netsum('name') returns the name of this function. netsum('type') returns the type of this function. netsum('fpnames') returns the names of the function parameters. netsum('fpdefaults') returns default function parameter values. netsum('fpcheck', FP) throws an error for illegal function parameters.</pre>	
	netsum('fullderiv derivative is S-by-Q	<pre>/') returns 0 or 1, depending on whether the or N-by-S-by-Q.</pre>
Examples		nes two sets of weighted input vectors and a bias. Ir to make B the same dimensions as Z1 and Z2.
	z1 = [1 2 4; 3 4 z2 = [-1 2 2; -5 b = [0; -1]	

#### netsum

	<pre>n = netsum({z1,z2,concur(b,3)})</pre>	
	Assign this net input function to layer i of a network.	
	<pre>net.layers(i).netFcn = 'compet';</pre>	
	Use feedforwardnet or cascadeforwardnet to create a standard network that uses netsum.	
See Also	cascadeforwardnet   feedforwardnet   netprod   netinv	

Create custom neural network		
<pre>net = network net = network(numInputs,numLayers,biasConnect,inputConnect,</pre>		
Type help network/network.		
network creates new custom networks. It is used to create networks that are then customized by functions such as feedforwardnet and narxnet.		
<b>net</b> = <b>network</b> without arguments returns a new neural network with no inputs, layers or outputs.		
<pre>net = network(numInputs,numLayers,biasConnect,inputConnect,layerConnect,out takes these optional arguments (shown with default values):</pre>		
numInputs	Number of inputs, 0	
numLayers	Number of layers, 0	
biasConnect	numLayers-by-1 Boolean vector, zeros	
inputConnect	numLayers-by-numInputs Boolean matrix, zeros	
layerConnect numLayers-by-numLayers Boolean mat zeros		
outputConnect	1-by-numLayers Boolean vector, zeros	
and returns		
net	New network with the given property values	
	<pre>net = network net = network(numInp layerConnect,outp Type help network/ne network creates new cu that are then customize narxnet. net = network without no inputs, layers or out net = network(numInputs,nut takes these optional arg numInputs numLayers biasConnect inputConnect layerConnect outputConnect and returns</pre>	

#### **Properties** Architecture Properties

net.numInputs	0 or a positive integer	Number of inputs.
net.numLayers	0 or a positive integer	Number of layers.
net.biasConnect	numLayer-by-1 Boolean vector	If net.biasConnect(i) is 1, then layer i has a bias, and net.biases{i} is a structure describing that bias.
net.inputConnect	numLayer-by-nu Boolean vector	<pre>mlfnpertsinputConnect(i,j) is 1, then layer i has a weight coming from input j, and net.inputWeights{i,j} is a structure describing that weight.</pre>
net.layerConnect	numLayer-by-nu Boolean vector	<pre>mLfayerslayerConnect(i,j) is 1, then layer i has a weight coming from layer j, and net.layerWeights{i,j} is a structure describing that weight.</pre>
net.numInputs	0 or a positive integer	Number of inputs.
net.numLayers	0 or a positive integer	Number of layers.
net.biasConnect	numLayer-by-1 Boolean vector	If net.biasConnect(i) is 1, then layer i has a bias, and net.biases{i} is a structure describing that bias.

net.inputConnect	numLayer-by-nur Boolean vector	<pre>IffmpertsinputConnect(i,j) is 1, then layer i has a weight coming from input j, and net.inputWeights{i,j} is a structure describing that weight.</pre>
net.layerConnect	numLayer-by-nur Boolean vector	<pre>https://www.section.com/s</pre>
net.outputConnect	1-by-numLayers Boolean vector	If net.outputConnect(i) is 1, then the network has an output from layer i, and net.outputs{i} is a structure describing that output.
net.numOutputs	0 or a positive integer (read only)	Number of network outputs according to net.outputConnect.
net.numInputDelay	<b>'9</b> or a positive integer (read only)	Maximum input delay according to all net.inputWeight{i,j}.delays.
net.numLayerDelay	<b>/9</b> or a positive number (read only)	Maximum layer delay according to all net.layerWeight{i,j}.delays.

#### Subobject Structure Properties

net.inputs	numInputs-by- cell array	<pre>1net.inputs{i} is a structure   defining input i.</pre>
net.layers	numLayers-by- cell array	<pre>1net.layers{i} is a structure   defining layer i.</pre>

### network

net.biases	numLayers-by- cell array	<pre>1If net.biasConnect(i) is 1, then net.biases{i} is a structure defining the bias for layer i.</pre>
net.inputWeights	numLayers-by- cell array	<pre>nLfm1eputsputConnect(i,j) is 1, then net.inputWeights{i,j} is a structure defining the weight to layer i from input j.</pre>
net.layerWeights	numLayers-by- cell array	<pre>nlfhtetyetayerConnect(i,j) is 1, then net.layerWeights{i,j} is a structure defining the weight to layer i from layer j.</pre>
net.outputs	1-by-numLayer cell array	<pre>sIf net.outputConnect(i) is 1, then net.outputs{i} is a structure defining the network output from layer i.</pre>

#### **Function Properties**

net.adaptFcn	Name of a network adaption function or ''
net.initFcn	Name of a network initialization function or ''
net.performFcn	Name of a network performance function or ''
net.trainFcn	Name of a network training function or ' '
Parameter Properties	

net.adaptParam	Network adaption parameters
net.initParam	Network initialization
	parameters

net.performParam	Network performance parameters
net.trainParam	Network training parameters

#### Weight and Bias Value Properties

net.IW	numLayers-by-numInputs cell array of input weight values
net.LW	numLayers-by-numLayers cell array of layer weight values
net.b	numLayers-by-1 cell array of bias values

#### **Other Properties**

Structure you can use to store useful values

## **Examples** Here is the code to create a network without any inputs and layers, and then set its numbers of inputs and layers to 1 and 2 respectively.

net = network
net.numInputs = 1
net.numLayers = 2

Here is the code to create the same network with one line of code.

```
net = network(1,2)
```

Here is the code to create a one-input, two-layer, feed-forward network. Only the first layer has a bias. An input weight connects to layer 1 from input 1. A layer weight connects to layer 2 from layer 1. Layer 2 is a network output and has a target.

```
net = network(1,2,[1;0],[1; 0],[0 0; 1 0],[0 1])
```

You can see the properties of subobjects as follows:

```
net.inputs{1}
net.layers{1}, net.layers{2}
net.biases{1}
net.inputWeights{1,1}, net.layerWeights{2,1}
net.outputs{2}
```

You can get the weight matrices and bias vector as follows:

net.iw.{1,1}, net.iw{2,1}, net.b{1}

You can alter the properties of any of these subobjects. Here you change the transfer functions of both layers:

net.layers{1}.transferFcn = 'tansig'; net.layers{2}.transferFcn = 'logsig';

Here you change the number of elements in input 1 to 2 by setting each element's range:

net.inputs{1}.range = [0 1; -1 1];

Next you can simulate the network for a two-element input vector:

p = [0.5; -0.1]; y = sim(net,p)

sim

#### See Also

Purpose	Design generalized regression neural network		
Syntax	<pre>net = newgrnn(P,T,spread)</pre>		
Description	Generalized regression neural networks (grnns) are a kind of radial basis network that is often used for function approximation. grnns can be designed very quickly.		
	net = newgrnn	n(P,T,spread) takes three inputs,	
	Р	R-by-Q matrix of Q input vectors	
	Т	S-by-Q matrix of Q target class vectors	
	spread	Spread of radial basis functions (default = $1.0$ )	
	and returns a n	new generalized regression neural network.	
	The larger the spread, the smoother the function approximation. To fit data very closely, use a spread smaller than the typical distance between input vectors. To fit the data more smoothly, use a larger spread.		
Properties	newgrnn creates a two-layer network. The first layer has radbas neurons, and calculates weighted inputs with dist and net input with netprod. The second layer has purelin neurons, calculates weighted input with normprod, and net inputs with netsum. Only the first layer has biases.		
	<code>newgrnn</code> sets the first layer weights to P', and the first layer biases are all set to $0.8326/spread$ , resulting in radial basis functions that cross 0.5 at weighted inputs of +/- spread. The second layer weights W2 are set to T.		
Examples	Here you desig	n a radial basis network, given inputs P and targets T.	
	P = [1 2 3]; T = [2.0 4.1 5.9];		

	<pre>net = newgrnn(P,T);</pre>
	The network is simulated for a new input.
	P = 1.5; Y = sim(net,P)
References	Wasserman, P.D., <i>Advanced Methods in Neural Computing</i> , New York, Van Nostrand Reinhold, 1993, pp. 155–61
See Also	sim   newrb   newrbe   newpnn

## newlind

Purpose	Design linear layer			
Syntax	<pre>net = newlind(P,T,Pi)</pre>			
Description	<pre>net = newlind(P,T,Pi) takes these input arguments,</pre>			
	Р	R-by-Q matrix of	Q input vectors	
	т	S-by-Q matrix of	Q target class vectors	
	Pi	1-by-ID cell arra	y of initial input delay states	
	where each element Pi{i,k} is an Ri-by-Q matrix, and the default = []; and returns a linear layer designed to output T (with minimum sum square error) given input P.			
	newlind(P,T,Pi) can also solve for linear networks with input delays and multiple inputs and layers by supplying input and target data in cell array form:			
	P Ni-by-TS cell Each element P{i,ts} is an array Ri-by-Q input matrix			
	т	Nt-by-TS cell array	Each element P{i,ts} is a Vi-by-Q matrix	
	Pi	Ni-by-ID cell array	Each element Pi{i,k} is an Ri-by-Q matrix, default = []	
Examples	<pre>and returns a linear network with ID input delays, Ni network inputs, and N1 layers, designed to output T (with minimum sum square error) given input P. You want a linear layer that outputs T given P for the following definitions: P = [1 2 3]; T = [2.0 4.1 5.9];</pre>			

Use newlind to design such a network and check its response.

net = newlind(P,T); Y = sim(net,P)

You want another linear layer that outputs the sequence T given the sequence P and two initial input delay states  $\mathsf{Pi.}$ 

P = {1 2 1 3 3 2}; Pi = {1 3}; T = {5.0 6.1 4.0 6.0 6.9 8.0}; net = newlind(P,T,Pi); Y = sim(net,P,Pi)

You want a linear network with two outputs Y1 and Y2 that generate sequences T1 and T2, given the sequences P1 and P2, with three initial input delay states Pi1 for input 1 and three initial delays states Pi2 for input 2.

```
P1 = {1 2 1 3 3 2}; Pi1 = {1 3 0};
P2 = {1 2 1 1 2 1}; Pi2 = {2 1 2};
T1 = {5.0 6.1 4.0 6.0 6.9 8.0};
T2 = {11.0 12.1 10.1 10.9 13.0 13.0};
net = newlind([P1; P2],[T1; T2],[Pi1; Pi2]);
Y = sim(net,[P1; P2],[Pi1; Pi2]);
Y1 = Y(1,:)
Y2 = Y(2,:)
```

## **Algorithms** newlind calculates weight W and bias B values for a linear layer from inputs P and targets T by solving this linear equation in the least squares sense:

[W b] \* [P; ones] = T

sim

#### See Also

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Purpose	Design probabilistic neural network	
Syntax	net = newpnn(P,T,spread)	
Description	Probabilistic neural networks (PNN) are a kind of radial basis network suitable for classification problems.	
	<pre>net = newpnn(P,T,</pre>	spread) takes two or three arguments,
	Р	R-by-Q matrix of Q input vectors
	Т	S-by-Q matrix of Q target class vectors
	spread Spread of radial basis functions (defa	
	-	probabilistic neural network.
	If spread is near zero, the network acts as a nearest neighbor classifier. As spread becomes larger, the designed network takes into account several nearby design vectors.	
Examples	<pre>Here a classification problem is defined with a set of inputs P and class indices Tc. P = [1 2 3 4 5 6 7]; Tc = [1 2 3 2 2 3 1]; The class indices are converted to target vectors, and a PNN is designed and tested. T = ind2vec(Tc) net = newpnn(P,T); Y = sim(net,P) Yc = vec2ind(Y)</pre>	
Algorithms	neurons, and calcula	vo-layer network. The first layer has radbas ates its weighted inputs with dist and its net input second layer has compet neurons, and calculates its

	weighted input with dotprod and its net inputs with netsum. Only the first layer has biases.		
	newpnn sets the first-layer weights to P', and the first-layer biases are all set to $0.8326/spread$ , resulting in radial basis functions that cross 0.5 at weighted inputs of +/- spread. The second-layer weights W2 are set to T.		
References	Wasserman, P.D., <i>Advanced Methods in Neural Computing</i> , New York, Van Nostrand Reinhold, 1993, pp. 35–55		
See Also	sim   ind2vec   vec2ind   newrb   newrbe   newgrnn		

Purpose	Design radial basis network		
Syntax	<pre>net = newrb(P,T,goal,spread,MN,DF)</pre>		
Description	Radial basis networks can be used to approximate functions. newrb adds neurons to the hidden layer of a radial basis network until it meets the specified mean squared error goal.		
	<pre>net = newrb(P,T,</pre>	goal,spread,MN,DF) takes two of these arguments,	
	Р	R-by-Q matrix of Q input vectors	
	Т	S-by-Q matrix of Q target class vectors	
	goal	Mean squared error goal (default = $0.0$ )	
	spread	Spread of radial basis functions (default = $1.0$ )	
	MN	Maximum number of neurons (default is Q)	
	DF	Number of neurons to add between displays (default = 25)	
	and returns a new radial basis network.		
	The larger <b>spread</b> is, the smoother the function approximation. Too large a spread means a lot of neurons are required to fit a fast-changing function. Too small a spread means many neurons are required to fit a smooth function, and the network might not generalize well. Call <b>newrb</b> with different spreads to find the best value for a given problem.		
Examples	Here you design a radial basis network, given inputs P and targets T.		
	_	P = [1 2 3]; T = [2.0 4.1 5.9]; net = newrb(P,T);	
	The network is simulated for a new input.		
	P = 1.5;		

Υ	=	sim(	(net	,P)
---	---	------	------	-----

# **Algorithms** newrb creates a two-layer network. The first layer has radbas neurons, and calculates its weighted inputs with dist and its net input with netprod. The second layer has purelin neurons, and calculates its weighted input with dotprod and its net inputs with netsum. Both layers have biases.

Initially the radbas layer has no neurons. The following steps are repeated until the network's mean squared error falls below goal.

- 1 The network is simulated.
- **2** The input vector with the greatest error is found.
- **3** A radbas neuron is added with weights equal to that vector.
- **4** The purelin layer weights are redesigned to minimize error.
- See Also sim | newrbe | newgrnn | newpnn

Purpose	Design exact radial basis network		
Syntax	<pre>net = newrbe(P,T,spread)</pre>		
Description	Radial basis networks can be used to approximate functions. newrbe very quickly designs a radial basis network with zero error on the design vectors.		
	net = newrbe(P,T	, spread) takes two or three arguments,	
	Р	RxQ matrix of Q R-element input vectors	
	т	$\mathbf{SxQ}\xxptox$ matrix of $\mathbf{Q}\xptox$ S-element target class vectors	
	spread	Spread of radial basis functions (default = $1.0$ )	
	and returns a new exact radial basis network. The larger the <b>spread</b> is, the smoother the function approximation will be. Too large a spread can cause numerical problems.		
Examples	Here you design a radial basis network given inputs P and targets T.		
	]; );		
	The network is simulated for a new input.		
	P = 1.5; Y = sim(net,P)		
Algorithms	newrbe creates a two-layer network. The first layer has radbas neurons, and calculates its weighted inputs with dist and its net input with netprod. The second layer has purelin neurons, and calculates its weighted input with dotprod and its net inputs with netsum. Both layers have biases.		

newrbe sets the first-layer weights to P', and the first-layer biases are all set to 0.8326/spread, resulting in radial basis functions that cross 0.5 at weighted inputs of +/- spread.

The second-layer weights  $IW{2,1}$  and biases  $b{2}$  are found by simulating the first-layer outputs  $A{1}$  and then solving the following linear expression:

[W{2,1} b{2}] \* [A{1}; ones] = T

See Also sim | newrb | newgrnn | newpnn

Purpose	Neural network fitting tool
Syntax	nftool
Description	nftool opens the neural network fitting tool GUI.
Algorithms	nftool leads you through solving a data fitting problem, solving it with a two-layer feed-forward network trained with Levenberg-Marquardt.
See Also	nntool

## nncell2mat

Purpose	Combine neural network cell data into matrix		
Syntax	[y,i,j] nncell2mat(x)		
Description	[y,i,j] nncell2mat(x) takes a cell array of matrices and returns,		
	y Cell array formed by concatenating matrices		
	i	Array of row sizes	
	ji	Array of column sizes	
Examples	<pre>The row and column sizes returned by nncell2mat can be used to convert the returned matrix back into a cell of matrices with mat2cell. Here neural network data is converted to a matrix and back. c = {rands(2,3) rands(2,3); rands(5,3) rands(5,3)}; [m,i,j] = nncell2mat(c) c3 = mat2cell(m,i,j)</pre>		
See Also	nndata   nnsize		

Purpose	Crross correlation between neural network time series		
Syntax	<pre>nncorr(a,b,maxlag,'flag')</pre>		
Description	<pre>nncorr(a,b,maxlag,'flag') takes these arguments,</pre>		
	<ul> <li>a Matrix or cell array, with columns interpreted a timesteps, and having a total number of matrix rows of N.</li> <li>b Matrix or cell array, with columns interpreted a timesteps, and having a total number of matrix rows of M.</li> <li>maxlag Maximum number of time lags</li> </ul>		
	flag	Type of normalization (default = 'none')	
	<ul> <li>and returns an N-by-M cell array where each {i,j} element is a 2*maxlag+1 length row vector formed from the correlations of a elements (i.e., matrix row) i and b elements (i.e., matrix column) j.</li> <li>If a and b are specified with row vectors, the result is returned in matrix form.</li> <li>The options for the normalization <i>flag</i> are: <ul> <li>'biased' — scales the raw cross-correlation by 1/N.</li> <li>'unbiased' — scales the raw correlation by 1/(N-abs(k)), where k is the index into the result.</li> </ul> </li> </ul>		
	<ul> <li>'coeff' — normalizes the sequence so that the correlations at zero lag are 1.0.</li> <li>'none' — no scaling. This is the default.</li> </ul>		
Examples		ation of a random 1-element, 1-sample, 20-timestep with a maximum lag of 10.	

a = nndata(1,1,20) aa = nncorr(a,a,10)

Here the cross-correlation of the first signal with another random 2-element signal are found, with a maximum lag of 8.

b = nndata(2,1,20)ab = nncorr(a,b,8)

See Also confusion | regression

Create neural network data		
nndata(N,Q,TS,v)		
nndata(N,Q,TS,v) takes these arguments,		
Ν	Vector of M element sizes	
Q	Number of samples	
TS	Number of timesteps	
v	Scalar value	
and returns an M-by-TS cell array where each row i has $N(i)$ -by-Q sized matrices of value v. If v is not specified, random values are returned.		
	subsets of neural network data with getelements, ttimesteps, and getsignals.	
You can set subsets of neural network data with setelements, setsamples, settimesteps, and setsignals. You can concatenate subsets of neural network data with catelements, catsamples, cattimesteps, and catsignals.		
To create random data with the same dimensions: x = nndata(2,4,5)		
	nndata(N,Q,TS nndata(N,Q,TS N Q TS V and returns an matrices of value You can access getsamples, ge You can set sub setsamples, se You can concate catsamples, ca Here four samp of zero values is x = nndata(2, To create rando x = nndata(2, Here static (1 ti	

## nndata

See Also nnsize | tonndata | fromnndata | nndata2sim | sim2nndata

Purpose	Format neural data for efficient GPU training or simulation
Syntax	nndata2gpu(x) [Y,Q,N,TS] = nndata2gpu(X) nndata2gpu(X,PRECISION)
Description	nndata2gpu requires Parallel Computing Toolbox™.
	nndata2gpu(x) takes an N-by-Q matrix X of Q N-element column vectors, and returns it in a form for neural network training and simulation on the current GPU device.
	The N-by-Q matrix becomes a QQ-by-N gpuArray where QQ is Q rounded up to the next multiple of 32. The extra rows (Q+1):QQ are filled with NaN values. The gpuArray has the same precision ('single' or 'double') as X.
	<pre>[Y,Q,N,TS] = nndata2gpu(X) can also take an M-by-TS cell array of M signals over TS time steps. Each element of X{i,ts} should be an Ni-by-Q matrix of Q Ni-element vectors, representing the ith signal vector at time step ts, across all Q time series. In this case, the gpuArray Y returned is QQ-by-(sum(Ni)*TS). Dimensions Ni, Q, and TS are also returned so they can be used with gpu2nndata to perform the reverse formatting.</pre>
	nndata2gpu(X,PRECISION) specifies the default precision of the gpuArray, which can be 'double' or 'single'.
Examples	Copy a matrix to the GPU and back:
	x = rand(5,6) [y,q] = nndata2gpu(x) x2 = gpu2nndata(y,q)
	Copy neural network cell array data, representing four time series, each consisting of five time steps of 2-element and 3-element signals:
	x = nndata([2;3],4,5) [y,q,n,ts] = nndata2gpu(x)

x2 = gpu2nndata(y,q,n,ts)

See Also gpu2nndata

## nndata2sim

Purpose	Convert neural network data to Simulink time series		
Syntax	nndata2sim(x,i,q)		
Description	<pre>nndata2sim(x,i,q) takes these arguments,</pre>		
	x	Neural network data	
	i	Index of signal (default = $1$ )	
	q	Index of sample (default = $1$ )	
Examples	and returns time series <b>q</b> of signal <b>i</b> as a Simulink time series structure. Here random neural network data is created with two signals having 4 and 3 elements respectively, over 10 timesteps. Three such series		
	are created. x = nndata([4;3],3,10);		
	x = muata([4, 5], 5, 10),		
	Now the second signal of the first series is converted to Simulink form.		
	$y_2_1 = nndata2sim(x,2,1)$		
See Also	nndata   sim2n	ndata   nnsize	

### nnsize

Purpose	Number of neural data elements, samples, timesteps, and signals	
Syntax	[N,Q,TS,M] = nnsize(X)	
Description	[N,Q,TS,M] = nnsize(X) takes neural network data x and returns,	
	Ν	Vector containing the number of element sizes for each of M signals
	Q	Number of samples
	TS	Number of timesteps
	Μ	Number of signals
	If X is a matrix, N is the number of rows of X, Q is the number of columns, and both TS and M are 1. If X is a cell array, N is an Sx1 vector, where M is the number of rows in X, and N(i) is the number of rows in X{i,1}. Q is the number of columns in the matrices in X.	
Examples	This code gets t	he dimensions of matrix data:
	x = [1 2 3; 4 7 4] [n,q,ts,s] = nnsize(x)	
	This code gets the dimensions of cell array data:	
	x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} [n,q,ts,s] = nnsize(x)	
See Also	nndata   numel	ements   numsamples   numsignals   numtimesteps

Purpose	Neural network getting started GUI
Syntax	nnstart
Description	nnstart opens a window with launch buttons for neural network fitting, pattern recognition, clustering and time series wizards. It also provides links to lists of data sets, examples, and other useful information for getting started.
See Also	nctool   nftool   nprtool   ntstool

## nntool

Purpose	Open Network/Data Manager
Syntax	nntool
Description	nntool opens the Network/Data Manager window, which allows you to import, create, use, and export neural networks and data.

#### nntraintool

Syntax nntraintool

**Description** nntraintool opens the neural network training GUI.

This function can be called to make the training GUI visible before training has occurred, after training if the window has been closed, or just to bring the training GUI to the front.

Network training functions handle all activity within the training window.

To access additional useful plots, related to the current or last network trained, during or after training, click their respective buttons in the training window.

## noloop

Purpose	Remove neural network open- and closed-loop feedback
Syntax	<pre>net = noloop(net)</pre>
Description	<pre>net = noloop(net) takes a neural network and returns the network with open- and closed-loop feedback removed.</pre>
	For outputs i, where net.outputs{i}.feedbackMode is 'open', the feedback mode is set to 'none', outputs{i}.feedbackInput is set to the empty matrix, and the associated network input is deleted.
	For outputs i, where net.outputs{i}.feedbackMode is 'closed', the feedback mode is set to 'none'.
Examples	Here a NARX network is designed. The NARX network has a standard input and an open-loop feedback output to an associated feedback input.
	<pre>[X,T] = simplenarx_dataset; net = narxnet(1:2,1:2,20); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts,Xi,Ai); view(net) Y = net(Xs,Xi,Ai)</pre>
	Now the network is converted to no loop form. The output and second input are no longer associated.
	net = noloop(net); view(net) [Xs,Xi,Ai] = preparets(net,X,T); Y = net(Xs,Xi,Ai)
See Also	closeloop   openloop

Purpose	Normalize columns	of matrix
Syntax	normc(M)	
Description	normc(M) normalize	es the columns of M to a length of 1.
Examples	m = [1 2; 3 4]; normc(m) ans =	
		0.4472
	0.9487 (	0.8944
See Also	normr	

## normprod

Purpose	Normalized dot product	weight function
Syntax	Z = normprod(W,P,FP) dim = normprod('size dw = normprod('dz_dw	
Description	normprod is a weight function. Weight functions apply weights to an input to get weighted inputs.	
	<pre>Z = normprod(W,P,FP)</pre>	takes these inputs,
	W	S-by-R weight matrix
	Р	R-by-Q matrix of Q input (column) vectors
	FP	Row cell array of function parameters (optional, ignored)
	and returns the S-by-Q n	natrix of normalized dot products.
	dim = normprod('size',S,R,FP) takes the layer dimension S, input dimension R, and function parameters, and returns the weight size [S-by-R].	
	<pre>dw = normprod('dz_dw',W,P,Z,FP) returns the derivative of Z w respect to W.</pre>	
Examples	Here you define a rando calculate the correspond	m weight matrix W and input vector P and ing weighted input Z.
	<pre>W = rand(4,3); P = rand(3,1); Z = normprod(W,P)</pre>	

Network Use	You can create a standard network that uses normprod by calling newgrnn.
	To change a network so an input weight uses normprod, set net.inputWeight{i,j}.weightFcn to 'normprod'. For a layer weight, set net.layerWeight{i,j}.weightFcn to 'normprod'.
	In either case, call <b>sim</b> to simulate the network with <b>normprod</b> . See <b>newgrnn</b> for simulation examples.
Algorithms	<b>normprod</b> returns the dot product normalized by the sum of the input vector elements.
	z = w*p/sum(p)
See Also	dotprod

#### normr

Purpose	Normalize rows of matrix
Syntax	normr(M)
Description	normr(M) normalizes the rows of M to a length of 1.
Examples	<pre>m = [1 2; 3 4]; normr(m) ans =</pre>
See Also	normc

Purpose	Neural network pattern recognition tool
Syntax	nprtool
Description	nprtool opens the neural network pattern-recognition GUI.
Algorithms	nprtool leads you through solving a pattern-recognition classification problem using a two-layer feed-forward patternnet network with sigmoid output neurons.
See Also	nctool   nftool   ntstool

### ntstool

Purpose	Neural network time series tool
Syntax	ntstool ntstool('close')
Description	ntstool opens the neural network time series wizard and leads you through solving a fitting problem using a two-layer feed-forward network.
	<pre>ntstool('close') closes the wizard.</pre>
See Also	nctool   nftool   nprtool

Purpose	Numeric two-point network derivative function
Syntax	num2deriv('dperf_dwb',net,X,T,Xi,Ai,EW) num2deriv('de_dwb',net,X,T,Xi,Ai,EW)
Description	This function calculates derivatives using the two-point numeric derivative rule.

$$\frac{dy}{dx} = \frac{y(x+dx) - y(x)}{dx}$$

This function is much slower than the analytical (non-numerical) derivative functions, but is provided as a means of checking the analytical derivative functions. The other numerical function, num5deriv, is slower but more accurate.

num2deriv('dperf\_dwb',net,X,T,Xi,Ai,EW) takes these arguments,

net	Neural network
Х	Inputs, an RxQ matrix (or NxTS cell array of RixQ matrices)
Т	Targets, an SxQ matrix (or MxTS cell array of SixQ matrices)
Xi	Initial input delay states (optional)
Ai	Initial layer delay states (optional)
EW	Error weights (optional)

and returns the gradient of performance with respect to the network's weights and biases, where R and S are the number of input and output elements and Q is the number of samples (and N and M are the number of input and output signals, Ri and Si are the number of each input and outputs elements, and TS is the number of timesteps).

num2deriv('de\_dwb',net,X,T,Xi,Ai,EW) returns the Jacobian of errors with respect to the network's weights and biases.

