Statistics Toolbox

For Use with MATLAB®

Computation

Visualization

Programming



User's Guide

Version 4

How to Contact The MathWorks:

	www.mathworks.com comp.soft-sys.matlab	Web Newsgroup
@	support@mathworks.com suggest@mathworks.com bugs@mathworks.com doc@mathworks.com service@mathworks.com info@mathworks.com	Technical support Product enhancement suggestions Bug reports Documentation error reports Order status, license renewals, passcodes Sales, pricing, and general information
1	508-647-7000	Phone
	508-647-7001	Fax
	The MathWorks, Inc. 3 Apple Hill Drive Natick, MA 01760-2098	Mail

For contact information about worldwide offices, see the MathWorks Web site.

Statistics Toolbox User's Guide

© COPYRIGHT 1993 - 2002 by The MathWorks, Inc.

The software described in this document is furnished under a license agreement. The software may be used or copied only under the terms of the license agreement. No part of this manual may be photocopied or reproduced in any form without prior written consent from The MathWorks, Inc.

FEDERAL ACQUISITION: This provision applies to all acquisitions of the Program and Documentation by or for the federal government of the United States. By accepting delivery of the Program, the government hereby agrees that this software qualifies as "commercial" computer software within the meaning of FAR Part 12.212, DFARS Part 227.7202-1, DFARS Part 27.7202-3, DFARS Part 252.227-7013, and DFARS Part 252.227-7014. The terms and conditions of The MathWorks, Inc. Software License Agreement shall pertain to the government's use and disclosure of the Program and Documentation, and shall supersede any conflicting contractual terms or conditions. If this license fails to meet the government's minimum needs or is inconsistent in any respect with federal procurement law, the government agrees to return the Program and Documentation, unused, to MathWorks.

MATLAB, Simulink, Stateflow, Handle Graphics, and Real-Time Workshop are registered trademarks, and TargetBox is a trademark of The MathWorks, Inc.

Other product or brand names are trademarks or registered trademarks of their respective holders.

Printing History:	September 1993	First printing	Version 1
	March 1996	Second printing	Version 2
	January 1997	Third printing	For MATLAB 5
	January 1999	Online only	Revised for Version 2.1.2 (Release 11)
	November 2000	Fourth printing	Revised for Version 3.0 (Release 12)
	May 2001	Fifth printing	Minor revisions
	July 2002	Sixth printing	Revised for Version 4.0 (Release 13)

Contents

Preface

How to Use This Guide	ĸ
Related Products Listxi	i
Typographical Conventions xiv	v

Introduction

1

What Is the Statistics Toolbox?	1-2
Primary Topic Areas	1-3
Random Numbers in Examples	1-5
Mathematical Notation	1-6

Probability Distributions

2

Introduction	2-2
Overview of the Functions	2-4
Probability Density Function (pdf)	2-4
Cumulative Distribution Function (cdf)	2-5
Inverse Cumulative Distribution Function	2-5
Random Number Generator	2-7

]	Mean and Variance as a Function of Parameters	2-9
Ov	erview of the Distributions	2-11
]	Beta Distribution	2-11
]	Binomial Distribution	2-13
(Chi-Square Distribution	2-16
]	Noncentral Chi-Square Distribution	2-17
]	Discrete Uniform Distribution	2-19
]	Exponential Distribution	2-20
]	F Distribution	2-22
]	Noncentral F Distribution	2-23
(Gamma Distribution	2-25
(Geometric Distribution	2-27
]	Hypergeometric Distribution	2-28
	Lognormal Distribution	
	Negative Binomial Distribution	
	Normal Distribution	
]	Poisson Distribution	2-36
	Rayleigh Distribution	
	Student's t Distribution	
	Noncentral t Distribution	
	Weibull Distribution	

Descriptive Statistics

Introduction	3-2
Measures of Central Tendency (Location)	3-3
Measures of Dispersion	3-5
Functions for Data with Missing Values (NaNs)	3-7
Function for Grouped Data	3-9
Percentiles and Graphical Descriptions 3	-11

3

Percentiles	3-11
Probability Density Estimation	3-13
Empirical Cumulative Distribution Function	3-16
The Bootstrap	3-19

Linear Models

Introduction
One-Way Analysis of Variance (ANOVA)
Example: One-Way ANOVA
Multiple Comparisons
Example: Multiple Comparisons
Two-Way Analysis of Variance (ANOVA)
Example: Two-Way ANOVA 4
N-Way Analysis of Variance 4
Example: N-Way ANOVA with Small Data Set 4
Example: N-Way ANOVA with Large Data Set 4
Multiple Linear Regression 4
Mathematical Foundations of Multiple Linear Regression 4
Example: Multiple Linear Regression 4
Quadratic Response Surface Models 4
Exploring Graphs of Multidimensional Polynomials 4
Stepwise Regression 4
Example: Stepwise Regression 4
Stepwise Regression Plot 4
Stepwise Regression Diagnostics Table 4

seneralized Linear Models		4-20
Example: Generalized Linea	r Models	4-28

Robust and Nonparametric Methods		4-32
Robust Regression		4-32
Kruskal-Wallis Test		4-34
Friedman's Test	••	4-34

Nonlinear Regression Models

5

6

Introduction	5-2
Nonlinear Least Squares	5-3
Example: Nonlinear Modeling	5-3
An Interactive GUI for Nonlinear Fitting and Prediction	5-7
Regression and Classification Trees	5-8

Hypothesis Tests

Introduction	6-2
Hypothesis Test Terminology	6-3
Hypothesis Test Assumptions	6-4
Example: Hypothesis Testing	6-5
Available Hypothesis Tests	6-9

Introduction	. 7-2
Principal Components Analysis	7-3
Example: Principal Components Analysis	
The Principal Components (First Output)	
The Component Scores (Second Output)	
The Component Variances (Third Output)	
Hotelling's T^2 (Fourth Output)	
Factor Analysis	7-13
Example: Finding Common Factors Affecting Stock Prices	7-13
Factor Rotation	7-16
Predicting Factor Scores	7-17
Comparison of Factor Analysis and Principal Components	
Analysis	7-19
Multivariate Analysis of Variance (MANOVA)	7-20
Example: Multivariate Analysis of Variance	7-20
Cluster Analysis	7-26
Hierarchical Clustering	7-26
K-Means Clustering	7-40
Classical Multidimensional Scaling	7-47
Overview	7-47
Reconstructing a Map from Inter-City Distances	7-49

Statistical Plots

8

7

Introduction	••	••	•••	•	 • •	•	•	•	•	 •	•	•	•	 •	•	•	 •	•	•	•	 •	•	•	• •	•	• •	8	3-2
Box Plots					 								•														٤	3-3

Distribution Plots	3-4
Normal Probability Plots 8	3-4
Quantile-Quantile Plots 8	3-6
Weibull Probability Plots 8	3-7
Empirical Cumulative Distribution Function (CDF) 8	3-8
Scatter Plots	·10

Statistical Process Control

9

Introduction	•••		• •		• •	•	•••	•	•••	•	• •		•	 •	•		•	•	•		•	•	•	9-2
Control Charts																								9-3
Xbar Charts						•															•			9-3
S Charts																					•			9-4
EWMA Charts		••	•••	• •	• •	•	• •	•	•••	•	• •	•••	•		•	• •	•	•	•	•••	•	•	•	9-5
Capability Studies						•				•			•		•						•			9-6

Design of Experiments

10 [

Introduction	10-2
Full Factorial Designs	10-4
Fractional Factorial Designs	10-6
Response Surface Designs Central Composite Designs Box-Behnken Designs	10-8
D-Optimal Designs	10-11

Generating D-Optimal Designs	10-11
Augmenting D-Optimal Designs	10-14
Designing Experiments with Uncontrolled Inputs	10-16
Controlling Candidate Points	10-16
Including Categorical Factors	10-17

Demos

11 [

Introduction 11-2	2
The disttool Demo	3
The polytool Demo 11-5	
Confidence Bounds 11-7 Overfitting 11-8	
The aoctool Demo 11-10 Example: aoctool with Sample Data 11-11	
The randtool Demo 11-18	3
The rsmdemo Demo 11-19	•
Part 1 11-19)
Part 2 11-20)
The glmdemo Demo 11-21	L
The robustdemo Demo 11-22	2

Reference

12 [

Functions – By Category 12-3

Probability Distributions	12-4
Descriptive Statistics	12-9
Statistical Plotting	12-10
Statistical Process Control	12-11
Linear Models	12-11
Nonlinear Regression	12-12
Design of Experiments	12-13
Multivariate Statistics	12-13
Decision Tree Techniques	12-15
Hypothesis Tests	12-15
Distribution Testing	12-15
Nonparametric Testing	12-15
File I/O	12-16
Demonstrations	12-16
Data	12-16
Functions — Alphabetical List	12-18

Selected Bibliography



Index

Preface

How to Use This Guide	•	•	•	•	•	•	•	•	•	•	•	•	•	. x
Related Products List	•			•	•	•			•		•	•	•	.xii
Typographical Conventions						•						•		xiv

How to Use This Guide

This guide introduces the MATLAB statistics environment through the toolbox functions. It describes the functions with regard to particular areas of interest, such as probability distributions, linear and nonlinear models, principal components analysis, design of experiments, statistical process control, and descriptive statistics. It also describes use of the graphical tools.

"Introduction"	Introduces the toolbox, and explains the mathematical notation it uses.
"Probability Distributions"	Describes the distributions and the distribution-related functions supported by the toolbox.
"Descriptive Statistics"	Explores toolbox features for working with descriptive statistics such as measures of location and spread, percentile estimates, and data with missing values.
"Linear Models"	Describes toolbox support for one-way, two-way, and higher-way analysis of variance (ANOVA), analysis of covariance (ANOCOVA), multiple linear regression, stepwise regression, response surface prediction, ridge regression, and one-way multivariate analysis of variance (MANOVA). It also describes support for nonparametric versions of one- and two-way ANOVA, and multiple comparisons of the estimates produced by ANOVA and ANOCOVA functions.
"Nonlinear Regression Models"	Discusses parameter estimation, interactive prediction and visualization of multidimensional nonlinear fits, and confidence intervals for parameters and predicted values. It also introduces classification and regression trees as a way to approximate a regression relationship.
"Hypothesis Tests"	Describes support for common tests of hypothesis – t-tests, Z-tests, nonparametric tests, and distribution tests.
"Multivariate Statistics"	Explores toolbox features that support methods in multivariate statistics, including principal components analysis, factor analysis, one-way multivariate analysis of variance, cluster analysis, and classical multidimensional scaling.

"Statistical Plots"	Describes box plots, normal probability plots, Weibull probability plots, control charts, and quantile-quantile plots which the toolbox adds to the arsenal of graphs in MATLAB. It also discusses extended support for polynomial curve fitting and prediction, creation of scatter plots or matrices of scatter plots for grouped data, interactive identification of points on such plots, and interactive exploration of a fitted regression model.
"Statistical Process Control"	Discusses the plotting of common control charts and the performing of process capability studies.
"Design of Experiments"	Discusses toolbox support for full and fractional factorial designs, response surface designs, and D-optimal designs. It also describes functions for generating designs, augmenting designs, and optimally assigning units with fixed covariates.
"Demos"	Describes GUIs that enable you to explore the probability distributions, random number generation, curve fitting, and design of experiments functions.
"Reference"	Lists the functions for each area supported by the toolbox, and provides a complete description of each function.
"Selected Bibliography"	Lists published materials that support concepts described in this guide.
PDF versi description	on about specific functions and tools is available online and in the on of this document. For functions and graphical tools, reference ns include a synopsis of the syntax, as well as a complete explanation and operation. Many reference descriptions also include examples, a

description of the function's algorithm, and references to additional reading material. "Demos" on page 11-1 further describes the use of the graphical tools.

Related Products List

The MathWorks provides several products that may be relevant to the kinds of tasks you can perform with the Statistics Toolbox.

For more information about any of these products, see either:

- The online documentation for that product if it is installed or if you are reading the documentation from the CD
- The MathWorks Web site, at http://www.mathworks.com; see the "products" section

Note The toolboxes listed below all include functions that extend the MATLAB capabilities. The blocksets all include blocks that extend Simulink's capabilities.

Product	Description	
Data Acquisition Toolbox	Acquire and send out data from plug-in data acquisition boards	
Database Toolbox	Exchange data with relational databases	
Financial Time Series Toolbox	Analyze and manage financial time series data	
Financial Toolbox	Model financial data and develop financial analysis algorithms	
GARCH Toolbox	Analyze financial volatility using univariate GARCH models	
Image Processing Toolbox	Perform image processing, analysis, and algorithm development	
Mapping Toolbox	Analyze and visualize geographically based information	

Product	Description
Neural Network Toolbox	Design and simulate neural networks
Optimization Toolbox	Solve standard and large-scale optimization problems
Signal Processing Toolbox	Perform signal processing, analysis, and algorithm development
System Identification Toolbox	Create linear dynamic models from measured input-output data

Typographical Conventions

This manual uses some or all of these conventions.

ltem	Convention	Example
Example code	Monospace font	To assign the value 5 to A, enter A = 5
Function names, syntax, filenames, directory/folder names, and user input	Monospace font	The cos function finds the cosine of each array element. Syntax line example is MLGetVar ML_var_name
Buttons and keys	Boldface with book title caps	Press the Enter key.
Literal strings (in syntax descriptions in reference chapters)	Monospace bold for literals	<pre>f = freqspace(n,'whole')</pre>
Mathematical expressions	<i>Italics</i> for variables Standard text font for functions, operators, and constants	This vector represents the polynomial $p = x^2 + 2x + 3$.
MATLAB output	Monospace font	MATLAB responds with A = 5
Menu and dialog box titles	Boldface with book title caps	Choose the File Options menu.
New terms and for emphasis	Italics	An <i>array</i> is an ordered collection of information.
Omitted input arguments	() ellipsis denotes all of the input/output arguments from preceding syntaxes.	[c,ia,ib] = union()
String variables (from a finite list)	Monospace italics	<pre>sysc = d2c(sysd,'method')</pre>

Introduction

What Is the Statistics Toolbox?	•	•	•	•	•	·	•	•	·	•	•	1-2
Primary Topic Areas	•		•				•			•		1-3
Random Numbers in Examples	•	•	•		•	•	•	•	•	•	•	1-5
Mathematical Notation												1-6

1

What Is the Statistics Toolbox?

The Statistics Toolbox, for use with MATLAB, supplies basic statistics capability on the level of a first course in engineering or scientific statistics. The statistics functions it provides are building blocks suitable for use inside other analytical tools.

The Statistics Toolbox is a collection of tools built on the MATLAB[®] numeric computing environment. The toolbox supports a wide range of common statistical tasks, from random number generation, to curve fitting, to design of experiments and statistical process control. The toolbox provides two categories of tools:

- Building-block probability and statistics functions
- Graphical, interactive tools

The first category of tools is made up of functions that you can call from the command line or from your own applications. Many of these functions are MATLAB M-files, series of MATLAB statements that implement specialized statistics algorithms. You can view the MATLAB code for these functions using the statement

```
type function_name
```

You can change the way any toolbox function works by copying and renaming the M-file, then modifying your copy. You can also extend the toolbox by adding your own M-files.

Secondly, the toolbox provides a number of interactive tools that let you access many of the functions through a graphical user interface (GUI). Together, the GUI-based tools provide an environment for polynomial fitting and prediction, as well as probability function exploration.

Primary Topic Areas

The Statistics Toolbox has more than 200 M-files, supporting work in these topical areas:

Probability Distributions

The Statistics Toolbox supports 20 probability distributions. For each distribution there are five associated functions. They are

- Probability density function (pdf)
- Cumulative distribution function (cdf)
- Inverse of the cumulative distribution function
- Random number generator
- Mean and variance as a function of the parameters

For data-driven distributions (beta, binomial, exponential, gamma, normal, Poisson, uniform, and Weibull), the Statistics Toolbox has functions for computing parameter estimates and confidence intervals.

Descriptive Statistics

The Statistics Toolbox provides functions for describing the features of a data sample. These descriptive statistics include measures of location and spread, percentile estimates and functions for dealing with data having missing values.

Linear Models

In the area of linear models, the Statistics Toolbox supports one-way, two-way, and higher-way analysis of variance (ANOVA), analysis of covariance (ANOCOVA), multiple linear regression, stepwise regression, response surface prediction, ridge regression, and one-way multivariate analysis of variance (MANOVA). It supports nonparametric versions of one- and two-way ANOVA. It also supports multiple comparisons of the estimates produced by ANOVA and ANOCOVA functions.

Nonlinear Models

For nonlinear models, the Statistics Toolbox provides functions for parameter estimation, interactive prediction and visualization of multidimensional nonlinear fits, and confidence intervals for parameters and predicted values. It

provides functions for using classification and regression trees to approximate regression relationships.

Hypothesis Tests

The Statistics Toolbox also provides functions that do the most common tests of hypothesis — t-tests, Z-tests, nonparametric tests, and distribution tests.

Multivariate Statistics

The Statistics Toolbox supports methods in multivariate statistics, including principal components analysis, factor analysis, one-way multivariate analysis of variance, cluster analysis, and classical multidimensional scaling.

Statistical Plots

The Statistics Toolbox adds box plots, normal probability plots, Weibull probability plots, control charts, and quantile-quantile plots to the arsenal of graphs in MATLAB. There is also extended support for polynomial curve fitting and prediction. There are functions to create scatter plots or matrices of scatter plots for grouped data, and to identify points interactively on such plots. There is a function to interactively explore a fitted regression model.

Statistical Process Control (SPC)

For SPC, the Statistics Toolbox provides functions for plotting common control charts and performing process capability studies.

Design of Experiments (DOE)

The Statistics Toolbox supports full and fractional factorial designs, response surface designs, and D-optimal designs. There are functions for generating designs, augmenting designs, and optimally assigning units with fixed covariates.

Random Numbers in Examples

The random number generation functions for various probability distributions are based on the primitive functions, randn and rand. There are many examples that start by generating data using random numbers. To duplicate the results in these examples, first execute the commands below.

```
seed = 931316785;
rand('seed',seed);
randn('seed',seed);
```

You might want to save these commands in an M-file script called initseed.m. Then, instead of three separate commands, you need only type initseed.

Note For rand and randn, the 'seed' syntax is replaced by the 'state' syntax in MATLAB Version 5. Although use of the 'seed' syntax is backward compatible in MATLAB Version 6.5, you should avoid its use. This book will be updated to use the 'state' syntax in a future release.

1

Mathematical Notation

This manual and the Statistics Toolbox functions use the following mathematical notation conventions.

β	Parameters in a linear model.			
E(x)	Expected value of x. $E(x) = \int t f(t) dt$			
$f(x \mid a, b)$	Probability density function. x is the independent variable; a and b are fixed parameters.			
$F(x \mid a, b)$	Cumulative distribution function.			
$I([a, b])$ or $I_{[a, b]}$	Indicator function. In this example the function takes the value 1 on the closed interval from a to b and is 0 elsewhere.			
p and q	p is the probability of some event. q is the probability of $\sim p$, so $q = 1-p$.			

2

Probability Distributions

Introduction	2-2
Overview of the Functions	2-4
Probability Density Function (pdf)	2-4
Cumulative Distribution Function (cdf)	
Inverse Cumulative Distribution Function	
Random Number Generator	
Mean and Variance as a Function of Parameters \ldots \ldots \ldots	
Overview of the Distributions	-11
Beta Distribution	
Binomial Distribution	
Chi-Square Distribution	
Noncentral Chi-Square Distribution	
Discrete Uniform Distribution	
Exponential Distribution	
F Distribution	
Noncentral F Distribution	
Gamma Distribution	
Geometric Distribution	
Hypergeometric Distribution	-28
Lognormal Distribution	
Negative Binomial Distribution	
Normal Distribution	
Poisson Distribution	
Rayleigh Distribution	-38
Student's t Distribution	
Noncentral t Distribution	
Weibull Distribution	

Introduction

Probability distributions arise from experiments where the outcome is subject to chance. The nature of the experiment dictates which probability distributions may be appropriate for modeling the resulting random outcomes. There are two types of probability distributions – *continuous* and *discrete*.

Continuous (data)	Continuous (statistics)	Discrete
Beta	Chi-square	Binomial
Exponential	Noncentral Chi-square	Discrete Uniform
Gamma	F	Geometric
Lognormal	Noncentral F	Hypergeometric
Normal	t	Negative Binomial
Rayleigh	Noncentral t	Poisson
Uniform		
Weibull		

Suppose you are studying a machine that produces videotape. One measure of the quality of the tape is the number of visual defects per hundred feet of tape. The result of this experiment is an integer, since you cannot observe 1.5 defects. To model this experiment you should use a discrete probability distribution.

A measure affecting the cost and quality of videotape is its thickness. Thick tape is more expensive to produce, while variation in the thickness of the tape on the reel increases the likelihood of breakage. Suppose you measure the thickness of the tape every 1000 feet. The resulting numbers can take a continuum of possible values, which suggests using a continuous probability distribution to model the results.

Using a probability model does not allow you to predict the result of any individual experiment but you can determine the probability that a given outcome will fall inside a specific range of values. This following two sections provide more information about the available distributions:

- "Overview of the Functions" on page 2-4
- "Overview of the Distributions" on page 2-11

Overview of the Functions

The Statistics Toolbox provides five functions for each distribution. They are discussed in the following sections:

- "Probability Density Function (pdf)" on page 2-4
- "Cumulative Distribution Function (cdf)" on page 2-5
- "Inverse Cumulative Distribution Function" on page 2-5
- "Random Number Generator" on page 2-7
- "Mean and Variance as a Function of Parameters" on page 2-9

Probability Density Function (pdf)

The probability density function (pdf) has a different meaning depending on whether the distribution is discrete or continuous.

For discrete distributions, the pdf is the probability of observing a particular outcome. In our videotape example, the probability that there is exactly one defect in a given hundred feet of tape is the value of the pdf at 1.

Unlike discrete distributions, the pdf of a continuous distribution at a value is not the probability of observing that value. For continuous distributions the probability of observing any particular value is zero. To get probabilities you must integrate the pdf over an interval of interest. For example the probability of the thickness of a videotape being between one and two millimeters is the integral of the appropriate pdf from one to two.

A pdf has two theoretical properties:

- The pdf is zero or positive for every possible outcome.
- The integral of a pdf over its entire range of values is one.

A pdf is not a single function. Rather a pdf is a family of functions characterized by one or more parameters. Once you choose (or estimate) the parameters of a pdf, you have uniquely specified the function.

The pdf function call has the same general format for every distribution in the Statistics Toolbox. The following commands illustrate how to call the pdf for the normal distribution.

```
x = [-3:0.1:3];
f = normpdf(x,0,1);
```

The variable f contains the density of the normal pdf with parameters $\mu=0$ and $\sigma=1$ at the values in x. The first input argument of every pdf is the set of values for which you want to evaluate the density. Other arguments contain as many parameters as are necessary to define the distribution uniquely. The normal distribution requires two parameters; a location parameter (the mean, μ) and a scale parameter (the standard deviation, σ).

Cumulative Distribution Function (cdf)

If f is a probability density function for random variable X, the associated cumulative distribution function (cdf) F is

$$F(x) = P(X \le x) = \int_{-\infty}^{x} f(t) dt$$

The cdf of a value x, F(x), is the probability of observing any outcome less than or equal to x.

A cdf has two theoretical properties:

- The cdf ranges from 0 to 1.
- If y > x, then the cdf of y is greater than or equal to the cdf of x.

The cdf function call has the same general format for every distribution in the Statistics Toolbox. The following commands illustrate how to call the cdf for the normal distribution.

```
x = [-3:0.1:3];
p = normcdf(x,0,1);
```

The variable p contains the probabilities associated with the normal cdf with parameters $\mu=0$ and $\sigma=1$ at the values in x. The first input argument of every cdf is the set of values for which you want to evaluate the probability. Other arguments contain as many parameters as are necessary to define the distribution uniquely.

Inverse Cumulative Distribution Function

The inverse cumulative distribution function returns critical values for hypothesis testing given significance probabilities. To understand the relationship between a continuous cdf and its inverse function, try the following:

```
x = [-3:0.1:3];
xnew = norminv(normcdf(x,0,1),0,1);
```

How does xnew compare with x? Conversely, try this:

```
p = [0.1:0.1:0.9];
pnew = normcdf(norminv(p,0,1),0,1);
```

How does pnew compare with p?

Calculating the cdf of values in the domain of a continuous distribution returns probabilities between zero and one. Applying the inverse cdf to these probabilities yields the original values.

For discrete distributions, the relationship between a cdf and its inverse function is more complicated. It is likely that there is no x value such that the cdf of x yields p. In these cases the inverse function returns the first value x such that the cdf of x equals or exceeds p. Try this:

```
x = [0:10];
y = binoinv(binocdf(x,10,0.5),10,0.5);
```

How does x compare with y?

The commands below illustrate the problem with reconstructing the probability p from the value x for discrete distributions.

```
p = [0.1:0.2:0.9];
pnew = binocdf(binoinv(p,10,0.5),10,0.5)
pnew =
        0.1719        0.3770        0.6230        0.8281        0.9453
```

The inverse function is useful in hypothesis testing and production of confidence intervals. Here is the way to get a 99% confidence interval for a normally distributed sample.

```
p = [0.005 0.995];
x = norminv(p,0,1)
x =
-2.5758 2.5758
```

The variable x contains the values associated with the normal inverse function with parameters $\mu=0$ and $\sigma=1$ at the probabilities in p. The difference p(2) - p(1) is 0.99. Thus, the values in x define an interval that contains 99% of the standard normal probability.

The inverse function call has the same general format for every distribution in the Statistics Toolbox. The first input argument of every inverse function is the set of probabilities for which you want to evaluate the critical values. Other arguments contain as many parameters as are necessary to define the distribution uniquely.

Random Number Generator

The methods for generating random numbers from any distribution all start with uniform random numbers. Once you have a uniform random number generator, you can produce random numbers from other distributions either directly or by using inversion or rejection methods, described below. See "Syntax for Random Number Functions" on page 2-8 for details on using generator functions.

Direct

Direct methods flow from the definition of the distribution.

As an example, consider generating binomial random numbers. You can think of binomial random numbers as the number of heads in n tosses of a coin with probability p of a heads on any toss. If you generate n uniform random numbers and count the number that are greater than p, the result is binomial with parameters n and p.

Inversion

The inversion method works due to a fundamental theorem that relates the uniform distribution to other continuous distributions.

If *F* is a continuous distribution with inverse F^{-1} , and U is a uniform random number, then $F^{-1}(U)$ has distribution *F*.

So, you can generate a random number from a distribution by applying the inverse function for that distribution to a uniform random number. Unfortunately, this approach is usually not the most efficient.

Rejection

The functional form of some distributions makes it difficult or time consuming to generate random numbers using direct or inversion methods. Rejection methods can sometimes provide an elegant solution in these cases.

Suppose you want to generate random numbers from a distribution with pdf f. To use rejection methods you must first find another density, g, and a constant, c, so that the inequality below holds.

 $f(x) \le cg(x) \forall x$

You then generate the random numbers you want using the following steps:

- 1 Generate a random number *x* from distribution *G* with density *g*.
- **2** Form the ratio $r = \frac{cg(x)}{f(x)}$.
- **3** Generate a uniform random number *u*.
- **4** If the product of *u* and *r* is less than one, return *x*.
- 5 Otherwise repeat steps one to three.

For efficiency you need a cheap method for generating random numbers from G, and the scalar c should be small. The expected number of iterations is c.

Syntax for Random Number Functions

You can generate random numbers from each distribution. This function provides a single random number or a matrix of random numbers, depending on the arguments you specify in the function call.

For example, here is the way to generate random numbers from the beta distribution. Four statements obtain random numbers: the first returns a single number, the second returns a 2-by-2 matrix of random numbers, and the third and fourth return 2-by-3 matrices of random numbers.

```
a = 1;
b = 2;
c = [.1 .5; 1 2];
d = [.25 .75; 5 10];
m = [2 3];
```

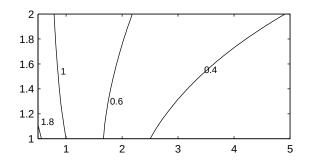
```
nrow = 2;
ncol = 3;
r1 = betarnd(a,b)
r1 =
    0.4469
r2 = betarnd(c,d)
r2 =
    0.8931
               0.4832
    0.1316
               0.2403
r3 = betarnd(a, b, m)
r3 =
    0.4196
               0.6078
                         0.1392
    0.0410
               0.0723
                         0.0782
r4 = betarnd(a,b,nrow,ncol)
r4 =
    0.0520
               0.3975
                         0.1284
    0.3891
               0.1848
                         0.5186
```

Mean and Variance as a Function of Parameters

The mean and variance of a probability distribution are generally simple functions of the parameters of the distribution. The Statistics Toolbox functions ending in "stat" all produce the mean and variance of the desired distribution for the given parameters.

The example below shows a contour plot of the mean of the Weibull distribution as a function of the parameters.

```
x = (0.5:0.1:5);
y = (1:0.04:2);
[X,Y] = meshgrid(x,y);
Z = weibstat(X,Y);
[c,h] = contour(x,y,Z,[0.4 0.6 1.0 1.8]);
clabel(c);
```



Overview of the Distributions

The following sections describe the available probability distributions:

- "Beta Distribution" on page 2-11
- "Binomial Distribution" on page 2-13
- "Chi-Square Distribution" on page 2-16
- "Noncentral Chi-Square Distribution" on page 2-17
- "Discrete Uniform Distribution" on page 2-19
- "Exponential Distribution" on page 2-20
- "F Distribution" on page 2-22
- "Noncentral F Distribution" on page 2-23
- "Gamma Distribution" on page 2-25
- "Geometric Distribution" on page 2-27
- "Hypergeometric Distribution" on page 2-28
- "Lognormal Distribution" on page 2-29
- "Negative Binomial Distribution" on page 2-31
- "Normal Distribution" on page 2-34
- "Poisson Distribution" on page 2-36
- "Rayleigh Distribution" on page 2-38
- "Student's t Distribution" on page 2-39
- "Noncentral t Distribution" on page 2-40
- "Uniform (Continuous) Distribution" on page 2-42
- "Weibull Distribution" on page 2-43

Beta Distribution

The following sections provide an overview of the beta distribution.

Background on the Beta Distribution

The beta distribution describes a family of curves that are unique in that they are nonzero only on the interval $(0\ 1)$. A more general version of the function assigns parameters to the end-points of the interval.

The beta cdf is the same as the incomplete beta function.

The beta distribution has a functional relationship with the t distribution. If Y is an observation from Student's t distribution with v degrees of freedom, then the following transformation generates X, which is beta distributed.

$$X = \frac{1}{2} + \frac{1}{2} \frac{Y}{\sqrt{\nu + Y^2}}$$

if
$$Y \sim t(v)$$
 then $X \sim \beta\left(\frac{v}{2}, \frac{v}{2}\right)$

The Statistics Toolbox uses this relationship to compute values of the t cdf and inverse function as well as generating t distributed random numbers.

Definition of the Beta Distribution

The beta pdf is

$$y = f(x|a,b) = \frac{1}{B(a,b)} x^{a-1} (1-x)^{b-1} I_{(0,1)}(x)$$

where $B(\cdot)$ is the Beta function. The indicator function $I_{(0,1)}(x)$ ensures that only values of x in the range (0 1) have nonzero probability.

Parameter Estimation for the Beta Distribution

Suppose you are collecting data that has hard lower and upper bounds of zero and one respectively. Parameter estimation is the process of determining the parameters of the beta distribution that fit this data best in some sense.

One popular criterion of goodness is to maximize the likelihood function. The likelihood has the same form as the beta pdf. But for the pdf, the parameters are known constants and the variable is *x*. The likelihood function reverses the roles of the variables. Here, the sample values (the *x*'s) are already observed. So they are the fixed constants. The variables are the unknown parameters. Maximum likelihood estimation (MLE) involves calculating the values of the parameters that give the highest likelihood given the particular set of data.

The function betafit returns the MLEs and confidence intervals for the parameters of the beta distribution. Here is an example using random numbers from the beta distribution with a = 5 and b = 0.2.

```
r = betarnd(5,0.2,100,1);
[phat, pci] = betafit(r)
```

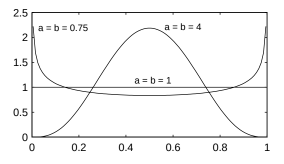
```
phat =
    4.5330 0.2301
pci =
    2.8051 0.1771
    6.2610 0.2832
```

The MLE for parameter a is 4.5330, compared to the true value of 5. The 95% confidence interval for a goes from 2.8051 to 6.2610, which includes the true value.

Similarly the MLE for parameter b is 0.2301, compared to the true value of 0.2. The 95% confidence interval for b goes from 0.1771 to 0.2832, which also includes the true value. Of course, in this made-up example we know the "true value." In experimentation we do not.

Example and Plot of the Beta Distribution

The shape of the beta distribution is quite variable depending on the values of the parameters, as illustrated by the plot below.



The constant pdf (the flat line) shows that the standard uniform distribution is a special case of the beta distribution.

Binomial Distribution

The following sections provide an overview of the binomial distribution.

Background of the Binomial Distribution

The binomial distribution models the total number of successes in repeated trials from an infinite population under the following conditions:

- Only two outcomes are possible on each of *n* trials.
- The probability of success for each trial is constant.
- All trials are independent of each other.

James Bernoulli derived the binomial distribution in 1713 (*Ars Conjectandi*). Earlier, Blaise Pascal had considered the special case where p = 1/2.

Definition of the Binomial Distribution

The binomial pdf is

$$y = f(x|n,p) = {\binom{n}{x}} p^{x} q^{(1-x)} I_{(0,1,...,n)}(x)$$

where
$$\binom{n}{x} = \frac{n!}{x!(n-x)!}$$
 and $q = 1-p$.

The binomial distribution is discrete. For zero and for positive integers less than n, the pdf is nonzero.

Parameter Estimation for the Binomial Distribution

Suppose you are collecting data from a widget manufacturing process, and you record the number of widgets within specification in each batch of 100. You might be interested in the probability that an individual widget is within specification. Parameter estimation is the process of determining the parameter, p, of the binomial distribution that fits this data best in some sense.

One popular criterion of goodness is to maximize the likelihood function. The likelihood has the same form as the binomial pdf above. But for the pdf, the parameters (n and p) are known constants and the variable is x. The likelihood function reverses the roles of the variables. Here, the sample values (the x's) are already observed. So they are the fixed constants. The variables are the unknown parameters. MLE involves calculating the value of p that give the highest likelihood given the particular set of data.

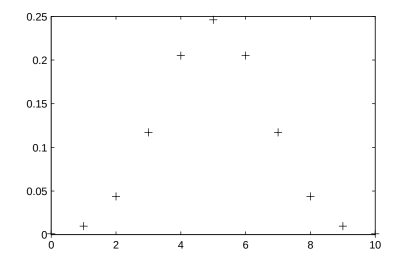
The function binofit returns the MLEs and confidence intervals for the parameters of the binomial distribution. Here is an example using random numbers from the binomial distribution with n = 100 and p = 0.9.

The MLE for parameter p is 0.8800, compared to the true value of 0.9. The 95% confidence interval for p goes from 0.7998 to 0.9364, which includes the true value. Of course, in this made-up example we know the "true value" of p. In experimentation we do not.

Example and Plot of the Binomial Distribution

The following commands generate a plot of the binomial pdf for n = 10 and p = 1/2.

x = 0:10; y = binopdf(x,10,0.5); plot(x,y,'+')



Chi-Square Distribution

The following sections provide an overview of the χ^2 distribution.

Background of the Chi-Square Distribution

The χ^2 distribution is a special case of the gamma distribution where b = 2 in the equation for gamma distribution below.

$$y = f(x|a, b) = \frac{1}{b^{a}\Gamma(a)}x^{a-1}e^{-\frac{x}{b}}$$

The χ^2 distribution gets special attention because of its importance in normal sampling theory. If a set of *n* observations is normally distributed with variance σ^2 , and s^2 is the sample standard deviation, then

$$\frac{(n-1)s^2}{\sigma^2} \sim \chi^2(n-1)$$

The Statistics Toolbox uses the above relationship to calculate confidence intervals for the estimate of the normal parameter σ^2 in the function normfit.

Definition of the Chi-Square Distribution

The χ^2 pdf is

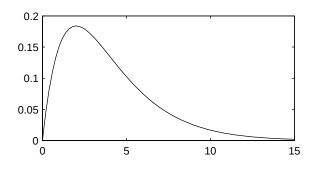
$$y = f(x|v) = \frac{x^{(v-2)/2}e^{-x/2}}{2^{\frac{v}{2}}\Gamma(v/2)}$$

where $\Gamma(\ \cdot\)$ is the Gamma function, and ν is the degrees of freedom.

Example and Plot of the Chi-Square Distribution

The χ^2 distribution is skewed to the right especially for few degrees of freedom (v). The plot shows the χ^2 distribution with four degrees of freedom.

```
x = 0:0.2:15;
y = chi2pdf(x,4);
plot(x,y)
```



Noncentral Chi-Square Distribution

The following sections provide an overview of the noncentral χ^2 distribution.

Background of the Noncentral Chi-Square Distribution

The χ^2 distribution is actually a simple special case of the noncentral chi-square distribution. One way to generate random numbers with a χ^2 distribution (with ν degrees of freedom) is to sum the squares of ν standard normal random numbers (mean equal to zero.)

What if we allow the normally distributed quantities to have a mean other than zero? The sum of squares of these numbers yields the noncentral chi-square

distribution. The noncentral chi-square distribution requires two parameters; the degrees of freedom and the noncentrality parameter. The noncentrality parameter is the sum of the squared means of the normally distributed quantities.

The noncentral chi-square has scientific application in thermodynamics and signal processing. The literature in these areas may refer to it as the Ricean or generalized Rayleigh distribution.

Definition of the Noncentral Chi-Square Distribution

There are many equivalent formulas for the noncentral chi-square distribution function. One formulation uses a modified Bessel function of the first kind. Another uses the generalized Laguerre polynomials. The Statistics Toolbox computes the cumulative distribution function values using a weighted sum of χ^2 probabilities with the weights equal to the probabilities of a Poisson distribution. The Poisson parameter is one-half of the noncentrality parameter of the noncentral chi-square.

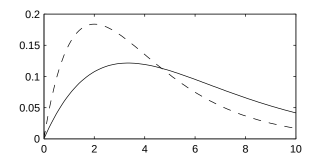
$$F(x|v, \delta) = \sum_{j=0}^{\infty} \left(\frac{\left(\frac{1}{2}\delta\right)^{j}}{j!} e^{-\frac{\delta}{2}} \right) Pr[\chi_{v+2j}^{2} \le x]$$

where $\boldsymbol{\delta}$ is the noncentrality parameter.

Example of the Noncentral Chi-Square Distribution

The following commands generate a plot of the noncentral chi-square pdf.

```
x = (0:0.1:10)';
p1 = ncx2pdf(x,4,2);
p = chi2pdf(x,4);
plot(x,p,'--',x,p1,'-')
```



Discrete Uniform Distribution

The following sections provide an overview of the discrete uniform distribution.

Background of the Discrete Uniform Distribution

The discrete uniform distribution is a simple distribution that puts equal weight on the integers from one to N.

Definition of the Discrete Uniform Distribution

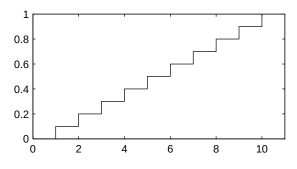
The discrete uniform pdf is

$$y = f(x|N) = \frac{1}{N}I_{(1,...,N)}(x)$$

Example and Plot of the Discrete Uniform Distribution

As for all discrete distributions, the cdf is a step function. The plot shows the discrete uniform cdf for N = 10.

x = 0:10; y = unidcdf(x,10); stairs(x,y) set(gca,'Xlim',[0 11])



To pick a random sample of 10 from a list of 553 items:

```
numbers = unidrnd(553, 1, 10)
```

numbers =

293 372 5 213 37 231 380 326 515 468

Exponential Distribution

The following sections provide an overview of the exponential distribution.

Background of the Exponential Distribution

Like the chi-square distribution, the exponential distribution is a special case of the gamma distribution (obtained by setting a = 1)

$$y = f(x|a,b) = \frac{1}{b^a \Gamma(a)} x^{a-1} e^{-\frac{x}{b}}$$

where $\Gamma(\cdot)$ is the Gamma function.

The exponential distribution is special because of its utility in modeling events that occur randomly over time. The main application area is in studies of lifetimes.

Definition of the Exponential Distribution

The exponential pdf is

$$y = f(x|\mu) = \frac{1}{\mu}e^{-\frac{x}{\mu}}$$

Parameter Estimation for the Exponential Distribution

Suppose you are stress testing light bulbs and collecting data on their lifetimes. You assume that these lifetimes follow an exponential distribution. You want to know how long you can expect the average light bulb to last. Parameter estimation is the process of determining the parameters of the exponential distribution that fit this data best in some sense.

One popular criterion of goodness is to maximize the likelihood function. The likelihood has the same form as the exponential pdf above. But for the pdf, the parameters are known constants and the variable is x. The likelihood function reverses the roles of the variables. Here, the sample values (the x's) are already observed. So they are the fixed constants. The variables are the unknown parameters. MLE involves calculating the values of the parameters that give the highest likelihood given the particular set of data.

The function expfit returns the MLEs and confidence intervals for the parameters of the exponential distribution. Here is an example using random numbers from the exponential distribution with $\mu = 700$.

```
lifetimes = exprnd(700,100,1);
[muhat, muci] = expfit(lifetimes)
muhat =
    672.8207
muci =
    547.4338
    810.9437
```

The MLE for parameter μ is 672, compared to the true value of 700. The 95% confidence interval for μ goes from 547 to 811, which includes the true value.