## num2deriv

Examples	Here a feedforward network is trained and both the gradient and Jacobian are calculated.
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); y = net(x); perf = perform(net,t,y); dwb = num2deriv('dperf_dwb',net,x,t)</pre>
See Also	bttderiv   defaultderiv   fpderiv   num5deriv   staticderiv

Purpose	Numeric five-point stencil neural network derivative function
Syntax	num5deriv('dperf_dwb',net,X,T,Xi,Ai,EW) num5deriv('de_dwb',net,X,T,Xi,Ai,EW)
Description	This function calculates derivatives using the five-point numeric derivative rule.
	$y_{1} = y(x + 2dx)$ $y_{2} = y(x + dx)$ $y_{3} = y(x - dx)$ $y_{4} = y(x - 2dx)$ $\frac{dy}{dy} = \frac{y1 - 8y_{2} + 8y_{3} - y_{4}}{y_{4}}$

dx

dx

This function is much slower than the analytical (non-numerical) derivative functions, but is provided as a means of checking the analytical derivative functions. The other numerical function, num2deriv, is faster but less accurate.

num5deriv('dperf\_dwb',net,X,T,Xi,Ai,EW) takes these arguments,

net	Neural network
Х	Inputs, an RxQ matrix (or NxTS cell array of RixQ matrices)
Т	Targets, an SxQ matrix (or MxTS cell array of SixQ matrices)
Xi	Initial input delay states (optional)
Ai	Initial layer delay states (optional)
EW	Error weights (optional)

and returns the gradient of performance with respect to the network's weights and biases, where R and S are the number of input and output

	elements and Q is the number of samples (and N and M are the number of input and output signals, Ri and Si are the number of each input and outputs elements, and TS is the number of timesteps). num5deriv('de_dwb',net,X,T,Xi,Ai,EW) returns the Jacobian of errors with respect to the network's weights and biases.
Examples	Here a feedforward network is trained and both the gradient and Jacobian are calculated.
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); y = net(x); perf = perform(net,t,y); dwb = num5deriv('dperf_dwb',net,x,t)</pre>
See Also	bttderiv   defaultderiv   fpderiv   num2deriv   staticderiv

## numelements

Purpose	Number of elements in neural network data
Syntax	numelements(x)
Description	numelements(x) takes neural network data x in matrix or cell array form, and returns the number of elements in each signal.
	If x is a matrix the result is the number of rows of x.
	If x is a cell array the result is an S-by-1 vector, where S is the number of signals (i.e., rows of X), and each element $S(i)$ is the number of elements in each signal i (i.e., rows of $x{i,1}$ ).
Examples	This code calculates the number of elements represented by matrix data:
	x = [1 2 3; 4 7 4] n = numelements(x)
	This code calculates the number of elements represented by cell data:
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} n = numelements(x)</pre>
See Also	nndata   nnsize   getelements   setelements   catelements   numsamples   numsignals   numtimesteps

## numfinite

Purpose	Number of finite values in neural network data
Syntax	numfinite(x)
Description	numfinite(x) takes a matrix or cell array of matrices and returns the number of finite elements in it.
Examples	x = [1 2; 3 NaN] n = numfinite(x)
	x = {[1 2; 3 NaN] [5 NaN; NaN 8]} n = numfinite(x)
See Also	numnan   nndata   nnsize

#### numnan

Purpose	Number of NaN values in neural network data
Syntax	numnan(x)
Description	<pre>numnan(x) takes a matrix or cell array of matrices and returns the number of NaN elements in it.</pre>
Examples	x = [1 2; 3 NaN] n = numnan(x)
	x = {[1 2; 3 NaN] [5 NaN; NaN 8]} n = numnan(x)
See Also	numnan   nndata   nnsize

# numsamples

Purpose	Number of samples in neural network data
Syntax	numsamples(x)
Description	<pre>numsamples(x) takes neural network data x in matrix or cell array form, and returns the number of samples.</pre>
	If x is a matrix, the result is the number of columns of x.
	If ${\boldsymbol x}$ is a cell array, the result is the number of columns of the matrices in ${\boldsymbol x}.$
Examples	This code calculates the number of samples represented by matrix data:
	x = [1 2 3; 4 7 4] n = numsamples(x)
	This code calculates the number of samples represented by cell data:
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} n = numsamples(x)</pre>
See Also	nndata   nnsize   getsamples   setsamples   catsamples   numelements   numsignals   numtimesteps

## numsignals

Purpose	Number of signals in neural network data
Syntax	numsignals(x)
Description	numsignals(x) takes neural network data x in matrix or cell array form, and returns the number of signals.
	If x is a matrix, the result is 1.
	If x is a cell array, the result is the number of rows in x.
Examples	This code calculates the number of signals represented by matrix data:
	x = [1 2 3; 4 7 4] n = numsignals(x)
	This code calculates the number of signals represented by cell data:
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} n = numsignals(x)</pre>
See Also	nndata   nnsize   getsignals   setsignals   catsignals   numelements   numsamples   numtimesteps

## numtimesteps

Purpose	Number of time steps in neural network data	
Syntax	numtimesteps(x)	
Description	numtimesteps(x) takes neural network data x in matrix or cell array form, and returns the number of signals.	
	If x is a matrix, the result is 1.	
	If x is a cell array, the result is the number of columns in x.	
Examples	This code calculates the number of time steps represented by matrix data:	
	x = [1 2 3; 4 7 4] n = numtimesteps(x)	
	This code calculates the number of time steps represented by cell data:	
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} n = numtimesteps(x)</pre>	
See Also	nndata   nnsize   gettimesteps   settimesteps   cattimesteps   numelements   numsamples   numsignals	

Purpose	Convert neural network closed-loop feedback to open loop
Syntax	<pre>net = openloop(net)</pre>
Description	<pre>net = openloop(net) takes a neural network and opens any closed-loop feedback. For each feedback output i whose property net.outputs{i}.feedbackMode is 'closed', it replaces its associated feedback layer weights with a new input and input weight connections. The net.outputs{i}.feedbackMode property is set to 'open', and the net.outputs{i}.feedbackInput property is set to the index of the new input. Finally, the value of net.outputs{i}.feedbackDelays is subtracted from the delays of the feedback input weights (i.e., to the delays values of the replaced layer weights).</pre>
Examples	<pre>Here a NARX network is designed in open-loop form and then converted to closed-loop form, then converted back. [X,T] = simplenarx_dataset; net = narxnet(1:2,1:2,10); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts,Xi,Ai); view(net) Yopen = net(Xs,Xi,Ai) net = closeloop(net) view(net) [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); Yclosed = net(Xs,Xi,Ai); net = openloop(net) view(net) [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); Yopen = net(Xs,Xi,Ai)</pre>
See Also	closeloop   noloop

## patternnet

Purpose	Pattern recognition network		
Syntax	patternnet(hiddenSiz	patternnet(hiddenSizes,trainFcn)	
Description	Pattern recognition networks are feedforward networks that can be trained to classify inputs according to target classes. The target data for pattern recognition networks should consist of vectors of all zero values except for a 1 in element i, where i is the class they are to represent.		
	patternnet(hiddenSiz	es,trainFcn) takes these arguments,	
	hiddenSizes	Row vector of one or more hidden layer sizes (default = 10)	
	trainFcn	Training function (default = 'trainscg')	
and returns a pattern recognition neural netw	ecognition neural network.		
Examples	Here a pattern recognition network is designed to classify iris flowers.		
	<pre>[x,t] = iris_dataset net = patternnet(10) net = train(net,x,t) view(net) y = net(x); perf = perform(net,t classes = vec2ind(y)</pre>	;	
See Also	lvqnet   competlayer	selforgmap   nprtool	

### perceptron

Purpose	Perceptron		
Syntax	perceptron(hardlimitTF,perceptronLF)		
Syntax			
Description	<ul> <li>Perceptrons are simple single-layer binary classifiers, which divide the input space with a linear decision boundary.</li> <li>Perceptrons are provide for historical interest. For much better results use patternnet, which can solve non-linearly separable problems. Sometimes when people refer to perceptrons they are referring to feed-forward pattern recognition networks, such as patternnet. But the original perceptron, described here, can solve only very simple problems.</li> </ul>		
	Perceptrons can learn to solve a narrow class of classification problems. Their significance is they have a simple learning rule and were one of the first neural networks to reliably solve a given class of problems.		
	perceptron(hardlimitTF,perceptronLF) takes these arguments,		
	hardlimitTF	Hard limit transfer function (default = 'hardlim')	
	perceptronLF	<pre>Perceptron learning rule (default = 'learnp')</pre>	
	and returns a perceptron		
	In addition to the default hard limit transfer functions, perceptrons can be created with the hardlims transfer function. The other option for the perceptron learning rule is learnpn.		
Examples	<b>Examples</b> Here a perceptron is used to solve a very simple classificatio problem.		
	x = [0 0 1 1; 0 1 0 1	];	

```
t = [0 1 1 1];
net = perceptron;
net = train(net,x,t);
view(net)
y = net(x);
```

#### See Also preparets | removedelay | timedelaynet | narnet | narxnet

Calculate network performance	
perform(net,t,y,ew)	
<pre>perform(net,t,y,ew) takes these arguments,</pre>	
net	Neural network
t	Target data
У	Output data
ew	Error weights (default = {1})
	perform(net,t,y,ev perform(net,t,y,ev net t y

and returns network performance calculated according to the net.performFcn and net.performParam property values.

The target and output data must have the same dimensions. The error weights may be the same dimensions as the targets, in the most general case, but may also have any of its dimension be 1. This gives the flexibility of defining error weights across any dimension desired.

Error weights can be defined by sample, output element, time step, or network output:

ew = [1.0 0.5 0.7 0.2]; % Across 4 samples ew = [0.1; 0.5; 1.0]; % Across 3 elements ew = {0.1 0.2 0.3 0.5 1.0}; % Across 5 timesteps ew = {1.0; 0.5}; % Across 2 outputs

The may also be defined across any combination, such as across two time-series (i.e. two samples) over four timesteps.

ew = {[0.5 0.4],[0.3 0.5],[1.0 1.0],[0.7 0.5]};

In the general case, error weights may have exactly the same dimensions as targets, in which case each target value will have an associated error weight.

# perform

	The default error weight treats all errors the same. ew = {1}
Examples	Here a simple fitting problem is solved with a feed-forward network and its performance calculated.
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); y = net(x); perf = perform(net,t,y)</pre>
See Also	train   configure   init

Purpose	Plot classification confusion matrix
Syntax	<pre>plotconfusion(targets,outputs) plotconfusion(targets1,outputs1,'name1',)</pre>
Description	<pre>plotconfusion(targets,outputs) displays the classification confusion grid.</pre>
	<pre>plotconfusion(targets1,outputs1, 'name1',) displays a series of plots.</pre>
Examples	<pre>load simpleclass_dataset net = patternnet(20); net = train(net,simpleclassInputs,simpleclassTargets); simpleclassOutputs = sim(net,simpleclassInputs); plotconfusion(simpleclassTargets,simpleclassOutputs);</pre>

# plotep

Purpose	Plot weight-bias position on error surface	
Syntax	H= plotep(W,B,E) H = plotep(W,B,E,	Н)
Description	plotep is used to show network learning on a plot already created by plotes.	
	H= plotep(W,B,E)	takes these arguments,
	W	Current weight value
	В	Current bias value
	E	Current error
	<ul> <li>and returns a vector H, containing information for continuing the plot.</li> <li>H = plotep(W,B,E,H) continues plotting using the vector H returned by the last call to plotep.</li> <li>H contains handles to dots plotted on the error surface, so they can be deleted next time, as well as points on the error contour, so they can be connected.</li> </ul>	
See Also	errsurf   plotes	

Purpose	Plot autocorrelation of error time series
Syntax	ploterrcorr(error) ploterrcorr(errors,'outputIndex',outIdx)
Description	<pre>ploterrcorr(error) takes an error time series and plots the autocorrelation of errors across varying lags. ploterrcorr(errors, 'outputIndex',outIdx) uses the optional property name/value pair to define which output error autocorrelation is plotted. The default is 1.</pre>
Examples	<pre>Here a NARX network is used to solve a time series problem. [X,T] = simplenarx_dataset; net = narxnet(1:2,20); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts,Xi,Ai); Y = net(Xs,Xi,Ai); E = gsubtract(Ts,Y); ploterrcorr(E)</pre>
See Also	plotinerrcorr   plotresponse

# ploterrhist

Purpose	Plot error histogram
Syntax	ploterrhist(e) ploterrhist(e1,'name1',e2,'name2',) ploterrhist(,'bins',bins)
Description	ploterrhist(e) plots a histogram of error values e.
	ploterrhist(e1, 'name1',e2, 'name2',) takes any number of errors and names and plots each pair.
	ploterrhist(, 'bins',bins) takes an optional property name/value pair which defines the number of bins to use in the histogram plot. The default is 20.
Examples	Here a feedforward network is used to solve a simple fitting problem:
	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); y = net(x); e = t - y; ploterrhist(e,'bins',30)</pre>
See Also	plotconfusion   ploterrcorr   plotinerrcorr

Purpose	Plot error surface of single-input neuron		
Syntax	<pre>plotes(WV,BV,ES,V)</pre>		
Description	<pre>plotes(WV,BV,ES,V) takes these arguments,</pre>		
	WV	1-by-N row vector of values of W	
	BV	1-by-M row vector of values of $B$	
	ES	M-by-N matrix of error vectors	
	V	View (default = $[-37.5, 30]$ )	
	-	surface with a contour underneath. surface ES with errsurf.	
Examples	<pre>p = [3 2]; t = [0.4 0.8]; wv = -4:0.4:4; bv = wv; ES = errsurf(p,t,wv,bv,'logsig'); plotes(wv,bv,ES,[60 30])</pre>		
See Also	errsurf		

# <u>p</u>lotfit

Purpose	Plot function fit
Syntax	plotfit(NET,INPUTS,TARGETS) plotfit(targets1,inputs1,'name1',)
Description	plotfit(NET, INPUTS, TARGETS) plots the output function of a network across the range of the inputs INPUTS and also plots target TARGETS and output data points associated with values in INPUTS. Error bars show the difference between outputs and INPUTS.
	The plot appears only for networks with one input.
	Only the first output/targets appear if the network has more than one output.
	plotfit(targets1,inputs1,'name1',) displays a series of plots.
Examples	<pre>load simplefit_dataset net = fitnet(20); [net,tr] = train(net,simplefitInputs,simplefitTargets); plotfit(net,simplefitInputs,simplefitTargets);</pre>
See Also	plottrainstate

Purpose	Plot input to error time-series cross correlation	
Syntax	plotinerrcorr(x,e) plotinerrcorr(,'inputIndex',inputIndex) plotinerrcorr(,'outputIndex',outputIndex)	
Description	plotinerrcorr(x,e) takes an input time series x and an error time series e, and plots the autocorrelation of inputs to errors across varying lags.	
	plotinerrcorr(,'inputIndex',inputIndex) optionally defines which input element is being correlated and plotted. The default is 1.	
	plotinerrcorr(, 'outputIndex', outputIndex) optionally defines which error element is being correlated and plotted. The default is 1.	
Examples	Here a NARX network is used to solve a time series problem.	
	<pre>[X,T] = simplenarx_dataset; net = narxnet(1:2,20); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts,Xi,Ai); Y = net(Xs,Xi,Ai); E = gsubtract(Ts,Y); ploterrcorr(E) plotinerrcorr(Xs,E)</pre>	
See Also	ploterrcorr   plotresponse   ploterrhist	

# plotpc

Purpose	Plot classification line on perceptron vector plot		
Syntax	plotpc(W,B) plotpc(W,B,H)		
Description	plotpc(W,B) takes these inputs,		
	W	S-by-R weight matrix (R must be 3 or less)	
	В	S-by-1 bias vector	
	and returns a h	andle to a plotted classification line.	
	plotpc(W,B,H)	takes an additional input,	
	Н	Handle to last plotted line	
	and deletes the	last line before plotting the new one.	
	This function do called after plo	bes not change the current axis and is intended to be tpv.	
Examples	The code below defines and plots the inputs and targets for a perceptron:		
<pre>p = [0 0 1 1; 0 1 0 1]; t = [0 0 0 1]; plotpv(p,t)</pre>			
	The following code creates a perceptron with inputs ranging over the values in P, assigns values to its weights and biases, and plots the resulting classification line.		
<pre>net = newp(mi net.iw{1,1} = net.b{1} = 1; plotpc(net.iv</pre>			

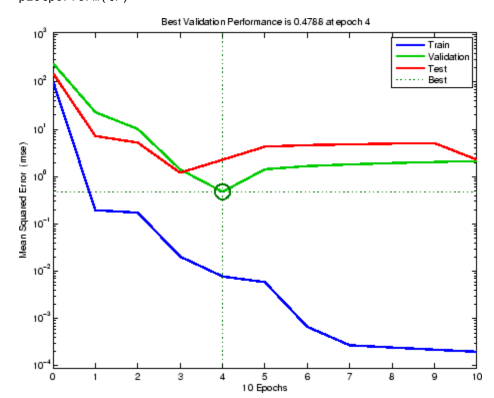
See Also plotpv

#### plotperform

- Purpose Plot network performance
- Syntax plotperform(TR)

**Description** plotperform(TR) plots the training, validation, and test performances given the training record TR returned by the function train.

Examples
 [x,t] = house\_dataset;
 net = feedforwardnet(10);
 [net,tr] = train(net,x,t);
 plotperform(tr)

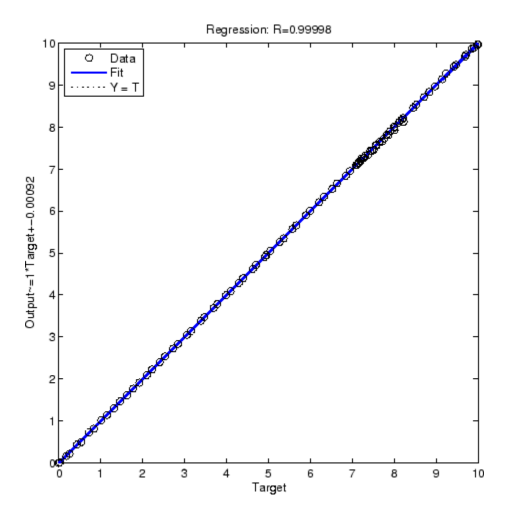


See Also plottrainstate

# <u>plot</u>pv

Purpose	Plot perceptron input/target vectors		
Syntax	plotpv(P,T) plotpv(P,T,V)		
Description	plotpv(P,T) takes these inputs,		
	Р	R-by-Q matrix of input vectors (R must be 3 or less)	
	Т	S-by-Q matrix of binary target vectors (S must be 3 or less)	
	and plots column vectors in P with markers based on T.		
	plotpv(P,T,V)	takes an additional input,	
	V	Graph limits = [x_min x_max y_min y_max]	
	and plots the co	lumn vectors with limits set by V.	
Examples	<b>es</b> The code below defines and plots the inputs and targets for a pe		
	<pre>p = [0 0 1 1; 0 1 0 1]; t = [0 0 0 1]; plotpv(p,t)</pre>		
	The following code creates a perceptron with inputs ranging over the values in P, assigns values to its weights and biases, and plots the resulting classification line.		
	<pre>net = newp(minmax(p),1); net.iw{1,1} = [-1.2 -0.5]; net.b{1} = 1; plotpc(net.iw{1,1},net.b{1})</pre>		
See Also	plotpc		

Purpose	Plot linear regression
Syntax	plotregression(targets,outputs) plotregression(targs1,outs1,'name1',targs2,outs2,'name2',)
Description	plotregression(targets,outputs) plots the linear regression of targets relative to outputs.
	<pre>plotregression(targs1,outs1,'name1',targs2,outs2,'name2',) generates multiple plots.</pre>
Examples	<pre>[x,t] = simplefit_dataset; net = feedforwardnet(10); net = train(net,x,t); y = net(x); plotregression(t,y,'Regression')</pre>



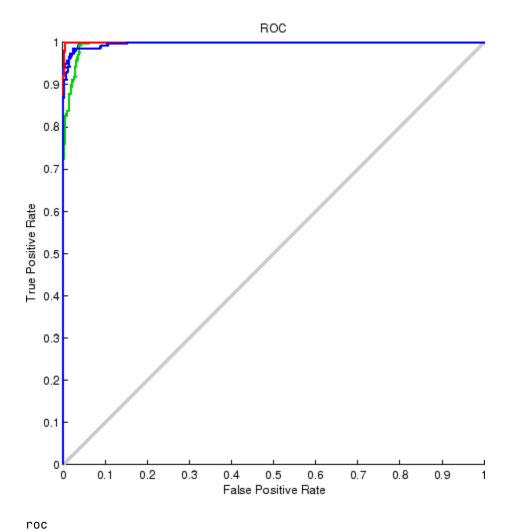


plottrainstate

Purpose	Plot dynamic network time series response
Syntax	plotresponse(t,y) plotresponse(t1,'name',t2,'name2',,y) plotresponse(,'outputIndex',outputIndex)
Description	plotresponse(t,y) takes a target time series t and an output time series y, and plots them on the same axis showing the errors between them.
	plotresponse(t1, 'name',t2, 'name2',,y) takes multiple target/name pairs, typically defining training, validation and testing targets, and the output. It plots the responses with colors indicating the different target sets.
	plotresponse(, outputIndex', outputIndex) optionally defines which error element is being correlated and plotted. The default is 1.
Examples	Here a NARX network is used to solve a time series problem.
	<pre>[X,T] = simplenarx_dataset; net = narxnet(1:2,20); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts,Xi,Ai); Y = net(Xs,Xi,Ai); plotresponse(Ts,Y)</pre>
See Also	ploterrcorr   plotinerrcorr   ploterrhist

# plotroc

Purpose	Plot receiver operating characteristic
Syntax	<pre>plotroc(targets,outputs) plotroc(targets1,outputs2,'name1',)</pre>
Description	plotroc(targets,outputs) plots the receiver operating characteristic for each output class. The more each curve hugs the left and top edges of the plot, the better the classification.
	<pre>plotroc(targets1,outputs2, 'name1',) generates multiple plots.</pre>
Examples	<pre>load simplecluster_dataset net = patternnet(20); net = train(net,simpleclusterInputs,simpleclusterTargets); simpleclusterOutputs = sim(net,simpleclusterInputs); plotroc(simpleclusterTargets,simpleclusterOutputs);</pre>



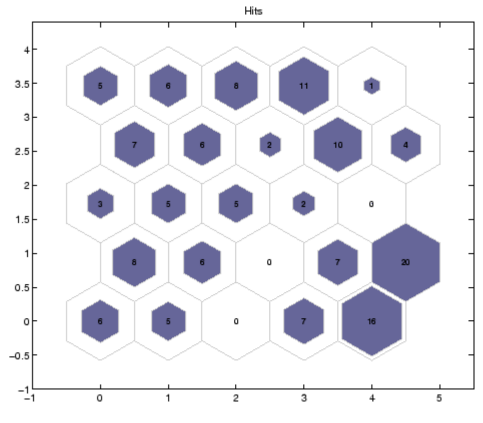
See Also

# plotsom

Purpose	Plot self-organizing map		
Syntax	plotsom(pos) plotsom(W,D,ND)		
Description	plotsom(pos) takes one argument,		
	POS	N-by-S matrix of S N-dimension neural positions	
	and plots the neuron positions with red dots, linking the neurons within a Euclidean distance of 1. plotsom(W,D,ND) takes three arguments,		
	W	S-by-R weight matrix	
	D	S-by-S distance matrix	
	ND	Neighborhood distance (default = $1$ )	
	and plots the neuron's weight vectors with connections between weight vectors whose neurons are within a distance of 1.		
Examples	<pre>This code generates plots of various layer topologies. pos = hextop(5,6); plotsom(pos) pos = gridtop(4,5); plotsom(pos) pos = randtop(18,12); plotsom(pos) pos = gridtop(4,5,2); plotsom(pos) pos = hextop(4,4,3); plotsom(pos) See newsom for an example of plotting a layer's weight vectors with the input vectors they map.</pre>		
See Also	initsompc   learnsom		

Purpose	Plot self-organizing map sample hits
Syntax	plotsomhits(net,inputs)
Description	plotsomhits(net,inputs) plots a SOM layer, with each neuron showing the number of input vectors that it classifies. The relative number of vectors for each neuron is shown via the size of a colored patch.
Examples	<pre>x = iris_dataset; net = selforgmap([5 5]); net = train(net,x); plotsomhits(net,x);</pre>

# plotsomhits



See Also

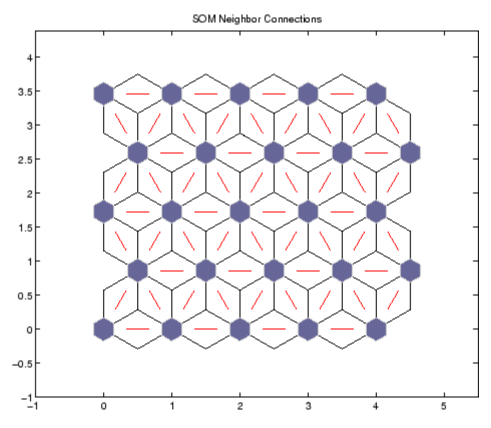
plotsomplanes

#### plotsomnc

- **Purpose** Plot self-organizing map neighbor connections
- Syntax plotsomnc(net)

**Description** plotsomnc(net) plots a SOM layer showing neurons as gray-blue patches and their direct neighbor relations with red lines.

Examples x = iris\_dataset; net = selforgmap([8 8]); net = train(net,x); plotsomnc(net)



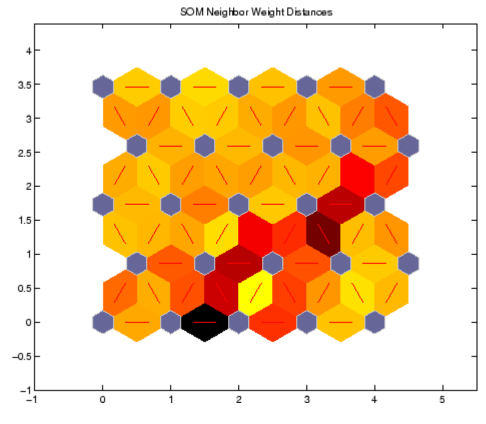
#### plotsomnc

See Also plotsomnd | plotsomplanes | plotsomhits

#### plotsomnd

Purpose	Plot self-organizing map neighbor distances
Syntax	plotsomnd(net)
Description	plotsomnd(net) plots a SOM layer showing neurons as gray-blue patches and their direct neighbor relations with red lines. The neighbor patches are colored from black to yellow to show how close each neuron's weight vector is to its neighbors.
Examples	<pre>x = iris_dataset; net = selforgmap([5 5]); net = train(net,x); plotsomnd(net);</pre>

## plotsomnd

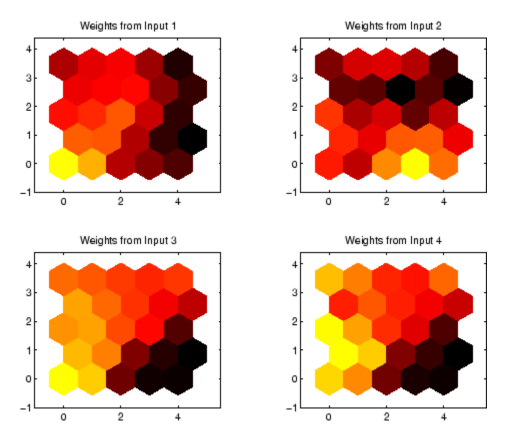


See Also

plotsomhits | plotsomnc | plotsomplanes

Purpose	Plot self-organizing map weight planes		
Syntax	plotsomplanes(net)		
Description	plotsomplanes(net) generates a set of subplots. Each ith subplot shows the weights from the ith input to the layer's neurons, with the most negative connections shown as blue, zero connections as black, and the strongest positive connections as red.		
	The plot is only shown for layers organized in one or two dimensions.		
	This function can also be called with standardized plotting function arguments used by the function train.		
Examples	<pre>x = iris_dataset; net = selforgmap([5 5]); net = train(net,x); plotsomplanes(net)</pre>		

### plotsomplanes

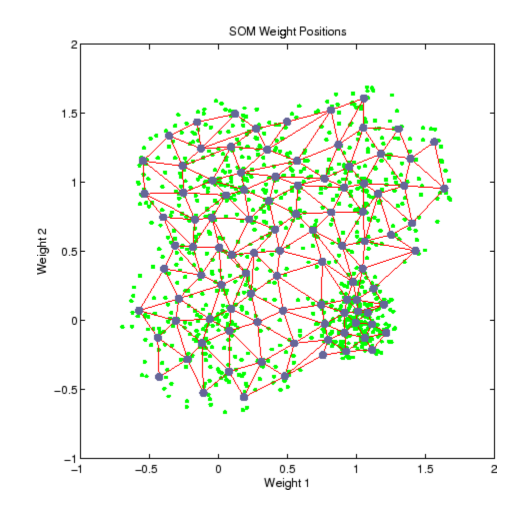


See Also

plotsomhits | plotsomnc | plotsomnd

Purpose	Plot self-organizing map weight positions
Syntax	plotsompos(net) plotsompos(net,inputs)
Description	plotsompos(net) plots the input vectors as green dots and shows how the SOM classifies the input space by showing blue-gray dots for each neuron's weight vector and connecting neighboring neurons with red lines.
	plotsompos(net, inputs) plots the input data alongside the weights.
Examples	<pre>x = iris_dataset; net = selforgmap([10 10]); net = train(net,x); plotsompos(net,x)</pre>

### plotsompos



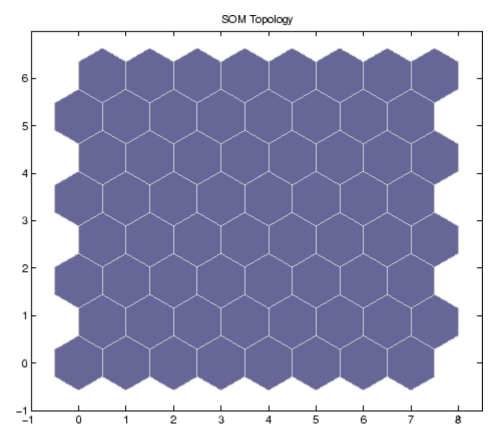


plotsomnd | plotsomplanes | plotsomhits

#### plotsomtop

- Purpose
   Plot self-organizing map topology
- Syntax plotsomtop(net)
- **Description** plotsomtop(net) plots the topology of a SOM layer.