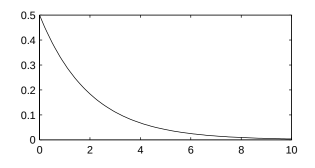
In our life tests we do not know the true value of μ so it is nice to have a confidence interval on the parameter to give a range of likely values.

Example and Plot of the Exponential Distribution

For exponentially distributed lifetimes, the probability that an item will survive an extra unit of time is independent of the current age of the item. The example shows a specific case of this special property.

The plot below shows the exponential pdf with its parameter (and mean), $\mu,$ set to 2.

x = 0:0.1:10; y = exppdf(x,2); plot(x,y)



F Distribution

The following sections provide an overview of the F distribution.

Background of the F distribution

The F distribution has a natural relationship with the chi-square distribution. If χ_1 and χ_2 are both chi-square with v_1 and v_2 degrees of freedom respectively, then the statistic *F* below is F distributed.

$$F(v_1, v_2) = \frac{\frac{\chi_1}{v_1}}{\frac{\chi_2}{v_2}}$$

The two parameters, v_1 and v_2 , are the numerator and denominator degrees of freedom. That is, v_1 and v_2 are the number of independent pieces information used to calculate χ_1 and χ_2 respectively.

Definition of the F distribution

The pdf for the F distribution is

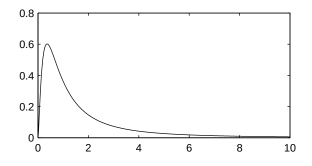
$$y = f(x|v_1, v_2) = \frac{\Gamma\left[\frac{(v_1 + v_2)}{2}\right]}{\Gamma\left(\frac{v_1}{2}\right)\Gamma\left(\frac{v_2}{2}\right)} \left(\frac{v_1}{v_2}\right)^{\frac{v_1}{2}} \frac{x^{\frac{v_1 - 2}{2}}}{\left[1 + \left(\frac{v_1}{v_2}\right)x\right]^{\frac{v_1 + v_2}{2}}}$$

where $\Gamma(\cdot)$ is the Gamma function.

Example and Plot of the F distribution

The most common application of the F distribution is in standard tests of hypotheses in analysis of variance and regression.

The plot shows that the F distribution exists on the positive real numbers and is skewed to the right.



Noncentral F Distribution

The following sections provide an overview of the noncentral F distribution.

Background of the Noncentral F Distribution

As with the χ^2 distribution, the F distribution is a special case of the noncentral F distribution. The F distribution is the result of taking the ratio of two χ^2 random variables each divided by its degrees of freedom.

If the numerator of the ratio is a noncentral chi-square random variable divided by its degrees of freedom, the resulting distribution is the noncentral F distribution.

The main application of the noncentral F distribution is to calculate the power of a hypothesis test relative to a particular alternative.

Definition of the Noncentral F Distribution

Similar to the noncentral χ^2 distribution, the toolbox calculates noncentral F distribution probabilities as a weighted sum of incomplete beta functions using Poisson probabilities as the weights.

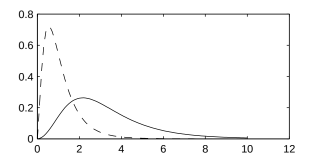
$$F(x|\mathbf{v}_1,\mathbf{v}_2,\boldsymbol{\delta}) = \sum_{j=0}^{\infty} \left(\frac{\left(\frac{1}{2}\boldsymbol{\delta}\right)^j}{j!} e^{-\frac{\boldsymbol{\delta}}{2}} \right) I\left(\frac{\mathbf{v}_1 \cdot x}{\mathbf{v}_2 + \mathbf{v}_1 \cdot x} \middle| \frac{\mathbf{v}_1}{2} + j, \frac{\mathbf{v}_2}{2} \right)$$

I(x | a, b) is the incomplete beta function with parameters a and b, and δ is the noncentrality parameter.

Example and Plot of the Noncentral F Distribution

The following commands generate a plot of the noncentral F pdf.

x = (0.01:0.1:10.01)'; p1 = ncfpdf(x,5,20,10); p = fpdf(x,5,20); plot(x,p,'--',x,p1,'-')



Gamma Distribution

The following sections provide an overview of the gamma distribution.

Background of the Gamma Distribution

The gamma distribution is a family of curves based on two parameters. The chi-square and exponential distributions, which are children of the gamma distribution, are one-parameter distributions that fix one of the two gamma parameters.

The gamma distribution has the following relationship with the incomplete Gamma function.

 $\Gamma(x|a,b) = \operatorname{gammainc}\left(\frac{x}{b},a\right)$

For b = 1 the functions are identical.

When a is large, the gamma distribution closely approximates a normal distribution with the advantage that the gamma distribution has density only for positive real numbers.

Definition of the Gamma Distribution

The gamma pdf is

$$y = f(x|a, b) = \frac{1}{b^{a}\Gamma(a)}x^{a-1}e^{-\frac{x}{b}}$$

where $\Gamma(\ \cdot\)$ is the Gamma function.

Parameter Estimation for the Gamma Distribution

Suppose you are stress testing computer memory chips and collecting data on their lifetimes. You assume that these lifetimes follow a gamma distribution. You want to know how long you can expect the average computer memory chip to last. Parameter estimation is the process of determining the parameters of the gamma distribution that fit this data best in some sense.

One popular criterion of goodness is to maximize the likelihood function. The likelihood has the same form as the gamma pdf above. But for the pdf, the parameters are known constants and the variable is x. The likelihood function reverses the roles of the variables. Here, the sample values (the x's) are already observed. So they are the fixed constants. The variables are the unknown parameters. MLE involves calculating the values of the parameters that give the highest likelihood given the particular set of data.

The function gamfit returns the MLEs and confidence intervals for the parameters of the gamma distribution. Here is an example using random numbers from the gamma distribution with a = 10 and b = 5.

```
lifetimes = gamrnd(10,5,100,1);
[phat, pci] = gamfit(lifetimes)
phat =
    10.9821    4.7258
pci =
    7.4001    3.1543
    14.5640    6.2974
```

Note phat(1) = \hat{a} and phat(2) = b. The MLE for parameter a is 10.98, compared to the true value of 10. The 95% confidence interval for a goes from 7.4 to 14.6, which includes the true value.

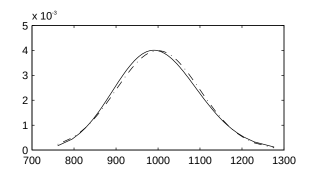
Similarly the MLE for parameter b is 4.7, compared to the true value of 5. The 95% confidence interval for b goes from 3.2 to 6.3, which also includes the true value.

In our life tests we do not know the true value of a and b so it is nice to have a confidence interval on the parameters to give a range of likely values.

Example and Plot of the Gamma Distribution

In the example the gamma pdf is plotted with the solid line. The normal pdf has a dashed line type.

```
x = gaminv((0.005:0.01:0.995),100,10);
y = gampdf(x,100,10);
y1 = normpdf(x,1000,100);
plot(x,y,'-',x,y1,'-.')
```



Geometric Distribution

The following sections provide an overview of the geometric distribution.

Background of the Geometric Distribution

The geometric distribution is discrete, existing only on the nonnegative integers. It is useful for modeling the runs of consecutive successes (or failures) in repeated independent trials of a system.

The geometric distribution models the number of successes before one failure in an independent succession of tests where each test results in success or failure.

Definition of the Geometric Distribution

The geometric pdf is

$$y = f(x|p) = pq^{x}I_{(0, 1, ...)}(x)$$

where q = 1 - p.

Example and Plot of the Geometric Distribution

Suppose the probability of a five-year-old battery failing in cold weather is 0.03. What is the probability of starting 25 consecutive days during a long cold snap?

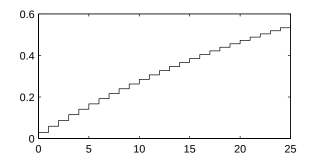
```
1 - geocdf(25, 0.03)
```

```
ans =
```

0.4530

The plot shows the cdf for this scenario.

x = 0:25; y = geocdf(x,0.03); stairs(x,y)



Hypergeometric Distribution

The following sections provide an overview of the hypergeometric distribution.

Background of the Hypergeometric Distribution

The hypergeometric distribution models the total number of successes in a fixed size sample drawn without replacement from a finite population.

The distribution is discrete, existing only for nonnegative integers less than the number of samples or the number of possible successes, whichever is greater. The hypergeometric distribution differs from the binomial only in that the population is finite and the sampling from the population is without replacement.

The hypergeometric distribution has three parameters that have direct physical interpretations. M is the size of the population. K is the number of items with the desired characteristic in the population. n is the number of samples drawn. Sampling "without replacement" means that once a particular sample is chosen, it is removed from the relevant population for all subsequent selections.

Definition of the Hypergeometric Distribution

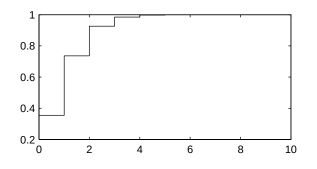
The hypergeometric pdf is

$$y = f(x|M, K, n) = \frac{\binom{K}{x}\binom{M-K}{n-x}}{\binom{M}{n}}$$

Example and Plot of the Hypergeometric Distribution

The plot shows the cdf of an experiment taking 20 samples from a group of 1000 where there are 50 items of the desired type.

x = 0:10; y = hygecdf(x,1000,50,20); stairs(x,y)



Lognormal Distribution

The following sections provide an overview of the lognormal distribution.

Background of the Lognormal Distribution

The normal and lognormal distributions are closely related. If X is distributed lognormal with parameters μ and σ^2 , then lnX is distributed normal with parameters μ and σ^2 .

The lognormal distribution is applicable when the quantity of interest must be positive, since lnX exists only when the random variable X is positive. Economists often model the distribution of income using a lognormal distribution.

Definition of the Lognormal Distribution

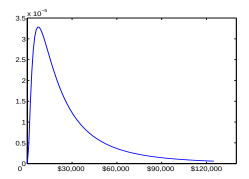
The lognormal pdf is

$$y = f(x|\mu, \sigma) = \frac{1}{x\sigma\sqrt{2\pi}}e^{\frac{-(\ln x - \mu)^2}{2\sigma^2}}$$

Example and Plot of the Lognormal Distribution

Suppose the income of a family of four in the United States follows a lognormal distribution with $\mu = \log(20,000)$ and $\sigma^2 = 1.0$. Plot the income density.

```
x = (10:1000:125010)';
y = lognpdf(x,log(20000),1.0);
plot(x,y)
set(gca,'xtick',[0 30000 60000 90000 120000])
set(gca,'xticklabel',str2mat('0','$30,000','$60,000',...
'$90,000','$120,000'))
```



Negative Binomial Distribution

The following sections provide an overview of the negative binomial distribution.

- "Background of the Negative Binomial Distribution" on page 2-31
- "Definition of the Negative Binomial Distribution" on page 2-31
- "Parameter Estimation for the Negative Binomial Distribution" on page 2-32
- "Example and Plot of the Negative Binomial Distribution" on page 2-33

Background of the Negative Binomial Distribution

In its simplest form, the negative binomial distribution models the number of successes before a specified number of failures is reached in an independent series of repeated identical trials. It can also be thought of as modelling the total number of trials required before a specified number of successes, thus motivating its name as the "inverse" of the binomial distribution. Its parameters are the probability of success in a single trial, p, and the number of failures, r. A special case of the negative binomial distribution, when r = 1, is the geometric distribution (also known as the Pascal distribution), which models the number of successes before the first failure.

More generally, the r parameter can take on non-integer values. This form of the negative binomial has no interpretation in terms of repeated trials, but, like the Poisson distribution, it is useful in modelling count data. It is, however, more general than the Poisson, because the negative binomial has a variance that is greater than its mean, often making it suitable for count data that do not meet the assumptions of the Poisson distribution. In the limit, as the parameter r increases to infinity, the negative binomial distribution approaches the Poisson distribution.

Definition of the Negative Binomial Distribution

When the r parameter is an integer, the negative binomial pdf is

$$y = f(x|r,p) = {r+x-1 \choose x} p^r q^x I_{(0, 1, ...)}(x)$$

where q = 1 - p. When *r* is non-integer, the binomial coefficient in the definition of the pdf is replaced by the equivalent expression

 $\frac{\Gamma(r+x)}{\Gamma(r)\Gamma(x+1)}$

Parameter Estimation for the Negative Binomial Distribution

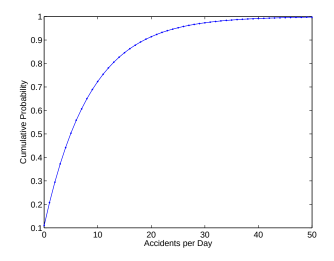
Suppose you are collecting data on the number of auto accidents on a busy highway, and would like to be able to model the number of accidents per day. Because these are count data, and because there are a very large number of cars and a small probability of an accident for any specific car, you might think to use the Poisson distribution. However, the probability of having an accident is likely to vary from day to day as the weather and amount of traffic change, and so the assumptions needed for the Poisson distribution are not met. In particular, the variance of this type of count data sometimes exceeds the mean by a large amount. The data below exhibit this effect: most days have few or no accidents, and a few days have a large number.

The negative binomial distribution is more general than the Poisson, and is often suitable for count data when the Poisson is not. The function nbinfit returns the maximum likelihood estimates (MLEs) and confidence intervals for the parameters of the negative binomial distribution. Here are the results from fitting the accident data above:

It's difficult to give a physical interpretation in this case to the individual parameters. However, the estimated parameters can be used in a model for the number of daily accidents. For example, a plot of the estimated cumulative probability function shows that while there is an estimated 10% chance of no

accidents on a given day, there is also about a 10% chance that there will be 20 or more accidents.

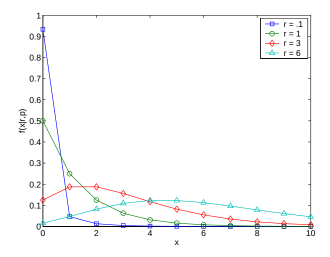
```
plot(0:50,nbincdf(0:50,phat(1),phat(2)),'.-');
xlabel('Accidents per Day')
ylabel('Cumulative Probability')
```



Example and Plot of the Negative Binomial Distribution

The negative binomial distribution can take on a variety of shapes ranging from very skewed to nearly symmetric. This example plots the probability function for different values of r, the desired number of successes: 1, 1, 3, 6.

```
x = 0:10;
plot(x,nbinpdf(x,.1,.5),'s-', ...
x,nbinpdf(x,1,.5),'o-', ...
x,nbinpdf(x,3,.5),'d-', ...
x,nbinpdf(x,6,.5),'^-');
legend({'r = .1' 'r = 1' 'r = 3' 'r = 6'})
xlabel('x')
ylabel('f(x|r,p')
```



Normal Distribution

The following sections provide an overview of the normal distribution.

Background of the Normal Distribution

The normal distribution is a two parameter family of curves. The first parameter, μ , is the mean. The second, σ , is the standard deviation. The standard normal distribution (written $\Phi(x)$) sets μ to 0 and σ to 1.

 $\Phi(x)$ is functionally related to the error function, *erf*.

 $erf(x) = 2\Phi(x\sqrt{2}) - 1$

The first use of the normal distribution was as a continuous approximation to the binomial.

The usual justification for using the normal distribution for modeling is the Central Limit Theorem, which states (roughly) that the sum of independent samples from any distribution with finite mean and variance converges to the normal distribution as the sample size goes to infinity.

Definition of the Normal Distribution

The normal pdf is

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

Parameter Estimation for the Normal Distribution

One of the first applications of the normal distribution in data analysis was modeling the height of school children. Suppose we want to estimate the mean, μ , and the variance, σ^2 , of all the 4th graders in the United States.

We have already introduced MLEs. Another desirable criterion in a statistical estimator is unbiasedness. A statistic is unbiased if the expected value of the statistic is equal to the parameter being estimated. MLEs are not always unbiased. For any data sample, there may be more than one unbiased estimator of the parameters of the parent distribution of the sample. For instance, every sample value is an unbiased estimate of the parameter μ of a normal distribution. The Minimum Variance Unbiased Estimator (MVUE) is the statistic that has the minimum variance of all unbiased estimators of a parameter.

The MVUEs of parameters μ and σ^2 for the normal distribution are the sample average and variance. The sample average is also the MLE for μ . There are two common textbook formulas for the variance.

They are

1)
$$s^{2} = \frac{1}{n} \sum_{i=1}^{n} (x_{i} - \bar{x})^{2}$$

2) $s^{2} = \frac{1}{n-1} \sum_{i=1}^{n} (x_{i} - \bar{x})^{2}$

where

$$\bar{x} = \sum_{i=1}^{n} \frac{x_i}{n}$$

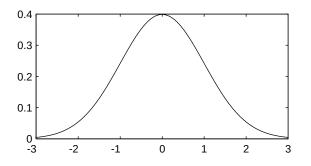
Equation 1 is the maximum likelihood estimator for $\sigma^2,$ and equation 2 is the MVUE.

The function normfit returns the MVUEs and confidence intervals for μ and σ^2 . Here is a playful example modeling the "heights" (inches) of a randomly chosen 4th grade class.

```
height = normrnd(50,2,30,1); % Simulate heights.
[mu,s,muci,sci] = normfit(height)
mu =
    50.2025
s =
    1.7946
muci =
    49.5210
    50.8841
sci =
    1.4292
    2.4125
```

Example and Plot of the Normal Distribution

The plot shows the "bell" curve of the standard normal pdf, with μ = 0 and σ = 1.



Poisson Distribution

The following sections provide an overview of the Poisson distribution.

Background of the Poisson Distribution

The Poisson distribution is appropriate for applications that involve counting the number of times a random event occurs in a given amount of time, distance, area, etc. Sample applications that involve Poisson distributions include the number of Geiger counter clicks per second, the number of people walking into a store in an hour, and the number of flaws per 1000 feet of video tape.

The Poisson distribution is a one parameter discrete distribution that takes nonnegative integer values. The parameter, λ , is both the mean and the variance of the distribution. Thus, as the size of the numbers in a particular sample of Poisson random numbers gets larger, so does the variability of the numbers.

As Poisson (1837) showed, the Poisson distribution is the limiting case of a binomial distribution where *N* approaches infinity and *p* goes to zero while $Np = \lambda$.

The Poisson and exponential distributions are related. If the number of counts follows the Poisson distribution, then the interval between individual counts follows the exponential distribution.

Definition of the Poisson Distribution

The Poisson pdf is

$$y = f(x|\lambda) = \frac{\lambda^x}{x!} e^{-\lambda} I_{(0, 1, \dots)}(x)$$

Parameter Estimation for the Poisson Distribution

The MLE and the MVUE of the Poisson parameter, λ , is the sample mean. The sum of independent Poisson random variables is also Poisson distributed with the parameter equal to the sum of the individual parameters. The Statistics Toolbox makes use of this fact to calculate confidence intervals on λ . As λ gets large the Poisson distribution can be approximated by a normal distribution with $\mu = \lambda$ and $\sigma^2 = \lambda$. The Statistics Toolbox uses this approximation for calculating confidence intervals for values of λ greater than 100.

Example and Plot of the Poisson Distribution

The plot shows the probability for each nonnegative integer when $\lambda = 5$.

Rayleigh Distribution

The following sections provide an overview of the Rayleigh distribution.

Background of the Rayleigh Distribution

The Rayleigh distribution is a special case of the Weibull distribution. If A and B are the parameters of the Weibull distribution, then the Rayleigh distribution with parameter b is equivalent to the Weibull distribution with parameters $A = 1/(2b^2)$ and B = 2.

If the component velocities of a particle in the x and y directions are two independent normal random variables with zero means and equal variances, then the distance the particle travels per unit time is distributed Rayleigh.

Definition of the Rayleigh Distribution

The Rayleigh pdf is

$$y = f(x|b) = \frac{x}{b^2} e^{\left(\frac{-x^2}{2b^2}\right)}$$

Parameter Estimation for the Rayleigh Distribution

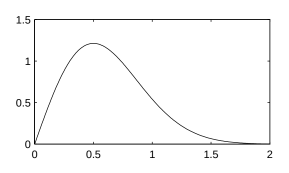
The raylfit function returns the MLE of the Rayleigh parameter. This estimate is

$$b = \sqrt{\frac{1}{2n} \sum_{i=1}^{n} x_i^2}$$

Example and Plot of the Rayleigh Distribution

The following commands generate a plot of the Rayleigh pdf.

```
x = [0:0.01:2];
p = raylpdf(x,0.5);
plot(x,p)
```



Student's t Distribution

The following sections provide an overview of Student's t distribution.

Background of Student's t Distribution

The t distribution is a family of curves depending on a single parameter ν (the degrees of freedom). As ν goes to infinity, the t distribution converges to the standard normal distribution.