Examples x = iris\_dataset; net = selforgmap([8 8]); plotsomtop(net);



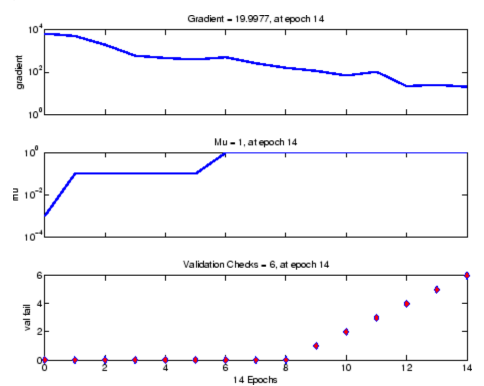
See Also plotsomnd | plotsomplanes | plotsomhits

#### plottrainstate

- Purpose Plot training state values
- **Syntax** plottrainstate(tr)

**Description** plottrainstate(tr) plots the training state from a training record tr returned by train.

Examples
 [x,t] = house\_dataset;
 net = feedforwardnet(10);
 [net,tr] = train(net,x,t);
 plottrainstate(tr)



# plottrainstate

See Also plotfit | plotperform | plotregression

Purpose	Plot vectors as lines from origin		
Syntax	plotv(M,T)		
Description	plotv(M,T) takes two inputs,		
	М	$R\text{-}\mathrm{by-}Q$ matrix of $Q$ column vectors with $R$ elements	
	т	The line plotting type (optional; default = '-')	
	and plots the column vectors of M. R must be 2 or greater. If R is greater than 2, only the first two rows of M are used for the plot.		
Examples	plotv([4 0.7 .2; -0.5 .1 0.5],'-')		

# plotvec

Purpose	Plot vectors with different colors	
Syntax	<pre>plotvec(X,C,M)</pre>	
Description	plotvec(X,C,M) takes these inputs,	
	х	Matrix of (column) vectors
	С	Row vector of color coordinates
	Μ	Marker (default = '+')
	and plots each ith vector in $X$ with a marker $M,$ using the ith value in ${\bf C}$ as the color coordinate.	
	plotvec(X) only takes a matrix X and plots each ith vector in X with marker '+' using the index i as the color coordinate.	
Examples	<pre>x = [0 1 0.5 0.7; -1 2 0.5 0.1]; c = [1 2 3 4]; plotvec(x,c)</pre>	

Purpose	Plot Hinton diagram of weight and bias values
Syntax	<pre>plotwb(net) plotwb(IW,LW,B) plotwb(,'toLayers',toLayers) plotwb(,'fromInputs',fromInputs) plotwb(,'fromLayers',fromLayers) plotwb(,'root',root)</pre>
Description	plotwb(net) takes a neural network and plots all its weights and biases.
	plotwb(IW,LW,B) takes a neural networks input weights, layer weights and biases and plots them.
	plotwb(,'toLayers',toLayers) optionally defines which destination layers whose input weights, layer weights and biases will be plotted.
	plotwb(, 'fromInputs', fromInputs) optionally defines which inputs will have their weights plotted.
	plotwb(,'fromLayers',fromLayers) optionally defines which layers will have weights coming from them plotted.
	<pre>plotwb(, 'root', root) optionally defines the root used to scale the weight/bias patch sizes. The default is 2, which makes the 2-dimensional patch sizes scale directly with absolute weight and bias sizes. Larger values of root magnify the relative patch sizes of smaller weights and biases, making differences in smaller values easier to see.</pre>
Examples	Here a cascade-forward network is configured for particular data and its weights and biases are plotted in several ways.
	<pre>[x,t] = simplefit_dataset; net = cascadeforwardnet([15 5]); net = configure(net,x,t); plotwb(net) plotwb(net,'root',3)</pre>

### plotwb

```
plotwb(net,'root',4)
plotwb(net,'toLayers',2)
plotwb(net,'fromLayers',1)
plotwb(net,'toLayers',2,'fromInputs',1)
```

#### See Also plotsomplanes

Purpose	Pseudonormalize columns of matrix		
Syntax	pnormc(X,R)		
Description	pnormc(X,R) takes these arguments,		
	х	M-by-N matrix	
	R	(Optional) radius to normalize columns to (default = 1)	
	and returns X with an additional row of elements, which results in new column vector lengths of R.		
	<b>Caution</b> For this function to work properly, the columns of X must originally have vector lengths less than R.		
Examples	x = [0.1 0.6; y = pnormc(x)	0.3 0.1];	
See Also	normc   normr		

# <u>p</u>oslin

Purpose	Positive linear transfer function		
Graph and Symbol	a +1: 0 1 -1 a = poslin(n) Positive Linear T	$\rightarrow n$ $\square$	
Syntax	A = poslin(N, info = poslin		
Description	<pre>poslin is a neural transfer function. Transfer functions calculate a layer's output from its net input. A = poslin(N,FP) takes N and optional function parameters,</pre>		
	Ν	S-by-Q matrix of net input (column) vectors	
	FP	Struct of function parameters (ignored)	
	and returns A, the S-by-Q matrix of N's elements clipped to [O, inf].		
	<pre>info = poslin('code') returns information about this function. The following codes are supported:</pre>		
	poslin('name') returns the name of this function. poslin('output',FP) returns the [min max] output range. poslin('active',FP) returns the [min max] active range.		
	poslin('fullderiv') returns 1 or 0, depending on whether dA_dN is S-by-S-by-Q or S-by-Q.		
	poslin('fpnam	es') returns the names of the function parameters.	
	poslin('fpdefaults') returns the default function parameters.		

Examples	Here is the code to create a plot of the poslin transfer function.		
	n = -5:0.1:5; a = poslin(n); plot(n,a)		
	Assign this transfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'poslin';</pre>		
Network Use	To change a network so that a layer uses poslin, set net.layers{i}.transferFcn to 'poslin'.		
	Call sim to simulate the network with poslin.		
Algorithms	The transfer function <b>poslin</b> returns the output <b>n</b> if <b>n</b> is greater than or equal to zero and 0 if <b>n</b> is less than or equal to zero.		
	poslin(n) = n, if n >= 0 = 0, if n <= 0		
See Also	sim   purelin   satlin   satlins		

#### preparets

Purpose	Prepare input and target time series data for network simulation or training	
Syntax	[Xs,Xi,Ai,Ts,EWs,shift] = preparets(net,Xnf,Tnf,Tf,EW)	
Description	<pre>This function simplifies the normally complex and error prone task of reformatting input and target time series. It automatically shifts input and target time series as many steps as are needed to fill the initial input and layer delay states. If the network has open loop feedback, then it copies feedback targets into the inputs as needed to define the open loop inputs.</pre> Each time a new network is designed, with different numbers of delays or feedback settings, preparets can be called to reformat input and target data accordingly. Also, each time a network is transformed with openloop, closeloop, removedelay or adddelay, this function can reformat the data accordingly. [Xs,Xi,Ai,Ts,EWs,shift] = preparets(net,Xnf,Tnf,Tf,EW) takes these arguments,	
	net	Neural network
	Xnf	Non-feedback inputs
	Tnf	Non-feedback targets
	Tf	Feedback targets
	EW	Error weights (default = $\{1\}$ )
	and returns,	
	Xs	Shifted inputs
	Xi	Initial input delay states
	Ai	Initial layer delay states
	Ts	Shifted targets

	EWs	Shifted error weights	
	shift	The number of timesteps truncated from the front of X and T in order to properly fill Xi and Ai.	
Examples	Here a time-delay network with 20 hidden neurons is created, trained and simulated.		
	<pre>net = timedelaynet(20); view(net) [X,T] = simpleseries_dataset; [Xs,Xi,Ai,Ts] = preparets(net,X,T); net = train(net,Xs,Ts); Y = net(Xs,Xi,Ai)</pre>		
	Here a NARX network is designed. The NARX network has a standard input and an open-loop feedback output to an associated feedback input.		
	<pre>[X,T] = simplenarx net = narxnet(1:2, view(net) [Xs,Xi,Ai,Ts] = pr net = train(net,Xs y = net(Xs,Xi,Ai);</pre>	1:2,20); reparets(net,X,{},T); s,Ts,Xi,Ai);	
	Now the network is converted to closed loop, and the data is reformatted to simulate the network's closed-loop response.		
	<pre>net = closeloop(ne view(net) [Xs,Xi,Ai] = prepa y = net(Xs,Xi,Ai);</pre>	arets(net,X,{},T);	
See Also	adddelay   closeloc   timedelaynet	op   narnet   narxnet   openloop   removedelay	

Purpose	Process columns of matrix with principal component analysis	
Syntax	[Y,PS] = processp [Y,PS] = processp Y = processpca('a X = processpca('r name = processpca fp = processpca(' names = processpc processpca('pchec	ca(X,FP) pply',X,PS) everse',Y,PS) ('name') pdefaults') a('pdesc')
Description	processpca processes matrices using principal component analysis so that each row is uncorrelated, the rows are in the order of the amount they contribute to total variation, and rows whose contribution to total variation are less than maxfrac are removed.	
	[Y,PS] = processpca(X,maxfrac) takes X and an optional param	
	Х	N-by-Q matrix or a 1-by-TS row cell array of N-by-Q matrices
	maxfrac	Maximum fraction of variance for removed rows (default is 0)
	and returns	
	Υ	Each N-by-Q matrix with N - M rows deleted (optional)
	PS	Process settings that allow consistent processing of values
	[Y,PS] = processpca(X,FP) takes parameters as a struct: FP.maxfrac. Y = processpca('apply',X,PS) returns Y, given X and settings PS.	

	X = processpca('reverse',Y,PS) returns X, given Y and settings PS.
	<pre>name = processpca('name') returns the name of this process method.</pre>
	<pre>fp = processpca('pdefaults') returns default process parameter structure.</pre>
	<pre>names = processpca('pdesc') returns the process parameter descriptions.</pre>
	processpca('pcheck',fp); throws an error if any parameter is illegal.
Examples	Here is how to format a matrix with an independent row, a correlated row, and a completely redundant row so that its rows are uncorrelated and the redundant row is dropped.
	<pre>x1_independent = rand(1,5) x1_correlated = rand(1,5) + x_independent; x1_redundant = x_independent + x_correlated x1 = [x1_independent; x1_correlated; x1_redundant] [y1,ps] = processpca(x1)</pre>
	Next, apply the same processing settings to new values.
	<pre>x2_independent = rand(1,5) x2_correlated = rand(1,5) + x_independent; x2_redundant = x_independent + x_correlated x2 = [x2_independent; x2_correlated; x2_redundant]; y2 = processpca('apply',x2,ps)</pre>
	Reverse the processing of y1 to get x1 again.
	x1_again = processpca('reverse',y1,ps)
Algorithms	Values in rows whose elements are not all the same value are set to
	y = 2*(x-minx)/(maxx-minx) - 1;
	Values in rows with all the same value are set to 0.

#### Definitions

In some situations, the dimension of the input vector is large, but the components of the vectors are highly correlated (redundant). It is useful in this situation to reduce the dimension of the input vectors. An effective procedure for performing this operation is principal component analysis. This technique has three effects: it orthogonalizes the components of the input vectors (so that they are uncorrelated with each other), it orders the resulting orthogonal components (principal components) so that those with the largest variation come first, and it eliminates those components that contribute the least to the variation in the data set. The following code illustrates the use of processpca, which performs a principal-component analysis using the processing setting maxfrac of 0.02.

[pn,ps1] = mapstd(p); [ptrans,ps2] = processpca(pn,0.02);

The input vectors are first normalized, using mapstd, so that they have zero mean and unity variance. This is a standard procedure when using principal components. In this example, the second argument passed to processpca is 0.02. This means that processpca eliminates those principal components that contribute less than 2% to the total variation in the data set. The matrix ptrans contains the transformed input vectors. The settings structure ps2 contains the principal component transformation matrix. After the network has been trained, these settings should be used to transform any future inputs that are applied to the network. It effectively becomes a part of the network, just like the network weights and biases. If you multiply the normalized input vectors pn by the transformation matrix transMat, you obtain the transformed input vectors ptrans.

If processpca is used to preprocess the training set data, then whenever the trained network is used with new inputs, you should preprocess them with the transformation matrix that was computed for the training set, using ps2. The following code applies a new set of inputs to a network already trained.

```
pnewn = mapstd('apply',pnew,ps1);
pnewtrans = processpca('apply',pnewn,ps2);
```

a = sim(net,pnewtrans);

Principal component analysis is not reliably reversible. Therefore it is only recommended for input processing. Outputs require reversible processing functions.

Principal component analysis is not part of the default processing for feedforwardnet. If you wish to add this, you can use the following command:

net.inputs{1}.processFcns{end+1} = 'processpca';

See Also fixunknowns | mapminmax | mapstd

#### prune

Purpose	Delete neural inputs, layers, and outputs with sizes of zero			
Syntax	[net,pi,pl,po] = prune(net)			
Description	This function removes zero-sized inputs, layers, and outputs from a network. This leaves a network which may have fewer inputs and outputs, but which implements the same operations, as zero-sized inputs and outputs do not convey any information.			
		plification is to prepare a network with zero sized ink, where zero sized signals are not supported.		
	The companion func with the transforme	tion prunedata can prune data to remain consistent d network.		
	[net,pi,pl,po] =	prune(net) takes a neural network and returns		
	net	The same network with zero-sized subobjects removed		
	pi	Indices of pruned inputs		
	pl	Indices of pruned layers		
	ро	Indices of pruned outputs		
Examples	Here a NARX dynamic network is created which has one external input and a second input which feeds back from the output.			
	<pre>net = narxnet(20); view(net) The network is then trained on a single random time-series problem with 50 timesteps. The external input happens to have no elements.</pre>			
	X = nndata(0,1,50); T = nndata(1,1,50); [Xs,Xi,Ai,Ts] = preparets(net,X,{},T); net = train(net,Xs,Ts);			

The network and data are then pruned before generating a Simulink diagram and initializing its input and layer states.

```
[net2,pi,pl,po] = prune(net);
view(net)
[Xs2,Xi2,Ai2,Ts2] = prunedata(net,pi,pl,po,Xs,Xi,Ai,Ts)
[sysName,netName] = gensim(net);
setsiminit(sysName,netName,Xi2,Ai2)
```

See Also prunedata | gensim

# prunedata

Purpose	Purpose			
Syntax	[Xp,Xip,Aip,Tp] =	[Xp,Xip,Aip,Tp] = prunedata(pi,pl,po,X,Xi,Ai,T)		
Description	-	es data to be consistent with a network whose ayers, and outputs have been removed with prune.		
		plification is to prepare a network with zero-sized link, where zero-sized signals are not supported.		
	[Xp,Xip,Aip,Tp] = arguments,	= prunedata(pi,pl,po,X,Xi,Ai,T) takes these		
	pi	Indices of pruned inputs		
	pl	Indices of pruned layers		
	ро	Indices of pruned outputs		
	Х	Input data		
	Xi	Initial input delay states		
	Ai Initial layer delay states			
	т	Target data		
	and returns the prus	ned inputs, input and layer delay states, and targets.		
Examples		mic network is created which has one external input which feeds back from the output.		
	<pre>net = narxnet(20); view(net) The network is then trained on a single random time-series problem with 50 timesteps. The external input happens to have no elements. X = nndata(0,1,50); T = nndata(1,1,50);</pre>			

```
[Xs,Xi,Ai,Ts] = preparets(net,X,{},T);
net = train(net,Xs,Ts);
The network and data are then pruned before generating a Simulink
diagram and initializing its input and layer states.
[net2,pi,pl,po] = prune(net);
view(net)
[Xs2,Xi2,Ai2,Ts2] = prunedata(net,pi,pl,po,Xs,Xi,Ai,Ts)
[sysName,netName] = gensim(net);
setsiminit(sysName,netName,Xi2,Ai2)
See Also prune | gensim
```

# purelin

Purpose	Linear transfer function			
Graph and Symbol	a +1 0 -1 a = purclin(n) Linear Transfer	Function		
Syntax	A = purelin(N info = pureli			
Description	•	ural transfer function. Transfer functions calculate a rom its net input.		
	A = purelin(N	,FP) takes N and optional function parameters,		
	Ν	S-by-Q matrix of net input (column) vectors		
	FP	Struct of function parameters (ignored)		
	and returns A, a	an S-by-Q matrix equal to N.		
	<pre>info = purelin('code') returns useful information for each supported code string:</pre>			
	purelin('name	') returns the name of this function.		
	purelin('outp	ut',FP) returns the [min max] output range.		
	purelin('acti	ve', FP) returns the [min max] active input range.		
	purelin('fullderiv') returns 1 or 0, depending on whether dA_dN is S-by-S-by-Q or S-by-Q.			
	purelin('fpna	mes') returns the names of the function parameters.		
	purelin('fpde	faults') returns the default function parameters.		

Examples	Here is the code to create a plot of the purelin transfer function.		
	n = -5:0.1:5; a = purelin(n); plot(n,a)		
	Assign this transfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'purelin';</pre>		
Algorithms	a = purelin(n) = n		
See Also	sim   satlin   satlins		

### quant

Purpose	Discretize values as multiples of quantity		
Syntax	quant(X,Q)		
Description	quant(X,Q) takes two inputs,		
	х	Matrix, vector, or scalar	
	Q	Minimum value	
	and returns val	ues from X rounded to nearest multiple of Q.	
Examples	<pre>x = [1.333 4.756 -3.897]; y = quant(x,0.1)</pre>		

Purpose	Radial basis transfer function		
Graph and Symbol	1.0 0.5 0.0 -0.833 +0.833 a = radbas(n) Radial Basis Fun	n	
Syntax	A = radbas(N,FP)		
Description	radbas is a neural transfer function. Transfer functions calculate a layer's output from its net input.		
	A = radbas(N,FP) takes one or two inputs,		
	Ν	S-by-Q matrix of net input (column) vectors	
	FP	Struct of function parameters (ignored)	
	and returns A, ar to each element o	n S-by-Q matrix of the radial basis function applied of N.	
Examples	Here you create a	a plot of the radbas transfer function.	
	n = -5:0.1:5; a = radbas(n); plot(n,a)		
	Assign this trans	fer function to layer i of a network.	
	net.layers{i}.	transferFcn = 'radbas';	

### radbas

Algorithms	а	=	radbas(n)	=	exp(-n^2)
------------	---	---	-----------	---	-----------

See Also sim | radbasn | tribas

Purpose	Normalized radial basis transfer function		
Graph and Symbol	a = radbas(n)		
Syntax	A = radbasn(N,FP)		
Description	<pre>radbasn is a neural transfer function. Transfer functions calculate a layer's output from its net input. This function is equivalent to radbas, except that output vectors are normalized by dividing by the sum of the pre-normalized values. A = radbasn(N,FP) takes one or two inputs, N S-by-Q matrix of net input (column) vectors</pre>		
Examples	<pre>FP Struct of function parameters (ignored) and returns A, an S-by-Q matrix of the radial basis function applied to each element of N. Here six random 3-element vectors are passed through the radial basis transform and normalized. n = rand(3,6) a = radbasn(n) Assign this transfer function to layer i of a network. net.layers{i}.transferFcn = 'radbasn';</pre>		

### radbasn

Algorithms	$a = radbasn(n) = exp(-n^2) / sum(exp(-n^2))$
See Also	sim   radbas   tribas

#### randnc

Purpose	Normalized column weight initialization function			
Syntax	W = randnc(S,	PR)		
Description	<pre>randnc is a weight initialization function. W = randnc(S,PR) takes two inputs,</pre>			
	S	Number	of rows (neu	cons)
	PR	R-by-2 m	atrix of input	value ranges = [Pmin Pmax]
	and returns an S-by-R random matrix with normalized columns. You can also call this in the form randnc(S,R).			
Examples	A random matrix of four normalized three-element columns is generated:			
		-0.4715 -0.6967	-0.2724 -0.9172 -0.2907	0.7819
See Also	randnr	0.3400	-0.2307	0.2747

#### randnr

Purpose	Normalized row weight initialization function			
Syntax	W = randnr(S,	PR)		
Description	<pre>randnr is a weight initialization function. W = randnr(S,PR) takes two inputs,</pre>			
	S	Number of rows (ne	urons)	
	PR	R-by-2 matrix of inp	ut value ranges = [Pmin Pmax]	
	and returns an S-by-R random matrix with normalized rows. You can also call this in the form randnr(S,R).			
Examples	A matrix of three normalized four-element rows is generated:			
			0.5381	
See Also	randnc			

Purpose	Symmetric rand	om weight/bias initialization function
Syntax	W = rands(S,PR) M = rands(S,R) v = rands(S)	
Description	rands is a weigh	nt/bias initialization function.
	W = rands(S,PH	3) takes
	S	Number of neurons
	PR	R-by-2 matrix of R input ranges
	M = rands(S,R)	S-by-R weight matrix of random values between –1 and 1. ) returns an S-by-R matrix of random values. v = ns an S-by-1 vector of random values.
Examples	Here, three sets	of random values are generated with rands.
	rands(4,[0 1; rands(4) rands(2,3)	-2 2])
Network Use	To prepare the v be initialized wi	weights and the bias of layer i of a custom network to th rands,
	Set net.initFcn to 'initlay'. (net.initParam automatically becomes initlay's default parameters.)	
	<ol> <li>2 Set net.layers{i}.initFcn to 'initwb'.</li> <li>3 Set each net.inputWeights{i,j}.initFcn to 'rands'.</li> <li>4 Set each net.layerWeights{i,j}.initFcn to 'rands'.</li> <li>5 Set each net.biases{i}.initFcn to 'rands'.</li> </ol>	

### rands

To initialize the network, call init.

See Also randsmall | randnr | randnc | initwb | initlay | init

Purpose	Small random weight/bias initialization function	
Syntax	<pre>W = randsmall(S,PR) M = rands(S,R) v = rands(S)</pre>	
Description	randsmall is a	weight/bias initialization function.
	W = randsmall	(S,PR) takes
	S	Number of neurons
	PR	R-by-2 matrix of R input ranges
	and returns an –0.1 and 0.1.	S-by-R weight matrix of small random values between
	M = rands(S,R) returns an S-by-R matrix of random values. v = rands(S) returns an S-by-1 vector of random values.	
Examples	Here three sets	of random values are generated with rands.
	randsmall(4,[ randsmall(4) randsmall(2,3	
Network Use	To prepare the be initialized with	weights and the bias of layer i of a custom network to ith rands,
		Fcn to 'initlay'. (net.initParam automatically tlay's default parameters.)
	<b>2</b> Set net.layers{i}.initFcn to 'initwb'.	
	3 Set each net	.inputWeights{i,j}.initFcn to 'randsmall'.
	4 Set each net	layerWeights{i,j}.initFcn to 'randsmall'.

## randsmall

5	Set each	net.biase	es{i}.initFcr	nto ' <mark>rands</mark> m	all'.

To initialize the network, call init.

See Also rands | randnr | randnc | initwb | initlay | init

Purpose	Random layer topology function	
Syntax	<pre>pos = randtop(dim1,dim2,,dimN)</pre>	
Description	randtop calculates the neuron positions for layers whose neurons are arranged in an N-dimensional random pattern.	
	<pre>pos = randtop(dim1,dim2,,dimN) takes N arguments,</pre>	
	dimi Length of layer in dimension i	
	and returns an N-by-S matrix of N coordinate vectors, where S is the product of dim1*dim2**dimN.	
Examples	This code creates and displays a two-dimensional layer with neurons arranged in a random pattern.	
	<pre>pos = randtop(8,5); net = selforgmap([8 5],'topologyFcn','randtop'); plotsomtop(net)</pre>	
See Also	gridtop   hextop   tritop	

# regression

Purpose	Linear regression	
Syntax	[r,m,b] = regression(t,y) [r,m,b] = regression(t,y,'one')	
Description	[r,m,b] = regress	sion(t,y) takes these arguments,
	t	Target matrix or cell array data with a total of N matrix rows
	У	Output matrix or cell array data of the same size
	and returns these outputs, r Regression values for each of the N matrix rows	
	m	Slope of regression fit for each of the N matrix rows $% \left( {{{\rm{N}}_{\rm{N}}}} \right)$
	b	Offset of regression fit for each of the $N\ matrix$ rows
		sion(t,y,'one') combines all matrix rows before g single scalar regression, slope and offset values.
Examples	<pre>Here a feedforward network is trained and regression performed on its targets and outputs. [x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); y = net(x); [r,m,b] = regression(t,y) plotregression(t,y)</pre>	
See Also	plotregression   confusion	

Purpose	Process matrices by removing rows with constant values		
Syntax	<pre>[Y,PS] = removeconstantrows(X,max_range) [Y,PS] = removeconstantrows(X,FP) Y = removeconstantrows('apply',X,PS) X = removeconstantrows('reverse',Y,PS)</pre>		
Description	<pre>removeconstantrows processes matrices by removing rows with constant values. [Y,PS] = removeconstantrows(X,max_range) takes X and an optional parameter, X Single N-by-Q matrix or a 1-by-TS row cell array of N-by-Q matrices</pre>		
	max_range	Maximum range of values for row to be removed (default is 0)	
	and returns		
	Y	Each M-by-Q matrix with N - M rows deleted (optional)	
	PS	Process settings that allow consistent processing of values	
	<pre>[Y,PS] = removeconstantrows(X,FP) takes parameters as a struct: FP.max_range. Y = removeconstantrows('apply',X,PS) returns Y, given X and settings PS. X = removeconstantrows('reverse',Y,PS) returns X, given Y and settings PS.</pre>		

#### removeconstantrows

Examples	Here is how to format a matrix so that the rows with constant values are removed.	
	x1 = [1 2 4; 1 1 1; 3 2 2; 0 0 0] [y1,PS] = removeconstantrows(x1)	
	Next, apply the same processing settings to new values.	
	x2 = [5 2 3; 1 1 1; 6 7 3; 0 0 0] y2 = removeconstantrows('apply',x2,PS)	
	Reverse the processing of $y1$ to get $x1$ again.	
	<pre>x1_again = removeconstantrows('reverse',y1,PS)</pre>	
See Also	fixunknowns   mapminmax   mapstd   processpca	

Purpose	Remove delay to neural network's response		
Syntax	<pre>net = removedelay(net,n)</pre>		
Description	<pre>net = removedelay(net,n) takes these arguments,</pre>		
	net Neural network		
	n Number of delays		
	and returns the network with input delay connections decreased, and output feedback delays increased, by the specified number of delays n. The result is a network which behaves identically, except that outputs are produced n timesteps later.		
	If the number of delays n is not specified, a default of one delay is used.		
Examples	Here a time delay network is created, trained and simulated in its original form on an input time series X and target series T. It is then with a delay removed and then added back. These first and third outputs will be identical, while the second will be shifted by one timestep.		
	<pre>[X,T] = simpleseries_dataset; net = timedelaynet(1:2,20); [Xs,Xi,Ai,Ts] = preparets(net,X,T); net = train(net,Xs,Ts,Xi); y1 = net(Xs) net2 = removedelay(net); [Xs,Xi,Ai,Ts] = preparets(net2,X,T); y2 = net2(Xs,Xi) net3 = adddelay(net2) [Xs,Xi,Ai,Ts] = preparets(net3,X,T); y3 = net3(Xs,Xi)</pre>		
See Also	adddelay   closeloop   openloop		