W. S. Gossett (1908) discovered the distribution through his work at the Guinness brewery. At that time, Guinness did not allow its staff to publish, so Gossett used the pseudonym Student.

If *x* and *s* are the mean and standard deviation of an independent random sample of size *n* from a normal distribution with mean μ and $\sigma^2 = n$, then

$$t(v) = \frac{x-\mu}{s}$$
$$v = n-1$$

Definition of Student's t Distribution

Student's t pdf is

$$y = f(x|v) = \frac{\Gamma\left(\frac{v+1}{2}\right)}{\Gamma\left(\frac{v}{2}\right)} \frac{1}{\sqrt{v\pi}} \frac{1}{\left(1 + \frac{x^2}{v}\right)^{\frac{v+1}{2}}}$$

where $\Gamma(\cdot)$ is the Gamma function.

Example and Plot of Student's t Distribution

The plot compares the t distribution with v = 5 (solid line) to the shorter tailed, standard normal distribution (dashed line).

$$x = -5:0.1:5;$$

$$y = tpdf(x,5);$$

$$z = normpdf(x,0,1);$$

$$plot(x,y,'-',x,z,'-.')$$

0.4
0.3
0.2
0.1
0.5
0
5

Noncentral t Distribution

The following sections provide an overview of the noncentral t distribution.

Background of the Noncentral t Distribution

The noncentral t distribution is a generalization of the familiar Student's t distribution.

If *x* and *s* are the mean and standard deviation of an independent random sample of size *n* from a normal distribution with mean μ and $\sigma^2 = n$, then

$$t(v) = \frac{x - \mu}{s}$$
$$v = n - 1$$

Suppose that the mean of the normal distribution is not μ . Then the ratio has the noncentral t distribution. The noncentrality parameter is the difference between the sample mean and μ .

The noncentral t distribution allows us to determine the probability that we would detect a difference between x and μ in a t test. This probability is the *power* of the test. As x- μ increases, the power of a test also increases.

Definition of the Noncentral t Distribution

The most general representation of the noncentral t distribution is quite complicated. Johnson and Kotz (1970) give a formula for the probability that a noncentral t variate falls in the range [-t, t].

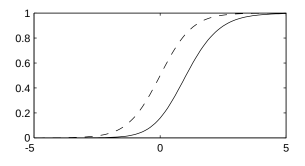
$$Pr((-t) < x < t | (v, \delta)) = \sum_{j=0}^{\infty} \left(\frac{\left(\frac{1}{2}\delta^{2}\right)^{j}}{j!} e^{-\frac{\delta^{2}}{2}} \right) I\left(\frac{x^{2}}{v+x^{2}} \left| \frac{1}{2} + j, \frac{v}{2} \right)$$

I(x | a, b) is the incomplete beta function with parameters a and b, δ is the noncentrality parameter, and v is the degrees of freedom.

Example and Plot of the Noncentral t Distribution

The following commands generate a plot of the noncentral t pdf.

```
x = (-5:0.1:5)';
p1 = nctcdf(x,10,1);
p = tcdf(x,10);
plot(x,p,'--',x,p1,'-')
```



Uniform (Continuous) Distribution

The following sections provide an overview of the uniform distribution.

Background of the Uniform Distribution

The uniform distribution (also called rectangular) has a constant pdf between its two parameters a (the minimum) and b (the maximum). The standard uniform distribution (a = 0 and b = 1) is a special case of the beta distribution, obtained by setting both of its parameters to 1.

The uniform distribution is appropriate for representing the distribution of round-off errors in values tabulated to a particular number of decimal places.

Definition of the Uniform Distribution

The uniform cdf is

 $p = F(x|a, b) = \frac{x-a}{b-a}I_{[a, b]}(x)$

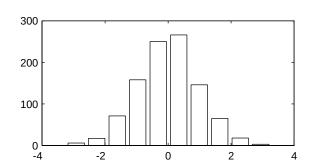
Parameter Estimation for the Uniform Distribution

The sample minimum and maximum are the MLEs of a and b respectively.

Example and Plot of the Uniform Distribution

The example illustrates the inversion method for generating normal random numbers using rand and norminy. Note that the MATLAB function, randn, does not use inversion since it is not efficient for this case.

```
u = rand(1000,1);
x = norminv(u,0,1);
hist(x)
```



Weibull Distribution

The following sections provide an overview of the Weibull distribution.

Background of the Weibull Distribution

Waloddi Weibull (1939) offered the distribution that bears his name as an appropriate analytical tool for modeling the breaking strength of materials. Current usage also includes reliability and lifetime modeling. The Weibull distribution is more flexible than the exponential for these purposes.

To see why, consider the hazard rate function (instantaneous failure rate). If f(t) and F(t) are the pdf and cdf of a distribution, then the hazard rate is

$$h(t) = \frac{f(t)}{1 - F(t)}$$

Substituting the pdf and cdf of the exponential distribution for f(t) and F(t) above yields a constant. The example below shows that the hazard rate for the Weibull distribution can vary.

Definition of the Weibull Distribution

The Weibull pdf is

$$y = f(x|a, b) = abx^{b-1}e^{-ax^{b}}I_{(0,\infty)}(x)$$

Parameter Estimation for the Weibull Distribution

Suppose we want to model the tensile strength of a thin filament using the Weibull distribution. The function weibfit gives MLEs and confidence intervals for the Weibull parameters.

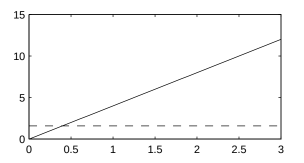
The default 95% confidence interval for each parameter contains the true value.

Example and Plot of the Weibull Distribution

The exponential distribution has a constant hazard function, which is not generally the case for the Weibull distribution.

The plot shows the hazard functions for exponential (dashed line) and Weibull (solid line) distributions having the same mean life. The Weibull hazard rate here increases with age (a reasonable assumption).

```
t = 0:0.1:3;
h1 = exppdf(t,0.6267) ./ (1-expcdf(t,0.6267));
h2 = weibpdf(t,2,2) ./ (1-weibcdf(t,2,2));
plot(t,h1,'--',t,h2,'-')
```



Descriptive Statistics

Introduction	•	•	•	. 3-2
Measures of Central Tendency (Location)				. 3-3
Measures of Dispersion				. 3-5
Functions for Data with Missing Values (NaNs)	•			. 3-7
Function for Grouped Data				. 3-9
Percentiles and Graphical Descriptions				
Percentiles	•			. 3-11
Probability Density Estimation				. 3-13
Empirical Cumulative Distribution Function	•	•	•	. 3-16
The Bootstrap				. 3-19

Introduction

Data samples can have thousands (even millions) of values. Descriptive statistics are a way to summarize this data into a few numbers that contain most of the relevant information. The following sections explore the features provided by the Statistics Toolbox for working with descriptive statistics:

- "Measures of Central Tendency (Location)" on page 3-3
- "Measures of Dispersion" on page 3-5
- "Functions for Data with Missing Values (NaNs)" on page 3-7
- "Function for Grouped Data" on page 3-9
- "Percentiles and Graphical Descriptions" on page 3-11
- "The Bootstrap" on page 3-19

Measures of Central Tendency (Location)

The purpose of measures of central tendency is to locate the data values on the number line. Another term for these statistics is *measures of location*.

The table gives the function names and descriptions.

Measures of Location		
geomean	Geometric mean	
harmmean	Harmonic mean	
mean	Arithmetic average (in MATLAB)	
median	50th percentile (in MATLAB)	
trimmean	Trimmed mean	

The average is a simple and popular estimate of location. If the data sample comes from a normal distribution, then the sample average is also optimal (MVUE of μ).

Unfortunately, outliers, data entry errors, or glitches exist in almost all real data. The sample average is sensitive to these problems. One bad data value can move the average away from the center of the rest of the data by an arbitrarily large distance.

The median and trimmed mean are two measures that are resistant (robust) to outliers. The median is the 50th percentile of the sample, which will only change slightly if you add a large perturbation to any value. The idea behind the trimmed mean is to ignore a small percentage of the highest and lowest values of a sample when determining the center of the sample.

The geometric mean and harmonic mean, like the average, are not robust to outliers. They are useful when the sample is distributed lognormal or heavily skewed.

The example below shows the behavior of the measures of location for a sample with one outlier.

```
x = [ones(1,6) \ 100]
x =
     1
           1
                 1
                        1
                              1
                                    1
                                         100
locate = [geomean(x) harmmean(x) mean(x) median(x)...
          trimmean(x,25)]
locate =
    1.9307
              1.1647
                        15.1429
                                   1.0000
                                              1.0000
```

You can see that the mean is far from any data value because of the influence of the outlier. The median and trimmed mean ignore the outlying value and describe the location of the rest of the data values.

Measures of Dispersion

The purpose of measures of dispersion is to find out how spread out the data values are on the number line. Another term for these statistics is measures of spread.

The table gives the function names and descriptions.

Measures of Dispersion		
iqr	Interquartile Range	
mad	Mean Absolute Deviation	
range	Range	
std	Standard deviation (in MATLAB)	
var	Variance (in MATLAB)	

The range (the difference between the maximum and minimum values) is the simplest measure of spread. But if there is an outlier in the data, it will be the minimum or maximum value. Thus, the range is not robust to outliers.

The standard deviation and the variance are popular measures of spread that are optimal for normally distributed samples. The sample variance is the MVUE of the normal parameter σ^2 . The standard deviation is the square root of the variance and has the desirable property of being in the same units as the data. That is, if the data is in meters, the standard deviation is in meters as well. The variance is in meters², which is more difficult to interpret.

Neither the standard deviation nor the variance is robust to outliers. A data value that is separate from the body of the data can increase the value of the statistics by an arbitrarily large amount.

The Mean Absolute Deviation (MAD) is also sensitive to outliers. But the MAD does not move quite as much as the standard deviation or variance in response to bad data.

The Interquartile Range (IQR) is the difference between the 75th and 25th percentile of the data. Since only the middle 50% of the data affects this measure, it is robust to outliers.

The example below shows the behavior of the measures of dispersion for a sample with one outlier.

Functions for Data with Missing Values (NaNs)

Most real-world data sets have one or more missing elements. It is convenient to code missing entries in a matrix as NaN (Not a Number).

Here is a simple example.

```
m = magic(3);
m([1 5]) = [NaN NaN]
m =
NaN 1 6
3 NaN 7
4 9 2
```

Any arithmetic operation that involves the missing values in this matrix yields NaN, as below.

```
sum(m)
ans =
NaN NaN 15
```

Removing cells with NaN would destroy the matrix structure. Removing whole rows that contain NaN would discard real data. Instead, the Statistics Toolbox has a variety of functions that are similar to other MATLAB functions, but that treat NaN values as missing and therefore ignore them in the calculations.

```
nansum(m)
ans =
7 10
```

13

NaN Functions	
nanmax	Maximum ignoring NaNs
nanmean	Mean ignoring NaNs
nanmedian	Median ignoring NaNs
nanmin	Minimum ignoring NaNs

NaN Functions	
nanstd	Standard deviation ignoring NaNs
nansum	Sum ignoring NaNs

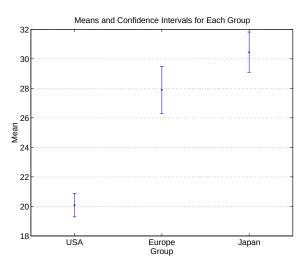
In addition, other Statistics Toolbox functions operate only on the numeric values, ignoring NaNs. These include iqr, kurtosis, mad, prctile, range, skewness, and trimmean.

Function for Grouped Data

As we saw in the previous section, the descriptive statistics functions can compute statistics on each column in a matrix. Sometimes, however, you may have your data arranged differently so that measurements appear in one column or variable, and a grouping code appears in a second column or variable. Although the MATLAB syntax makes it simple to apply functions to a subset of an array, in this case it is simpler to use the grpstats function.

The grpstats function can compute the mean, standard error of the mean, and count (number of observations) for each group defined by one or more grouping variables. If you supply a significance level, it also creates a graph of the group means with confidence intervals.

As an example, load the larger car data set. We can look at the average value of MPG (miles per gallon) for cars grouped by org (location of the origin of the car).



We can also get the complete set of statistics for MPG grouped by three variables: org, cyl4 (the engine has four cylinders or not), and when (when the car was made).

```
[m,s,c,n] = grpstats(MPG,{org cyl4 when});
[n num2cell([m s c])]
```

```
ans =
```

'USA'	'Other'	'Early'	[14.896]	[0.33306]	[77]
'USA'	'Other'	'Mid'	[17.479]	[0.30225]	[75]
'USA'	'Other'	'Late'	[21.536]	[0.97961]	[25]
'USA'	'Four'	'Early'	[23.333]	[0.87328]	[12]
'USA'	'Four'	'Mid'	[27.027]	[0.75456]	[22]
'USA'	'Four'	'Late'	[29.734]	[0.71126]	[38]
'Europe'	'Other'	'Mid'	[17.5]	[0.9478]	[4]
'Europe'	'Other'	'Late'	[30.833]	[3.1761]	[3]
'Europe'	'Four'	'Early'	[24.714]	[0.73076]	[21]
'Europe'	'Four'	'Mid'	[26.912]	[1.0116]	[26]
'Europe'	'Four'	'Late'	[35.7]	[1.4265]	[16]
'Japan'	'Other'	'Early'	[19]	[0.57735]	[3]
'Japan'	'Other'	'Mid'	[20.833]	[0.92796]	[3]
'Japan'	'Other'	'Late'	[26.5]	[2.0972]	[4]
'Japan'	'Four'	'Early'	[26.083]	[1.1772]	[12]
'Japan'	'Four'	'Mid'	[29.5]	[0.86547]	[25]
'Japan'	'Four'	'Late'	[35.3]	[0.68346]	[32]

Percentiles and Graphical Descriptions

Trying to describe a data sample with two numbers, a measure of location and a measure of spread, is frugal but may be misleading. Here are some other approaches:

- "Percentiles" on page 3-11
- "Probability Density Estimation" on page 3-13
- "Empirical Cumulative Distribution Function" on page 3-16

Percentiles

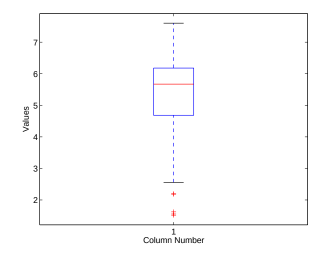
Another option is to compute a reasonable number of the sample percentiles. This provides information about the shape of the data as well as its location and spread.

The example shows the result of looking at every quartile of a sample containing a mixture of two distributions.

Compare the first two quantiles to the rest.

The box plot is a graph for descriptive statistics. The graph below is a box plot of the data above.

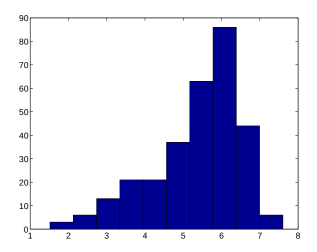
boxplot(x)



The long lower tail and plus signs show the lack of symmetry in the sample values. For more information on box plots, see "Statistical Plots" on page 8-1.

The histogram is a complementary graph.

hist(x)

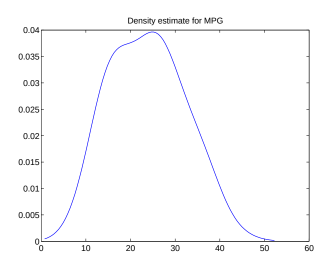


Probability Density Estimation

You can also describe a data sample by estimating its density in a nonparametric way. The ksdensity function does this by using a kernel smoothing function and an associated bandwidth to estimate the density.

This example uses the carsmall data set to estimate the probability density of the MPG (miles per gallon) measurements for 94 cars. It uses the default kernel function, a normal distribution, and its default bandwidth.

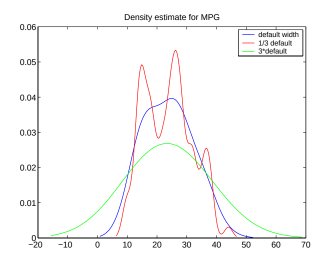
```
cars = load('carsmall','MPG','Origin');
MPG = cars.MPG;
Origin = cars.Origin;
[f,x] = ksdensity(MPG);
plot(x,f);
title('Density estimate for MPG')
```



Kernel Bandwidth

The choice of kernel bandwidth controls the smoothness of the probability density curve. The graph below shows the density estimate for the same mileage data using different bandwidths. The default bandwidth is in blue and looks like the graph above. Estimates for smaller and larger bandwidths are in red and green. The first call to ksdensity returns the default bandwidth, u, of the kernel smoothing function. Subsequent calls, modify this bandwidth.

```
[f,x,u] = ksdensity(MPG);
plot(x,f)
title('Density estimate for MPG')
hold on
[f,x] = ksdensity(MPG,'width',u/3);
plot(x,f,'r');
[f,x] = ksdensity(MPG,'width',u*3);
plot(x,f,'g');
legend('default width','1/3 default','3*default')
hold off
```



The default bandwidth seems to be doing a good job -- reasonably smooth, but not so smooth as to obscure features of the data. This bandwidth is the one that is theoretically optimal for estimating densities for the normal distribution.

The green curve shows a density with the kernel bandwidth set too high. This curve smooths out the data so much that the end result looks just like the kernel function. The red curve has a smaller bandwidth and is rougher-looking than the blue curve. It may be too rough, but it does provide an indication that there may be two major peaks rather than the single peak of the blue curve. A

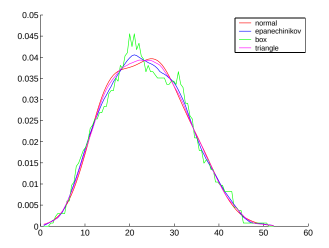
reasonable choice of width might lead to a curve that is intermediate between the red and blue curves.

Kernel Smoothing Function

You can also specify a kernel function by supplying either the function name or a function handle. The four pre-selected functions, 'normal', 'epanechinikov', 'box', and 'triangle', are all scaled to have standard deviation equal to one, so the "bandwidth" parameter means roughly the same thing regardless of kernel function.

Using default bandwidths, we now plot the same mileage data, using each of the available kernel smoothers.

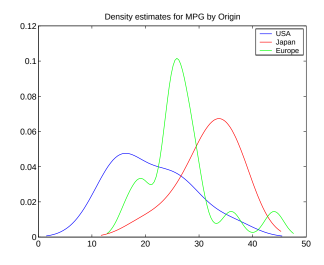
```
hname = {'normal' 'epanechinikov' 'box' 'triangle'};
hold on;
colors = {'r' 'b' 'g' 'm'};
for j=1:4
    [f,x] = ksdensity(MPG,'kernel',hname{j});
    plot(x,f,colors{j});
end
legend(hname{:});
hold off
```



The density estimates are roughly comparable, but the box kernel produces a density that is rougher than the others.

Usefulness of Smooth Density Estimates

In addition to the aesthetic appeal of the smooth density estimate, there are other appeals as well. While it is difficult to overlay two histograms to compare them, you can easily overlay smooth density estimates. For example, the following graph shows the MPG distributions for cars from different countries of origin.



Empirical Cumulative Distribution Function

The ksdensity function described in the last section produces an empirical version of a probability density function (pdf). That is, instead of selecting a density with a particular parametric form and estimating the parameters, it produces a nonparametric density estimate that tries to adapt itself to the data.

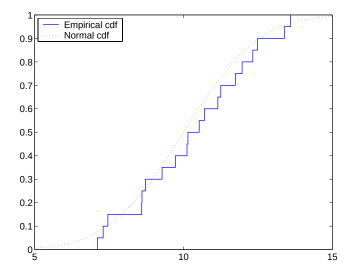
Similarly, it is possible to produce an empirical version of the cumulative distribution function (cdf). The ecdf function computes this empirical cdf. It

returns the values of a function F such that F(x) represents the proportion of observations in a sample less than or equal to x.

The idea behind the empirical cdf is simple. It is a function that assigns probability 1/n to each of n observations in a sample. Its graph has a stair-step appearance. If a sample comes from a distribution in a parametric family (such as a normal distribution), its empirical cdf is likely to resemble the parametric distribution. If not, its empirical distribution still gives an estimate of the cdf for the distribution that generated the data.

In the following example, we generate 20 observations from a normal distribution with mean 10 and standard deviation 2. We use ecdf to calculate the empirical cdf and stairs to plot it. Then we overlay the normal distribution curve on the empirical function.

```
x = normrnd(10,2,20,1);
[f,xf] = ecdf(x);
stairs(xf,f)
xx=linspace(5,15,100);
yy = normcdf(xx,10,2);
hold on; plot(xx,yy,'r:'); hold off
legend('Empirical cdf','Normal cdf',2)
```



The empirical cdf is especially useful in survival analysis applications. In such applications the data may be censored, that is, not observed exactly. Some individuals may fail during a study, and we observe their failure time exactly. Other individuals may drop out of the study, or may not fail until after the study is complete. The ecdf function has arguments for dealing with censored data.

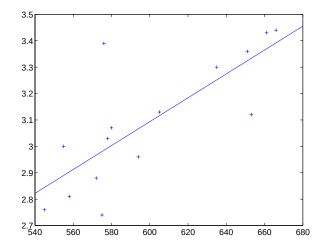
The Bootstrap

In recent years the statistical literature has examined the properties of resampling as a means to acquire information about the uncertainty of statistical estimators.

The bootstrap is a procedure that involves choosing random samples *with replacement* from a data set and analyzing each sample the same way. Sampling *with replacement* means that every sample is returned to the data set after sampling. So a particular data point from the original data set could appear multiple times in a given bootstrap sample. The number of elements in each bootstrap sample equals the number of elements in the original data set. The range of sample estimates we obtain allows us to establish the uncertainty of the quantity we are estimating.

Here is an example taken from Efron and Tibshirani (1993) comparing Law School Admission Test (LSAT) scores and subsequent law school grade point average (GPA) for a sample of 15 law schools.

```
load lawdata
plot(lsat,gpa,'+')
lsline
```



The least squares fit line indicates that higher LSAT scores go with higher law school GPAs. But how sure are we of this conclusion? The plot gives us some intuition but nothing quantitative.

We can calculate the correlation coefficient of the variables using the corrcoef function.

```
rhohat = corrcoef(lsat,gpa)
rhohat =
    1.0000    0.7764
    0.7764    1.0000
```

Now we have a number, 0.7764, describing the positive connection between LSAT and GPA, but though 0.7764 may seem large, we still do not know if it is statistically significant.

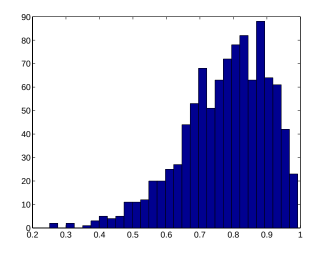
Using the bootstrp function we can resample the lsat and gpa vectors as many times as we like and consider the variation in the resulting correlation coefficients.

Here is an example.

```
rhos1000 = bootstrp(1000, 'corrcoef',lsat,gpa);
```

This command resamples the 1sat and gpa vectors 1000 times and computes the corrcoef function on each sample. Here is a histogram of the result.

```
hist(rhos1000(:,2),30)
```



Nearly all the estimates lie on the interval [0.4 1.0].

This is strong quantitative evidence that LSAT and subsequent GPA are positively correlated. Moreover, it does not require us to make any strong assumptions about the probability distribution of the correlation coefficient.



4

Linear Models

Introduction
One-Way Analysis of Variance (ANOVA)
Multiple Comparisons 4-6 Example: Multiple Comparisons 4-6
Two-Way Analysis of Variance (ANOVA)
N-Way Analysis of Variance
Multiple Linear Regression
Quadratic Response Surface Models
Stepwise Regression
Generalized Linear Models<
Robust and Nonparametric Methods
Friedman's Test

Introduction

Linear models represent the relationship between a continuous response variable and one or more predictor variables (either continuous or categorical) in the form

$$y = X\beta + \varepsilon$$

where:

- *y* is an *n*-by-1 vector of observations of the response variable.
- *X* is the *n*-by-*p* design matrix determined by the predictors.
- β is a *p*-by-1 vector of parameters.
- ε is an *n*-by-1 vector of random disturbances, independent of each other and usually having a normal distribution.

MATLAB uses this general form of the linear model to solve a variety of specific regression and analysis of variance (ANOVA) problems. For example, for polynomial and multiple regression problems, the columns of *X* are predictor variable values or powers of such values. For one-way, two-way, and higher-way ANOVA models, the columns of *X* are dummy (or indicator) variables that encode the predictor categories. For analysis of covariance (ANOCOVA) models, *X* contains values of a continuous predictor and codes for a categorical predictor.

The following sections describe a number of functions for fitting various types of linear models:

- "One-Way Analysis of Variance (ANOVA)" on page 4-4
- "Two-Way Analysis of Variance (ANOVA)" on page 4-9
- "N-Way Analysis of Variance" on page 4-12
- "Multiple Linear Regression" on page 4-18
- "Quadratic Response Surface Models" on page 4-22
- "Stepwise Regression" on page 4-24
- "Generalized Linear Models" on page 4-28
- "Robust and Nonparametric Methods" on page 4-32

See the sections below for a tour of some of the related graphical tools:

- "The polytool Demo" on page 11-5
- "The aoctool Demo" on page 11-10
- "The rsmdemo Demo" on page 11-19

Note See "Linear Models" on page 4-1 for information on fitting nonlinear models.

One-Way Analysis of Variance (ANOVA)

The purpose of one-way ANOVA is to find out whether data from several groups have a common mean. That is, to determine whether the groups are actually different in the measured characteristic.

One-way ANOVA is a simple special case of the linear model. The one-way ANOVA form of the model is

$$y_{ij} = \alpha_{j} + \varepsilon_{ij}$$

where:

- y_{ij} is a matrix of observations in which each column represents a different group.
- $\alpha_{,j}$ is a matrix whose columns are the group means. (The "dot j" notation means that α applies to all rows of the *j*th column. That is, the value α_{ij} is the same for all *i*.)
- ε_{ij} is a matrix of random disturbances.

The model posits that the columns of y are a constant plus a random disturbance. You want to know if the constants are all the same.

The following sections explore one-way ANOVA in greater detail:

- "Example: One-Way ANOVA" on page 4-4
- "Multiple Comparisons" on page 4-6

Example: One-Way ANOVA

The data below comes from a study by Hogg and Ledolter (1987) of bacteria counts in shipments of milk. The columns of the matrix hogg represent different shipments. The rows are bacteria counts from cartons of milk chosen randomly from each shipment. Do some shipments have higher counts than others?

```
load hogg
hogg
hogg =
24 14 11 7 19
```

[p,tbl,stats] = anova1(hogg); р p = 1.1971e-04

The standard ANOVA table has columns for the sums of squares, degrees of freedom, mean squares (SS/df), F statistic, and p-value.

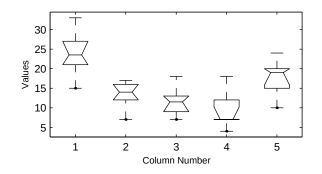
🚺 Figure I	No. 1: One-way	ANOVA	۱		_ 🗆 ×
<u>F</u> ile <u>E</u> dit	$\underline{T}ools$ $\underline{W}indow$	<u>H</u> elp			
		ANO	VA Table		
Source	SS	df	MS	F	Prob>F 🛌
Columns Error Total	803 557.17 1360.17	4 25 29	200.75 22.287	9.01	0.0001

You can use the F statistic to do a hypothesis test to find out if the bacteria counts are the same. anoval returns the p-value from this hypothesis test.

In this case the p-value is about 0.0001, a very small value. This is a strong indication that the bacteria counts from the different tankers are not the same. An F statistic as extreme as the observed F would occur by chance only once in 10,000 times if the counts were truly equal.

The p-value returned by anoval depends on assumptions about the random disturbances ϵ_{ij} in the model equation. For the p-value to be correct, these disturbances need to be independent, normally distributed, and have constant variance. See "Robust and Nonparametric Methods" on page 4-32 for a nonparametric function that does not require a normal assumption.

You can get some graphical assurance that the means are different by looking at the box plots in the second figure window displayed by anova1.



Multiple Comparisons

Sometimes you need to determine not just if there are any differences among the means, but specifically which pairs of means are significantly different. It is tempting to perform a series of t tests, one for each pair of means, but this procedure has a pitfall.

In a t test, we compute a t statistic and compare it to a critical value. The critical value is chosen so that when the means are really the same (any apparent difference is due to random chance), the probability that the t statistic will exceed the critical value is small, say 5%. When the means are different, the probability that the statistic will exceed the critical value is larger.

In this example there are five means, so there are 10 pairs of means to compare. It stands to reason that if all the means are the same, and if we have a 5% chance of incorrectly concluding that there is a difference in one pair, then the probability of making at least one incorrect conclusion among all 10 pairs is much larger than 5%.

Fortunately, there are procedures known as *multiple comparison procedures* that are designed to compensate for multiple tests.

Example: Multiple Comparisons

You can perform a multiple comparison test using the multcompare function and supplying it with the stats output from anova1.

[c,m] = multcompare(stats)
c =

1.0000	2.0000	2.4953	10.5000	18.5047
1.0000	3.0000	4.1619	12.1667	20.1714
1.0000	4.0000	6.6619	14.6667	22.6714
1.0000	5.0000	-2.0047	6.0000	14.0047
2.0000	3.0000	-6.3381	1.6667	9.6714
2.0000	4.0000	-3.8381	4.1667	12.1714
2.0000	5.0000	-12.5047	-4.5000	3.5047
3.0000	4.0000	-5.5047	2.5000	10.5047
3.0000	5.0000	-14.1714	-6.1667	1.8381
4.0000	5.0000	-16.6714	-8.6667	-0.6619
=				
23.8333	1.9273			
13.3333	1.9273			
11.6667	1.9273			
9.1667	1.9273			

m

17.8333

The first output from mult compare has one row for each pair of groups, with an estimate of the difference in group means and a confidence interval for that group. For example, the second row has the values

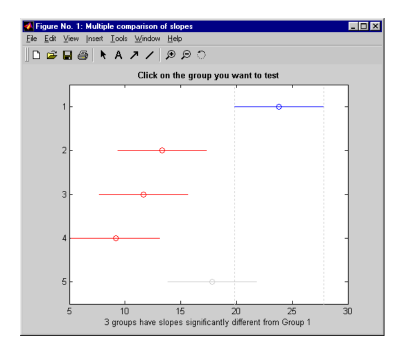
1.0000 3.0000 4.1619 12.1667 20.1714

1.9273

indicating that the mean of group 1 minus the mean of group 3 is estimated to be 12.1667, and a 95% confidence interval for this difference is [4.1619, 20.1714]. This interval does not contain 0, so we can conclude that the means of groups 1 and 3 are different.

The second output contains the mean and its standard error for each group.

It is easier to visualize the difference between group means by looking at the graph that multcompare produces.



The graph shows that group 1 is significantly different from groups 2, 3, and 4. By using the mouse to select group 4, you can determine that it is also significantly different from group 5. Other pairs are not significantly different.

Two-Way Analysis of Variance (ANOVA)

The purpose of two-way ANOVA is to find out whether data from several groups have a common mean. One-way ANOVA and two-way ANOVA differ in that the groups in two-way ANOVA have two categories of defining characteristics instead of one.

Suppose an automobile company has two factories, and each factory makes the same three models of car. It is reasonable to ask if the gas mileage in the cars varies from factory to factory as well as from model to model. We use two predictors, factory and model, to explain differences in mileage.

There could be an overall difference in mileage due to a difference in the production methods between factories. There is probably a difference in the mileage of the different models (irrespective of the factory) due to differences in design specifications. These effects are called *additive*.

Finally, a factory might make high mileage cars in one model (perhaps because of a superior production line), but not be different from the other factory for other models. This effect is called an *interaction*. It is impossible to detect an interaction unless there are duplicate observations for some combination of factory and car model.

Two-way ANOVA is a special case of the linear model. The two-way ANOVA form of the model is

 $y_{ijk} = \mu + \alpha_{.j} + \beta_{i.} + \gamma_{ij} + \varepsilon_{ijk}$

where, with respect to the automobile example above:

- y_{ijk} is a matrix of gas mileage observations (with row index *i*, column index *j*, and repetition index *k*).
- μ is a constant matrix of the overall mean gas mileage.
- α_j is a matrix whose columns are the deviations of each car's gas mileage (from the mean gas mileage μ) that are attributable to the car's *model*. All values in a given column of α_j are identical, and the values in each row of α_j sum to 0.
- $\beta_{i.}$ is a matrix whose rows are the deviations of each car's gas mileage (from the mean gas mileage μ) that are attributable to the car's *factory*. All values in a given row of $\beta_{i.}$ are identical, and the values in each column of $\beta_{i.}$ sum to 0.

- γ_{ij} is a matrix of interactions. The values in each row of γ_{ij} sum to 0, and the values in each column of γ_{ij} sum to 0.
- ε_{ijk} is a matrix of random disturbances.

The next section provides an example of a two-way analysis.

Example: Two-Way ANOVA

The purpose of the example is to determine the effect of car model and factory on the mileage rating of cars.

```
load mileage
mileage
mileage =
   33.3000
              34.5000
                        37.4000
   33.4000
              34.8000
                        36.8000
   32.9000
                        37.6000
              33.8000
   32.6000
              33.4000
                        36.6000
   32.5000
              33.7000
                        37.0000
                        36.7000
   33.0000
              33.9000
cars = 3;
[p,tbl,stats] = anova2(mileage,cars);
р
p =
    0.0000
               0.0039
                         0.8411
```

There are three models of cars (columns) and two factories (rows). The reason there are six rows in mileage instead of two is that each factory provides three cars of each model for the study. The data from the first factory is in the first three rows, and the data from the second factory is in the last three rows.

The standard ANOVA table has columns for the sums of squares, degrees-of-freedom, mean squares (SS/df), F statistics, and p-values.

🛃 Figure No. 1:	Two-way ANOV	/Α			_ 🗆 ×
<u>File E</u> dit <u>T</u> ools	<u>W</u> indow <u>H</u> elp				
	A	NOV	A Table		
Source	SS	df	MS	F	Prob>F 🛌
Columns Rows Interaction Error Total	53.3511 1.445 0.04 1.3667 56.2028	2 1 12 12 17	26.6756 1.445 0.02 0.1139	234.22 12.69 0.18	0 0.0039 0.8411

You can use the F statistics to do hypotheses tests to find out if the mileage is the same across models, factories, and model-factory pairs (after adjusting for the additive effects). anova2 returns the p-value from these tests.

The p-value for the model effect is zero to four decimal places. This is a strong indication that the mileage varies from one model to another. An F statistic as extreme as the observed F would occur by chance less than once in 10,000 times if the gas mileage were truly equal from model to model. If you used the multcompare function to perform a multiple comparison test, you would find that each pair of the three models is significantly different.

The p-value for the factory effect is 0.0039, which is also highly significant. This indicates that one factory is out-performing the other in the gas mileage of the cars it produces. The observed p-value indicates that an F statistic as extreme as the observed F would occur by chance about four out of 1000 times if the gas mileage were truly equal from factory to factory.

There does not appear to be any interaction between factories and models. The p-value, 0.8411, means that the observed result is quite likely (84 out 100 times) given that there is no interaction.

The p-values returned by anova2 depend on assumptions about the random disturbances ϵ_{ijk} in the model equation. For the p-values to be correct these disturbances need to be independent, normally distributed, and have constant variance. See "Robust and Nonparametric Methods" on page 4-32 for nonparametric methods that do not require a normal distribution.

In addition, anova2 requires that data be *balanced*, which in this case means there must be the same number of cars for each combination of model and factory. The next section discusses a function that supports unbalanced data with any number of predictors.

N-Way Analysis of Variance

You can use N-way ANOVA to determine if the means in a set of data differ when grouped by multiple factors. If they do differ, you can determine which factors or combinations of factors are associated with the difference.

N-way ANOVA is a generalization of two-way ANOVA. For three factors, the model can be written

$$y_{ijkl} = \mu + \alpha_{.j.} + \beta_{i..} + \gamma_{..k} + (\alpha\beta)_{ij.} + (\alpha\gamma)_{i.k} + (\beta\gamma)_{.jk} + (\alpha\beta\gamma)_{ijk} + \varepsilon_{ijkl}$$

In this notation parameters with two subscripts, such as $(\alpha\beta)_{ij}$, represent the interaction effect of two factors. The parameter $(\alpha\beta\gamma)_{ijk}$ represents the three-way interaction. An ANOVA model can have the full set of parameters or any subset, but conventionally it does not include complex interaction terms unless it also includes all simpler terms for those factors. For example, one would generally not include the three-way interaction without also including all two-way interactions.

The anovan function performs N-way ANOVA. Unlike the anova1 and anova2 functions, anovan does not expect data in a tabular form. Instead, it expects a vector of response measurements and a separate vector (or text array) containing the values corresponding to each factor. This input data format is more convenient than matrices when there are more than two factors or when the number of measurements per factor combination is not constant.

The following examples explore anovan in greater detail:

- "Example: N-Way ANOVA with Small Data Set" on page 4-12
- "Example: N-Way ANOVA with Large Data Set" on page 4-14

Example: N-Way ANOVA with Small Data Set

Consider the following two-way example using anova2.

```
m = [23 15 20;27 17 63;43 3 55;41 9 90]
m =
23 15 20
27 17 63
43 3 55
41 9 90
```

anova2(m,2) ans = 0.0197 0.2234 0.2663

The factor information is implied by the shape of the matrix m and the number of measurements at each factor combination (2). Although anova2 does not actually require arrays of factor values, for illustrative purposes we could create them as follows.

```
cfactor = repmat(1:3,4,1)
cfactor =
     1
            2
                   3
     1
            2
                   3
     1
            2
                   3
     1
            2
                   3
rfactor = [ones(2,3); 2*ones(2,3)]
rfactor =
     1
            1
                   1
     1
            1
                   1
     2
            2
                   2
     2
            2
                   2
```

The cfactor matrix shows that each column of m represents a different level of the column factor. The rfactor matrix shows that the top two rows of m represent one level of the row factor, and bottom two rows of m represent a second level of the row factor. In other words, each value m(i,j) represents an observation at column factor level cfactor(i,j) and row factor level rfactor(i,j).

To solve the above problem with anovan, we need to reshape the matrices ${\tt m},$ cfactor, and rfactor to be vectors.

```
m = m(:);
cfactor = cfactor(:);
rfactor = rfactor(:);
[m cfactor rfactor]
ans =
```

4 Linear Models

23	1	1
27	1	1
43	1	2
41	1	2
15	2	1
17	2	1
3	2	2
9	2	2
20	3	1
63	3	1
55	3	2
90	3	2
anovan(m,{	cfacto	r rfactor},2)

ans =

0.0197 0.2234 0.2663

Example: N-Way ANOVA with Large Data Set

In the previous example we used anova2 to study a small data set measuring car mileage. Now we study a larger set of car data with mileage and other information on 406 cars made between 1970 and 1982. First we load the data set and look at the variable names.

load carbig whos			
Name	Size	Bytes	Class
Acceleration	406x1	3248	double array
Cylinders	406x1	3248	double array
Displacement	406x1	3248	double array
Horsepower	406x1	3248	double array
MPG	406x1	3248	double array
Model	406x36	29232	char array
Model_Year	406x1	3248	double array
Origin	406x7	5684	char array
Weight	406x1	3248	double array

cyl4	406x5	4060	char array
org	406x7	5684	char array
when	406x5	4060	char array

We will focus our attention on four variables. MPG is the number of miles per gallon for each of 406 cars (though some have missing values coded as NaN). The other three variables are factors: cyl4 (four-cylinder car or not), org (car originated in Europe, Japan, or the USA), and when (car was built early in the period, in the middle of the period, or late in the period).

First we fit the full model, requesting up to three-way interactions and Type 3 sums-of-squares.

```
varnames = {'Origin';'4Cyl';'MfgDate'};
anovan(MPG,{org cyl4 when},3,3,varnames)
ans =
    0.0000
    NaN
    0
0.7032
0.0001
0.2072
0.6990
```

<u>File E</u> dit ⊻iew <u>I</u> nsert <u>T</u> ools <u>W</u> i	indow <u>H</u> elp				
	Analysis	of Vari	iance		
Source	Sum Sq.	d.f.	Mean Sq.	F	Prob>F
<pre># Origin # 4Cyl # MfgDate # Origin*4Cyl # Origin*MfgDate # 4Cyl*MfgDate # 0rigin*4Cyl*MfgDate # Origin*4Cyl*MfgDate Error Total</pre>	416.8 0 1112.3 2.1 301.2 22.7 20.3 5411.8 24252.6	1 0 1 3 1 381 397	$\begin{array}{r} 416.77\\0\\1112.27\\2.07\\100.41\\22.68\\6.77\\14.2\end{array}$	29.34 0 78.31 0.15 7.07 1.6 0.48	

Note that many terms are marked by a "#" symbol as not having full rank, and one of them has zero degrees of freedom and is missing a p-value. This can happen when there are missing factor combinations and the model has higher-order terms. In this case, the cross-tabulation below shows that there are no cars made in Europe during the early part of the period with other than four cylinders, as indicated by the 0 in table(2,1,1).

```
[table,factorvals] = crosstab(org,when,cyl4)
table(:,:,1) =
    82
                  25
           75
     0
            4
                  3
     3
            3
                  4
table(:,:,2) =
    12
           22
                  38
    23
           26
                  17
    12
           25
                  32
factorvals =
    'USA'
                  'Early'
                              'Other'
    'Europe'
                  'Mid'
                              'Four'
    'Japan'
                  'Late'
                                    []
```

Consequently it is impossible to estimate the three-way interaction effects, and including the three-way interaction term in the model makes the fit singular.