#### removerows

Purpose	Process matrices by removing rows with specified indices	
Syntax	<pre>[Y,PS] = removerows(X,'ind',ind) [Y,PS] = removerows(X,FP) Y = removerows('apply',X,PS) X = removerows('reverse',Y,PS) dx_dy = removerows('dx',X,Y,PS) dx_dy = removerows('dx',X,[],PS) name = removerows('name') fp = removerows('pdefaults') names = removerows('pdesc') removerows('pcheck',FP)</pre>	
Description	<pre>removerows processes matrices by removing rows with the specified indices. [Y,PS] = removerows(X,'ind',ind) takes X and an optional parameter,</pre>	
	Х	N-by-Q matrix or a 1-by-TS row cell array of N-by-Q matrices
	ind	Vector of row indices to remove (default is [])
	and returns Y Each M-by-Q matrix, where M == N-length (optional)	
	PS	Process settings that allow consistent processing of values
	[Y,PS] = removerows(X,FP) takes parameters as a struct: F	
	Y = removerows('	<code>apply',X,PS)</code> returns Y, given X and settings PS.
	X = removerows('reverse', Y, PS) returns X, given Y and settings PS	

	$dx_dy = removerows('dx',X,Y,PS)$ returns the M-by-N-by-Q derivative of Y with respect to X.		
	<pre>dx_dy = removerows('dx',X,[],PS) returns the derivative, less efficiently.</pre>		
	<pre>name = removerows('name') returns the name of this process method.</pre>		
	<pre>fp = removerows('pdefaults') returns the default process parameter structure.</pre>		
	<pre>names = removerows('pdesc') returns the process parameter descriptions.</pre>		
	removerows('pcheck',FP) throws an error if any parameter is illegal.		
Examples	Here is how to format a matrix so that rows 2 and 4 are removed:		
	x1 = [1 2 4; 1 1 1; 3 2 2; 0 0 0] [y1,ps] = removerows(x1,'ind',[2 4])		
	Next, apply the same processing settings to new values.		
	x2 = [5 2 3; 1 1 1; 6 7 3; 0 0 0] y2 = removerows('apply',x2,ps)		
	Reverse the processing of y1 to get x1 again.		
	x1_again = removerows('reverse',y1,ps)		
Algorithms	In the reverse calculation, the unknown values of replaced rows are represented with NaN values.		
See Also	fixunknowns   mapminmax   mapstd   processpca		

#### revert

Purpose	Change network weights and biases to previous initialization values
Syntax	net = revert (net)
Description	<pre>net = revert (net) returns neural network net with weight and bias values restored to the values generated the last time the network was initialized.</pre>
	If the network is altered so that it has different weight and bias connections or different input or layer sizes, then revert cannot set the weights and biases to their previous values and they are set to zeros instead.
Examples	Here a perceptron is created with input size set to 2 and number of neurons to 1.
	<pre>net = perceptron; net.inputs{1}.size = 2; net.layers{1}.size = 1;</pre>
	The initial network has weights and biases with zero values.
	net.iw{1,1}, net.b{1}
	Change these values as follows:
	net.iw{1,1} = [1 2]; net.b{1} = 5; net.iw{1,1}, net.b{1}
	You can recover the network's initial values as follows:
	<pre>net = revert(net); net.iw{1,1}, net.b{1}</pre>
See Also	init   sim   adapt   train

Purpose	Receiver operating characteristic	
Syntax	[tpr,fpr,threshol	.ds] = roc(targets,outputs)
Description	The <i>receiver operating characteristic</i> is a metric used to check the quality of classifiers. For each class of a classifier, roc applies threshold values across the interval [0,1] to outputs. For each threshold, two values are calculated, the True Positive Ratio (the number of outputs greater or equal to the threshold, divided by the number of one targets), and the False Positive Ratio (the number of outputs less than the threshold, divided by the number of zero targets).	
	You can visualize the results of this function with plotroc.	
	<pre>[tpr,fpr,thresholds] = roc(targets,outputs) takes these arguments:</pre>	
	targets	<ul><li>S-by-Q matrix, where each column vector contains a single 1 value, with all other elements</li><li>O. The index of the 1 indicates which of S categories that vector represents.</li></ul>
	outputs	S-by-Q matrix, where each column contains values in the range [0,1]. The index of the largest element in the column indicates which of S categories that vector presents. Alternately, 1-by-Q vector, where values greater or equal to 0.5 indicate class membership, and values below 0.5, nonmembership.

and returns these values:

	tpr	1-by-S cell array of 1-by-N true-positive/positive ratios.
	fpr	1-by-S cell array of 1-by-N false-positive/negative ratios.
	thresholds	1-by-S cell array of 1-by-N thresholds over interval [0,1].
	roc(targets,outpu	ts) takes these arguments:
	targets	1-by-Q matrix of Boolean values indicating class membership.
	outputs	S-by-Q matrix, of values in [0,1] interval, where values greater than or equal to 0.5 indicate class membership.
	and returns these va	alues:
	tpr	1-by-N vector of true-positive/positive ratios.
	fpr	1-by-N vector of false-positive/negative ratios.
	thresholds	1-by-N vector of thresholds over interval [0,1].
Examples	irisOutputs = sim	20); risInputs,irisTargets);
See Also	plotroc   confusic	on

Purpose	Sum absolute error	performance function
Syntax	[] = sae(,'	regularization',regularization) normalization',normalization) squaredWeighting',squaredWeighting)
Description	<pre>sae is a network performance function. It measures performance according to the sum of squared errors. perf = sae(net,t,y,ew) takes these input arguments and optional function parameters,</pre>	
	net	Neural network
	t	Matrix or cell array of target vectors
	У	Matrix or cell array of output vectors
	ew	Error weights (default = {1})

and returns the sum squared error.

This function has three optional function parameters that can be defined with parameter name/pair arguments, or as a structure FP argument with fields having the parameter name and assigned the parameter values:

[]	<pre>= sae(,'regularization',regularization)</pre>
[]	<pre>= sae(,'normalization',normalization)</pre>
[]	<pre>= sae(,'squaredWeighting',squaredWeighting)</pre>
[]	= sae(,FP)

• regularization — can be set to any value between the default of 0 and 1. The greater the regularization value, the more squared weights and biases are taken into account in the performance calculation.

	<ul> <li>normalization — can be set to the default 'absolute', or 'normalized' (which normalizes errors to the [+2 -2] range consistent with normalized output and target ranges of [-1 1]) or 'percent' (which normalizes errors to the range [-1 +1]).</li> <li>squaredWeighting — can be set to the default false, for applying error weights to absolute errors, or false for applying error weights to the squared errors before squaring.</li> </ul>
Examples	Here a network is trained to fit a simple data set and its performance calculated
	<pre>[x,t] = simplefit_dataset; net = fitnet(10,'trainscg'); net.performFcn = 'sae'; net = train(net,x,t) y = net(x) e = t-y perf = sae(net,t,y)</pre>
Network Use	To prepare a custom network to be trained with sae, set net.performFcn to 'sae'. This automatically sets net.performParam to the default function parameters.
	Then calling train, adapt or perform will result in sae being used to calculate performance.

Purpose	Saturating linear transfer function	
Graph and Symbol	a $-1$ $0$ $+1$ $a = satlin(n)$ Satlin Transfer Function	
Syntax	A = satlin(N,FP)	
Description	<pre>satlin is a neural transfer function. Transfer functions calculate a layer's output from its net input. A = satlin(N,FP) takes one input,</pre>	
	NS-by-Q matrix of net input (column) vectorsFPStruct of function parameters (ignored)	
	and returns A, the S-by-Q matrix of N's elements clipped to $[0, 1]$ .	
	<pre>info = satlin('code') returns useful information for each supported code string:</pre>	
	<pre>satlin('name') returns the name of this function.</pre>	
	<pre>satlin('output',FP) returns the [min max] output range.</pre>	
	<pre>satlin('active',FP) returns the [min max] active input range.</pre>	
	<pre>satlin('fullderiv') returns 1 or 0, depending on whether dA_dN is S-by-S-by-Q or S-by-Q.</pre>	
	<pre>satlin('fpnames') returns the names of the function parameters.</pre>	
	<pre>satlin('fpdefaults') returns the default function parameters.</pre>	

## satlin

Examples	Here is the code to create a plot of the satlin transfer function.	
	n = -5:0.1:5; a = satlin(n); plot(n,a)	
	Assign this transfer function to layer i of a network.	
	<pre>net.layers{i}.transferFcn = 'satlin';</pre>	
Algorithms	a = satlin(n) = 0, if n <= 0 n, if 0 <= n <= 1 1, if 1 <= n	
See Also	sim   poslin   satlins   purelin	

Purpose	Symmetric saturating linear transfer function	
Graph and Symbol	a + 1 $-1$ $a = satlins(n)$ Satlins Transfer Function	
Syntax	A = satlins(N,FP)	
Description	satlins is a neural transfer function. Transfer functions calculate a layer's output from its net input.	
	A = satlins(N,FP) takes N and an optional argument,	
	N S-by-Q matrix of net input (column) vectors	
	FP         Struct of function parameters (optional, ignored)	)
	and returns A, the S-by-Q matrix of N's elements clipped to $[-1, 1]$ .	
	<pre>info = satlins('code') returns useful information for each supported code string:</pre>	
	<pre>satlins('name') returns the name of this function.</pre>	
	<pre>satlins('output',FP) returns the [min max] output range.</pre>	
	<pre>satlins('active',FP) returns the [min max] active input range.</pre>	
	<pre>satlins('fullderiv') returns 1 or 0, depending on whether dA_dN i S-by-S-by-Q or S-by-Q.</pre>	is
	<pre>satlins('fpnames') returns the names of the function parameters.</pre>	
	<pre>satlins('fpdefaults') returns the default function parameters.</pre>	

## satlins

Examples	Here is the code to create a plot of the satlins transfer function.	
	n = -5:0.1:5; a = satlins(n); plot(n,a)	
Algorithms	satlins(n) = -1, if n <= -1 n, if -1 <= n <= 1 1, if 1 <= n	
See Also	sim   satlin   poslin   purelin	

Purpose	Scalar product weight function	
Syntax	Z = scalprod(W,P) dim = scalprod('size',S,R,FP) dw = scalprod('dw',W,P,Z,FP)	
Description	scalprod is the scalar product weight function. Weight functions apply weights to an input to get weighted inputs.	
	Z = scalprod(W,P)	takes these inputs,
	W	1-by-1 weight matrix
	Р	R-by-Q matrix of Q input (column) vectors
	<pre>and returns the R-by-Q scalar product of W and P defined by Z = w*P. dim = scalprod('size',S,R,FP) takes the layer dimension S, input dimension R, and function parameters, and returns the weight size [1-by-1]. dw = scalprod('dw',W,P,Z,FP) returns the derivative of Z with respect to W.</pre>	
Examples	<pre>Here you define a random weight matrix W and input vector P and calculate the corresponding weighted input Z. W = rand(1,1); P = rand(3,1); Z = scalprod(W,P)</pre>	
Network Use	To change a network so an input weight uses scalprod, set net.inputWeight{i,j}.weightFcn to 'scalprod'. For a layer weight, set net.layerWeight{i,j}.weightFcn to 'scalprod'. In either case, call sim to simulate the network with scalprod.	

### scalprod

See help newp and help newlin for simulation examples.

See Also dotprod | sim | dist | negdist | normprod

## selforgmap

Purpose	Self-organizing map		
Syntax	<pre>selforgmap(dimensions,coverSteps,initNeighbor,topologyFcn,</pre>		
Description	Self-organizing maps learn to cluster data based on similarity, topology, with a preference (but no guarantee) of assigning the same number of instances to each class.		
	Self-organizing maps are used both to cluster data and to reduce the dimensionality of data. They are inspired by the sensory and motor mappings in the mammal brain, which also appear to automatically organizing information topologically.		
	<pre>selforgmap(dimensions,coverSteps,initNeighbor,topologyFcn,distanceFcn) takes these arguments,</pre>		
	dimensions	Row vector of dimension sizes (default = [8 8])	
	coverSteps	Number of training steps for initial covering of the input space (default = 100)	
	initNeighbor	Initial neighborhood size (default = 3)	
	topologyFcn	Layer topology function (default = 'hextop')	
	distanceFcn	Neuron distance function (default = 'linkdist')	
	and returns a self-organizing map.		
Examples	Here a self-organizing map is used to cluster a simple set of data.		
	<pre>x = simplecluster_dataset; net = selforgmap([8 8]) net = train(net,x); view(net) y = net(x);</pre>		

### selforgmap

classes = vec2ind(y)

See Also lvqnet | competlayer | nctool

Purpose	Separate biases and weight values from weight/bias vector	
Syntax	[b,IW,LW] = separatewb(net,wb)	
Description	[b,IW,LW] = separatewb(net,wb) takes two arguments,	
	net	Neural network
	wb	Weight/bias vector
	and returns	
	b	Cell array of bias vectors
	IW	Cell array of input weight matrices
	LW	Cell array of layer weight matrices
Examples	<pre>Here a feedforward network is trained to fit some data, then its bias and weight values formed into a vector. The single vector is then redivided into the original biases and weights. [x,t] = simplefit_dataset; net = feedforwardnet(20); net = train(net,x,t); wb = formwb(net,net.b,net.iw,net.lw) [b,iw,lw] = separatewb(net,wb)</pre>	
See Also	getwb   formwb   setwb	

#### seq2con

Purpose	Convert sequential vectors to concurrent vectors		
Syntax	b = seq2con(s)	b = seq2con(s)	
Description	Neural Network Toolbox software represents batches of vectors with a matrix, and sequences of vectors with multiple columns of a cell array.		
	seq2con and con2se sequential vectors, a	eq allow concurrent vectors to be converted to and back again.	
	b = seq2con(s) tal	xes one input,	
	S	N-by-TS cell array of matrices with ${\tt M}$ columns	
	and returns		
	b	N-by-1 cell array of matrices with $M*TS$ columns	
Examples	Here three sequenti	al values are converted to concurrent values.	
	p1 = {1 4 2} p2 = seq2con(p1)		
	Here two sequences concurrent vectors.	of vectors over three time steps are converted to	
	p1 = {[1; 1] [5; 4] [1; 2]; [3; 9] [4; 1] [9; 8]} p2 = seq2con(p1)		
See Also	con2seq   concur		

Purpose	Set neural network data elements		
Syntax	<pre>setelements(x,i,v)</pre>		
Description	<pre>setelements(x,i,v) takes these arguments,</pre>		
	x Neural network matrix or cell array data		
	i	Indices	
	v Neural network data to store into x		
Examples	and returns the original data x with the data v stored in the element indicated by the indices i. This code sets elements 1 and 3 of matrix data: x = [1 2; 3 4; 7 4]		
$v = [10 \ 11; \ 12 \ 13];$			
y = setelements		x,[1 3],V)	
	This code sets elements 1 and 3 of cell array data:		
	<pre>x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} v = {[20 21 22; 23 24 25] [26 27 28; 29 30 31]} y = setelements(x,[1 3],v)</pre>		
See Also	nndata   numeleme setsignals   sett	nts   getelements   catelements   setsamples   imesteps	

### setsamples

Purpose	Set neural network data samples	
Syntax	<pre>setsamples(x,i,v)</pre>	
Description	<pre>setsamples(x,i,v) takes these arguments,</pre>	
	x Neural network matrix or cell array data	
	i	Indices
	v	Neural network data to store into x
	and returns the original data x with the data v stored in the samp indicated by the indices i.	
Examples	$x = [1 \ 2 \ 3; \ 4 \ 7 \ 4]$	
	v = [10 11; 12 13]; y = setsamples(x,[1 3],v)	
	y = 3ec3ampres(x,[1, 0],v)	
	This code sets samples 1 and 3 of cell array data:	
	x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} v = {[20 21; 22 23] [24 25; 26 27]; [28 29] [30 31]} y = setsamples(x,[1 3],v)	
See Also	nndata   numsampl setsignals   sett	es   getsamples   catsamples   setelements   imesteps

Purpose	Set neural network data signals	
Syntax	<pre>setsignals(x,i,v)</pre>	
Description	<pre>setsignals(x,i,v) takes these arguments,</pre>	
	× Neural network matrix or cell array data	
	i	Indices
	v	Neural network data to store into x
	<pre>and returns the original data x with the data v stored in the signals indicated by the indices i. This code sets signal 2 of cell array data: x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} v = {[20:22] [23:25]} y = setsignals(x,2,v)</pre>	
Examples		
See Also	nndata   numsignals   getsignals   catsignals   setelements   setsamples   settimesteps	

### setsiminit

Purpose	Set neural network Simulink block initial conditions	
Syntax	<pre>setsiminit(sysName,netName,net,xi,ai,Q)</pre>	
Description	<pre>setsiminit(sysName,netName,net,xi,ai,Q) takes these arguments,</pre>	
	sysName	The name of the Simulink system containing the neural network block
	netName	The name of the Simulink neural network block
	net	The original neural network
	xi	Initial input delay states
	ai	Initial layer delay states
	Q	Sample number (default is 1)
Examples	<ul><li>es and sets the Simulink neural network blocks initial conditions as specified.</li><li>es Here a NARX network is designed. The NARX network has a stan input and an open loop feedback output to an associated feedback i</li></ul>	
<pre>[x,t] = simplenarx_dataset; net = narxnet(1:2,1:2,20); view(net) [xs,xi,ai,ts] = preparets(net,x,{},t); net = train(net,xs,ts,xi,ai); y = net(xs,xi,ai); Now the network is converted to closed loop, and the approximately appro</pre>		et(1:2,1:2,20); s] = preparets(net,x,{},t); (net,xs,ts,xi,ai); xi,ai);
	<pre>to simulate the network's closed loop response. net = closeloop(net); view(net) [xs,xi,ai,ts] = preparets(net,x,{},t);</pre>	

y = net(xs,xi,ai);

Here the network is converted to a Simulink system with workspace input and output ports. Its delay states are initialized, inputs X1 defined in the workspace, and it is ready to be simulated in Simulink.

```
[sysName,netName] = gensim(net,'InputMode','Workspace',...
'OutputMode','WorkSpace','SolverMode','Discrete');
setsiminit(sysName,netName,net,xi,ai,1);
x1 = nndata2sim(x,1,1);
```

Finally the initial input and layer delays are obtained from the Simulink model. (They will be identical to the values set with setsiminit.)

[xi,ai] = getsiminit(sysName,netName,net);

See Also gensim | getsiminit | nndata2sim | sim2nndata

### settimesteps

Purpose	Set neural network data timesteps	
Syntax	<pre>settimesteps(x,i,v)</pre>	
Description	<pre>settimesteps(x,i,v) takes these arguments,</pre>	
	x Neural network matrix or cell array data	
	i	Indices
	v	Neural network data to store into x
Examples	<pre>and returns the original data x with the data v stored in the timesteps indicated by the indices i. This code sets timestep 2 of cell array data: x = {[1:3; 4:6] [7:9; 10:12]; [13:15] [16:18]} v = {[20:22; 23:25]; [25:27]} y = settimesteps(x,2,v)</pre>	
See Also	nndata   numtimesteps   gettimesteps   cattimesteps   setelements   setsamples   setsignals	

Purpose	Set all network weight and bias values with single vector		
Syntax	<pre>net = setwb(net,wb)</pre>		
Description	This function sets a network's weight and biases to a vector of values. net = setwb(net,wb) takes the following inputs:		
	net Neural network		
	wb	Vector of weight and bias values	
Examples	<pre>Here you create a network with a two-element input and one layer of three neurons. net = feedforwardnet(3); net = configure(net,[0;0],0);</pre>		
	The network has six weights (3 neurons * 2 input elements) and three biases (3 neurons) for a total of nine weight and bias values. You can set them to random values as follows:		
	<pre>net = setwb(net,rand(9,1));</pre>		
	You can then view the weight and bias values as follows:		
	net.iw{1,1} net.b{1}		
See Also	getwb   formwb   separatewb		

Purpose	Simulate neural network	
Syntax	<pre>[Y,Xf,Af] = sim(net,X,Xi,Ai,T) [Y,Xf,Af] = sim(net,{Q TS},Xi,Ai) [Y,] = sim(net,,'useParallel',) [Y,] = sim(net,,'useGPU',) [Y,] = sim(net,,'showResources',) [Ycomposite,] = sim(net,Xcomposite,) [Ygpu,] = sim(net,Xgpu,)</pre>	
To Get Help	Type help network/sim.	
Description	<pre>sim simulates n [Y,Xf,Af] = s: net X Xi Ai T and returns</pre>	eural networks. im(net,X,Xi,Ai,T) takes Network Network inputs Initial input delay conditions (default = zeros) Initial layer delay conditions (default = zeros) Network targets (default = zeros)
	Y	Network outputs
	Xf	Final input delay conditions
	Xf	Final layer delay conditions

sim is usually called implicitly by calling the neural network as a function. For instance, these two expressions return the same result:

y = sim(net,x,xi,ai)
y = net(x,xi,ai)

Note that arguments Xi, Ai, Xf, and Af are optional and need only be used for networks that have input or layer delays.

The signal arguments can have two formats: cell array or matrix.

The cell array format is easiest to describe. It is most convenient for networks with multiple inputs and outputs, and allows sequences of inputs to be presented:

Х	Ni-by-TS cell array	Each element X{i,ts} is an Ri-by-Q matrix.
Xi	Ni-by-ID cell array	Each element Xi{i,k} is an Ri-by-Q matrix.
Ai	N1-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.
Т	No-by-TS cell array	Each element X{i,ts} is a Ui-by-Q matrix.
Y	No-by-TS cell array	Each element Y{i,ts} is a Ui-by-Q matrix.
Xf	Ni-by-ID cell array	Each element Xf{i,k} is an Ri-by-Q matrix.
Af	N1-by-LD cell array	Each element Af{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
No	=	net.numOutputs
D	=	net.numInputDelays

LD	=	net.numLayerDelays
TS	=	Number of time steps
Q	=	Batch size
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Ui	=	net.outputs{i}.size

The columns of Xi, Ai, Xf, and Af are ordered from oldest delay condition to most recent:

Xi{i,k}	=	Input i at time $ts = k - ID$
Xf{i,k}	=	Input i at time ts = $TS + k - ID$
Ai{i,k}	=	Layer output i at time ts = $k - LD$
Af{i,k}	=	Layer output i at time ts = TS + k - LD

The matrix format can be used if only one time step is to be simulated (TS = 1). It is convenient for networks with only one input and output, but can also be used with networks that have more.

Each matrix argument is found by storing the elements of the corresponding cell array argument in a single matrix:

Х	(sum	of	Ri)-by-Q matrix
Xi	(sum	of	Ri)-by-(ID*Q) matrix
Ai	(sum	of	Si)-by-(LD*Q) matrix

Т	(sum	of	Ui)-by-Q matrix
Y	(sum	of	Ui)-by-Q matrix

```
Xf(sum of Ri)-by-(ID*Q) matrixAf(sum of Si)-by-(LD*Q) matrix
```

[Y,Xf,Af] = sim(net, {Q TS},Xi,Ai) is used for networks that do not have an input, such as Hopfield networks, when cell array notation is used.

```
[Y,...] = sim(net,...,'useParallel',...),
[Y,...] = sim(net,...,'useGPU',...), or [Y,...] =
sim(net,...,'showResources',...) (or the network called as a
function) accepts optional name/value pair arguments to control how
calculations are performed. Two of these options allow training to
happen faster or on larger datasets using parallel workers or GPU
devices if Parallel Computing Toolbox is available. These are the
optional name/value pairs:
```

```
'useParallel', Catculations occur on normal MATLAB thread. This is the default 'useParallel' setting.
```

- 'useParallel', Cytesulations occur on parallel workers if a MATLAB pool is open. Otherwise calculations occur on the normal MATLAB thread.
- 'useGPU', 'no' Calculations occur on the CPU. This is the default 'useGPU' setting.
- 'useGPU', 'yes'Calculations occur on the current gpuDevice if it is a supported GPU (See Parallel Computing Toolbox for GPU requirements.) If the current gpuDevice is not supported, calculations remain on the CPU. If 'useParallel' is also 'yes' and a MATLAB pool is open, then each worker with a unique GPU uses that GPU, other workers run calculations on their respective CPU cores.

```
'useGPU', 'onlyIf no MATLAB pool is open, then this setting is the
                                    same as 'yes'. If a MATLAB pool is open, then only
                                    workers with unique GPUs are used. However, if a
                                    MATLAB pool is open, but no supported GPUs are
                                    available, then calculations revert to performing on
                                    all worker CPUs.
                     'showResourcesDo'mot' display computing resources used at the
                                    command line. This is the default setting.
                     'showResourcesShowest the command line a summary of the
                                    computing resources actually used. The actual
                                    resources may differ from the requested resources,
                                    if parallel or GPU computing is requested but a
                                    MATLAB pool is not open or a supported GPU is
                                    not available. When parallel workers are used, each
                                    worker's computation mode is described, including
                                    workers in the pool that are not used.
                    [Ycomposite,...] = sim(net,Xcomposite,...) takes Composite
                    data and returns Composite results. If Composite data is used, then
                    'useParallel' is automatically set to 'yes'.
                    [Ygpu,...] = sim(net,Xgpu,...) takes gpuArray data and
                   returns gpuArray results. If gpuArray data is used, then 'useGPU' is
                    automatically set to 'yes'.
Examples
                   In the following examples, the sim function is called implicitly by calling
                   the neural network object (net) as a function.
                   Simulate Feedforward Networks
                   This example loads a dataset that maps neighborhood characteristics,
```

x, to median house prices, t. A feedforward network with 10 neurons is created and trained on that data, then simulated.

```
[x,t] = house_dataset;
net = feedforwardnet(10);
net = train(net,x,t);
```

```
y = net(x);
```

#### Simulate NARX Time Series Networks

This example trains an open-loop nonlinear-autoregressive network with external input, to model a levitated magnet system defined by a control current x and the magnet's vertical position response t, then simulates the network. The function preparets prepares the data before training and simulation. It creates the open-loop network's combined inputs x0, which contains both the external input x and previous values of position t. It also prepares the delay states x1.

```
[x,t] = maglev_dataset;
net = narxnet(10);
[xo,xi,~,to] = preparets(net,x,{},t);
net = train(net,xo,to,xi);
y = net(xo,xi)
```

This same system can also be simulated in closed-loop form.

```
netc = closeloop(net);
view(netc)
[xc,xi,ai,tc] = preparets(netc,x,{},t);
yc = netc(xc,xi,ai);
```

#### Simulate in Parallel on a MATLAB Pool

Parallel Computing Toolbox allows Neural Network Toolbox to simulate and train networks faster and on larger datasets than can fit on one PC. Here training and simulation happens across parallel MATLAB workers.

```
matlabpool open
[X,T] = vinyl_dataset;
net = feedforwardnet(10);
net = train(net,X,T,'useParallel','yes','showResources','yes');
Y = net(X,'useParallel','yes');
```

#### Simulate on GPUs

Use Composite values to distribute the data manually, and get back the results as a Composite value. If the data is loaded as it is distributed, then while each piece of the dataset must fit in RAM, the entire dataset is limited only by the total RAM of all the workers.