Using even the limited information available in the ANOVA table, we can see that the three-way interaction has a p-value of 0.699, so it is not significant. We decide to request only two-way interactions this time.

```
[p,tbl,stats,termvec] = anovan(MPG,{org cyl4 when},2,3,varnames);
termvec'
```

ans =

1 2 4 3 5 6

e <u>E</u> dit ⊻iew <u>I</u> nsert]	ools <u>W</u> indow	<u>H</u> elp			
	Ana	alysis d	of Variance		
Source	Sum Sq.	d.f.	Mean Sq.	F	Prob>F
Origin 4Cyl MfgDate Origin*4Cyl Origin*MfgDate 4Cyl*MfgDate Error Total	532.6 1769.8 2887.1 12.5 350.4 31 5432.1 24252.6	2 1 2 4 2 384 397	266.29 1769.85 1443.55 6.27 87.59 15.52 14.15	18.82 125.11 102.05 0.44 6.19 1.1	0 0 0.6422 0.0001 0.3348

Now all terms are estimable. The p-values for interaction term 4 (Origin*4Cyl) and interaction term 6 (4Cyl*MfgDate) are much larger than a typical cutoff value of 0.05, indicating these terms are not significant. We could choose to omit these terms and pool their effects into the error term. The output termvec variable returns a vector of codes, each of which is a bit pattern representing a term. We can omit terms from the model by deleting their entries from termvec and running anovan again, this time supplying the resulting vector as the model argument.

```
termvec([4 6]) = []
termvec =
    1
    2
    4
    5
```

anovan(MPG,{org cyl4 when},termvec,3,varnames)

Figure No. 2: N-Way /	ANOVA					_ []
ile <u>E</u> dit <u>V</u> iew <u>I</u> nsert <u>T</u>	ools <u>W</u> indow	<u>H</u> elp				
	Ana	alysis o	f Variance			
Source	Sum Sq.	d.f.	Mean Sq.	F	Prob>F	
Origin 4Cyl MfgDate Origin*MfgDate Error Total	686.7 4206.2 3590.7 336.8 5473 24252.6	2 1 2 4 388 397	343.36 4206.17 1795.34 84.19 14.11	24.34 298.19 127.28 5.97	0 0 0 0.0001	
	Constra	ined (Type	III) sums of squares			

Now we have a more parsimonious model indicating that the mileage of these cars seems to be related to all three factors, and that the effect of the manufacturing date depends on where the car was made.

Multiple Linear Regression

The purpose of multiple linear regression is to establish a quantitative relationship between a group of predictor variables (the columns of X) and a response, y. This relationship is useful for:

- Understanding which predictors have the greatest effect.
- Knowing the direction of the effect (i.e., increasing *x* increases/decreases *y*).
- Using the model to predict future values of the response when only the predictors are currently known.

The following sections explain multiple linear regression in greater detail:

- "Mathematical Foundations of Multiple Linear Regression" on page 4-18
- "Example: Multiple Linear Regression" on page 4-20

Mathematical Foundations of Multiple Linear Regression

The linear model takes its common form

 $y = X\beta + \varepsilon$

where:

- *y* is an *n*-by-1 vector of observations.
- *X* is an *n*-by-*p* matrix of regressors.
- β is a *p*-by-1 vector of parameters.
- ε is an *n*-by-1 vector of random disturbances.

The solution to the problem is a vector, b, which estimates the unknown vector of parameters, β . The least squares solution is

$$b = \hat{\beta} = (X^T X)^{-1} X^T y$$

This equation is useful for developing later statistical formulas, but has poor numeric properties. regress uses QR decomposition of X followed by the backslash operator to compute b. The QR decomposition is not necessary for computing b, but the matrix R is useful for computing confidence intervals.

You can plug *b* back into the model formula to get the predicted *y* values at the data points.

$$\hat{y} = Xb = Hy$$

 $H = X(X^TX)^{-1}X^T$

Statisticians use a hat (circumflex) over a letter to denote an estimate of a parameter or a prediction from a model. The projection matrix H is called the *hat matrix*, because it puts the "hat" on y.

The residuals are the difference between the observed and predicted *y* values.

$$r = y - \hat{y} = (I - H)y$$

The residuals are useful for detecting failures in the model assumptions, since they correspond to the errors, ε , in the model equation. By assumption, these errors each have independent normal distributions with mean zero and a constant variance.

The residuals, however, are correlated and have variances that depend on the locations of the data points. It is a common practice to scale ("Studentize") the residuals so they all have the same variance.

In the equation below, the scaled residual, t_i , has a Student's t distribution with (n-p-1) degrees of freedom

$$t_i = \frac{r_i}{\hat{\sigma}_{(i)} \sqrt{1 - h_i}}$$

where

$$\hat{\sigma}^{2}(i) = \frac{\|r\|^{2}}{n-p-1} - \frac{r_{i}^{2}}{(n-p-1)(1-h_{i})}$$

and:

- t_i is the scaled residual for the *i*th data point.
- r_i is the raw residual for the *i*th data point.
- *n* is the sample size.
- *p* is the number of parameters in the model.

• h_i is the *i*th diagonal element of *H*.

The left-hand side of the second equation is the estimate of the variance of the errors excluding the *i*th data point from the calculation.

A hypothesis test for outliers involves comparing t_i with the critical values of the t distribution. If t_i is large, this casts doubt on the assumption that this residual has the same variance as the others.

A confidence interval for the mean of each error is

$$c_i = r_i \pm t_{\left(1-\frac{\alpha}{2},\nu\right)} \hat{\sigma}_{(i)} \sqrt{1-h_i}$$

Confidence intervals that do not include zero are equivalent to rejecting the hypothesis (at a significance probability of α) that the residual mean is zero. Such confidence intervals are good evidence that the observation is an outlier for the given model.

Example: Multiple Linear Regression

The example comes from Chatterjee and Hadi (1986) in a paper on regression diagnostics. The data set (originally from Moore (1975)) has five predictor variables and one response.

```
load moore
X = [ones(size(moore,1),1) moore(:,1:5)];
```

Matrix X has a column of ones, and then one column of values for each of the five predictor variables. The column of ones is necessary for estimating the *y*-intercept of the linear model.

```
y = moore(:,6);
[b,bint,r,rint,stats] = regress(y,X);
```

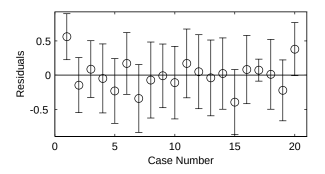
The *y*-intercept is b(1), which corresponds to the column index of the column of ones.

```
stats
stats =
0.8107 11.9886 0.0001
```

The elements of the vector stats are the regression \mathbb{R}^2 statistic, the F statistic (for the hypothesis test that all the regression coefficients are zero), and the p-value associated with this F statistic.

 R^2 is 0.8107 indicating the model accounts for over 80% of the variability in the observations. The F statistic of about 12 and its p-value of 0.0001 indicate that it is highly unlikely that all of the regression coefficients are zero.

```
rcoplot(r,rint)
```



The plot shows the residuals plotted in case order (by row). The 95% confidence intervals about these residuals are plotted as error bars. The first observation is an outlier since its error bar does not cross the zero reference line.

In problems with just a single predictor, it is simpler to use the polytool function (see "The polytool Demo" on page 11-5). This function can form an X matrix with predictor values, their squares, their cubes, and so on.

Quadratic Response Surface Models

Response Surface Methodology (RSM) is a tool for understanding the quantitative relationship between multiple input variables and one output variable.

Consider one output, *z*, as a polynomial function of two inputs, *x* and *y*. The function z = f(x,y) describes a two-dimensional surface in the space (x,y,z). Of course, you can have as many input variables as you want and the resulting surface becomes a hypersurface. You can have multiple output variables with a separate hypersurface for each one.

For three inputs (x_1, x_2, x_3) , the equation of a quadratic response surface is

 $y = b_0 + b_1 x_1 + b_2 x_2 + b_3 x_3 + \dots$ (linear terms) + $b_{12} x_1 x_2 + b_{13} x_1 x_3 + b_{23} x_2 x_3 + \dots$ (interaction terms) + $b_{11} x_1^2 + b_{22} x_2^2 + b_{33} x_3^2$ (quadratic terms)

It is difficult to visualize a k-dimensional surface in k+1 dimensional space for k>2. The function rstool is a graphical user interface (GUI) designed to make this visualization more intuitive, as is discussed in the next section.

Exploring Graphs of Multidimensional Polynomials

The function rstool is useful for fitting response surface models. The purpose of rstool is larger than just fitting and prediction for polynomial models. This GUI provides an environment for exploration of the graph of a multidimensional polynomial.

You can learn about rstool by trying the commands below. The chemistry behind the data in reaction.mat deals with reaction kinetics as a function of the partial pressure of three chemical reactants: hydrogen, n-pentane, and isopentane.

```
load reaction
rstool(reactants,rate,'quadratic',0.01,xn,yn)
```

You will see a "vector" of three plots. The dependent variable of all three plots is the reaction rate. The first plot has hydrogen as the independent variable. The second and third plots have n-pentane and isopentane respectively. Each plot shows the fitted relationship of the reaction rate to the independent variable at a fixed value of the other two independent variables. The fixed value of each independent variable is in an editable text box below each axis. You can change the fixed value of any independent variable by either typing a new value in the box or by dragging any of the three vertical lines to a new position.

When you change the value of an independent variable, all the plots update to show the current picture at the new point in the space of the independent variables.

Note that while this example only uses three inputs (reactants) and one output (rate), rstool can accommodate an arbitrary number of inputs and outputs. Interpretability may be limited by the size of the monitor for large numbers of inputs or outputs.

The GUI also has two pop-up menus. The **Export** menu facilitates saving various important variables in the GUI to the base workspace. Below the **Export** menu there is another menu that allows you to change the order of the polynomial model from within the GUI. If you used the commands above, this menu will have the string **Full Quadratic**. Other choices are:

- Linear has the constant and first order terms only.
- Pure Quadratic includes constant, linear and squared terms.
- Interactions includes constant, linear, and cross product terms.

The rstool GUI is used by the rsmdemo function to visualize the results of a designed experiment for studying a chemical reaction. See "The rsmdemo Demo" on page 11-19.

Stepwise Regression

Stepwise regression is a technique for choosing the variables to include in a multiple regression model. Forward stepwise regression starts with no model terms. At each step it adds the most statistically significant term (the one with the highest F statistic or lowest p-value) until there are none left. Backward stepwise regression starts with all the terms in the model and removes the least significant terms until all the remaining terms are statistically significant. It is also possible to start with a subset of all the terms and then add significant terms or remove insignificant terms.

An important assumption behind the method is that some input variables in a multiple regression do not have an important explanatory effect on the response. If this assumption is true, then it is a convenient simplification to keep only the statistically significant terms in the model.

One common problem in multiple regression analysis is multicollinearity of the input variables. The input variables may be as correlated with each other as they are with the response. If this is the case, the presence of one input variable in the model may mask the effect of another input. Stepwise regression used as a canned procedure is a dangerous tool because the resulting model may include different variables depending on the choice of starting model and inclusion strategy.

The following example explores an interactive tool for stepwise regression.

Example: Stepwise Regression

The Statistics Toolbox provides an interactive graphical user interface (GUI) to make comparison of competing models more understandable. You can explore the GUI using the Hald (1960) data set. Here are the commands to get started.

```
load hald
stepwise(ingredients,heat)
```

The Hald data come from a study of the heat of reaction of various cement mixtures. There are four components in each mixture, and the amount of heat produced depends on the amount of each ingredient in the mixture.

The interface consists of three interactively linked figure windows. Two of these are discussed in the following sections:

• "Stepwise Regression Plot" on page 4-25

• "Stepwise Regression Diagnostics Table" on page 4-25

All three windows have *hot* regions. When your mouse is above one of these regions, the pointer changes from an arrow to a circle. Clicking on this point initiates some activity in the interface.

Stepwise Regression Plot

This plot shows the regression coefficient and confidence interval for every term (in or out of the model). The green lines represent terms in the model while red lines indicate terms that are not currently in the model.

Statistically significant terms are solid lines. Dotted lines show that the fitted coefficient is not significantly different from zero.

Clicking on a line in this plot toggles its state. That is, a term currently in the model (green line) is removed (turns red), and a term currently not in the model (red line) is added (turns green).

The coefficient for a term out of the model is the coefficient resulting from adding that term to the current model.

Scale Inputs

Pressing this button centers and normalizes the columns of the input matrix to have a standard deviation of one.

Export

This pop-up menu allows you to export variables from the stepwise function to the base workspace.

Close

The Close button removes all the figure windows.

Stepwise Regression Diagnostics Table

This table is a quantitative view of the information in the Stepwise Regression Plot. The table shows the Hald model with the second and third terms removed.

		Confidence	Intervals
Column #	Parameter	Lower	Upper
1	1.44	1.02	1.86
2	0.4161	-0.1602	0.9924
3	-0.41	-1.029	0.2086
4	-0.614	-0.7615	-0.4664
RMSE	R-square	F	P
2.734	0.9725	176.6	1.581e-08

Coefficients and Confidence Intervals

The table at the top of the figure shows the regression coefficient and confidence interval for every term (in or out of the model.) The green rows in the table (on your monitor) represent terms in the model while red rows indicate terms not currently in the model.

Clicking on a row in this table toggles the state of the corresponding term. That is, a term currently in the model (green row) is removed (turns red), and a term currently not in the model (red row) is added to the model (turns green).

The coefficient for a term out of the model is the coefficient resulting from adding that term to the current model.

Additional Diagnostic Statistics

There are also several diagnostic statistics at the bottom of the table:

- RMSE the root mean squared error of the current model.
- R-square the amount of response variability explained by the model.
- F the overall F statistic for the regression.
- P the associated significance probability.

Close Button

Shuts down all windows.

Help Button

Activates online help.

Stepwise History

This plot shows the RMSE and a confidence interval for every model generated in the course of the interactive use of the other windows.

Recreating a Previous Model

Clicking on one of these lines recreates the current model at that point in the analysis using a *new* set of windows. You can thus compare the two candidate models directly.

Generalized Linear Models

So far, the functions in this section have dealt with models that have a linear relationship between the response and one or more predictors. Sometimes you may have a nonlinear relationship instead. To fit nonlinear models you can use the functions described in "Nonlinear Regression Models" on page 5-1.

There are some nonlinear models, known as generalized linear models, that you can fit using simpler linear methods. To understand generalized linear models, first let's review the linear models we have seen so far. Each of these models has the following three characteristics:

- The response has a normal distribution with mean μ .
- A coefficient vector *b* defines a linear combination *X***b* of the predictors *X*.
- The model equates the two as $\mu = X^*b$.

In generalized linear models, these characteristics are generalized as follows:

- The response has a distribution that may be normal, binomial, Poisson, gamma, or inverse Gaussian, with parameters including a mean μ .
- A coefficient vector *b* defines a linear combination *X***b* of the predictors *X*.
- A link function $f(\cdot)$ defines the link between the two as $f(\mu) = X^*b$.

The following example explores this in greater detail.

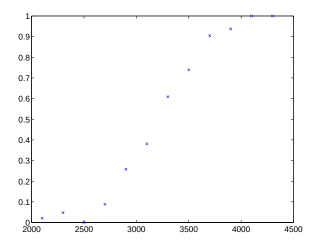
Example: Generalized Linear Models

For example, consider the following data derived from the carbig data set. We have cars of various weights, and we record the total number of cars of each weight and the number qualifying as poor-mileage cars because their miles per gallon value is below some target. (Suppose we don't know the miles per gallon for each car, only the number passing the test.) It might be reasonable to assume that the value of the variable poor follows a binomial distribution with parameter N=total and with a p parameter that depends on the car weight. A plot shows that the proportion of poor-mileage cars follows a nonlinear S-shape.

```
w = [2100 2300 2500 2700 2900 3100 3300 3500 3700 3900 4100 4300]';
poor = [1 2 0 3 8 8 14 17 19 15 17 21]';
total = [48 42 31 34 31 21 23 23 21 16 17 21]';
```

[w poor ans =	total]			
	2100	1	48	
	2300	2	42	
	2500	0	31	
	2700	3	34	
	2900	8	31	
	3100	8	21	
	3300	14	23	
	3500	17	23	
	3700	19	21	
	3900	15	16	
	4100	17	17	
	4300	21	21	

plot(w,poor./total,'x')



This shape is typical of graphs of proportions, as they have natural boundaries at 0.0 and 1.0.

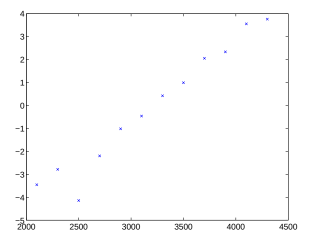
A linear regression model would not produce a satisfactory fit to this graph. Not only would the fitted line not follow the data points, it would produce invalid proportions less than 0 for light cars, and higher than 1 for heavy cars.

There is a class of regression models for dealing with proportion data. The logistic model is one such model. It defines the relationship between proportion p and weight w to be

$$\log\left(\frac{p}{1-p}\right) = b_1 + b_2 w$$

Is this a good model for our data? It would be helpful to graph the data on this scale, to see if the relationship appears linear. However, some of our proportions are 0 and 1, so we cannot explicitly evaluate the left-hand-side of the equation. A useful trick is to compute adjusted proportions by adding small increments to the poor and total values — say a half observation to poor and a full observation to total. This keeps the proportions within range. A graph now shows a more nearly linear relationship.

```
padj = (poor+.5) ./ (total+1);
plot(w,log(padj./(1-padj)),'x')
```



We can use the glmfit function to fit this logistic model.

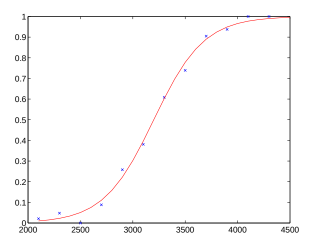
```
b = glmfit(w,[poor total],'binomial')
b =
    -13.3801
    0.0042
```

To use these coefficients to compute a fitted proportion, we have to invert the logistic relationship. Some simple algebra shows that the logistic equation can also be written as

$$p = \frac{1}{1 + \exp(-b_1 - b_2 w)}$$

Fortunately, the function glmval can decode this link function to compute the fitted values. Using this function we can graph fitted proportions for a range of car weights, and superimpose this curve on the original scatter plot.

```
x = 2100:100:4500;
y = glmval(b,x, logit );
plot(w,poor./total,'x',x,y,'r-')
```



Generalized linear models can fit a variety of distributions with a variety of relationships between the distribution parameters and the predictors. A full description is beyond the scope of this document. For more information see Dobson (1990), or McCullagh and Nelder (1990). Also see the reference material for glmfit.

Robust and Nonparametric Methods

As mentioned in the previous sections, regression and analysis of variance procedures depend on certain assumptions, such as a normal distribution for the error term. Sometimes such an assumption is not warranted. For example, if the distribution of the errors is asymmetric or prone to extreme outliers, that is a violation of the assumption of normal errors.

The Statistics Toolbox has a robust regression function that is useful when there may be outliers. Robust methods are designed to be relatively insensitive to large changes in a small part of the data.

The Statistics Toolbox also has nonparametric versions of the one-way and two-way analysis of variance functions. Unlike classical tests, nonparametric tests make only mild assumptions about the data, and are appropriate when the distribution of the data is not normal. On the other hand, they are less powerful than classical methods for normally distributed data.

The following sections describe the robust regression and nonparametric functions in greater detail:

- "Robust Regression" on page 4-32
- "Kruskal-Wallis Test" on page 4-34
- "Friedman's Test" on page 4-34

Both of the nonparametric functions described here can return a stats structure that you can use as input to the multcompare function to perform multiple comparisons.

Note See for information on another type of nonparametric regression.

Robust Regression

In "Example: Multiple Linear Regression" on page 4-20 we found an outlier when we used ordinary least squares regression to model a response as a function of five predictors. How did that outlier affect the results?

Let's estimate the coefficients using the robustfit function.

load moore

```
x = moore(:,1:5);
y = moore(:,6);
[br,statsr] = robustfit(x,y);
br
br =
    -1.7742
    0.0000
    0.0009
    0.0002
    0.0002
    0.0062
    0.0001
```

Compare these estimates to those we obtained from the regress function.

```
b
b =
-2.1561
-0.0000
0.0013
0.0001
0.0079
0.0001
```

To understand why the two differ, it is helpful to look at the weight variable from the robust fit. It measures how much weight was given to each point during the fit. In this case, the first point had a very low weight so it was effectively ignored.

```
statsr.w'
ans =
 Columns 1 through 7
   0.0577
            0.9977
                      0.9776
                                0.9455
                                         0.9687
                                                   0.8734
                                                             0.9177
 Columns 8 through 14
   0.9990
            0.9653
                      0.9679
                                0.9768
                                         0.9882
                                                   0.9998
                                                             0.9979
 Columns 15 through 20
    0.8185
              0.9757
                         0.9875
                                   0.9991
                                              0.9021
                                                        0.6953
```

For another example illustrating robust fitting, see "The robustdemo Demo" on page 11-22.