Networks can be simulated using the current GPU device, if it is supported by Parallel Computing Toolbox.

```
gpuDevice % Check if there is a supported GPU
Y = net(X,'useGPU','yes','showResources','yes');
```

To put the data on a GPU manually, and get the results on the GPU:

```
Xgpu = gpuArray(X);
Ygpu = net(Xgpu,'showResources','yes');
Y = gather(Ygpu);
```

To run in parallel, with workers associated with unique GPUs taking advantage of that hardware, while the rest of the workers use CPUs:

```
Y = net(X,'useParallel','yes','useGPU','yes','showResources','yes');
```

Using only workers with unique GPUs might result in higher speeds, as CPU workers might not keep up.

```
Y = net(X,'useParallel','yes','useGPU','only','showResources','yes');
```

#### **Algorithms** sim uses these properties to simulate a network net.

net.numInputs, net.numLayers
net.outputConnect, net.biasConnect
net.inputConnect, net.layerConnect

These properties determine the network's weight and bias values and the number of delays associated with each weight:

net.IW{i,j}
net.LW{i,j}
net.b{i}
net.inputWeights{i,j}.delays
net.layerWeights{i,j}.delays

These function properties indicate how sim applies weight and bias values to inputs to get each layer's output:

```
net.inputWeights{i,j}.weightFcn
net.layerWeights{i,j}.weightFcn
net.layers{i}.netInputFcn
net.layers{i}.transferFcn
```

See Also init | adapt | train | revert

### sim2nndata

Purpose	Convert Simulink time series to neural network data
Syntax	<pre>sim2nndata(x)</pre>
Description	<pre>sim2nndata(x) takes either a column vector of values or a Simulink time series structure and converts it to a neural network data time series.</pre>
Examples	Here a random Simulink 20-step time series is created and converted.
	simts = rands(20,1); nnts = sim2nndata(simts)
	Here a similar time series is defined with a Simulink structure and converted.
	<pre>simts.time = 0:19 simts.signals.values = rands(20,1); simts.dimensions = 1; nnts = sim2nndata(simts)</pre>
See Also	nndata   nndata2sim

Purpose	Soft max transfer function		
Graph	Inputn Outputa		
and Symbol	$\begin{array}{c c c c c c c c c c c c c c c c c c c $		
Syntax	A = softmax(N,FP)		
Description	<b>softmax</b> is a neural transfer function. Transfer functions calculate a layer's output from its net input.		
	A = softmax(N,FP) takes N and optional function parameters,		
	N S-by-Q matrix of net input (column) vectors		
	FP Struct of function parameters (ignored)		
	and returns A, the S-by-Q matrix of the softmax competitive function applied to each column of N.		
	<pre>info = softmax('code') returns information about this function. The following codes are defined:</pre>		
	<pre>softmax('name') returns the name of this function.</pre>		
	<pre>softmax('output',FP) returns the [min max] output range.</pre>		
	<pre>softmax('active',FP) returns the [min max] active input range.</pre>		
	<pre>softmax('fullderiv') returns 1 or 0, depending on whether dA_dN is S-by-S-by-Q or S-by-Q.</pre>		
	<pre>softmax('fpnames') returns the names of the function parameters.</pre>		
	<pre>softmax('fpdefaults') returns the default function parameters.</pre>		

### softmax

Examples	Here you define a net input vector N, calculate the output, and plot both with bar graphs.		
	<pre>n = [0; 1; -0.5; 0.5]; a = softmax(n); subplot(2,1,1), bar(n), ylabel('n') subplot(2,1,2), bar(a), ylabel('a')</pre>		
	Assign this transfer function to layer i of a network.		
	<pre>net.layers{i}.transferFcn = 'softmax';</pre>		
Algorithms	a = softmax(n) = exp(n)/sum(exp(n))		
See Also	sim   compet		

Purpose	1-D minimization using backtracking		
Syntax		ccode,delta,tol] = srchbac(net,X,Pd,Tl,Ai,Q,TS, dperf,delta,TOL,ch_perf)	
Description	srchbac is a linear search routine. It searches in a given direction to locate the minimum of the performance function in that direction. It uses a technique called backtracking.		
	[a,gX,perf,retcode,delta,tol] = srchbac(net,X,Pd,Tl,Ai,Q,TS,dX,gX,perf,dperf,delta,TOL,ch_perf) takes these inputs,		
	net	Neural network	
	X Vector containing current values of weights and biases		
	Pd Delayed input vectors		
	T1   Layer target vectors		
	Ai	Initial input delay conditions	
	Q Batch size		
	TS Time steps		
	dX	Search direction vector	
	gX Gradient vector		
	perf Performance value at current X		
	dperf Slope of performance value at current X in direction of dX		
	delta Initial step size		
	tol Tolerance on search		
	ch_perf	Change in performance on previous step	

### srchbac

#### and returns

а	Step size that minimizes performance
gX	Gradient at new minimum point
perf	Performance value at new minimum point
retcode	Return code that has three elements. The first two elements correspond to the number of function evaluations in the two stages of the search. The third element is a return code. These have different meanings for different search algorithms. Some might not be used in this function.
	0 Normal
	1 Minimum step taken
	2 Maximum step taken
	<b>3</b> Beta condition not met
delta	New initial step size, based on the current step size
tol	New tolerance on search

Parameters used for the backstepping algorithm are

alpha	Scale factor that determines sufficient reduction in perf
beta	Scale factor that determines sufficiently large step size
low_lim	Lower limit on change in step size
up_lim	Upper limit on change in step size
maxstep	Maximum step length

minstep	Minimum step length
scale_tol	Parameter that relates the tolerance tol to the initial step size delta, usually set to 20

The defaults for these parameters are set in the training function that calls them. See traincgf, traincgb, traincgp, trainbfg, and trainoss.

Dimensions for these variables are

Pd	No-by-Ni-by-TS cell array	Each element P{i,j,ts} is a Dij-by-Q matrix.
Tl	N1-by-TS cell array	Each element P{i,ts} is a Vi-by-Q matrix.
V	Nl-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
LD	=	net.numLayerDelays
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Vi	=	net.targets{i}.size
Dij	=	Ri * length(net.inputWeights{i,j}.delays)

**Examples** Here is a problem consisting of inputs **p** and targets **t** to be solved with a network.

 $p = [0 \ 1 \ 2 \ 3 \ 4 \ 5];$ t = [0 \ 0 \ 0 \ 1 \ 1 \ 1];

A two-layer feed-forward network is created. The network's input ranges from [0 to 10]. The first layer has two tansig neurons, and the second layer has one logsig neuron. The traincgf network training function and the srchbac search function are to be used.

#### **Create and Test a Network**

```
net = newff([0 5],[2 1],{'tansig','logsig'},'traincgf');
a = sim(net,p)
```

#### **Train and Retest the Network**

```
net.trainParam.searchFcn = 'srchbac';
net.trainParam.epochs = 50;
net.trainParam.show = 10;
net.trainParam.goal = 0.1;
net = train(net,p,t);
a = sim(net,p)
```

#### Network Use

You can create a standard network that uses srchbac with newff, newcf, or newelm.

To prepare a custom network to be trained with traincgf, using the line search function srchbac,

- 1 Set net.trainFcn to 'traincgf'. This sets net.trainParam to traincgf's default parameters.
- 2 Set net.trainParam.searchFcn to 'srchbac'.

The srchbac function can be used with any of the following training functions: traincgf, traincgb, traincgp, trainbfg, trainoss.

Algorithms	srchbac locates the minimum of the performance function in the search direction dX, using the backtracking algorithm described on page 126 and 328 of Dennis and Schnabel's book, noted below.
References	Dennis, J.E., and R.B. Schnabel, Numerical Methods for Unconstrained Optimization and Nonlinear Equations, Englewood Cliffs, NJ, Prentice-Hall, 1983
Definitions	The backtracking search routine srchbac is best suited to use with the quasi-Newton optimization algorithms. It begins with a step multiplier of 1 and then backtracks until an acceptable reduction in the performance is obtained. On the first step it uses the value of performance at the current point and a step multiplier of 1. It also uses the value of the derivative of performance at the current point to obtain a quadratic approximation to the performance function along the search direction. The minimum of the quadratic approximation becomes a tentative optimum point (under certain conditions) and the performance at this point is tested. If the performance is not sufficiently reduced, a cubic interpolation is obtained and the minimum of the cubic interpolation becomes the new tentative optimum point. This process is continued until a sufficient reduction in the performance is obtained. The backtracking algorithm is described in Dennis and Schnabel. It is used as the default line search for the quasi-Newton algorithms, otherwise is might and the best technique for all methans.
See Also	although it might not be the best technique for all problems. srchcha   srchgol   srchhyb

### srchbre

Purpose	1-D interval location using Brent's method		
Syntax	<pre>[a,gX,perf,retcode,delta,tol] = srchbre(net,X,Pd,Tl,Ai,Q,TS, dX,gX,perf,dperf,delta,tol,ch_perf)</pre>		
Description	srchbre is a linear search routine. It searches in a given direction to locate the minimum of the performance function in that direction. It uses a technique called Brent's technique.		
	<pre>[a,gX,perf,retcode,delta,tol] = srchbre(net,X,Pd,Tl,Ai,Q,TS,dX,gX,perf,dperf,delta,tol,ch_perf) takes these inputs,</pre>		
	net	Neural network	
	Х	Vector containing current values of weights and biases	
	Pd	Delayed input vectors	
	Tl	Layer target vectors	
	Ai	Initial input delay conditions	
	Q	Batch size	
	TS	Time steps	
	dX	Search direction vector	
	gX	Gradient vector	
	perf	Performance value at current X	
	dperf	Slope of performance value at current $\boldsymbol{X}$ in direction of $d\boldsymbol{X}$	
	delta	Initial step size	
	tol	Tolerance on search	
	ch_perf	Change in performance on previous step	

and :	returns
-------	---------

a	Step size that minimizes performance	
gX	Gradient at new minimum point	
perf	Performance value at new minimum point	
retcode	Return code that has three elements. The first two elements correspond to the number of function evaluations in the two stages of the search. The third element is a return code. These have different meanings for different search algorithms. Some might not be used in this function.	
	0 Normal	
	1 Minimum step taken	
	2 Maximum step taken	
	<b>3</b> Beta condition not met	
delta	New initial step size, based on the current step size	
tol	New tolerance on search	

Parameters used for the Brent algorithm are

alpha	Scale factor that determines sufficient reduction in perf
beta	Scale factor that determines sufficiently large step size
bmax	Largest step size
scale_tol	Parameter that relates the tolerance tol to the initial step size delta, usually set to 20

The defaults for these parameters are set in the training function that calls them. See traincgf, traincgb, traincgp, trainbfg, and trainoss.

Dimensions for these variables are

Pd	No-by-Ni-by-TS cell array	Each element P{i,j,ts} is a Dij-by-Q matrix.
Tl	N1-by-TS cell array	Each element P{i,ts} is a Vi-by-Q matrix.
Ai	N1-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
LD	=	net.numLayerDelays
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Vi	=	net.targets{i}.size
Dij	=	Ri * length(net.inputWeights{i,j}.delays)

# **Examples** Here is a problem consisting of inputs p and targets t to be solved with a network.

 $p = [0 \ 1 \ 2 \ 3 \ 4 \ 5];$ t = [0 0 0 1 1 1];

A two-layer feed-forward network is created. The network's input ranges from  $[0 \ to \ 10]$ . The first layer has two tansig neurons,

and the second layer has one logsig neuron. The traincgf network training function and the srchbac search function are to be used.

#### **Create and Test a Network**

```
net = newff([0 5],[2 1],{'tansig','logsig'},'traincgf');
a = sim(net,p)
```

#### Train and Retest the Network

```
net.trainParam.searchFcn = 'srchbre';
net.trainParam.epochs = 50;
net.trainParam.show = 10;
net.trainParam.goal = 0.1;
net = train(net,p,t);
a = sim(net,p)
```

- **Network** Use You can create a standard network that uses srchbre with newff, newcf, or newelm. To prepare a custom network to be trained with traincgf, using the line search function srchbre,
  - 1 Set net.trainFcn to 'traincgf'. This sets net.trainParam to traincgf's default parameters.
  - 2 Set net.trainParam.searchFcn to 'srchbre'.

The srchbre function can be used with any of the following training functions: traincgf, traincgb, traincgp, trainbfg, trainoss.

**Algorithms** srchbre brackets the minimum of the performance function in the search direction dX, using Brent's algorithm, described on page 46 of Scales (see reference below). It is a hybrid algorithm based on the golden section search and the quadratic approximation.

# **References** Scales, L.E., *Introduction to Non-Linear Optimization*, New York, Springer-Verlag, 1985

#### Definitions

Brent's search is a linear search that is a hybrid of the golden section search and a quadratic interpolation. Function comparison methods, like the golden section search, have a first-order rate of convergence, while polynomial interpolation methods have an asymptotic rate that is faster than superlinear. On the other hand, the rate of convergence for the golden section search starts when the algorithm is initialized, whereas the asymptotic behavior for the polynomial interpolation methods can take many iterations to become apparent. Brent's search attempts to combine the best features of both approaches.

For Brent's search, you begin with the same interval of uncertainty used with the golden section search, but some additional points are computed. A quadratic function is then fitted to these points and the minimum of the quadratic function is computed. If this minimum is within the appropriate interval of uncertainty, it is used in the next stage of the search and a new quadratic approximation is performed. If the minimum falls outside the known interval of uncertainty, then a step of the golden section search is performed.

See [Bren73] for a complete description of this algorithm. This algorithm has the advantage that it does not require computation of the derivative. The derivative computation requires a backpropagation through the network, which involves more computation than a forward pass. However, the algorithm can require more performance evaluations than algorithms that use derivative information.

See Also srchbac | srchcha | srchgol | srchhyb

Purpose	1-D minimization using Charalambous' method	
Syntax	[a,gX,perf,retcode,delta,tol] = srchcha(net,X,Pd,Tl,Ai,Q,TS, dX,gX,perf,dperf,delta,tol,ch_perf)	
Description	srchcha is a linear search routine. It searches in a given direction to locate the minimum of the performance function in that direction. It uses a technique based on Charalambous' method.	
	[a,gX,perf,retcode,delta,tol] = srchcha(net,X,Pd,Tl,Ai,Q,TS,dX,gX,perf,dperf,delta,tol,ch_perf) takes these inputs,	
	net	Neural network
	Х	Vector containing current values of weights and biases
	Pd	Delayed input vectors
	Tl	Layer target vectors
	Ai	Initial input delay conditions
	Q	Batch size
	TS	Time steps
	dX	Search direction vector
	gX	Gradient vector
	perf	Performance value at current X
	dperf	Slope of performance value at current $\boldsymbol{X}$ in direction of $d\boldsymbol{X}$
	delta	Initial step size
	tol	Tolerance on search
	ch_perf	Change in performance on previous step

### srchcha

and returns

а	Step size that minimizes performance
gX	Gradient at new minimum point
perf	Performance value at new minimum point
retcode	Return code that has three elements. The first two elements correspond to the number of function evaluations in the two stages of the search. The third element is a return code. These have different meanings for different search algorithms. Some might not be used in this function.
	0 Normal
	1 Minimum step taken
	2 Maximum step taken
	<b>3</b> Beta condition not met
delta	New initial step size, based on the current step size
tol	New tolerance on search

Parameters used for the Charalambous algorithm are

alpha	Scale factor that determines sufficient reduction in perf
beta	Scale factor that determines sufficiently large step size
gama	Parameter to avoid small reductions in performance, usually set to 0.1
scale_tol	Parameter that relates the tolerance tol to the initial step size delta, usually set to 20

The defaults for these parameters are set in the training function that calls them. See traincgf, traincgb, traincgp, trainbfg, and trainoss.

Dimensions for these variables are

Pd	No-by-Ni-by-TS cell array	Each element P{i,j,ts} is a Dij-by-Q matrix.
Tl	N1-by-TS cell array	Each element P{i,ts} is a Vi-by-Q matrix.
Ai	Nl-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
LD	=	net.numLayerDelays
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Vi	=	net.targets{i}.size
Dij	=	Ri * length(net.inputWeights{i,j}.delays)

# **Examples** Here is a problem consisting of inputs **p** and targets **t** to be solved with a network.

 $p = [0 \ 1 \ 2 \ 3 \ 4 \ 5]; \\ t = [0 \ 0 \ 0 \ 1 \ 1 \ 1];$ 

A two-layer feed-forward network is created. The network's input ranges from [0 to 10]. The first layer has two tansig neurons, and the second layer has one logsig neuron. The traincgf network training function and the srchcha search function are to be used.

### **Create and Test a Network**

```
net = newff([0 5],[2 1],{'tansig','logsig'},'traincgf');
a = sim(net,p)
```

### **Train and Retest the Network**

```
net.trainParam.searchFcn = 'srchcha';
net.trainParam.epochs = 50;
net.trainParam.show = 10;
net.trainParam.goal = 0.1;
net = train(net,p,t);
a = sim(net,p)
```

Network Use You can create a standard network that uses srchcha with newff, newcf, or newelm.

To prepare a custom network to be trained with traincgf, using the line search function srchcha,

1 Set net.trainFcn to 'traincgf'. This sets net.trainParam to traincgf's default parameters.

2 Set net.trainParam.searchFcn to 'srchcha'.

The srchcha function can be used with any of the following training functions: traincgf, traincgb, traincgp, trainbfg, trainoss.

# **Algorithms** srchcha locates the minimum of the performance function in the search direction dX, using an algorithm based on the method described in Charalambous (see reference below).

References	Charalambous, C., "Conjugate gradient algorithm for efficient training of artificial neural networks," <i>IEEE Proceedings</i> , Vol. 139, No. 3, June, 1992, pp. 301–310.
Definitions	The method of Charalambous, srchcha, was designed to be used in combination with a conjugate gradient algorithm for neural network training. Like srchbre and srchhyb, it is a hybrid search. It uses a cubic interpolation together with a type of sectioning.
	See [Char92] for a description of Charalambous' search. This routine is used as the default search for most of the conjugate gradient algorithms because it appears to produce excellent results for many different problems. It does require the computation of the derivatives (backpropagation) in addition to the computation of performance, but it overcomes this limitation by locating the minimum with fewer steps. This is not true for all problems, and you might want to experiment with other line searches.
See Also	srchbac   srchbre   srchgol   srchhyb

## srchgol

Purpose	1-D minimization using golden section search		
Syntax	<pre>[a,gX,perf,retcode,delta,tol] = srchgol(net,X,Pd,Tl,Ai,Q,TS, dX,gX,perf,dperf,delta,tol,ch_perf)</pre>		
Description	<pre>srchgol is a linear search routine. It searches in a given direction to locate the minimum of the performance function in that direction. It uses a technique called the golden section search. [a,gX,perf,retcode,delta,tol] = srchgol(net,X,Pd,Tl,Ai,Q,TS,dX,gX,perf,dperf,delta,tol,ch_perf) takes these inputs,</pre>		
	net	Neural network	
	Х	Vector containing current values of weights and biases	
	Pd	Delayed input vectors	
	Tl	Layer target vectors	
	Ai	Initial input delay conditions	
	Q	Batch size	
	TS	Time steps	
	dX	Search direction vector	
	gX	Gradient vector	
	perf	Performance value at current X	
	dperf	Slope of performance value at current $X$ in direction of $d X$	
	delta	Initial step size	
	tol	Tolerance on search	
	ch_perf	Change in performance on previous step	

and	returns
-----	---------

а	Step size that minimizes performance
gX	Gradient at new minimum point
perf	Performance value at new minimum point
retcode	Return code that has three elements. The first two elements correspond to the number of function evaluations in the two stages of the search. The third element is a return code. These have different meanings for different search algorithms. Some might not be used in this function.
	0 Normal
	1 Minimum step taken
	2 Maximum step taken
	<b>3</b> Beta condition not met
delta	New initial step size, based on the current step size
tol	New tolerance on search

Parameters used for the golden section algorithm are

alpha	Scale factor that determines sufficient reduction in $\ensuremath{perf}$
bmax	Largest step size
scale_tol	Parameter that relates the tolerance tol to the initial step size delta, usually set to 20

The defaults for these parameters are set in the training function that calls them. See traincgf, traincgb, traincgp, trainbfg, and trainoss.

Dimensions for these variables are

Pd	No-by-Ni-by-TS cell array	Each element P{i,j,ts} is a Dij-by-Q matrix.
Tl	Nl-by-TS cell array	Each element P{i,ts} is a Vi-by-Q matrix.
Ai	Nl-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
LD	=	net.numLayerDelays
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Vi	=	net.targets{i}.size
Dij	=	<pre>Ri * length(net.inputWeights{i,j}.delays)</pre>

# **Examples** Here is a problem consisting of inputs p and targets t to be solved with a network.

 $p = [0 \ 1 \ 2 \ 3 \ 4 \ 5];$ t = [0 0 0 1 1 1];

A two-layer feed-forward network is created. The network's input ranges from  $[0 \ to \ 10]$ . The first layer has two tansig neurons,

and the second layer has one logsig neuron. The traincgf network training function and the srchgol search function are to be used.

### **Create and Test a Network**

```
net = newff([0 5],[2 1],{'tansig','logsig'},'traincgf');
a = sim(net,p)
```

### **Train and Retest the Network**

```
net.trainParam.searchFcn = 'srchgol';
net.trainParam.epochs = 50;
net.trainParam.show = 10;
net.trainParam.goal = 0.1;
net = train(net,p,t);
a = sim(net,p)
```

NetworkYou can create a standard network that uses srchgol with newff,<br/>newcf, or newelm.

To prepare a custom network to be trained with traincgf, using the line search function srchgol,

- 1 Set net.trainFcn to 'traincgf'. This sets net.trainParam to traincgf's default parameters.
- 2 Set net.trainParam.searchFcn to 'srchgol'.

The srchgol function can be used with any of the following training functions: traincgf, traincgb, traincgp, trainbfg, trainoss.

**Algorithms** srchgol locates the minimum of the performance function in the search direction dX, using the golden section search. It is based on the algorithm as described on page 33 of Scales (see reference below).

# **References** Scales, L.E., *Introduction to Non-Linear Optimization*, New York, Springer-Verlag, 1985

## Definitions

The golden section search srchgol is a linear search that does not require the calculation of the slope. This routine begins by locating an interval in which the minimum of the performance function occurs. This is accomplished by evaluating the performance at a sequence of points, starting at a distance of delta and doubling in distance each step, along the search direction. When the performance increases between two successive iterations, a minimum has been bracketed. The next step is to reduce the size of the interval containing the minimum. Two new points are located within the initial interval. The values of the performance at these two points determine a section of the interval that can be discarded, and a new interior point is placed within the new interval. This procedure is continued until the interval of uncertainty is reduced to a width of tol, which is equal to delta/scale tol.

See [HDB96], starting on page 12-16, for a complete description of the golden section search. Try the *Neural Network Design* demonstration nnd12sd1 [HDB96] for an illustration of the performance of the golden section search in combination with a conjugate gradient algorithm.

See Also srchbac | srchbre | srchcha | srchhyb

Purpose	1-D minimization using a hybrid bisection-cubic search		
Syntax	<pre>[a,gX,perf,retcode,delta,tol] = srchhyb(net,X,Pd,Tl,Ai,Q,TS, dX,gX,perf,dperf,delta,tol,ch_perf)</pre>		
Description	<b>srchhyb</b> is a linear search routine. It searches in a given direction to locate the minimum of the performance function in that direction. It uses a technique that is a combination of a bisection and a cubic interpolation.		
	[a,gX,perf,retcode,delta,tol] = srchhyb(net,X,Pd,Tl,Ai,Q,TS,dX,gX,perf,dperf,delta,tol,ch_perf) takes these inputs,		
	net	Neural network	
	х	Vector containing current values of weights and biases	
	Pd	Delayed input vectors	
	Tl	Layer target vectors	
	Ai	Initial input delay conditions	
	Q	Batch size	
	TS	Time steps	
	dX	Search direction vector	
	gX	Gradient vector	
	perf	Performance value at current X	
	dperf	Slope of performance value at current $\boldsymbol{X}$ in direction of $\boldsymbol{d}\boldsymbol{X}$	
	delta	Initial step size	
	tol	Tolerance on search	
	ch_perf	Change in performance on previous step	

## srchhyb

### and returns

а	Step size that minimizes performance
gX	Gradient at new minimum point
perf	Performance value at new minimum point
retcode	Return code that has three elements. The first two elements correspond to the number of function evaluations in the two stages of the search. The third element is a return code. These have different meanings for different search algorithms. Some might not be used in this function.
	0 Normal
	1 Minimum step taken
	2 Maximum step taken
	<b>3</b> Beta condition not met
delta	New initial step size, based on the current step size
tol	New tolerance on search

Parameters used for the hybrid bisection-cubic algorithm are

alpha	Scale factor that determines sufficient reduction in perf
beta	Scale factor that determines sufficiently large step size
bmax	Largest step size
scale_tol	Parameter that relates the tolerance tol to the initial step size delta, usually set to 20

The defaults for these parameters are set in the training function that calls them. See traincgf, traincgb, traincgp, trainbfg, and trainoss.

Dimensions for these variables are

Pd	No-by-Ni-by-TS cell array	Each element P{i,j,ts} is a Dij-by-Q matrix.
Tl	Nl-by-TS cell array	Each element P{i,ts} is a Vi-by-Q matrix.
Ai	Nl-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
LD	=	net.numLayerDelays
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size
Vi	=	net.targets{i}.size
Dij	=	Ri * length(net.inputWeights{i,j}.delays)

# **Examples** Here is a problem consisting of inputs p and targets t to be solved with a network.

 $p = [0 \ 1 \ 2 \ 3 \ 4 \ 5]; \\ t = [0 \ 0 \ 0 \ 1 \ 1 \ 1];$ 

A two-layer feed-forward network is created. The network's input ranges from [0 to 10]. The first layer has two tansig neurons,

and the second layer has one logsig neuron. The traincgf network training function and the srchhyb search function are to be used.

## Create and Test a Network

```
net = newff([0 5],[2 1],{'tansig','logsig'},'traincgf');
a = sim(net,p)
```

### Train and Retest the Network

	<pre>net.trainParam.searchFcn = 'srchhyb'; net.trainParam.epochs = 50; net.trainParam.show = 10; net.trainParam.goal = 0.1; net = train(net,p,t); a = sim(net,p)</pre>
Network Use	You can create a standard network that uses srchhyb with newff, newcf, or newelm.
	To prepare a custom network to be trained with traincgf, using the line search function srchhyb,
	Set net.trainFcn to 'traincgf'. This sets net.trainParam to traincgf's default parameters.
	<b>2</b> Set net.trainParam.searchFcn to 'srchhyb'.
	The srchhyb function can be used with any of the following training functions: traincgf, traincgb, traincgp, trainbfg, trainoss.
Algorithms	srchhyb locates the minimum of the performance function in the search direction dX, using the hybrid bisection-cubic interpolation algorithm described on page 50 of Scales (see reference below).
References	Scales, L.E., <i>Introduction to Non-Linear Optimization</i> , New York Springer-Verlag, 1985

### **Definitions**

Like Brent's search, srchhyb is a hybrid algorithm. It is a combination of bisection and cubic interpolation. For the bisection algorithm, one point is located in the interval of uncertainty, and the performance and its derivative are computed. Based on this information, half of the interval of uncertainty is discarded. In the hybrid algorithm, a cubic interpolation of the function is obtained by using the value of the performance and its derivative at the two endpoints. If the minimum of the cubic interpolation falls within the known interval of uncertainty, then it is used to reduce the interval of uncertainty. Otherwise, a step of the bisection algorithm is used.

See [Scal85] for a complete description of the hybrid bisection-cubic search. This algorithm does require derivative information, so it performs more computations at each step of the algorithm than the golden section search or Brent's algorithm.

See Also srchbac | srchbre | srchcha | srchgol

Purpose	Sum squared error performance function	
Syntax	[] = sse(	., 'regularization', regularization) ., 'normalization', normalization) ., 'squaredWeighting', squaredWeighting)
Description	<b>sse</b> is a network performance function. It measures performance according to the sum of squared errors.	
	<pre>perf = sse(net,t,y,ew) takes these input arguments and optional function parameters,</pre>	
	net	Neural network
	t	Matrix or cell array of target vectors
	У	Matrix or cell array of output vectors
	ew	Error weights (default = $\{1\}$ )

and returns the sum squared error.

This function has three optional function parameters which can be defined with parameter name/pair arguments, or as a structure FP argument with fields having the parameter name and assigned the parameter values.

```
[...] = sse(...,'regularization',regularization)
[...] = sse(...,'normalization',normalization)
[...] = sse(...,'squaredWeighting',squaredWeighting)
[...] = sse(...,FP)
```

• regularization — can be set to any value between the default of 0 and 1. The greater the regularization value, the more squared weights and biases are taken into account in the performance calculation.

normalization — can be set to the default 'absolute', or 'normalized' (which normalizes errors to the [+2 -2] range consistent with normalized output and target ranges of [-1 1]) or 'percent' (which normalizes errors to the range [-1 +1]).
squaredWeighting — can be set to the default true, for applying error weights to squared errors; or false for applying error weights to the absolute errors before squaring.

# **Examples** Here a network is trained to fit a simple data set and its performance calculated

```
[x,t] = simplefit_dataset;
net = fitnet(10);
net.performFcn = 'sse';
net = train(net,x,t)
y = net(x)
e = t-y
perf = sse(net,t,y)
```

# NetworkTo prepare a custom network to be trained with sse, setUsenet.performFcn to 'sse'. This automatically sets net.performParam<br/>to the default function parameters.

Then calling train, adapt or perform will result in sse being used to calculate performance.

## staticderiv

Purpose	Static derivative function		
Syntax		staticderiv('dperf_dwb',net,X,T,Xi,Ai,EW) staticderiv('de_dwb',net,X,T,Xi,Ai,EW)	
Description	This function calculates derivatives using the chain rule from the networks performance or outputs back to its inputs. For time series data and dynamic networks this function ignores the delay connections resulting in a approximation (which may be good or not) of the actual derivative. This function is used by Elman networks (elmannet) which is a dynamic network trained by the static derivative approximation when full derivative calculations are not available. As full derivatives are calculated by all the other derivative functions, this function is not recommended for dynamic networks except for research into training algorithms. staticderiv('dperf_dwb',net,X,T,Xi,Ai,EW) takes these arguments,		
	net Neural network		
	Х	Inputs, an RxQ matrix (or NxTS cell array of RixQ matrices)	
	Т	Targets, an SxQ matrix (or MxTS cell array of SixQ matrices)	
	Xi	Initial input delay states (optional)	
	Ai	Initial layer delay states (optional)	
	EW	Error weights (optional)	

and returns the gradient of performance with respect to the network's weights and biases, where R and S are the number of input and output elements and Q is the number of samples (and N and M are the number of input and output signals, Ri and Si are the number of each input and outputs elements, and TS is the number of timesteps).

staticderiv('de\_dwb',net,X,T,Xi,Ai,EW) returns the Jacobian of errors with respect to the network's weights and biases.

**Examples** Here a feedforward network is trained and both the gradient and Jacobian are calculated.

```
[x,t] = simplefit_dataset;
net = feedforwardnet(20);
net = train(net,x,t);
y = net(x);
perf = perform(net,t,y);
gwb = staticderiv('dperf_dwb',net,x,t)
jwb = staticderiv('de_dwb',net,x,t)
See Also bttderiv | defaultderiv | fpderiv | num2deriv
```

## sumabs

Purpose	Sum of absolute elements of matrix or matrices	
Syntax	[s,n] = sumabs	5(X)
Description	[s,n] = sumabs(x) takes a matrix or cell array of matrices and returns,	
	S	Sum of all absolute finite values
	n	Number of finite values
	If x contains no	finite values, the sum returned is 0.
Examples	m = sumabs([1 [m,n] = sumabs	2;3 4]) s({[1 2; NaN 4], [4 5; 2 3]})
See Also	meanabs   meansqr   sumsqr	

Purpose	Sum of squared elements of matrix or matrices	
Syntax	[s,n] = sumsqr	^(X)
Description	[s,n] = sumsqr(x) takes a matrix or cell array of matrices and returns,	
	S	Sum of all squared finite values
	n	Number of finite values
	If x contains no	finite values, the sum returned is 0.
Examples	m = sumsqr([1 [m,n] = sumsqr	2;3 4]) ^({[1 2; NaN 4], [4 5; 2 3]})
See Also	meanabs   meansqr   sumabs	

## tansig

Purpose	Hyperbolic tangent sigmoid transfer function	
Graph and Symbol	a + 1 $a + 1$ $a = tansig(n)$ Tan-Sigmoid Transfer Function	
Syntax	A = tansig(N,FP)	
Description	tansig is a neural transfer function. Transfer functions calculate a layer's output from its net input.	
	A = tansig(N,FP) takes N and optional function parameters,	
	N S-by-Q matrix of net input (column) vectors	
	FP Struct of function parameters (ignored)	
	and returns A, the S-by-Q matrix of N's elements squashed into [ -1 1].	
Examples	Here is the code to create a plot of the tansig transfer function.	
	n = -5:0.1:5; a = tansig(n); plot(n,a)	
	Assign this transfer function to layer i of a network.	
	<pre>net.layers{i}.transferFcn = 'tansig';</pre>	
Algorithms	a = tansig(n) = 2/(1+exp(-2*n))-1	

	This is mathematically equivalent to tanh(N). It differs in that it runs faster than the MATLAB implementation of tanh, but the results can have very small numerical differences. This function is a good tradeoff for neural networks, where speed is important and the exact shape of the transfer function is not.
References	Vogl, T.P., J.K. Mangis, A.K. Rigler, W.T. Zink, and D.L. Alkon, "Accelerating the convergence of the backpropagation method," <i>Biological Cybernetics</i> , Vol. 59, 1988, pp. 257–263
See Also	sim   logsig

## tapdelay

Purpose	Shift neural network time series data for tap delay		
Syntax	tapdelay(x,i,	<pre>tapdelay(x,i,ts,delays)</pre>	
Description	tapdelay(x,i,	ts,delays) takes these arguments,	
	х	Neural network time series data	
	i	Signal index	
	ts	Timestep index	
	delays	Row vector of increasing zero or positive delays	
	specified tap de		
Examples	Here a random signal x consisting of eight timesteps is defined, and a tap delay with delays of [0 1 4] is simulated at timestep 6.		
	<pre>x = num2cell(rand(1,8)); y = tapdelay(x,1,6,[0 1 4])</pre>		
See Also	nndata   extendts   preparets		

Purpose	Time delay neural network		
Syntax	timedelaynet(inpu	<pre>timedelaynet(inputDelays,hiddenSizes,trainFcn)</pre>	
Description	Time delay networks are similar to feedforward networks, except that the input weight has a tap delay line associated with it. This allows the network to have a finite dynamic response to time series input data. This network is also similar to the distributed delay neural network (distdelaynet), which has delays on the layer weights in addition to the input weight.		
	<pre>timedelaynet(inputDelays,hiddenSizes,trainFcn) takes these arguments,</pre>		
	inputDelays Row vector of increasing 0 or positive delays (default = 1:2)		
	hiddenSizesRow vector of one or more hidden layer sizes (default = 10)trainFcnTraining function (default = 'trainlm')and returns a time delay neural network.		
Examples	Here a time delay n problem.	eural network is used to solve a simple time series	
	<pre>[X,T] = simpleseries_dataset; net = timedelaynet(1:2,10) [Xs,Xi,Ai,Ts] = preparets(net,X,T) net = train(net,Xs,Ts,Xi,Ai); view(net) Y = net(Xs,Xi,Ai); perf = perform(net,Ts,Y)</pre>		
See Also	preparets   removedelay   distdelaynet   narnet   narxnet		

## tonndata

Purpose	Convert data to standard neural network cell array form		
Syntax	[y,wasMatrix] = tonndata(x,columnSamples,cellTime)		
Description	<pre>[y,wasMatrix] = tonndata(x,columnSamples,cellTime) takes these arguments,</pre>		
	x	Matrix or cell array of matrices	
	columnSamples	True if original samples are oriented as columns, false if rows	
	cellTime True if original samples are columns of cell, fa if they are store in matrix		
	and returns		
	y Original data transformed into standard neural network cell array form		
	wasMatrix True if original data was a mat to cell array)	True if original data was a matrix (as apposed to cell array)	
	If columnSamples is false, then matrix x or matrices in cell array x will be transposed, so row samples will now be stored as column vectors. If cellTime is false, then matrix samples will be separated into columns of a cell array so time originally represented as vectors in a matrix will now be represented as columns of a cell array.		
	The returned value wasMatrix can be used by fromnndata to reverse the transformation.		
Examples	Here data consisting of six timesteps of 5-element vectors is originally represented as a matrix with six columns is converted to standard neural network representation and back.		

	x = rand(5,6)
	[y,wasMatrix] = tonndata(x,true,false)
	<pre>x2 = fromnndata(y,wasMatrix,columnSamples,cellTime)</pre>
See Also	nndata   fromnndata   nndata2sim   sim2nndata

## train

Purpose	Train neural network		
Syntax	<pre>[net,tr] = train(net,X,T,Xi,Ai) [net,] = train(,'useParallel',) [net,] = train(,'useGPU',) [net,] = train(,'showResources',) [net,] = train(Xcomposite,Tcomposite,) [net,] = train(Xgpu,Tgpu,)</pre>		
To Get Help	Type help network/train.		
Description	train trains a network net according to net.trainFcn and net.trainParam. [net,tr] = train(net,X,T,Xi,Ai) takes net Network		
	Х	Network inputs	
	т	Network targets (default = zeros)	
	Xi	Initial input delay conditions (default = zeros)	
	Ai Initial layer delay conditions (default = ze and returns		
	net	New network	
	Training record (epoch and perf)		

Note that T is optional and need only be used for networks that require targets. Xi is also optional and need only be used for networks that have input or layer delays.

train's signal arguments can have two formats: cell array or matrix.

The cell array format is easiest to describe. It is most convenient for networks with multiple inputs and outputs, and allows sequences of inputs to be presented.

Х	Ni-by-TS cell array	Each element X{i,j,ts} is an Ni-by-Q matrix.
Т	N1-by-TS cell array	Each element T{i,ts} is a Ui-by-Q matrix.
Xi	Ni-by-ID cell array	Each element Xi{i,k} is an Ri-by-Q matrix.
Ai	N1-by-LD cell array	Each element Ai{i,k} is an Si-by-Q matrix.

where

Ni	=	net.numInputs
Nl	=	net.numLayers
ID	=	net.numInputDelays
LD	=	net.numLayerDelays
TS	=	Number of time steps
Q	=	Batch size
Ri	=	net.inputs{i}.size
Si	=	net.layers{i}.size

The columns of Xi and Ai are ordered from the oldest delay condition to the most recent:

Xi{i,k}	=	Input i at time $ts = k - ID$
Ai{i,k}	=	Layer output i at time $ts = k - LD$

The matrix format can be used if only one time step is to be simulated (TS = 1). It is convenient for networks with only one input and output, but can be used with networks that have more.

Each matrix argument is found by storing the elements of the corresponding cell array argument in a single matrix:

Х	(sum of	Ri)-by-Q matrix
Т	(sum of	Ui)-by-Q matrix
Xi	(sum of	Ri)-by-(ID*Q) matrix
Ai	(sum of	Si)-by-(LD*Q) matrix

[net,...] = train(...,'useParallel',...), [net,...] = train(...,'useGPU',...), or [net,...] = train(...,'showResources',...) accepts optional name/value pair arguments to control how calculations are performed. Two of these options allow training to happen faster or on larger datasets using parallel workers or GPU devices if Parallel Computing Toolbox is available. These are the optional name/value pairs:

'useParallel',Catculations occur on normal MATLAB thread. This is the default 'useParallel' setting.
'useParallel',Cytallations occur on parallel workers if a MATLAB pool is open. Otherwise calculations occur on the normal MATLAB thread.
'useGPU', 'no' Calculations occur on the CPU. This is the default 'useGPU' setting.

- 'useGPU', 'yes'Calculations occur on the current gpuDevice if it is a supported GPU (See Parallel Computing Toolbox for GPU requirements.) If the current gpuDevice is not supported, calculations remain on the CPU. If 'useParallel' is also 'yes' and a MATLAB pool is open, then each worker with a unique GPU uses that GPU, other workers run calculations on their respective CPU cores.
- 'useGPU', 'onlyIf no MATLAB pool is open, then this setting is the same as 'yes'. If a MATLAB pool is open then only workers with unique GPUs are used. However, if a MATLAB pool is open, but no supported GPUs are available, then calculations revert to performing on all worker CPUs.
- 'showResourcesDo'mot' display computing resources used at the command line. This is the default setting.
- 'showResourcesShoyeat the command line a summary of the computing resources actually used. The actual resources may differ from the requested resources, if parallel or GPU computing is requested but a MATLAB pool is not open or a supported GPU is not available. When parallel workers are used, each worker's computation mode is described, including workers in the pool that are not used.
- 'reduction',N For most neural networks, the default CPU training computation mode is a compiled MEX algorithm. However, for large networks the calculations might occur with a MATLAB calculation mode. This can be confirmed using 'showResources'. If MATLAB is being used and memory is an issue, setting the reduction option to a value N greater than 1, reduces much of the temporary storage required to train by a factor of N, in exchange for longer training times.

[net,...] = train(Xcomposite,Tcomposite,...) takes Composite
data and returns Composite results. If Composite data is used, then
'useParallel' is automatically set to 'yes'.

[net,...] = train(Xgpu,Tgpu,...) takes gpuArray data and returns gpuArray results. If gpuArray data is used, then 'useGPU' is automatically set to 'yes'.

## **Examples** Train and Plot Networks

Here input x and targets t define a simple function that you can plot:

x = [0 1 2 3 4 5 6 7 8]; t = [0 0.84 0.91 0.14 -0.77 -0.96 -0.28 0.66 0.99]; plot(x,t, '0')

Here feedforwardnet creates a two-layer feed-forward network. The network has one hidden layer with ten neurons.

```
net = feedforwardnet(10);
net = configure(net,x,t);
y1 = net(x)
plot(x,t,'o',x,y1,'x')
```

The network is trained and then resimulated.

```
net = train(net,x,t);
y2 = net(x)
plot(x,t,'o',x,y1,'x',x,y2,'*')
```

### Train a NARX Time Series Network

This example trains an open-loop nonlinear-autoregressive network with external input, to model a levitated magnet system defined by a control current x and the magnet's vertical position response t, then simulates the network. The function preparets prepares the data before training and simulation. It creates the open-loop network's combined inputs xo, which contains both the external input x and previous values of position t. It also prepares the delay states xi.

```
[x,t] = maglev_dataset;
net = narxnet(10);
[xo,xi,~,to] = preparets(net,x,{},t);
net = train(net,xo,to,xi);
y = net(xo,xi)
```

This same system can also be simulated in closed-loop form.

```
netc = closeloop(net);
view(netc)
[xc,xi,ai,tc] = preparets(netc,x,{},t);
yc = netc(xc,xi,ai);
```

### Train a Network in Parallel on a MATLAB Pool

Parallel Computing Toolbox allows Neural Network Toolbox to simulate and train networks faster and on larger datasets than can fit on one PC. Here training and simulation happens across parallel MATLAB workers.

```
matlabpool open
[X,T] = vinyl_dataset;
net = feedforwardnet(10);
net = train(net,X,T,'useParallel','yes','showResources','yes');
Y = net(X);
```

### Train a Network on GPUs

Use Composite values to distribute the data manually, and get back the results as a Composite value. If the data is loaded as it is distributed then while each piece of the dataset must fit in RAM, the entire dataset is limited only by the total RAM of all the workers.

Yc = net(Xc);

Networks can be trained using the current GPU device, if it is supported by Parallel Computing Toolbox.

```
net = train(net,X,T,'useGPU','yes');
y = net(X);
```

To put the data on a GPU manually:

```
Xgpu = gpuArray(X);
Tgpu = gpuArray(T);
net = train(net,Xgpu,Tgpu);
Ygpu = net(Xgpu);
Y = gather(Ygpu);
```

To run in parallel, with workers associated with unique GPUs taking advantage of that hardware, while the rest of the workers use CPUs:

```
net = train(net,X,T,'useParallel','yes','useGPU','yes');
y = net(X);
```

Using only workers with unique GPUs might result in higher speed, as CPU workers might not keep up.

```
net = train(net,X,T,'useParallel','yes','useGPU','only');
Y = net(X);
```

### **Algorithms**

train calls the function indicated by net.trainFcn, using the training parameter values indicated by net.trainParam.

Typically one epoch of training is defined as a single presentation of all input vectors to the network. The network is then updated according to the results of all those presentations.

Training occurs until a maximum number of epochs occurs, the performance goal is met, or any other stopping condition of the function net.trainFcn occurs.

Some training functions depart from this norm by presenting only one input vector (or sequence) each epoch. An input vector (or sequence) is chosen randomly for each epoch from concurrent input vectors (or sequences). competlayer returns networks that use trainru, a training function that does this.

**See Also** init | revert | sim | adapt

## trainb

Purpose	Batch training with weight and bias learning rules			
Syntax	net.trainFcn = 'trainb' [net,tr] = train(net,)			
Description	trainb is not called directly. Instead it is called by train for networks whose net.trainFcn property is set to 'trainb', thus:			
	net.trainFcn = 'trainb'			
	[net,tr] = train(net,)			
	trainb trains a network with weight and bias learning rules with batch updates. The weights and biases are updated at the end of an entire pass through the input data. Training occurs according to trainb's training parameters, shown here with their default values:			
net.trainParam.epochs		100	Maximum number of epochs to train	
net.trainParam.goal		0	Performance goal	
net.trainParam.max_fail		5	Maximum validation failures	

	•	Maximum vanaabon fanares	
net.trainParam.show 25		Epochs between displays (NaN for no displays)	
net.trainParam.showCommandLinefalse		Generate command-line output	
net.trainParam.showWindow true		Show training GUI	
net.trainParam.time	inf	Maximum time to train in seconds	

Network Use	You can create a standard network that uses trainb by calling linearlayer.		
	To prepare a custom network to be trained with trainb,		
	1 Set net.trainFcn to 'trainb'. This sets net.trainParam to trainb's default parameters.		

	2 Set each net.inputWeights{i,j}.learnFcn to a learning function. Set each net.layerWeights{i,j}.learnFcn to a learning function. Set each net.biases{i}.learnFcn to a learning function. (Weight and bias learning parameters are automatically set to default values for the given learning function.) To train the network,
	1 Set net.trainParam properties to desired values.
	<b>2</b> Set weight and bias learning parameters to desired values.
	<b>3</b> Call train.
Algorithms	Each weight and bias is updated according to its learning function after each epoch (one pass through the entire set of input vectors).
	Training stops when any of these conditions is met:
	• The maximum number of epochs (repetitions) is reached.
	• Performance is minimized to the goal.
	• The maximum amount of time is exceeded.
	• Validation performance has increased more than max_fail times
	since the last time it decreased (when using validation).

## trainbfg

Purpose	BFGS quasi-Newton backpropagation			
Syntax	net.trainFcn = 'trainbfg' [net,tr] = train(net,)			
Description	trainbfg is a network training function that updates weight and bias values according to the BFGS quasi-Newton method.			
	net.trainFcn = 'tra:	inbfg'		
	[net,tr] = train(ne	t,)		
	Training occurs according to trainbfg's training parameters here with their default values:			
net.trainParam.epochs		100	Maximum number of epochs to train	
net.trainParam.sh	owWindow	0	Show training window	
net.trainParam.show		25	Epochs between displays (NaN for no displays)	
net.trainParam.showCommandLine		0	Generate command-line output	
net.trainParam.goal		0	Performance goal	
net.trainParam.time		inf	Maximum time to train in seconds	
net.trainParam.min_grad		1e-6	Minimum performance gradient	
net.trainParam.max_fail		5	Maximum validation failures	
net.trainParam.se	archFcn	'srchcha'	Name of line search routine to use	

Parameters related to line search methods (not all used for all methods):

### trainbfg

net.trainParam.scal_tol	20	Divide into delta to determine tolerance for linear search.
net.trainParam.alpha	0.001	Scale factor that determines sufficient reduction in perf
net.trainParam.beta	0.1	Scale factor that determines sufficiently large step size
net.trainParam.delta	0.01	Initial step size in interval location step
net.trainParam.gama	0.1	Parameter to avoid small reductions in performance, usually set to 0.1 (see srch_cha)
net.trainParam.low_lim	0.1	Lower limit on change in step size
net.trainParam.up_lim	0.5	Upper limit on change in step size
net.trainParam.maxstep	100	Maximum step length
net.trainParam.minstep	1.0e-6	Minimum step length
net.trainParam.bmax	26	Maximum step size
net.trainParam.batch_frag	0	In case of multiple batches, they are considered independent. Any nonzero value implies a fragmented batch, so the final layer's conditions

of a previous trained epoch are used as initial conditions for the next epoch.

### Network Use

You can create a standard network that uses trainbfg with feedfowardnet or cascadeforwardnet. To prepare a custom network to be trained with trainbfg:

- 1 Set NET.trainFcn to 'trainbfg'. This sets NET.trainParam to trainbfg's default parameters.
- 2 Set NET.trainParam properties to desired values.

In either case, calling train with the resulting network trains the network with trainbfg.

### trainbfg

```
Examples
                    Here a neural network is trained to predict median house prices.
                    [x,t] = house dataset;
                    net = feedforwardnet(10, 'trainbfg');
                    net = train(net,x,t);
                    y = net(x)
Algorithms
                    trainbfg can train any network as long as its weight, net input, and
                    transfer functions have derivative functions.
                    Backpropagation is used to calculate derivatives of performance perf
                    with respect to the weight and bias variables X. Each variable is
                    adjusted according to the following:
                    X = X + a*dX;
                    where dX is the search direction. The parameter a is selected to
                    minimize the performance along the search direction. The line search
                    function searchFcn is used to locate the minimum point. The first
                    search direction is the negative of the gradient of performance. In
                    succeeding iterations the search direction is computed according to
                    the following formula:
                    dX = -H \setminus gX;
                    where qX is the gradient and H is a approximate Hessian matrix. See
                    page 119 of Gill, Murray, and Wright (Practical Optimization, 1981) for
                    a more detailed discussion of the BFGS guasi-Newton method.
                    Training stops when any of these conditions occurs:
                    • The maximum number of epochs (repetitions) is reached.
                    • The maximum amount of time is exceeded.
                    • Performance is minimized to the goal.
                    • The performance gradient falls below min grad.

    Validation performance has increased more than max fail times

                       since the last time it decreased (when using validation).
```

#### **References** Gill, Murray, & Wright, *Practical Optimization*, 1981

### Definitions

Newton's method is an alternative to the conjugate gradient methods for fast optimization. The basic step of Newton's method is

$$\mathbf{x}_{k+1} = \mathbf{x}_k - \mathbf{A}_k^{-1} \mathbf{g}_k$$

where  $\mathbf{A}_{k}^{-1}$  is the Hessian matrix (second derivatives) of the performance index at the current values of the weights and biases. Newton's method often converges faster than conjugate gradient methods. Unfortunately, it is complex and expensive to compute the Hessian matrix for feedforward neural networks. There is a class of algorithms that is based on Newton's method, but which does not require calculation of second derivatives. These are called quasi-Newton (or secant) methods. They update an approximate Hessian matrix at each iteration of the algorithm. The update is computed as a function of the gradient. The quasi-Newton method that has been most successful in published studies is the Broyden, Fletcher, Goldfarb, and Shanno (BFGS) update. This algorithm is implemented in the trainbfg routine.

The BFGS algorithm is described in [DeSc83]. This algorithm requires more computation in each iteration and more storage than the conjugate gradient methods, although it generally converges in fewer iterations. The approximate Hessian must be stored, and its dimension is  $n \times n$ , where n is equal to the number of weights and biases in the network. For very large networks it might be better to use Rprop or one of the conjugate gradient algorithms. For smaller networks, however, trainbfg can be an efficient training function.

#### See Also cascadeforwardnet | feedforwardnet | traingdm | traingda | traingdx | trainlm | trainrp | traincgf | traincgb | trainscg | traincgp | trainoss

## trainbfgc

Purpose	BFGS quasi-Newton backpropagation for use with NN model reference adaptive controller			
Syntax	[net,TR,Y,E,Pf,Af,flag_stop] = trainbfgc(net,P,T,Pi,Ai,epochs, TS,Q) info = trainbfgc(code)			
Description	trainbfgc is a network training function that updates weight and bias values according to the BFGS quasi-Newton method. This function is called from nnmodref, a GUI for the model reference adaptive control Simulink block.			
	[net,TR,Y,E,Pf,Af,flag_stop] = trainbfgc(net,P,T,Pi,Ai,epochs,TS,Q) takes these inputs,			
	net Neural network			
	Р	PDelayed input vectorsTLayer target vectorsPiInitial input delay conditionsAiInitial layer delay conditions		
	т			
	Pi			
	Ai			
	epochsNumber of iterations for trainingTSTime steps			
	Q	Batch size		
	and returns			
	net		Trained network	
	TR	Training record of various values over eac epoch:		

TR.epoch Epoch number

TR.perf Training performance

	TR.vperf Validation performance
	TR.tperf Test performance
Υ	Network output for last epoch
E	Layer errors for last epoch
Pf	Final input delay conditions
Af	Collective layer outputs for last epoch
flag_stop	Indicates if the user stopped the training

Training occurs according to trainbfgc's training parameters, shown here with their default values:

net.trainParam.epochs	100	Maximum number of epochs to train
net.trainParam.show	25	Epochs between displays (NaN for no displays)
net.trainParam.goal	0	Performance goal
net.trainParam.time	inf	Maximum time to train in seconds
net.trainParam.min_grad	1e-6	Minimum performance gradient
net.trainParam.max_fail	5	Maximum validation failures
net.trainParam.searchFcn	'srchbac	×Name of line search routine to use

#### Parameters related to line search methods (not all used for all methods):

net.trainParam.scal_tol	20	Divide into delta to determine tolerance for linear search.
net.trainParam.alpha	0.001	Scale factor that determines sufficient reduction in perf
net.trainParam.beta	0.1	Scale factor that determines sufficiently large step size
net.trainParam.delta	0.01	Initial step size in interval location step

net.trainParam.gama	0.1	Parameter to avoid small reductions in performance, usually set to 0.1 (see srch_cha)
net.trainParam.low_lim	0.1	Lower limit on change in step size
net.trainParam.up_lim	0.5	Upper limit on change in step size
net.trainParam.maxstep	100	Maximum step length
net.trainParam.minstep	1.0e-6	Minimum step length
net.trainParam.bmax	26	Maximum step size

info = trainbfgc(code) returns useful information for each code
string:

'pnames'	Names of training parameters
'pdefaults'	Default training parameters

### Algorithms

trainbfgc can train any network as long as its weight, net input, and transfer functions have derivative functions. Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to the following:

X = X + a\*dX;

where dX is the search direction. The parameter a is selected to minimize the performance along the search direction. The line search function searchFcn is used to locate the minimum point. The first search direction is the negative of the gradient of performance. In succeeding iterations the search direction is computed according to the following formula:

 $dX = -H \setminus gX;$ 

where gX is the gradient and H is an approximate Hessian matrix. See page 119 of Gill, Murray, and Wright (*Practical Optimization*, 1981) for a more detailed discussion of the BFGS quasi-Newton method.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- Precision problems have occurred in the matrix inversion.
- **References** Gill, Murray, and Wright, *Practical Optimization*, 1981

## trainbr

Purpose	Bayesian regulation backpropagation
Syntax	net.trainFcn = 'trainbr' [net,tr] = train(net,)
Description	trainbr is a network training function that updates the weight and bias values according to Levenberg-Marquardt optimization. It minimizes a combination of squared errors and weights, and then determines the correct combination so as to produce a network that generalizes well. The process is called Bayesian regularization.
	net.trainFcn = 'trainbr'
	[net,tr] = train(net,)

Training occurs according to trainbr's training parameters, shown here with their default values:

net.trainParam.epochs	100	Maximum number of epochs to train
net.trainParam.goal	0	Performance goal
net.trainParam.mu	0.005	Marquardt adjustment parameter
net.trainParam.mu_dec	0.1	Decrease factor for mu
net.trainParam.mu_inc	10	Increase factor for mu
net.trainParam.mu_max	1e10	Maximum value for mu
net.trainParam.max_fail	5	Maximum validation failures
net.trainParam.mem_reduc	1	Factor to use for memory/speed tradeoff
net.trainParam.min_grad	1e-10	Minimum performance gradient
net.trainParam.show	25	Epochs between displays (NaN for no displays)
net.trainParam.showCommandLine	0	Generate command-line output
net.trainParam.showWindow	1	Show training GUI
net.trainParam.time	inf	Maximum time to train in seconds

Network Use	You can create a standard network that uses trainbr with feedforwardnet or cascadeforwardnet. To prepare a custom networ to be trained with trainbr,		
	1 Set NET.trainFcn to 'trainbr'. This sets NET.trainParam to trainbr's default parameters.		
	2 Set NET.trainParam properties to desired values.		
	In either case, calling train with the resulting network trains the network with trainbr. See feedforwardnet and cascadeforwardnet for examples.		
Examples	Here is a problem consisting of inputs <b>p</b> and targets <b>t</b> to be solved with a network. It involves fitting a noisy sine wave.		
	<pre>p = [-1:.05:1]; t = sin(2*pi*p)+0.1*randn(size(p));</pre>		
	A feed-forward network is created with a hidden layer of 2 neurons.		
	<pre>net = feedforwardnet(2,'trainbr');</pre>		
	Here the network is trained and tested.		
	<pre>net = train(net,p,t); a = net(p)</pre>		
Algorithms	trainbr can train any network as long as its weight, net input, and transfer functions have derivative functions.		
	Bayesian regularization minimizes a linear combination of squared errors and weights. It also modifies the linear combination so that at the end of training the resulting network has good generalization qualities. See MacKay ( <i>Neural Computation</i> , Vol. 4, No. 3, 1992, pp. 415 to 447) and Foresee and Hagan ( <i>Proceedings of the International</i>		

Joint Conference on Neural Networks, June, 1997) for more detailed discussions of Bayesian regularization.

This Bayesian regularization takes place within the Levenberg-Marquardt algorithm. Backpropagation is used to calculate the Jacobian jX of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to Levenberg-Marquardt,

jj = jX \* jX je = jX \* E dX = -(jj+I\*mu) ∖ je

where E is all errors and I is the identity matrix.

The adaptive value mu is increased by mu\_inc until the change shown above results in a reduced performance value. The change is then made to the network, and mu is decreased by mu\_dec.

The parameter mem\_reduc indicates how to use memory and speed to calculate the Jacobian jX. If mem\_reduc is 1, then trainlm runs the fastest, but can require a lot of memory. Increasing mem\_reduc to 2 cuts some of the memory required by a factor of two, but slows trainlm somewhat. Higher values continue to decrease the amount of memory needed and increase the training times.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- mu exceeds mu\_max.

# **Limitations** This function uses the Jacobian for calculations, which assumes that performance is a mean or sum of squared errors. Therefore networks

trained with this function must use either the  $\tt mse$  or  $\tt sse$  performance function.

References	MacKay, Neural Computation, Vol. 4, No. 3, 1992, pp. 415–447			
	Foresee and Hagan, Proceedings of the International Joint Conference on Neural Networks, June, 1997			
See Also	cascadeforwardnet   feedforwardnet   traingdm   traingda   traingdx   trainlm   trainrp   traincgf   traincgb   trainscg			

traincgp | trainbfg

### trainbu

Purpose	Batch unsupervised weight/bias training				
Syntax		net.trainFcn = 'trainbu' [net,tr] = train(net,)			
Description	trainbu trains a network with weight and bias learning rules with batch updates. Weights and biases updates occur at the end of an entire pass through the input data.				
			tly. Instead the train function calls it for nFcn property is set to 'trainbu', thus:		
	net.trainFcn =	'trainb	u'		
	[net,tr] = tra:	in(net,.	)		
	Training occurs according to trainbu training parameters, shown here with the following default values:				
net.trainParam.	epochs	100	Maximum number of epochs to train		
net.trainParam.show		25	Epochs between displays (NaN for no displays)		
net.trainParam.showCommandLine		false	Generate command-line output		
net.trainParam.showGUI		true	Show training GUI		
net.trainParam.time		inf	Maximum time to train in seconds		
Network Use	<ul> <li>Validation and test vectors have no impact on training for this function, but act as independent measures of network generalization.</li> <li>You can create a standard network that uses trainbu by calling selforgmap. To prepare a custom network to be trained with trainb:</li> <li>Set NET.trainFcn to 'trainbu'. (This option sets NET.trainParam to trainbu default parameters.)</li> </ul>				
	<b>2</b> Set each NET.inputWeights{i,j}.learnFcn to a learning function.				