Kruskal-Wallis Test

In "One-Way Analysis of Variance (ANOVA)" on page 4-4 we used one-way analysis of variance to determine if the bacteria counts of milk varied from shipment to shipment. Our one-way analysis rested on the assumption that the measurements were independent, and that each had a normal distribution with a common variance and with a mean that was constant in each column. We concluded that the column means were not all the same. Let's repeat that analysis using a nonparametric procedure.

The Kruskal-Wallis test is a nonparametric version of one-way analysis of variance. The assumption behind this test is that the measurements come from a continuous distribution, but not necessarily a normal distribution. The test is based on an analysis of variance using the ranks of the data values, not the data values themselves. Output includes a table similar to an anova table, and a box plot.

We can run this test as follows.

p = kruskalwallis(hogg)
p =
 0.0020

The low p-value means the Kruskal-Wallis test results agree with the one-way analysis of variance results.

Friedman's Test

In "Two-Way Analysis of Variance (ANOVA)" on page 4-9 we used two-way analysis of variance to study the effect of car model and factory on car mileage. We tested whether either of these factors had a significant effect on mileage, and whether there was an interaction between these factors. We concluded that there was no interaction, but that each individual factor had a significant effect. Now we will see if a nonparametric analysis will lead to the same conclusion.

Friedman's test is a nonparametric test for data having a two-way layout (data grouped by two categorical factors). Unlike two-way analysis of variance, Friedman's test does not treat the two factors symmetrically and it does not test for an interaction between them. Instead, it is a test for whether the columns are different after adjusting for possible row differences. The test is based on an analysis of variance using the ranks of the data across categories of the row factor. Output includes a table similar to an anova table. We can run Friedman's test as follows.

```
p = friedman(mileage, 3)
ans =
7.4659e-004
```

Recall the classical analysis of variance gave a p-value to test column effects, row effects, and interaction effects. This p-value is for column effects. Using either this p-value or the p-value from ANOVA (p < 0.0001), we conclude that there are significant column effects.

In order to test for row effects, we need to rearrange the data to swap the roles of the rows in columns. For a data matrix x with no replications, we could simply transpose the data and type

p = friedman(x)

With replicated data it is slightly more complicated. A simple way is to transform the matrix into a three-dimensional array with the first dimension representing the replicates, swapping the other two dimensions, and restoring the two-dimensional shape.

```
x = reshape(mileage, [3 2 3]);
x = permute(x, [1 3 2]);
x = reshape(x, [9 2])
x =
   33.3000
             32,6000
   33,4000
             32,5000
   32.9000
             33.0000
   34.5000
             33.4000
   34.8000
             33.7000
   33.8000
             33,9000
   37.4000
             36,6000
   36.8000
             37.0000
   37.6000
             36.7000
friedman(x, 3)
ans =
    0.0082
```

Again, the conclusion is similar to the conclusion from the classical analysis of variance. Both this p-value and the one from ANOVA (p = 0.0039) lead us to conclude there are significant row effects.

You cannot use Friedman's test to test for interactions between the row and column factors.

5

Nonlinear Regression Models

Introduction	•	•	5-2
Nonlinear Least Squares			5 - 3
Example: Nonlinear Modeling			5 - 3
An Interactive GUI for Nonlinear Fitting and Prediction	•		5-7
Regression and Classification Trees			5-8

Introduction

Not all relationships are well described by a linear regression model. The Statistics Toolbox provides two nonlinear regression techniques that may be preferable to linear regression in some cases.

- "Nonlinear Least Squares" on page 5-3 fits a model that has a known parametric form but unknown parameter values.
- "Regression and Classification Trees" on page 5-8 approximates a regression relationship using a decision tree. Such a tree seeks to partition the data set into regions, using values of the predictor variables, so that the response variables are roughly constant in each region.

Nonlinear Least Squares

Response Surface Methodology (RSM) is an empirical modeling approach using polynomials as local approximations to the true input/output relationship. This empirical approach is often adequate for process improvement in an industrial setting.

In scientific applications there is usually relevant theory that allows us to make a mechanistic model. Often such models are nonlinear in the unknown parameters. Nonlinear models are more difficult to fit, requiring iterative methods that start with an initial guess of the unknown parameters. Each iteration alters the current guess until the algorithm converges.

The Statistics Toolbox has functions for fitting nonlinear models of the form

$$y = f(X, \beta) + \varepsilon$$

where:

- *y* is an-*n* by-1 vector of observations.
- *f* is any function of *X* and β .
- *X* is an *n*-by-*p* matrix of input variables.
- β is a *p*-by-1 vector of unknown parameters to be estimated.
- ε is an *n*-by-1 vector of random disturbances.

This is explored further in the following sections:

- "Example: Nonlinear Modeling" on page 5-3
- "An Interactive GUI for Nonlinear Fitting and Prediction" on page 5-7

Example: Nonlinear Modeling

The Hougen-Watson model (Bates and Watts, [2]) for reaction kinetics is one specific example of this type. The form of the model is

$$rate = \frac{\beta_1 \cdot x_2 - x_3 / \beta_5}{1 + \beta_2 \cdot x_1 + \beta_3 \cdot x_2 + \beta_4 \cdot x_3}$$

where β_1 , β_2 , ..., β_5 are the unknown parameters, and x_1 , x_2 , and x_3 are the three input variables. The three inputs are hydrogen, n-pentane, and

isopentane. It is easy to see that the parameters do not enter the model linearly.

The file reaction.mat contains simulated data from this reaction.

load reaction who Your variables are: beta rate xn model reactants yn

The variables are as follows:

- rate is a 13-by-1 vector of observed reaction rates.
- reactants is a 13-by-3 matrix of reactants.
- beta is 5-by-1 vector of initial parameter estimates.
- model is a string containing the nonlinear function name.
- xn is a string matrix of the names of the reactants.
- yn is a string containing the name of the response.

The data and model are explored further in the following sections:

- "Fitting the Hougen-Watson Model" on page 5-4
- "Confidence Intervals on the Parameter Estimates" on page 5-6
- "Confidence Intervals on the Predicted Responses" on page 5-6
- "An Interactive GUI for Nonlinear Fitting and Prediction" on page 5-7

Fitting the Hougen-Watson Model

The Statistics Toolbox provides the function nlinfit for finding parameter estimates in nonlinear modeling. nlinfit returns the least squares parameter estimates. That is, it finds the parameters that minimize the sum of the squared differences between the observed responses and their fitted values. It uses the Gauss-Newton algorithm with Levenberg-Marquardt modifications for global convergence.

nlinfit requires the input data, the responses, and an initial guess of the unknown parameters. You must also supply the name of a function that takes the input data and the current parameter estimate and returns the predicted responses. In MATLAB terminology, nlinfit is called a "function" function.

Here is the hougen function.

function yhat = hougen(beta,x) %HOUGEN Hougen-Watson model for reaction kinetics. YHAT = HOUGEN(BETA, X) gives the predicted values of the % % reaction rate, YHAT, as a function of the vector of parameters, BETA, and the matrix of data, X. % % BETA must have five elements and X must have three % columns. % % The model form is: v = (b1*x2 - x3/b5)./(1+b2*x1+b3*x2+b4*x3)% b1 = beta(1);b2 = beta(2);b3 = beta(3);b4 = beta(4);b5 = beta(5);x1 = x(:,1); $x^2 = x(:,2);$ x3 = x(:,3);yhat = (b1*x2 - x3/b5)./(1+b2*x1+b3*x2+b4*x3);

To fit the reaction data, call the function nlinfit.

```
load reaction
betahat = nlinfit(reactants,rate,'hougen',beta)
betahat =
    1.2526
    0.0628
    0.0400
    0.1124
    1.1914
```

nlinfit has two optional outputs. They are the residuals and Jacobian matrix at the solution. The residuals are the differences between the observed and fitted responses. The Jacobian matrix is the direct analog of the matrix X in the standard linear regression model.

These outputs are useful for obtaining confidence intervals on the parameter estimates and predicted responses.

Confidence Intervals on the Parameter Estimates

Using nlparci, form 95% confidence intervals on the parameter estimates, betahat, from the reaction kinetics example.

```
[betahat,resid,J] = nlinfit(reactants,rate,'hougen',beta);
betaci = nlparci(betahat,resid,J)
```

betaci =

-0.7467	3.2519
-0.0377	0.1632
-0.0312	0.1113
-0.0609	0.2857
-0.7381	3.1208

Confidence Intervals on the Predicted Responses

Using nlpredci, form 95% confidence intervals on the predicted responses from the reaction kinetics example.

```
[yhat,delta] = nlpredci('hougen',reactants,betahat,resid,J);
opd = [rate yhat delta]
```

opd =

8.5500	8.2937	0.9178
3.7900	3.8584	0.7244
4.8200	4.7950	0.8267
0.0200	-0.0725	0.4775
2.7500	2.5687	0.4987
14.3900	14.2227	0.9666
2.5400	2.4393	0.9247
4.3500	3.9360	0.7327
13.0000	12.9440	0.7210
8.5000	8.2670	0.9459
0.0500	-0.1437	0.9537
11.3200	11.3484	0.9228
3.1300	3.3145	0.8418

Matrix opd has the observed rates in column 1 and the predictions in column 2. The 95% confidence interval is column $2\pm$ column 3. These are simultaneous confidence intervals for the estimated function at each input value. They are not intervals for new response observations at those inputs, even though most of the confidence intervals do contain the original observations.

An Interactive GUI for Nonlinear Fitting and Prediction

The function nlintool for nonlinear models is a direct analog of rstool for polynomial models. nlintool calls nlinfit and requires the same inputs.

The purpose of nlintool is larger than just fitting and prediction for nonlinear models. This GUI provides an environment for exploration of the graph of a multidimensional nonlinear function.

If you have already loaded reaction.mat, you can start nlintool.

```
nlintool(reactants,rate, 'hougen',beta,0.01,xn,yn)
```

You will see a "vector" of three plots. The dependent variable of all three plots is the reaction rate. The first plot has hydrogen as the independent variable. The second and third plots have n-pentane and isopentane respectively.

Each plot shows the fitted relationship of the reaction rate to the independent variable at a fixed value of the other two independent variables. The fixed value of each independent variable is in an editable text box below each axis. You can change the fixed value of any independent variable by either typing a new value in the box or by dragging any of the three vertical lines to a new position.

When you change the value of an independent variable, all the plots update to show the current picture at the new point in the space of the independent variables.

Note that while this example only uses three reactants, nlintool can accommodate an arbitrary number of independent variables. Interpretability may be limited by the size of the monitor for large numbers of inputs.

Regression and Classification Trees

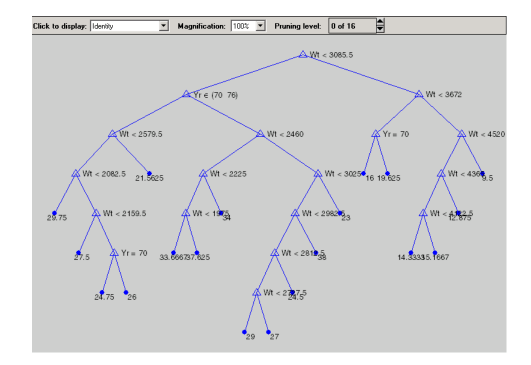
In nonlinear least squares we suppose that we know the form of the relationship between the response and predictor. Suppose instead that we don't know that relationship, and also that we are unwilling to assume the relationship can be well approximated by a linear model. We need a more nonparametric type of regression fitting approach. One such approach is based on "trees."

A regression tree is a sequence of questions that can be answered as yes or no, plus a set of fitted response values. Each question asks whether a predictor satisfies a given condition. Predictors can be continuous or discrete. Depending on the answers to one question, we either proceed to another question or we arrive at a fitted response value.

In this example we fit a regression tree to variables from the carsmall data set. We use the same variables as in the Analysis of Covariance example (see "The aoctool Demo" on page 11-10), so we have one continuous predictor (car weight) and one discrete predictor (model year).

Our goal is to model mileage (MPG) as a function of car weight and model year. First we load the data and then create a matrix x of predictor values and a vector y of response variables. We fit a regression tree, specifying the model year column as a categorical variable. In this data set there are cars from the three different model years 1970, 1976, and 1982.

```
load carsmall
x = [Weight,Model_Year];
y = MPG;
t = treefit(x,y,'catidx',2);
treedisp(t,'name',{'Wt' 'Yr'});
```



Now we want to use this model to determine the predicted mileage for a car weighing 3000 pounds from model year 1982. Start at the top node. The weight is less than the cutoff value of 3085.5, so we take the left branch. The model year is not 1970 or 1976, so we take the right branch. We continue moving down the tree until we arrive at a terminal node that gives the predicted value. In this case, the predicted value is 38 miles per gallon. We can use the treeval function to find the fitted value for any set of predictor values.

```
treeval(t,[3000 82])
ans =
38
```

With a tree like this one having many branches, there is a danger that it fits the current data set well but would not do a good job at predicting new values. Some of its lower branches may be strongly affected by outliers and other artifacts of the current data set. If possible we would prefer to find a simpler tree that avoids this problem of over-fitting.

We'll estimate the best tree size by cross validation. First we compute a "resubstitution" estimate of the error variance for this tree and a sequence of simpler trees and plot it as the lower (blue) line in the figure. This estimate probably under-estimates the true error variance. Then we compute a "cross-validation" estimate of the same quantity and plot it as the upper (red) line. The cross-validation procedure also provides us with an estimate of the pruning level, best, needed to achieve the best tree size.

```
[c,s,ntn] = treetest(t, 'resub');
[c2,s2,n2,best] = treetest(t, 'cross',x,y);
plot(ntn,c,'b-', n2,c2,'r-', n2(best+1),c2(best+1),'mo');
xlabel('Number of terminal nodes')
ylabel('Residual variance')
legend('Resubstitution error','Cross-validation
error', 'Estimated best tree size')
best
best =
     10
   70
                                 Resubstitution error
                                 Cross-validation error
                                 Estimated best tree size
   60
  50
 Residual variance
  40
   30
  20
   10
```

0^L 0

5

10

Number of terminal nodes

15

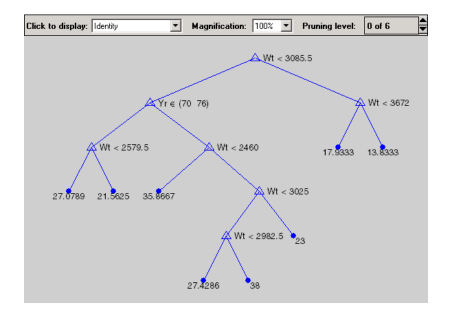
20

The best tree is the one that has a residual variance that is no more than one standard error above the minimum value along the cross-validation line. In this case the variance is just over 14. The output best takes on values starting with 0 (representing no pruning), so we need to add 1 to use it as an index into the other output arguments.

```
c2(best+1)
ans =
14.3440
```

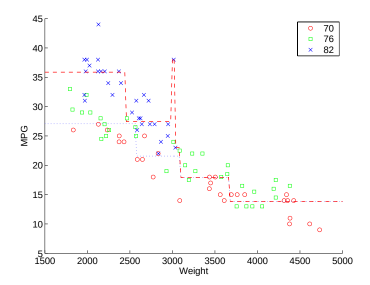
Use the output best to create a smaller tree that is pruned to our estimated best size.

```
t0 = treeprune(t, 'level', best);
treedisp(t0, 'name', {'Wt' 'Yr'})
```



Now plot the original data and overlay the fitted values that we get using this tree. Notice that this tree does not distinguish between cars from 1970 or 1976, so we'll create a vector yold containing fitted values for 1976 and another ynew for year 1982. Cars from 1970 have the same fitted values as those from 1976.

```
xx = (1500:20:5000)';
ynew = treeval(t0,[xx 82*ones(size(xx))]);
yold = treeval(t0,[xx 76*ones(size(xx))]);
gscatter(Weight,MPG,Model_Year,'rgb','osx');
hold on; plot(xx,yold,'b:', xx,ynew,'r--'); hold off
```



The tree functions (treedisp, treefit, treeprune, treetest, and treeval) can also accept a categorical response variable. In that case, the fitted value from the tree is the category with the highest predicted probability for the range of predictor values falling in a given node. The Statistics Toolbox demo entitled Classification of Fisher's Iris Data shows how to use decision trees for classification.

6

Hypothesis Tests

Introduction	•	•	•	•	•	•	•	•	•	•	•	•	6-2
Hypothesis Test Terminology	•	•	•	•	•			•	•	•	•	•	6-3
Hypothesis Test Assumptions								•	•	•	•	•	6-4
Example: Hypothesis Testing	•		•	•					•	•	•	•	6-5
Available Hypothesis Tests .									•			•	6-9

Introduction

A hypothesis test is a procedure for determining if an assertion about a characteristic of a population is reasonable.

For example, suppose that someone says that the average price of a gallon of regular unleaded gas in Massachusetts is \$1.15. How would you decide whether this statement is true? You could try to find out what every gas station in the state was charging and how many gallons they were selling at that price. That approach might be definitive, but it could end up costing more than the information is worth.

A simpler approach is to find out the price of gas at a small number of randomly chosen stations around the state and compare the average price to \$1.15.

Of course, the average price you get will probably not be exactly \$1.15 due to variability in price from one station to the next. Suppose your average price was \$1.18. Is this three cent difference a result of chance variability, or is the original assertion incorrect? A hypothesis test can provide an answer.

The following sections provide an overview of hypothesis testing with the Statistics Toolbox:

- "Hypothesis Test Terminology" on page 6-3
- "Hypothesis Test Assumptions" on page 6-4
- "Example: Hypothesis Testing" on page 6-5
- "Available Hypothesis Tests" on page 6-9

Hypothesis Test Terminology

To get started, there are some terms to define and assumptions to make:

- The *null hypothesis* is the original assertion. In this case the null hypothesis is that the average price of a gallon of gas is \$1.15. The notation is H_0 : $\mu = 1.15$.
- There are three possibilities for the *alternative hypothesis*. You might only be interested in the result if gas prices were actually higher. In this case, the alternative hypothesis is H₁: $\mu > 1.15$. The other possibilities are H₁: $\mu < 1.15$ and H₁: $\mu \neq 1.15$.
- The significance level is related to the degree of certainty you require in order to reject the null hypothesis in favor of the alternative. By taking a small sample you cannot be certain about your conclusion. So you decide in advance to reject the null hypothesis if the probability of observing your sampled result is less than the significance level. For a typical significance level of 5%, the notation is $\alpha = 0.05$. For this significance level, the probability of incorrectly rejecting the null hypothesis when it is actually true is 5%. If you need more protection from this error, then choose a lower value of α .
- The *p*-value is the probability of observing the given sample result under the assumption that the null hypothesis is true. If the p-value is less than α , then you reject the null hypothesis. For example, if $\alpha = 0.05$ and the p-value is 0.03, then you reject the null hypothesis.

The converse is not true. If the p-value is greater than α , you have insufficient evidence to reject the null hypothesis.

• The outputs for many hypothesis test functions also include *confidence intervals*. Loosely speaking, a confidence interval is a range of values that have a chosen probability of containing the true hypothesized quantity. Suppose, in our example, 1.15 is inside a 95% confidence interval for the mean, μ . That is equivalent to being unable to reject the null hypothesis at a significance level of 0.05. Conversely if the 100(1- α) confidence interval does not contain 1.15, then you reject the null hypothesis at the α level of significance.

Hypothesis Test Assumptions

The difference between hypothesis test procedures often arises from differences in the assumptions that the researcher is willing to make about the data sample. For example, the Z-test assumes that the data represents independent samples from the same normal distribution and that you know the standard deviation, σ . The t-test has the same assumptions except that you estimate the standard deviation using the data instead of specifying it as a known quantity.

Both tests have an associated signal-to-noise ratio

$$Z = \frac{\bar{x} - \mu}{\sigma} \quad or \quad T = \frac{\bar{x} - \mu}{s}$$

where $\bar{x} = \sum_{i=1}^{n} \frac{x_i}{n}$

The signal is the difference between the average and the hypothesized mean. The noise is the standard deviation posited or estimated.

If the null hypothesis is true, then Z has a standard normal distribution, N(0,1). T has a Student's t distribution with the degrees of freedom, ν , equal to one less than the number of data values.

Given the observed result for Z or T, and knowing the distribution of Z and T assuming the null hypothesis is true, it is possible to compute the probability (p-value) of observing this result. A very small p-value casts doubt on the truth of the null hypothesis. For example, suppose that the p-value was 0.001, meaning that the probability of observing the given Z or T was one in a thousand. That should make you skeptical enough about the null hypothesis that you reject it rather than believe that your result was just a lucky 999 to 1 shot.

There are also nonparametric tests that do not even require the assumption that the data come from a normal distribution. In addition, there are functions for testing whether the normal assumption is reasonable.