	<b>3</b> Set each NET.layerWeights{i,j}.learnFcn to a learning function.
	<b>4</b> Set each NET.biases{i}.learnFcn to a learning function. (Weight and bias learning parameters are automatically set to default values for the given learning function.)
	To train the network:
	1 Set NET.trainParam properties to desired values.
	<b>2</b> Set weight and bias learning parameters to desired values.
	<b>3</b> Call train.
	See selforgmap for training examples.
Algorithms	Each weight and bias updates according to its learning function after each epoch (one pass through the entire set of input vectors).
	Training stops when any of these conditions is met:
	• The maximum number of epochs (repetitions) is reached.
	• Performance is minimized to the goal.
	• The maximum amount of time is exceeded.
	• Validation performance has increased more than max_fail times since the last time it decreased (when using validation).

See Also train | trainb

### trainc

Purpose	Cyclical order weight/bias training		
Syntax	net.trainFcn = [net,tr] = tra		
Description	trainc is not called directly. Instead it is called by train for networks whose net.trainFcn property is set to 'trainc', thus:		
	net.trainFcn =	'trainc	1
	[net,tr] = tra	in(net,.	)
		ates after	rith weight and bias learning rules with each presentation of an input. Inputs are
	Training occurs a with their defaul		to trainc's training parameters, shown here
net.trainParam.e	pochs	100	Maximum number of epochs to train
net.trainParam.g	joal	0	Performance goal
net.trainParam.max_fail		5	Maximum validation failures
net.trainParam.s	show	25	Epochs between displays (NaN for no displays)
net.trainParam.s	howCommandLine	false	Generate command-line output
net.trainParam.s	showWindow	true	Show training GUI
net.trainParam.t	ime	inf	Maximum time to train in seconds

Network Use	You can create a standard network that uses trainc by calling competlayer. To prepare a custom network to be trained with trainc,				
	1 Set net.trainFcn to 'trainc'. This sets net.trainParam to trainc's default parameters.				

	2 Set each net.inputWeights{i,j}.learnFcn to a learning function. Set each net.layerWeights{i,j}.learnFcn to a learning function. Set each net.biases{i}.learnFcn to a learning function. (Weight and bias learning parameters are automatically set to default values for the given learning function.)
	To train the network,
	1 Set net.trainParam properties to desired values.
	<b>2</b> Set weight and bias learning parameters to desired values.
	<b>3</b> Call train.
	See newp for training examples.
Algorithms	For each epoch, each vector (or sequence) is presented in order to the network, with the weight and bias values updated accordingly after each individual presentation.
	Training stops when any of these conditions is met:
	• The maximum number of epochs (repetitions) is reached.
	• Performance is minimized to the goal.
	• The maximum amount of time is exceeded.
See Also	competlayer   train

## traincgb

Purpose	Conjugate gradien	t backpro	opagation with Powell-Beale restarts
Syntax	net.trainFcn = [net,tr] = train	-	
Description	traincgb is a network training function that updates weight and bias values according to the conjugate gradient backpropagation with Powell-Beale restarts.		
	net.trainFcn =	'traincg	b '
	[net,tr] = train	n(net,	.)
	Training occurs ac here with their de	-	o <b>traincgb</b> 's training parameters, shown ues:
net.trainParam.e	pochs	100	Maximum number of epochs to train
net.trainParam.s	how	25	Epochs between displays (NaN for no displays)
net.trainParam.s	howCommandLine	0	Generate command-line output
net.trainParam.showWindow		1	Show training GUI
net.trainParam.goal		0	Performance goal
net.trainParam.time		inf	Maximum time to train in seconds
net.trainParam.n	iin_grad	1e-6	Minimum performance gradient
net.trainParam.n	nax_fail	5	Maximum validation failures
net.trainParam.searchFcn		'srchch	Name of line search routine to use

Parameters related to line search methods (not all used for all methods):

net.trainParam.scal_tol	20	Divide into delta to determine tolerance for linear search.
net.trainParam.alpha	0.001	Scale factor that determines sufficient reduction in perf

## traincgb

net.trainParam.beta	0.1	Scale factor that determines sufficiently large step size
net.trainParam.delta	0.01	Initial step size in interval location step
net.trainParam.gama	0.1	Parameter to avoid small reductions in performance, usually set to 0.1 (see srch_cha)
net.trainParam.low_lim	0.1	Lower limit on change in step size
net.trainParam.up_lim	0.5	Upper limit on change in step size
net.trainParam.maxstep	100	Maximum step length
net.trainParam.minstep	1.0e-6	Minimum step length
net.trainParam.bmax	26	Maximum step size

Network Use	You can create a standard network that uses traincgb with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traincgb,
	Set net.trainFcn to 'traincgb'. This sets net.trainParam to traincgb's default parameters.
	2 Set net.trainParam properties to desired values.
	In either case, calling train with the resulting network trains the network with traincgb.
Examples	Here a neural network is trained to predict median house prices.
	<pre>[x,t] = house_dataset; net = feedforwardnet(10,'traincgb'); net = train(net,x,t); y = net(x)</pre>
Algorithms	traincgb can train any network as long as its weight, net input, and transfer functions have derivative functions.

Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to the following:

X = X + a \* dX;

where dX is the search direction. The parameter a is selected to minimize the performance along the search direction. The line search function searchFcn is used to locate the minimum point. The first search direction is the negative of the gradient of performance. In succeeding iterations the search direction is computed from the new gradient and the previous search direction according to the formula

 $dX = -gX + dX_old*Z;$ 

where gX is the gradient. The parameter Z can be computed in several different ways. The Powell-Beale variation of conjugate gradient is distinguished by two features. First, the algorithm uses a test to determine when to reset the search direction to the negative of the gradient. Second, the search direction is computed from the negative gradient, the previous search direction, and the last search direction before the previous reset. See Powell, *Mathematical Programming*, Vol. 12, 1977, pp. 241 to 254, for a more detailed discussion of the algorithm.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- Validation performance has increased more than max\_fail times since the last time it decreased (when using validation).

#### **References** Powell, M.J.D., "Restart procedures for the conjugate gradient method," *Mathematical Programming*, Vol. 12, 1977, pp. 241–254

#### Definitions

For all conjugate gradient algorithms, the search direction is periodically reset to the negative of the gradient. The standard reset point occurs when the number of iterations is equal to the number of network parameters (weights and biases), but there are other reset methods that can improve the efficiency of training. One such reset method was proposed by Powell [Powe77], based on an earlier version proposed by Beale [Beal72]. This technique restarts if there is very little orthogonality left between the current gradient and the previous gradient. This is tested with the following inequality:

$$\left|\mathbf{g}_{k-1}^{T}\mathbf{g}_{k}\right| \geq 0.2 \left\|\mathbf{g}_{k}\right\|^{2}$$

If this condition is satisfied, the search direction is reset to the negative of the gradient.

The traincgb routine has somewhat better performance than traincgp for some problems, although performance on any given problem is difficult to predict. The storage requirements for the Powell-Beale algorithm (six vectors) are slightly larger than for Polak-Ribiére (four vectors).

See Also traingdm | traingda | traingdx | trainlm | traincgp | traincgf | trainscg | trainoss | trainbfg

## traincgf

Purpose	Conjugate gradie	nt backpr	opagation with Fletcher-Reeves updates
Syntax	net.trainFcn = [net,tr] = tra:	-	
Description	traincgf is a network training function that updates weight and bias values according to conjugate gradient backpropagation with Fletcher-Reeves updates.		
	net.trainFcn =	'trainco	)f'
	[net,tr] = tra:	in(net,	)
	Training occurs a here with their d	0	to traincgf's training parameters, shown ues:
net.trainParam.e	pochs	100	Maximum number of epochs to train
net.trainParam.s	how	25	Epochs between displays (NaN for no displays)
net.trainParam.showCommandLine		0	Generate command-line output
net.trainParam.s	howWindow	1	Show training GUI
net.trainParam.goal		0	Performance goal
net.trainParam.time		inf	Maximum time to train in seconds
net.trainParam.m	in_grad	1e-6	Minimum performance gradient
net.trainParam.m	ax_fail	5	Maximum validation failures
net.trainParam.s	earchFcn	'srchcha	Name of line search routine to use

Parameters related to line search methods (not all used for all methods):

net.trainParam.scal_tol	20	Divide into delta to determine tolerance for linear search.
net.trainParam.alpha	0.001	Scale factor that determines sufficient reduction in perf

net.trainParam.beta	0.1	Scale factor that determines sufficiently large step size
net.trainParam.delta	0.01	Initial step size in interval location step
net.trainParam.gama	0.1	Parameter to avoid small reductions in performance, usually set to 0.1 (see srch_cha)
net.trainParam.low_lim	0.1	Lower limit on change in step size
net.trainParam.up_lim	0.5	Upper limit on change in step size
net.trainParam.maxstep	100	Maximum step length
net.trainParam.minstep	1.0e-6	Minimum step length
net.trainParam.bmax	26	Maximum step size

Network Use	You can create a standard network that uses traincgf with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traincgf,
	Set net.trainFcn to 'traincgf'. This sets net.trainParam to traincgf's default parameters.
	2 Set net.trainParam properties to desired values.
	In either case, calling train with the resulting network trains the network with traincgf.
Examples	Here a neural network is trained to predict median house prices.
	<pre>[x,t] = house_dataset; net = feedforwardnet(10,'traincgf'); net = train(net,x,t); y = net(x)</pre>
Algorithms	traincgf can train any network as long as its weight, net input, and transfer functions have derivative functions.

Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to the following:

X = X + a \* dX;

where dX is the search direction. The parameter a is selected to minimize the performance along the search direction. The line search function searchFcn is used to locate the minimum point. The first search direction is the negative of the gradient of performance. In succeeding iterations the search direction is computed from the new gradient and the previous search direction, according to the formula

 $dX = -gX + dX_old*Z;$ 

where gX is the gradient. The parameter Z can be computed in several different ways. For the Fletcher-Reeves variation of conjugate gradient it is computed according to

```
Z = normnew_sqr/norm_sqr;
```

where norm\_sqr is the norm square of the previous gradient and normnew\_sqr is the norm square of the current gradient. See page 78 of Scales (*Introduction to Non-Linear Optimization*) for a more detailed discussion of the algorithm.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- Validation performance has increased more than max\_fail times since the last time it decreased (when using validation).

# **References** Scales, L.E., *Introduction to Non-Linear Optimization*, New York, Springer-Verlag, 1985

**Definitions** All the conjugate gradient algorithms start out by searching in the steepest descent direction (negative of the gradient) on the first iteration.

 $\mathbf{p}_0 = -\mathbf{g}_0$ 

A line search is then performed to determine the optimal distance to move along the current search direction:

$$\mathbf{x}_{k+1} = \mathbf{x}_k \alpha_k \mathbf{p}_k$$

Then the next search direction is determined so that it is conjugate to previous search directions. The general procedure for determining the new search direction is to combine the new steepest descent direction with the previous search direction:

 $\mathbf{p}_k = -\mathbf{g}_k + \beta_k \mathbf{p}_{k-1}$ 

The various versions of the conjugate gradient algorithm are distinguished by the manner in which the constant  $\beta_k$  is computed. For the Fletcher-Reeves update the procedure is

$$\beta_k = \frac{\mathbf{g}_k^T \mathbf{g}_k}{\mathbf{g}_{k-1}^T \mathbf{g}_{k-1}}$$

This is the ratio of the norm squared of the current gradient to the norm squared of the previous gradient.

See [FlRe64] or [HDB96] for a discussion of the Fletcher-Reeves conjugate gradient algorithm.

The conjugate gradient algorithms are usually much faster than variable learning rate backpropagation, and are sometimes faster than trainrp, although the results vary from one problem to another. The conjugate gradient algorithms require only a little more storage

## traincgf

than the simpler algorithms. Therefore, these algorithms are good for networks with a large number of weights.

Try the *Neural Network Design* demonstration nnd12cg [HDB96] for an illustration of the performance of a conjugate gradient algorithm.

See Also traingdm | traingda | traingdx | trainlm | traincgb | trainscg | traincgp | trainoss | trainbfg

Purpose	Conjugate gradient backpropagation with Polak-Ribiére updates			
Syntax	net.trainFcn = 'traincgp' [net,tr] = train(net,)			
Description	traincgp is a network training function that updates weight and bias values according to conjugate gradient backpropagation with Polak-Ribiére updates.			
	net.trainFcn =	traincgp	1	
	[net,tr] = trair	[net,tr] = train(net,)		
	Training occurs ac here with their def	-	traincgp's training parameters, shown es:	
net.trainParam.e	epochs	100	Maximum number of epochs to train	
net.trainParam.show		25	Epochs between displays (NaN for no displays)	
net.trainParam.showCommandLine		0	Generate command-line output	
net.trainParam.showWindow		1	Show training GUI	
net.trainParam.goal		0	Performance goal	
net.trainParam.time		inf	Maximum time to train in seconds	
net.trainParam.min_grad		1e-6	Minimum performance gradient	
net.trainParam.max_fail		5	Maximum validation failures	
net.trainParam.searchFcn		'srchcha	Name of line search routine to use	

Parameters related to line search methods (not all used for all methods):

net.trainParam.scal_tol	20	Divide into delta to determine tolerance for linear search.
net.trainParam.alpha	0.001	Scale factor that determines sufficient reduction in perf

net.trainParam.beta	0.1	Scale factor that determines sufficiently large step size
net.trainParam.delta	0.01	Initial step size in interval location step
net.trainParam.gama	0.1	Parameter to avoid small reductions in performance, usually set to 0.1 (see srch_cha)
net.trainParam.low_lim	0.1	Lower limit on change in step size
net.trainParam.up_lim	0.5	Upper limit on change in step size
net.trainParam.maxstep	100	Maximum step length
net.trainParam.minstep	1.0e-6	Minimum step length
net.trainParam.bmax	26	Maximum step size

Network Use	You can create a standard network that uses traincgp with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traincgp,			
	1 Set net.trainFcn to 'traincgp'. This sets net.trainParam to traincgp's default parameters.			
	2 Set net.trainParam properties to desired values.			
	In either case, calling train with the resulting network trains the network with traincgp.			
Examples	Examples			
	Here a neural network is trained to predict median house prices.			
	<pre>[x,t] = house_dataset; net = feedforwardnet(10,'traincgp'); net = train(net,x,t);</pre>			

```
y = net(x)
```

# **Algorithms** traincgp can train any network as long as its weight, net input, and transfer functions have derivative functions.

Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to the following:

X = X + a \* dX;

where dX is the search direction. The parameter a is selected to minimize the performance along the search direction. The line search function searchFcn is used to locate the minimum point. The first search direction is the negative of the gradient of performance. In succeeding iterations the search direction is computed from the new gradient and the previous search direction according to the formula

 $dX = -gX + dX_old*Z;$ 

where gX is the gradient. The parameter Z can be computed in several different ways. For the Polak-Ribiére variation of conjugate gradient, it is computed according to

where norm\_sqr is the norm square of the previous gradient, and gX\_old is the gradient on the previous iteration. See page 78 of Scales (*Introduction to Non-Linear Optimization*, 1985) for a more detailed discussion of the algorithm.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- Validation performance has increased more than max\_fail times since the last time it decreased (when using validation).

### traincgp

References	Scales, L.E., Introduction to Non-Linear Optimization, New York,
	Springer-Verlag, 1985

**Definitions** Another version of the conjugate gradient algorithm was proposed by Polak and Ribiére. As with the Fletcher-Reeves algorithm, traincgf, the search direction at each iteration is determined by

$$\mathbf{p}_k = -\mathbf{g}_k + \beta_k \mathbf{p}_{k-1}$$

For the Polak-Ribiére update, the constant  $\beta_k$  is computed by

$$\beta_k = \frac{\Delta \mathbf{g}_{k-1}^T \mathbf{g}_k}{\mathbf{g}_{k-1}^T \mathbf{g}_{k-1}}$$

This is the inner product of the previous change in the gradient with the current gradient divided by the norm squared of the previous gradient. See [FlRe64] or [HDB96] for a discussion of the Polak-Ribiére conjugate gradient algorithm.

The traincgp routine has performance similar to traincgf. It is difficult to predict which algorithm will perform best on a given problem. The storage requirements for Polak-Ribiére (four vectors) are slightly larger than for Fletcher-Reeves (three vectors).

See Also traingdm | traingda | traingdx | trainlm | trainrp | traincgf | traincgb | trainscg | trainoss | trainbfg

Purpose	Gradient descent backpropagation		
Syntax	net.trainFcn = 'traingd' [net,tr] = train(net,)		
Description	traingd is a network training function that updates weight and bias values according to gradient descent.		
	net.trainFcn =	'traing	d '
	[net,tr] = tra	in(net,.	)
	Training occurs a here with their d		to traingd's training parameters, shown ues:
net.trainParam.epochs		10	Maximum number of epochs to train
net.trainParam.goal		0	Performance goal
net.trainParam.showCommandLine		0	Generate command-line output
net.trainParam.showWindow		1	Show training GUI
net.trainParam.lr		0.01	Learning rate
net.trainParam.max_fail		5	Maximum validation failures
net.trainParam.min_grad		1e-10	Minimum performance gradient
net.trainParam.show		25	Epochs between displays (NaN for no displays)
net.trainParam.time		inf	Maximum time to train in seconds

#### Network Use

You can create a standard network that uses traingd with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traingd,

1 Set net.trainFcn to 'traingd'. This sets net.trainParam to traingd's default parameters.

## traingd

	2 Set net.trainParam properties to desired values.
	In either case, calling train with the resulting network trains the network with traingd.
	See help feedforwardnet and help cascadeforwardnet for examples.
Algorithms	traingd can train any network as long as its weight, net input, and transfer functions have derivative functions.
	Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to gradient descent:
	dX = lr * dperf/dX
	Training stops when any of these conditions occurs:
	• The maximum number of epochs (repetitions) is reached.
	• The maximum amount of time is exceeded.
	• Performance is minimized to the goal.
	• The performance gradient falls below min_grad.
	• Validation performance has increased more than max_fail times since the last time it decreased (when using validation).
Definitions	The batch steepest descent training function is traingd. The weights and biases are updated in the direction of the negative gradient of the performance function. If you want to train a network using batch steepest descent, you should set the network trainFcn to traingd, and then call the function train. There is only one training function associated with a given network.
	There are seven training parameters associated with traingd:
	• epochs
	• show

- goal
- time
- min\_grad
- max\_fail
- lr

The learning rate 1r is multiplied times the negative of the gradient to determine the changes to the weights and biases. The larger the learning rate, the bigger the step. If the learning rate is made too large, the algorithm becomes unstable. If the learning rate is set too small, the algorithm takes a long time to converge. See page 12-8 of [HDB96] for a discussion of the choice of learning rate.

The training status is displayed for every show iterations of the algorithm. (If show is set to NaN, then the training status is never displayed.) The other parameters determine when the training stops. The training stops if the number of iterations exceeds epochs, if the performance function drops below goal, if the magnitude of the gradient is less than mingrad, or if the training time is longer than time seconds. max\_fail, which is associated with the early stopping technique, is discussed in Improving Generalization.

The following code creates a training set of inputs p and targets t. For batch training, all the input vectors are placed in one matrix.

 $p = [-1 \ -1 \ 2 \ 2; \ 0 \ 5 \ 0 \ 5]; \\ t = [-1 \ -1 \ 1 \ 1];$ 

Create the feedforward network.

net = feedforwardnet(3, 'traingd');

In this simple example, turn off a feature that is introduced later.

net.divideFcn = '';

At this point, you might want to modify some of the default training parameters.

net.trainParam.show = 50; net.trainParam.lr = 0.05; net.trainParam.epochs = 300; net.trainParam.goal = 1e-5;

If you want to use the default training parameters, the preceding commands are not necessary.

Now you are ready to train the network.

[net,tr] = train(net,p,t);

The training record tr contains information about the progress of training.

Now you can simulate the trained network to obtain its response to the inputs in the training set.

a = net(p) a = -1.0026 -0.9962 1.0010 0.9960

Try the *Neural Network Design* demonstration nnd12sd1 [HDB96] for an illustration of the performance of the batch gradient descent algorithm.

```
See Also traingdm | traingda | traingdx | trainlm
```

Purpose	Gradient descent with adaptive learning rate backpropagation			
Syntax	net.trainFcn = 'traingda' [net,tr] = train(net,)			
Description	traingda is a network training function that updates weight and bias values according to gradient descent with adaptive learning rate.			
	net.trainFcn = '	traingd	a'	
	[net,tr] = trair	[net,tr] = train(net,)		
	-	Training occurs according to traingda's training parameters, shown here with their default values:		
net.trainParam.epochs		10	Maximum number of epochs to train	
net.trainParam.	goal	0	Performance goal	
net.trainParam.	lr	0.01	Learning rate	
net.trainParam.	lr_inc	1.05	Ratio to increase learning rate	
net.trainParam.lr_dec		0.7	Ratio to decrease learning rate	
net.trainParam.max_fail		5	Maximum validation failures	
net.trainParam.max_perf_inc		1.04	Maximum performance increase	
net.trainParam.min_grad		1e-10	Minimum performance gradient	
net.trainParam.show		25	Epochs between displays (NaN for no displays)	
net.trainParam.	showCommandLine	0	Generate command-line output	
net.trainParam.showWindow		1	Show training GUI	

inf Maximum time to train in seconds

### Network Use

net.trainParam.time

You can create a standard network that uses traingda with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traingda,

## traingda

	Set net.trainFcn to 'traingda'. This sets net.trainParam to traingda's default parameters.				
	2 Set net.trainParam properties to desired values.				
	In either case, calling train with the resulting network trains the network with traingda.				
	See help feedforwardnet and help cascadeforwardnet for examples.				
Algorithms	traingda can train any network as long as its weight, net input, and transfer functions have derivative functions.				
	Backpropagation is used to calculate derivatives of performance dperf with respect to the weight and bias variables X. Each variable is adjusted according to gradient descent:				
	dX = lr*dperf/dX				
	At each epoch, if performance decreases toward the goal, then the learning rate is increased by the factor lr_inc. If performance increases by more than the factor max_perf_inc, the learning rate is adjusted by the factor lr_dec and the change that increased the performance is not made.				
	Training stops when any of these conditions occurs:				
	• The maximum number of epochs (repetitions) is reached.				
	• The maximum amount of time is exceeded.				
	• Performance is minimized to the goal.				
	• The performance gradient falls below min_grad.				
	• Validation performance has increased more than max_fail times since the last time it decreased (when using validation).				
Definitions	With standard steepest descent, the learning rate is held constant throughout training. The performance of the algorithm is very sensitive to the proper setting of the learning rate. If the learning rate is set				

too high, the algorithm can oscillate and become unstable. If the learning rate is too small, the algorithm takes too long to converge. It is not practical to determine the optimal setting for the learning rate before training, and, in fact, the optimal learning rate changes during the training process, as the algorithm moves across the performance surface.

You can improve the performance of the steepest descent algorithm if you allow the learning rate to change during the training process. An adaptive learning rate attempts to keep the learning step size as large as possible while keeping learning stable. The learning rate is made responsive to the complexity of the local error surface.

An adaptive learning rate requires some changes in the training procedure used by traingd. First, the initial network output and error are calculated. At each epoch new weights and biases are calculated using the current learning rate. New outputs and errors are then calculated.

As with momentum, if the new error exceeds the old error by more than a predefined ratio,  $max\_perf\_inc$  (typically 1.04), the new weights and biases are discarded. In addition, the learning rate is decreased (typically by multiplying by  $lr\_dec = 0.7$ ). Otherwise, the new weights, etc., are kept. If the new error is less than the old error, the learning rate is increased (typically by multiplying by  $lr\_inc = 1.05$ ).

This procedure increases the learning rate, but only to the extent that the network can learn without large error increases. Thus, a near-optimal learning rate is obtained for the local terrain. When a larger learning rate could result in stable learning, the learning rate is increased. When the learning rate is too high to guarantee a decrease in error, it is decreased until stable learning resumes.

Try the *Neural Network Design* demonstration nnd12v1 [HDB96] for an illustration of the performance of the variable learning rate algorithm.

Backpropagation training with an adaptive learning rate is implemented with the function traingda, which is called just like traingd, except for the additional training parameters max\_perf\_inc, lr\_dec, and lr\_inc. Here is how it is called to train the previous
two-layer network:

```
p = [-1 -1 2 2; 0 5 0 5];
t = [-1 -1 1 1];
net = feedforwardnet(3, 'traingda');
net.trainParam.lr = 0.05;
net.trainParam.lr_inc = 1.05;
net = train(net,p,t);
y = net(p)
```

See Also traingd | traingdm | traingdx | trainlm

Purpose	Gradient descent with momentum backpropagation		
Syntax	net.trainFcn = 'traingdm' [net,tr] = train(net,)		
Description	traingdm is a network training function that updates weight and bias values according to gradient descent with momentum.		
	net.trainFcn = '	traingdm	۱ <sup>۰</sup>
	[net,tr] = train	(net,	)
	Training occurs according to traingdm's training parameters, shown here with their default values:		
net.trainParam.epochs		10	Maximum number of epochs to train
net.trainParam.goal		0	Performance goal
net.trainParam.lr		0.01	Learning rate
net.trainParam.max_fail		5	Maximum validation failures
net.trainParam.mc		0.9	Momentum constant
net.trainParam.min_grad		1e-10	Minimum performance gradient
net.trainParam.show		25	Epochs between showing progress
net.trainParam.showCommandLine		0	Generate command-line output
net.trainParam.	showWindow	1	Show training GUI
net.trainParam.time		inf	Maximum time to train in seconds

Network	You can create a standard network that uses traingdm with
Use	feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traingdm,

1 Set net.trainFcn to 'traingdm'. This sets net.trainParam to traingdm's default parameters.

# traingdm

	2 Set net.trainParam properties to desired values.				
	In either case, calling train with the resulting network trains the network with traingdm.				
	See help feedforwardnet and help cascadeforwardnet for ${ m examples}.$				
Algorithms	traingdm can train any network as long as its weight, net input, and transfer functions have derivative functions.				
	Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to gradient descent with momentum,				
	dX = mc*dXprev + lr*(1-mc)*dperf/dX				
	where dXprev is the previous change to the weight or bias.				
	Training stops when any of these conditions occurs:				
	• The maximum number of <b>epochs</b> (repetitions) is reached.				
	• The maximum amount of time is exceeded.				
	• Performance is minimized to the goal.				
	• The performance gradient falls below min_grad.				
	• Validation performance has increased more than max_fail times since the last time it decreased (when using validation).				
Definitions	In addition to traingd, there are three other variations of gradient descent.				
	Gradient descent with momentum, implemented by traingdm, allows a network to respond not only to the local gradient, but also to recent trends in the error surface. Acting like a lowpass filter, momentum allows the network to ignore small features in the error surface. Without momentum a network can get stuck in a shallow local minimum. With momentum a network can slide through such a minimum. See page 12–9 of [HDB96] for a discussion of momentum.				

Gradient descent with momentum depends on two training parameters. The parameter 1r indicates the learning rate, similar to the simple gradient descent. The parameter mc is the momentum constant that defines the amount of momentum. mc is set between 0 (no momentum) and values close to 1 (lots of momentum). A momentum constant of 1 results in a network that is completely insensitive to the local gradient and, therefore, does not learn properly.)

```
p = [-1 -1 2 2; 0 5 0 5];
t = [-1 -1 1 1];
net = feedforwardnet(3, 'traingdm');
net.trainParam.lr = 0.05;
net.trainParam.mc = 0.9;
net = train(net,p,t);
y = net(p)
```

Try the *Neural Network Design* demonstration nnd12mo [HDB96] for an illustration of the performance of the batch momentum algorithm.

See Also traingd | traingda | traingdx | trainlm

# traingdx

Purpose	Gradient descent with momentum and adaptive learning rate backpropagation			
Syntax	net.trainFcn = 'traingdx' [net,tr] = train(net,)			
Description	traingdx is a network training function that updates weight and bias values according to gradient descent momentum and an adaptive learning rate.			
	net.trainFcn = 'traingdx'			
	[net,tr] = train(net,)			
	Training occurs according to traingdx's training parameters, shown			

here with their default values:

net.trainParam.epochs	10	Maximum number of epochs to train
net.trainParam.goal	0	Performance goal
net.trainParam.lr	0.01	Learning rate
net.trainParam.lr_inc	1.05	Ratio to increase learning rate
net.trainParam.lr_dec	0.7	Ratio to decrease learning rate
net.trainParam.max_fail	5	Maximum validation failures
net.trainParam.max_perf_inc	1.04	Maximum performance increase
net.trainParam.mc	0.9	Momentum constant
net.trainParam.min_grad	1e-10	Minimum performance gradient
net.trainParam.show	25	Epochs between displays (NaN for no displays)
net.trainParam.showCommandLine	0	Generate command-line output
net.trainParam.showWindow	1	Show training GUI
net.trainParam.time	inf	Maximum time to train in seconds

Network Use	You can create a standard network that uses traingdx with feedforwardnet or cascadeforwardnet. To prepare a custom network to be trained with traingdx,				
	1 Set net.trainFcn to 'traingdx'. This sets net.trainParam to traingdx's default parameters.				
	2 Set net.trainParam properties to desired values.				
	In either case, calling train with the resulting network trains the network with traingdx.				
	See help feedforwardnet and help cascadeforwardnet for examples.				
Algorithms	traingdx can train any network as long as its weight, net input, and transfer functions have derivative functions.				
	Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to gradient descent with momentum,				
	dX = mc*dXprev + lr*mc*dperf/dX				
	where dXprev is the previous change to the weight or bias.				
	For each epoch, if performance decreases toward the goal, then the learning rate is increased by the factor lr_inc. If performance increases by more than the factor max_perf_inc, the learning rate is adjusted by the factor lr_dec and the change that increased the performance is not made.				
	Training stops when any of these conditions occurs:				
	• The maximum number of <b>epochs</b> (repetitions) is reached.				
	• The maximum amount of time is exceeded.				
	• Performance is minimized to the goal.				
	• The performance gradient falls below min_grad.				

# traingdx

	• Validation performance has increased more than max_fail times since the last time it decreased (when using validation).
Definitions	The function traingdx combines adaptive learning rate with momentum training. It is invoked in the same way as traingda, except that it has the momentum coefficient mc as an additional training parameter.
See Also	traingd   traingda   traingdm   trainlm

Purpose	Levenberg-Marquardt backpropagation
Syntax	net.trainFcn = 'trainlm' [net,tr] = train(net,)
Description	trainlm is a network training function that updates weight and bias values according to Levenberg-Marquardt optimization.
	trainlm is often the fastest backpropagation algorithm in the toolbox, and is highly recommended as a first-choice supervised algorithm, although it does require more memory than other algorithms.
	net.trainFcn = 'trainlm'
	[net,tr] = train(net,)

Training occurs according to trainlm's training parameters, shown here with their default values:

net.trainParam.epochs	1000	Maximum number of epochs to train
net.trainParam.goal	0	Performance goal
net.trainParam.max_fail	6	Maximum validation failures
net.trainParam.min_grad	1e-7	Minimum performance gradient
net.trainParam.mu	0.001	Initial mu
net.trainParam.mu_dec	0.1	mu decrease factor
net.trainParam.mu_inc	10	mu increase factor
net.trainParam.mu_max	1e10	Maximum mu
net.trainParam.show	25	Epochs between displays (NaN for no displays)
net.trainParam.showCommandLine	0	Generate command-line output
net.trainParam.showWindow	1	Show training GUI
net.trainParam.time	inf	Maximum time to train in seconds

# trainlm

	Validation vectors are used to stop training early if the network performance on the validation vectors fails to improve or remains the same for max_fail epochs in a row. Test vectors are used as a further check that the network is generalizing well, but do not have any effect on training.
	trainlm is the default training function for several network creation functions including newcf, newdtdnn, newff, and newnarx.
Network Use	You can create a standard network that uses trainlm with feedforwardnet or cascadeforwardnet.
	To prepare a custom network to be trained with trainlm,
	1 Set net.trainFcn to 'trainlm'. This sets net.trainParam to trainlm's default parameters.
	<b>2</b> Set net.trainParam properties to desired values.
	In either case, calling train with the resulting network trains the network with trainlm.
	See help feedforwardnet and help cascadeforwardnet for examples.
Examples	Here a neural network is trained to predict median house prices.
	<pre>[x,t] = house_dataset; net = feedforwardnet(10,'trainlm'); net = train(net,x,t); y = net(x)</pre>
Algorithms	trainlm supports training with validation and test vectors if the network's NET.divideFcn property is set to a data division function. Validation vectors are used to stop training early if the network performance on the validation vectors fails to improve or remains the same for max_fail epochs in a row. Test vectors are used as a further check that the network is generalizing well, but do not have any effect on training.

trainlm can train any network as long as its weight, net input, and transfer functions have derivative functions.

Backpropagation is used to calculate the Jacobian jX of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to Levenberg-Marquardt,

jj = jX \* jX je = jX \* E dX = -(jj+I\*mu) ∖ je

where E is all errors and I is the identity matrix.

The adaptive value mu is increased by mu\_inc until the change above results in a reduced performance value. The change is then made to the network and mu is decreased by mu\_dec.

The parameter mem\_reduc indicates how to use memory and speed to calculate the Jacobian jX. If mem\_reduc is 1, then trainlm runs the fastest, but can require a lot of memory. Increasing mem\_reduc to 2 cuts some of the memory required by a factor of two, but slows trainlm somewhat. Higher states continue to decrease the amount of memory needed and increase training times.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- mu exceeds mu\_max.
- Validation performance has increased more than max\_fail times since the last time it decreased (when using validation).

# **Definitions** Like the quasi-Newton methods, the Levenberg-Marquardt algorithm was designed to approach second-order training speed without having to compute the Hessian matrix. When the performance function has

the form of a sum of squares (as is typical in training feedforward networks), then the Hessian matrix can be approximated as

 $\mathbf{H} = \mathbf{J}^T \mathbf{J}$ 

and the gradient can be computed as

 $\mathbf{g} = \mathbf{J}^T \mathbf{e}$ 

where **J** is the Jacobian matrix that contains first derivatives of the network errors with respect to the weights and biases, and **e** is a vector of network errors. The Jacobian matrix can be computed through a standard backpropagation technique (see [HaMe94]) that is much less complex than computing the Hessian matrix.

The Levenberg-Marquardt algorithm uses this approximation to the Hessian matrix in the following Newton-like update:

$$\mathbf{x}_{k+1} = \mathbf{x}_k - [\mathbf{J}^T \mathbf{J} + \mu \mathbf{I}]^{-1} \mathbf{J}^T \mathbf{e}$$

When the scalar  $\mu$  is zero, this is just Newton's method, using the approximate Hessian matrix. When  $\mu$  is large, this becomes gradient descent with a small step size. Newton's method is faster and more accurate near an error minimum, so the aim is to shift toward Newton's method as quickly as possible. Thus,  $\mu$  is decreased after each successful step (reduction in performance function) and is increased only when a tentative step would increase the performance function. In this way, the performance function is always reduced at each iteration of the algorithm.

The original description of the Levenberg-Marquardt algorithm is given in [Marq63]. The application of Levenberg-Marquardt to neural network training is described in [HaMe94] and starting on page 12-19 of [HDB96]. This algorithm appears to be the fastest method for training moderate-sized feedforward neural networks (up to several hundred weights). It also has an efficient implementation in MATLAB<sup>®</sup> software, because the solution of the matrix equation is a built-in function, so its attributes become even more pronounced in a MATLAB environment. Try the *Neural Network Design* demonstration nnd12m [HDB96] for an illustration of the performance of the batch Levenberg-Marquardt algorithm.

# **Limitations** This function uses the Jacobian for calculations, which assumes that performance is a mean or sum of squared errors. Therefore, networks trained with this function must use either the mse or sse performance function.

# trainoss

Purpose	One-step secant backpropagation		
Syntax	net.trainFcn = 'trainoss' [net,tr] = train(net,)		
Description	trainoss is a network training function that updates weight and bias values according to the one-step secant method.		
	net.trainFcn =	'trainos	S '
	[net,tr] = trai	n(net,	.)
	Training occurs according to trainoss's training parameters, shown here with their default values:		
net.trainParam.epochs		100	Maximum number of epochs to train
net.trainParam.show		25	Epochs between displays (NaN for no displays)
net.trainParam.showCommandLine		0	Generate command-line output
net.trainParam.showWindow		1	Show training GUI
net.trainParam.goal		0	Performance goal
net.trainParam.time		inf	Maximum time to train in seconds
net.trainParam.r	nin_grad	1e-6	Minimum performance gradient
net.trainParam.max_fail		5	Maximum validation failures
net.trainParam.searchFcn		'srchcha	a Name of line search routine to use

Parameters related to line search methods (not all used for all methods):

net.trainParam.scal_tol	20	Divide into delta to determine tolerance for linear search.
net.trainParam.alpha	0.001	Scale factor that determines sufficient reduction in perf

#### trainoss

net.trainParam.beta	0.1	Scale factor that determines sufficiently large step size
net.trainParam.delta	0.01	Initial step size in interval location step
net.trainParam.gama	0.1	Parameter to avoid small reductions in performance, usually set to 0.1 (see srch_cha)
net.trainParam.low_lim	0.1	Lower limit on change in step size
net.trainParam.up_lim	0.5	Upper limit on change in step size
net.trainParam.maxstep	100	Maximum step length
net.trainParam.minstep	1.0e-6	Minimum step length
net.trainParam.bmax	26	Maximum step size

Network	You can create a standard network that uses trainoss with
Use	feedforwardnet or cascadeforwardnet. To prepare a custom network
	to be trained with trainoss:

- 1 Set net.trainFcn to 'trainoss'. This sets net.trainParam to trainoss's default parameters.
- 2 Set net.trainParam properties to desired values.

In either case, calling train with the resulting network trains the network with trainoss.

#### **Examples** Here a neural network is trained to predict median house prices.

[x,t] = house\_dataset; net = feedforwardnet(10,'trainoss'); net = train(net,x,t); y = net(x)

**Algorithms** trainoss can train any network as long as its weight, net input, and transfer functions have derivative functions.

Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to the following:

X = X + a\*dX;

where dX is the search direction. The parameter a is selected to minimize the performance along the search direction. The line search function searchFcn is used to locate the minimum point. The first search direction is the negative of the gradient of performance. In succeeding iterations the search direction is computed from the new gradient and the previous steps and gradients, according to the following formula:

 $dX = -gX + Ac*X_step + Bc*dgX;$ 

where gX is the gradient,  $X\_step$  is the change in the weights on the previous iteration, and dgX is the change in the gradient from the last iteration. See Battiti (*Neural Computation*, Vol. 4, 1992, pp. 141–166) for a more detailed discussion of the one-step secant algorithm.

Training stops when any of these conditions occurs:

- The maximum number of epochs (repetitions) is reached.
- The maximum amount of time is exceeded.
- Performance is minimized to the goal.
- The performance gradient falls below min\_grad.
- Validation performance has increased more than max\_fail times since the last time it decreased (when using validation).
- **References** Battiti, R., "First and second order methods for learning: Between steepest descent and Newton's method," *Neural Computation*, Vol. 4, No. 2, 1992, pp. 141–166

# **Definitions** Because the BFGS algorithm requires more storage and computation in each iteration than the conjugate gradient algorithms, there is need

for a secant approximation with smaller storage and computation requirements. The one step secant (OSS) method is an attempt to bridge the gap between the conjugate gradient algorithms and the quasi-Newton (secant) algorithms. This algorithm does not store the complete Hessian matrix; it assumes that at each iteration, the previous Hessian was the identity matrix. This has the additional advantage that the new search direction can be calculated without computing a matrix inverse.

The one step secant method is described in [Batt92]. This algorithm requires less storage and computation per epoch than the BFGS algorithm. It requires slightly more storage and computation per epoch than the conjugate gradient algorithms. It can be considered a compromise between full quasi-Newton algorithms and conjugate gradient algorithms.

See Also traingdm | traingda | traingdx | trainlm | trainrp | traincgf | traincgb | trainscg | traincgp | trainbfg

## trainr

Purpose	Random order incremental training with learning functions		
Syntax	net.trainFcn = 'trainr' [net,tr] = train(net,)		
Description	trainr is not called directly. Instead it is called by train for networks whose net.trainFcn property is set to 'trainr', thus:		
	net.trainFcn =	'trainr'	
	[net,tr] = trai	n(net,	.)
		tes after e	th weight and bias learning rules with each presentation of an input. Inputs are
	Training occurs ac with their default		o trainr's training parameters, shown here
net.trainParam.e	epochs	100	Maximum number of epochs to train
net.trainParam.goal		0	Performance goal
net.trainParam.show		25	Epochs between displays (NaN for no displays)
net.trainParam.showCommandLine		0	Generate command-line output
net.trainParam.s	showWindow	1	Show training GUI
net.trainParam.t	ime	inf	Maximum time to train in seconds

Network Use	You can create a standard network that uses trainr by calling competlayer or selforgmap. To prepare a custom network to be trained with trainr,			
	1 Set net.trainFcn to 'trainr'. This sets net.trainParam to trainr's default parameters.			

2 Set each net.inputWeights{i,j}.learnFcn to a learning function.

	<b>3</b> Set each net.layerWeights{i,j}.learnFcn to a learning function.		
	<b>4</b> Set each net.biases{i}.learnFcn to a learning function. (Weight and bias learning parameters are automatically set to default values for the given learning function.)		
	To train the network,		
	1 Set net.trainParam properties to desired values.		
	<b>2</b> Set weight and bias learning parameters to desired values.		
	<b>3</b> Call train.		
	See help competlayer and help selforgmap for training examples.		
Algorithms	For each epoch, all training vectors (or sequences) are each presented once in a different random order, with the network and weight and bias values updated accordingly after each individual presentation.		
	Training stops when any of these conditions is met:		
	• The maximum number of epochs (repetitions) is reached.		
	• Performance is minimized to the goal.		
	• The maximum amount of time is exceeded.		
See Also	train		

# trainrp

Purpose	Resilient backpropagation		
Syntax	net.trainFcn = [net,tr] = tra		•
Description	trainrp is a network training function that updates weight and bias values according to the resilient backpropagation algorithm (Rprop).		
	net.trainFcn =	'trainr	'q
	[net,tr] = tra	in(net,.	)
	Training occurs a here with their d		to trainrp's training parameters, shown lues:
net.trainParam.	epochs	100	Maximum number of epochs to train
net.trainParam.	show	25	Epochs between displays (NaN for no displays)
net.trainParam.	showCommandLine	0	Generate command-line output
net.trainParam.	showWindow	1	Show training GUI
net.trainParam.	goal	0	Performance goal
net.trainParam.	time	inf	Maximum time to train in seconds
net.trainParam.	nin_grad	1e-6	Minimum performance gradient
net.trainParam.	nax_fail	5	Maximum validation failures
net.trainParam.	lr	0.01	Learning rate
net.trainParam.	delt_inc	1.2	Increment to weight change
net.trainParam.	delt_dec	0.5	Decrement to weight change
net.trainParam.	deltaO	0.07	Initial weight change
net.trainParam.	deltamax	50.0	Maximum weight change

Network Use	You can create a standard network that uses trainrp with feedforwardnet or cascadeforwardnet.
	To prepare a custom network to be trained with trainrp,
	Set net.trainFcn to 'trainrp'. This sets net.trainParam to trainrp's default parameters.
	2 Set net.trainParam properties to desired values.
	In either case, calling train with the resulting network trains the network with trainrp.
Examples	Here is a problem consisting of inputs ${\sf p}$ and targets ${\sf t}$ to be solved with a network.
	p = [0 1 2 3 4 5]; t = [0 0 0 1 1 1];
	A two-layer feed-forward network with two hidden neurons and this training function is created.
	Create and test a network.
	<pre>net = feedforwardnet(2,'trainrp');</pre>
	Here the network is trained and retested.
	net.trainParam.epochs = 50; net.trainParam.show = 10;
	net.trainParam.goal = 0.1;
	<pre>net = train(net,p,t); a = net(p)</pre>
	See help feedforwardnet and help cascadeforwardnet for other examples.

Algorithms	trainrp can train any network as long as its weight, net input, and transfer functions have derivative functions.			
	Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X. Each variable is adjusted according to the following:			
	dX = deltaX.*sign(gX);			
	where the elements of deltaX are all initialized to deltaO, and gX is the gradient. At each iteration the elements of deltaX are modified. If an element of gX changes sign from one iteration to the next, then the corresponding element of deltaX is decreased by delta_dec. If an element of gX maintains the same sign from one iteration to the next, then the corresponding element of deltaX is increased by delta_inc. See Riedmiller, <i>Proceedings of the IEEE International Conference on</i> <i>Neural Networks (ICNN)</i> , San Francisco, 1993, pp. 586 to 591.			
	Training stops when any of these conditions occurs:			
	• The maximum number of epochs (repetitions) is reached.			
	<ul><li>The maximum amount of time is exceeded.</li><li>Performance is minimized to the goal.</li></ul>			
	• The performance gradient falls below min_grad.			
	• Validation performance has increased more than max_fail times since the last time it decreased (when using validation).			
References	Riedmiller, Proceedings of the IEEE International Conference on Neural Networks (ICNN), San Francisco, 1993, pp. 586–591			
Definitions	Multilayer networks typically use sigmoid transfer functions in the hidden layers. These functions are often called "squashing" functions, because they compress an infinite input range into a finite output range. Sigmoid functions are characterized by the fact that their slopes must approach zero as the input gets large. This causes a problem when you use steepest descent to train a multilayer network with sigmoid			

functions, because the gradient can have a very small magnitude and, therefore, cause small changes in the weights and biases, even though the weights and biases are far from their optimal values.

The purpose of the resilient backpropagation (Rprop) training algorithm is to eliminate these harmful effects of the magnitudes of the partial derivatives. Only the sign of the derivative can determine the direction of the weight update; the magnitude of the derivative has no effect on the weight update. The size of the weight change is determined by a separate update value. The update value for each weight and bias is increased by a factor delt inc whenever the derivative of the performance function with respect to that weight has the same sign for two successive iterations. The update value is decreased by a factor delt dec whenever the derivative with respect to that weight changes sign from the previous iteration. If the derivative is zero, the update value remains the same. Whenever the weights are oscillating, the weight change is reduced. If the weight continues to change in the same direction for several iterations, the magnitude of the weight change increases. A complete description of the Rprop algorithm is given in [ReBr93].

The following code recreates the previous network and trains it using the Rprop algorithm. The training parameters for trainrp are epochs, show, goal, time, min\_grad, max\_fail, delt\_inc, delt\_dec, delta0, and deltamax. The first eight parameters have been previously discussed. The last two are the initial step size and the maximum step size, respectively. The performance of Rprop is not very sensitive to the settings of the training parameters. For the example below, the training parameters are left at the default values:

```
p = [-1 -1 2 2;0 5 0 5];
t = [-1 -1 1 1];
net = feedforwardnet(3,'trainrp');
net = train(net,p,t);
y = net(p)
```

rprop is generally much faster than the standard steepest descent algorithm. It also has the nice property that it requires only a modest

### trainrp

increase in memory requirements. You do need to store the update values for each weight and bias, which is equivalent to storage of the gradient.

See Also traingdm | traingda | traingdx | trainlm | traincgp | traincgf | traincgb | trainscg | trainoss | trainbfg

Purpose	Unsupervised random order weight/bias training		
Syntax	net.trainFcn = 'trainru' [net,tr] = train(net,)		
Description	trainru is not called directly. Instead it is called by train for networks whose net.trainFcn property is set to 'trainru', thus:		
	net.trainFcn =	'trainru	r
	[net,tr] = trai	n(net,	.)
	trainru trains a network with weight and bias learning rules with incremental updates after each presentation of an input. Inputs are presented in random order.		
	Training occurs according to trainr's training parameters, shown here with their default values:		
net.trainParam.	epochs	100	Maximum number of epochs to train
net.trainParam.	goal	0	Performance goal
net.trainParam.	show	25	Epochs between displays (NaN for no displays)
net.trainParam.	showCommandLine	0	Generate command-line output
net.trainParam.	showWindow	1	Show training GUI
net.trainParam.	time	Inf	Maximum time to train in seconds
Network	To prepare a custo	m netwo	rk to be trained with trainru,

_	_	-	_	-	
I	J	5	ρ		

- 1 Set net.trainFcn to 'trainr'. This sets net.trainParam to trainru's default parameters.
- 2 Set each net.inputWeights{i,j}.learnFcn to a learning function.
- **3** Set each net.layerWeights{i,j}.learnFcn to a learning function.

# trainru

	<b>4</b> Set each net.biases{i}.learnFcn to a learning function. (Weight and bias learning parameters are automatically set to default values for the given learning function.)			
	To train the network,			
	1 Set net.trainParam properties to desired values.			
	<b>2</b> Set weight and bias learning parameters to desired values.			
	<b>3</b> Call train.			
Algorithms	For each epoch, all training vectors (or sequences) are each presented once in a different random order, with the network and weight and bias values updated accordingly after each individual presentation.			
	Training stops when any of these conditions is met:			
	• The maximum number of epochs (repetitions) is reached.			
	• The maximum amount of time is exceeded.			
See Also	train   trainr			

Purpose	Sequential order incremental training with learning functions
Syntax	net.trainFcn = 'trains' [net,tr] = train(net,)
Description	trains is not called directly. Instead it is called by train for networks whose net.trainFcn property is set to 'trains', thus:
	net.trainFcn = 'trains'
	[net,tr] = train(net,)
	trains trains a network with weight and bias learning rules with sequential updates. The sequence of inputs is presented to the network with updates occurring after each time step.
	This incremental training algorithm is commonly used for adaptive applications.
Network Use	You can create a standard network that uses trains for adapting by calling perceptron or linearlayer.
	To prepare a custom network to adapt with trains,
	Set net.adaptFcn to 'trains'. This sets net.adaptParam to trains's default parameters.
	2 Set each net.inputWeights{i,j}.learnFcn to a learning function. Set each net.layerWeights{i,j}.learnFcn to a learning function. Set each net.biases{i}.learnFcn to a learning function. (Weight and bias learning parameters are automatically set to default values for the given learning function.)
	To allow the network to adapt,
	<b>1</b> Set weight and bias learning parameters to desired values.
	<b>2</b> Call adapt.

# trains

	See help perceptron and help linearlayer for adaption examples.
Algorithms	Each weight and bias is updated according to its learning function after each time step in the input sequence.
See Also	train   trainb   trainc   trainr

Purpose	Scaled conjugate g	gradient k	packpropagation
Syntax	net.trainFcn = [net,tr] = trai		-
Description	-		ning function that updates weight and bias led conjugate gradient method.
	net.trainFcn =	'trainsc	.a ,
	[net,tr] = trai	n(net,	.)
	Training occurs ac here with their de	-	o <b>trainscg</b> 's training parameters, shown aes:
net.trainParam.e	epochs	100	Maximum number of epochs to train
net.trainParam.	show	25	Epochs between displays (NaN for no displays)
net.trainParam.s	showCommandLine	0	Generate command-line output
net.trainParam.showWindow		1	Show training GUI
net.trainParam.goal		0	Performance goal
net.trainParam.time		inf	Maximum time to train in seconds
net.trainParam.min_grad		1e-6	Minimum performance gradient
net.trainParam.max_fail		5	Maximum validation failures
net.trainParam.sigma		5.0e-5	Determine change in weight for second derivative approximation
net.trainParam.]	Lambda	5.0e-7	Parameter for regulating the indefiniteness of the Hessian

# NetworkYou can create a standard network that uses trainscg withUsefeedforwardnet or cascadeforwardnet. To prepare a custom network<br/>to be trained with trainscg,

	1 Set net.trainFcn to 'trainscg'. This sets net.trainParam to trainscg's default parameters.
	<b>2</b> Set net.trainParam properties to desired values.
	In either case, calling train with the resulting network trains the network with trainscg.
Examples	Here is a problem consisting of inputs p and targets t to be solved with a network.
	p = [0 1 2 3 4 5]; t = [0 0 0 1 1 1];
	A two-layer feed-forward network with two hidden neurons and this training function is created.
	<pre>net = feedforwardnet(2,'trainscg');</pre>
	Here the network is trained and retested.
	net = train(net,p,t); a = net(p)
	See help feedforwardnet and help cascadeforwardnet for other examples.
Algorithms	trainscg can train any network as long as its weight, net input, and transfer functions have derivative functions. Backpropagation is used to calculate derivatives of performance perf with respect to the weight and bias variables X.
	The scaled conjugate gradient algorithm is based on conjugate directions, as in traincgp, traincgf, and traincgb, but this algorithm does not perform a line search at each iteration. See Moller ( <i>Neural Networks</i> , Vol. 6, 1993, pp. 525–533) for a more detailed discussion of the scaled conjugate gradient algorithm.
	Training stops when any of these conditions occurs:

The maximum number of epochs (repetitions) is reached.
 The maximum amount of time is exceeded.
 Performance is minimized to the goal.
 The performance gradient falls below min\_grad.
 Validation performance has increased more than max\_fail times since the last time it decreased (when using validation).
 References Moller, Neural Networks, Vol. 6, 1993, pp. 525–533
 See Also traingdm | traingda | traingdx | trainlm | trainrp | traincgf | traincgb | trainbfg | traincgp | trainoss

# tribas

Purpose	Triangular basis	transfer function	
Graph and Symbol	a $+1$ $-1$ $0$ $+1$ $-1$ $a = tribas(n)$ Triangular Basis	n 🔼	
Syntax	A = tribas(N,F	P)	
Description	tribas is a neural transfer function. Transfer functions calculate a layer's output from its net input.		
	A = tribas(N,FP) takes N and optional function parameters,		
	Ν	S-by-Q matrix of net input (column) vectors	
	FP	Struct of function parameters (ignored)	
	and returns A, an S-by-Q matrix of the triangular basis function app to each element of N.		
	<pre>info = tribas( information:</pre>	$^{\prime}\textit{code}^{\prime}$ ) can take the following forms to return specific	
	tribas('name')	returns the name of this function.	
tribas('output', FP) returns the [min max] output range.			
	tribas('active	', FP) returns the [min max] active input range.	
	tribas('fullde S-by-S-by-Q or S-	riv') returns 1 or 0, depending on whether dA_dN is by-Q.	
	tribas('fpname	<b>s</b> ') returns the names of the function parameters.	

tribas('fpdefaults') returns the default function parameters.

Examples	Here you create a plot of the tribas transfer function.	
	n = -5:0.1:5; a = tribas(n); plot(n,a)	
	Assign this transfer function to layer i of a network.	
	<pre>net.layers{i}.transferFcn = 'tribas';</pre>	
Algorithms	a = tribas(n) = 1 - abs(n), if -1 <= n <= 1 = 0, otherwise	
See Also	sim   radbas	

# tritop

Purpose	Triangle layer topology function	
Syntax	<pre>pos = triptop(dim1,dim2,,dimN)</pre>	
Description	tritop calculates neuron positions for layers whose neurons are arranged in an N-dimensional triangular grid. pos = triptop(dim1,dim2,,dimN) takes N arguments,	
	dimi Length of layer in dimension i	
	and returns an N-by-S matrix of N coordinate vectors, where S is the product of dim1*dim2**dimN.	
Examples	This code creates and displays a two-dimensional layer with 40 neurons arranged in an 8-by-5 triangular grid.	
	<pre>pos = tritop(8,5); net = selforgmap([8 5],'topologyFcn','tritop'); plotsomtop(net)</pre>	
See Also	gridtop   hextop   randtop	

Purpose	Unconfigure network inputs and outputs
Syntax	unconfigure(net) unconfigure(net, 'inputs', i) unconfigure(net, 'outputs', i)
Description	unconfigure(net) returns a network with its input and output sizes set to 0, its input and output processing settings and related weight initialization settings set to values consistent with zero-sized signals. The new network will be ready to be reconfigured for data of the same or different dimensions than it was previously configured for.
	unconfigure(net, 'inputs', i) unconfigures the inputs indicated by the indices i. If no indices are specified, all inputs are unconfigured.
	unconfigure(net, 'outputs', i) unconfigures the outputs indicated by the indices i. If no indices are specified, all outputs are unconfigured.
Examples	Here a network is configured for a simple fitting problem, and then unconfigured.
	<pre>[x,t] = simplefit_dataset; net = fitnet(10); view(net) net = configure(net,x,t); view(net) net = unconfigure(net) view(net)</pre>
See Also	configure   isconfigured

# vec2ind

Purpose	Convert vectors to indices	
Syntax	ind2vec vec2ind	
Description	ind2vec and vec2ind allow indices to be represented either by themselves or as vectors containing a 1 in the row of the index they represent.	
	vec2ind(vec) takes one argument,	
	vec Matrix of vectors, each containing a single 1	
	and returns the indices of the 1s.	
Examples	Here four vectors (each containing only one "1" element) are defined, and the indices of the 1s are found.	
	<pre>vec = [1 0 0 0; 0 0 1 0; 0 1 0 1] ind = vec2ind(vec)</pre>	
See Also	ind2vec	

- Syntax view(net)
- **Description** view(net) launches a window that shows your neural network (specified in net) as a graphical diagram.

## view

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