Example: Hypothesis Testing

This example uses the gasoline price data in gas.mat. There are two samples of 20 observed gas prices for the months of January and February, 1993.

```
load gas
prices = [price1 price2];
```

As a first step, you may want to test whether the samples from each month follow a normal distribution. As each sample is relatively small, you might choose to perform a Lilliefors test (rather than a Jarque-Bera test):

```
lillietest(price1)
ans =
     0
lillietest(price2)
ans =
     0
```

The result of the hypothesis test is a Boolean value that is 0 when you do not reject the null hypothesis, and 1 when you do reject that hypothesis. In each case, there is no need to reject the null hypothesis that the samples have a normal distribution.

Suppose it is historically true that the standard deviation of gas prices at gas stations around Massachusetts is four cents a gallon. The Z-test is a procedure for testing the null hypothesis that the average price of a gallon of gas in January (price1) is \$1.15.

The Boolean output is h = 0, so you do not reject the null hypothesis.

The result suggests that \$1.15 is reasonable. The 95% confidence interval [1.1340 1.1690] neatly brackets \$1.15.

What about February? Try a t-test with price2. Now you are not assuming that you know the standard deviation in price.

With the Boolean result h = 1, you can reject the null hypothesis at the default significance level, 0.05.

It looks like \$1.15 is not a reasonable estimate of the gasoline price in February. The low end of the 95% confidence interval is greater than 1.15.

The function ttest2 allows you to compare the means of the two data samples.

The confidence interval (ci above) indicates that gasoline prices were between one and six cents lower in January than February.

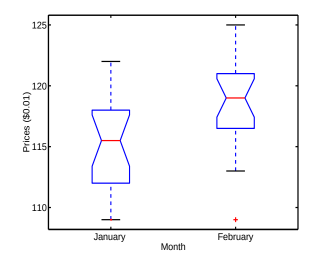
If the two samples were not normally distributed but had similar shape, it would have been more appropriate to use the nonparametric rank sum test in place of the t-test. We can still use the rank sum test with normally distributed data, but it is less powerful than the t-test.

As might be expected, the rank sum test leads to the same conclusion but it is less sensitive to the difference between samples (higher p-value).

The box plot below gives the same conclusion graphically. Note that the notches have little, if any, overlap. Refer to "Statistical Plots" on page 8-1 for more information about box plots.

```
boxplot(prices,1)
set(gca,'XtickLabel',str2mat('January','February'))
xlabel('Month')
ylabel('Prices ($0.01)')
```

6 Hypothesis Tests



Available Hypothesis Tests

The Statistics Toolbox has functions for performing the following tests.

Function	What it Tests	
jbtest	Normal distribution for one sample	
kstest	Any specified distribution for one sample	
kstest2	Equal distributions for two samples	
lillietest	Normal distribution for one sample	
ranksum	Median of two unpaired samples	
signrank	Median of two paired samples	
signtest	Median of two paired samples	
ttest	Mean of one normal sample	
ttest2	Mean of two normal samples	
ztest	Mean of normal sample with known standard deviation	



7

Multivariate Statistics

Introduction
Principal Components Analysis
Example: Principal Components Analysis
The Principal Components (First Output)
The Component Scores (Second Output) 7-7
The Component Variances (Third Output)
Hotelling's T ² (Fourth Output) $\ldots \ldots \ldots \ldots \ldots \ldots \ldots $
Factor Analysis
Example: Finding Common Factors Affecting Stock Prices 7-13
Factor Rotation
Predicting Factor Scores
Comparison of Factor Analysis and Principal Components
Analysis
Multivariate Analysis of Variance (MANOVA) 7-20
Example: Multivariate Analysis of Variance
Cluster Analysis
Hierarchical Clustering
K-Means Clustering
Classical Multidimensional Scaling
Overview
Reconstructing a Map from Inter-City Distances

Introduction

Multivariate statistics is an omnibus term for a number of different statistical methods. The defining characteristic of these methods is that they all aim to understand a data set by considering a group of variables together rather than focusing on only one variable at a time.

The Statistics Toolbox has functions for these methods:

- "Principal Components Analysis" on page 7-3
- "Factor Analysis" on page 7-13
- "Multivariate Analysis of Variance (MANOVA)" on page 7-20
- "Cluster Analysis" on page 7-26
- "Classical Multidimensional Scaling" on page 7-47

Principal Components Analysis

One of the difficulties inherent in multivariate statistics is the problem of visualizing multidimensionality. In MATLAB, the plot command displays a graph of the relationship between two variables. The plot3 and surf commands display different three-dimensional views. When there are more than three variables, it stretches the imagination to visualize their relationships.

Fortunately, in data sets with many variables, groups of variables often move together. One reason for this is that more than one variable may be measuring the same driving principle governing the behavior of the system. In many systems there are only a few such driving forces. But an abundance of instrumentation allows us to measure dozens of system variables. When this happens, we can take advantage of this redundancy of information. We can simplify our problem by replacing a group of variables with a single new variable.

Principal components analysis is a quantitatively rigorous method for achieving this simplification. The method generates a new set of variables, called *principal components*. Each principal component is a linear combination of the original variables. All the principal components are orthogonal to each other so there is no redundant information. The principal components as a whole form an orthogonal basis for the space of the data.

There are an infinite number of ways to construct an orthogonal basis for several columns of data. What is so special about the principal component basis?

The first principal component is a single axis in space. When you project each observation on that axis, the resulting values form a new variable. And the variance of this variable is the maximum among all possible choices of the first axis.

The second principal component is another axis in space, perpendicular to the first. Projecting the observations on this axis generates another new variable. The variance of this variable is the maximum among all possible choices of this second axis.

The full set of principal components is as large as the original set of variables. But it is commonplace for the sum of the variances of the first few principal components to exceed 80% of the total variance of the original data. By examining plots of these few new variables, researchers often develop a deeper understanding of the driving forces that generated the original data.

The function princomp is used to find the principal components. The following sections provide an example and explain the four outputs of princomp:

- "Example: Principal Components Analysis" on page 7-4
- "The Principal Components (First Output)" on page 7-7
- "The Component Scores (Second Output)" on page 7-7
- "The Component Variances (Third Output)" on page 7-10
- "Hotelling's T2 (Fourth Output)" on page 7-12

Example: Principal Components Analysis

Let us look at a sample application that uses nine different indices of the quality of life in 329 U.S. cities. These are climate, housing, health, crime, transportation, education, arts, recreation, and economics. For each index, higher is better; so, for example, a higher index for crime means a lower crime rate.

We start by loading the data in cities.mat.

load cities whos			
Name	Size	Bytes	Class
categories	9x14	252	char array
names	329x43	28294	char array
ratings	329x9	23688	double array

The whos command generates a table of information about all the variables in the workspace.

The cities data set contains three variables:

- categories, a string matrix containing the names of the indices.
- names, a string matrix containing the 329 city names.
- ratings, the data matrix with 329 rows and 9 columns.

Let's look at the value of the categories variable.

```
categories
```

```
categories =
   climate
   housing
   health
   crime
   transportation
   education
   arts
   recreation
   economics
```

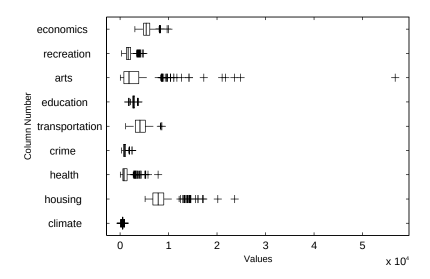
Now, let's look at the first several rows of names variable.

```
first5 = names(1:5,:)
first5 =
   Abilene, TX
   Akron, OH
   Albany, GA
   Albany-Troy, NY
   Albuquerque, NM
```

To get a quick impression of the ratings data, make a box plot.

```
boxplot(ratings,0,'+',0)
set(gca,'YTicklabel',categories)
```

These commands generate the plot below. Note that there is substantially more variability in the ratings of the arts and housing than in the ratings of crime and climate.



Ordinarily you might also graph pairs of the original variables, but there are 36 two-variable plots. Perhaps principal components analysis can reduce the number of variables we need to consider.

Sometimes it makes sense to compute principal components for raw data. This is appropriate when all the variables are in the same units. Standardizing the data is reasonable when the variables are in different units or when the variance of the different columns is substantial (as in this case).

You can standardize the data by dividing each column by its standard deviation.

```
stdr = std(ratings);
sr = ratings./repmat(stdr,329,1);
```

Now we are ready to find the principal components.

[pcs,newdata,variances,t2] = princomp(sr);

The following sections explain the four outputs from princomp:

- "The Principal Components (First Output)" on page 7-7
- "The Component Scores (Second Output)" on page 7-7
- "The Component Variances (Third Output)" on page 7-10
- "Hotelling's T2 (Fourth Output)" on page 7-12

The Principal Components (First Output)

The first output of the princomp function, pcs, contains the nine principal components. These are the linear combinations of the original variables that generate the new variables.

Let's look at the first three principal component vectors.

рЗ	= pcs(:,	1:3)	
рЗ	=		
	0.2064	0.2178	-0.6900
	0.3565	0.2506	-0.2082
	0.4602	-0.2995	-0.0073
	0.2813	0.3553	0.1851
	0.3512	-0.1796	0.1464
	0.2753	-0.4834	0.2297
	0.4631	-0.1948	-0.0265
	0.3279	0.3845	-0.0509
	0.1354	0.4713	0.6073

The largest weights in the first column (first principal component) are the third and seventh elements, corresponding to the variables health and arts. All the elements of the first principal component are the same sign, making it a weighted average of all the variables.

To show the orthogonality of the principal components, note that premultiplying them by their transpose yields the identity matrix.

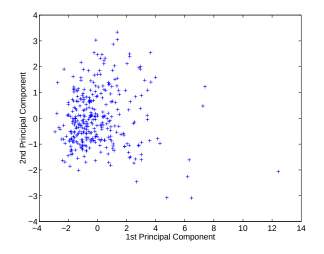
I = p3'*p3 I = 1.0000 -0.0000 -0.0000 -0.0000 1.0000 -0.0000 -0.0000 -0.0000 1.0000

The Component Scores (Second Output)

The second output, newdata, is the data in the new coordinate system defined by the principal components. This output is the same size as the input data matrix.

A plot of the first two columns of newdata shows the ratings data projected onto the first two principal components.

```
plot(newdata(:,1),newdata(:,2),'+')
xlabel('1st Principal Component');
ylabel('2nd Principal Component');
```



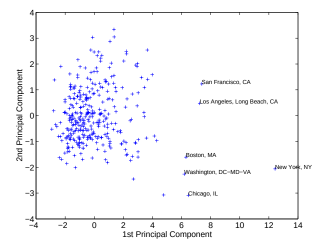
Note the outlying points in the lower right corner.

The function gname is useful for graphically identifying a few points in a plot like this. You can call gname with a string matrix containing as many case labels as points in the plot. The string matrix names works for labeling points with the city names.

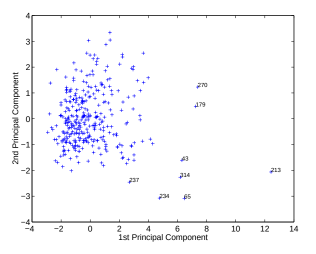
```
gname(names)
```

Move your cursor over the plot and click once near each point at the top right. As you click on each point, MATLAB labels it with the proper row from the names string matrix. When you are finished labeling points, press the **Return** key.

Here is the resulting plot.



The labeled cities are the biggest population centers in the United States. Perhaps we should consider them as a completely separate group. If we call gname without arguments, it labels each point with its row number.



We can create an index variable containing the row numbers of all the metropolitan areas we chose.

```
metro = [43 65 179 213 234 270 314];
names(metro,:)
ans =
Boston, MA
Chicago, IL
Los Angeles, Long Beach, CA
New York, NY
Philadelphia, PA-NJ
San Francisco, CA
Washington, DC-MD-VA
```

To remove these rows from the ratings matrix, type the following.

```
rsubset = ratings;
nsubset = names;
nsubset(metro,:) = [];
rsubset(metro,:) = [];
size(rsubset)
ans =
322 9
```

To practice, repeat the analysis using the variable rsubset as the new data matrix and nsubset as the string matrix of labels.

The Component Variances (Third Output)

The third output, variances, is a vector containing the variance explained by the corresponding column of newdata.

```
variances
variances =
    3.4083
    1.2140
    1.1415
    0.9209
    0.7533
    0.6306
    0.4930
```

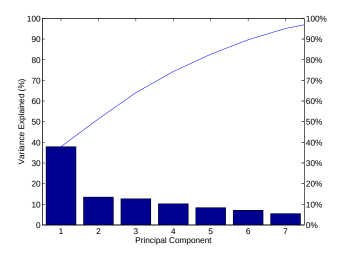
0.3180 0.1204

You can easily calculate the percent of the total variability explained by each principal component.

```
percent_explained = 100*variances/sum(variances)
percent_explained =
    37.8699
    13.4886
    12.6831
    10.2324
    8.3698
    7.0062
    5.4783
    3.5338
    1.3378
```

A "Scree" plot is a pareto plot of the percent variability explained by each principal component.

```
pareto(percent_explained)
xlabel('Principal Component')
ylabel('Variance Explained (%)')
```



We can see that the first three principal components explain roughly two thirds of the total variability in the standardized ratings.

Hotelling's T² (Fourth Output)

The last output of the princomp function, t2, is Hotelling's T^2 , a statistical measure of the multivariate distance of each observation from the center of the data set. This is an analytical way to find the most extreme points in the data.

```
[st2, index] = sort(t2); % Sort in ascending order.
st2 = flipud(st2); % Values in descending order.
index = flipud(index); % Indices in descending order.
extreme = index(1)
extreme = 213
names(extreme,:)
ans =
New York, NY
```

It is not surprising that the ratings for New York are the furthest from the average U.S. town.

Factor Analysis

Multivariate data often includes a large number of measured variables, and sometimes those variables "overlap" in the sense that groups of them may be dependent. For example, in a decathalon, each athlete competes in 10 events, but several of them can be thought of as "speed" events, while others can be thought of as "strength" events, etc. Thus, you can think of a competitor's 10 event scores as largely dependent on a smaller set of 3 or 4 types of athletic ability.

Factor Analysis is a way to fit a model to multivariate data to estimate just this sort of interdependence. In the Factor Analysis model, the measured variables depend on a smaller number of unobserved (latent) factors. Because each factor may affect several variables in common, they are known as "common factors". Each variable is assumed to be dependent on a linear combination of the common factors, and the coefficients are known as loadings. Each measured variable also includes a component due to independent random variability, known as "specific variance" because it is specific to one variable.

Specifically, Factor Analysis assumes that the covariance matrix of your data is of the form

$$\sigma_x = \Lambda \Lambda^T + \Psi$$

where Λ is the matrix of loadings, and the elements of the diagonal matrix Ψ are the specific variances. The function factoran fits the Factor Analysis model using maximum likelihood.

This section includes these topics:

- "Example: Finding Common Factors Affecting Stock Prices" on page 7-13
- "Factor Rotation" on page 7-16
- "Predicting Factor Scores" on page 7-17
- "Comparison of Factor Analysis and Principal Components Analysis" on page 7-19

Example: Finding Common Factors Affecting Stock Prices

Over the course of 100 weeks, the percent change in stock prices for ten companies has been recorded. Of the ten companies, the first four can be

classified as primarily technology, the next three as financial, and last three as retail. It seems reasonable that the stock prices for companies that are in the same sector might vary together as economic conditions change. Factor Analysis can provide quantitative evidence that companies within each sector do experience similar week-to-week changes in stock price.

First load the data, then call factoran and specify a model fit with three common factors. By default, factoran would compute rotated estimates of the loadings to try and make their interpretation simpler, but for the moment, specify an unrotated solution.

The first two factor an return arguments are the estimated loadings and the estimated specific variances. Each row of the loadings matrix represents one of the ten stocks, and each column corresponds to a common factor. With unrotated estimates, interpretation of the factors in this fit is difficult because most of the stocks contain fairly large coefficients for two or more factors.

```
Loadings
Loadings =
                  0.2367
       0.8885
                            -0.2354
       0.7126
                  0.3862
                             0.0034
       0.3351
                  0.2784
                            -0.0211
       0.3088
                  0.1113
                            -0.1905
       0.6277
                 -0.6643
                             0.1478
       0.4726
                 -0.6383
                             0.0133
       0.1133
                 -0.5416
                             0.0322
       0.6403
                  0.1669
                             0.4960
       0.2363
                  0.5293
                             0.5770
       0.1105
                  0.1680
                             0.5524
```

Note "Factor Rotation" on page 7-16 helps to simplify the structure in the loadings matrix, so that it will be easier to assign meaningful interpretations to the factors.

From the estimated specific variances, we can see that the model indicates that a particular stock price varies quite a lot beyond the variation due to the common factors.

A specific variance of 1 would indicate that there is *no* common factor component in that variable, while a specific variance of 0 would indicate that the variable is *entirely* determined by common factors. These data seem to fall somewhere in between.

The p-value returned in the stats structure fails to reject the null hypothesis of three common factors, suggesting that this model provides a satisfactory explanation of the covariation in these data.

To determine if fewer than three factors can provide an acceptable fit, you can try a model with two common factors. The p-value for this second fit is highly significant, and rejects the hypothesis of two factors, indicating that the simpler model is not sufficient to explain the pattern in these data.

Factor Rotation

As the results above illustrate, the estimated loadings from an unrotated Factor Analysis fit can have a complicated structure. The goal of factor rotation is to find a parameterization in which each variable has only a small number of large loadings, i.e., is affected by a small number of factors, preferably only one. This can often make it easier to interpret what the factors represent.

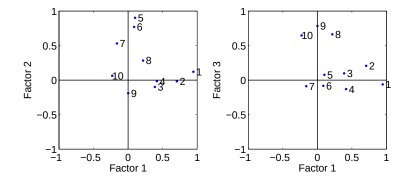
If you think of each row of the loadings matrix as coordinates of a point in M-dimensional space, then each factor corresponds to a coordinate axis. Factor rotation is equivalent to rotating those axes, and computing new loadings in the rotated coordinate system. There are various ways to do this. Some methods leave the axes orthogonal, while others are oblique methods that change the angles between them. For this example, rotate the estimated loadings by using the promax criterion, a common oblique method.

```
[LoadingsPM,specVarPM] = factoran(stocks,3,'rotate','promax');
LoadingsPM
LoadingsPM =
       0.9452
                  0.1214
                            -0.0617
       0.7064
                 -0.0178
                             0.2058
       0.3885
                 -0.0994
                             0.0975
       0.4162
                 -0.0148
                            -0.1298
       0.1021
                  0.9019
                             0.0768
       0.0873
                  0.7709
                            -0.0821
      -0.1616
                  0.5320
                            -0.0888
       0.2169
                  0.2844
                             0.6635
                 -0.1881
       0.0016
                             0.7849
      -0.2289
                  0.0636
                             0.6475
```

Promax rotation has created a simpler structure in the loadings, one in which most of the stocks have a large loading on only one factor. To see this more clearly, you can plot each stock using the factor loadings as coordinates. "Simple" structure in the loadings appears in this plot as stocks that fall along one of the factor axes. Because there are three factors, it's easier to see the loadings if you make a pair of two-dimensional plots.

```
subplot(1,2,1);
plot(LoadingsPM(:,1),LoadingsPM(:,2),'b.');
text(LoadingsPM(:,1),LoadingsPM(:,2), num2str((1:10)'));
line([-1 1 NaN 0 0 NaN 0 0],[0 0 NaN -1 1 NaN 0 0],...
'Color','black');
```

```
xlabel('Factor 1');
ylabel('Factor 2');
axis square;
subplot(1,2,2);
plot(LoadingsPM(:,1),LoadingsPM(:,3), 'b.');
text(LoadingsPM(:,1),LoadingsPM(:,3), num2str((1:10)'));
line([-1 1 NaN 0 0 NaN 0 0],[0 0 NaN -1 1 NaN 0 0],...
'Color', 'black');
xlabel('Factor 1');
ylabel('Factor 3');
axis square;
```



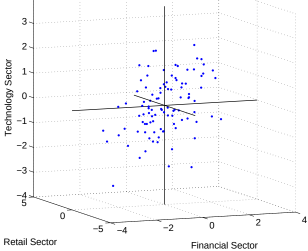
This plot shows that promax has succeeded in rotating the factor loadings to a simpler structure. Each stock depends primarily on only one factor, and it is possible to describe each factor in terms of the stocks that it affects. Based on which companies are near which axes, you could reasonably conclude that the first factor axis represents the financial sector, the second retail, and the third technology. The original conjecture, that stocks vary primarily within sector, is apparently supported by the data.

Predicting Factor Scores

Sometimes, it is useful to be able to classify an observation based on its factor scores. For example, if you accepted the three-factor model and the interpretation of the rotated factors, you might want to categorize each week in terms of how favorable it was for each of the three stock sectors, based on the data from the ten observed stocks.

Since the data in this example are the raw stock price changes, and not just their correlation matrix, you can have factoran return estimates of the value of each of the three rotated common factors for each week. You can then plot the estimated scores to see how the different stock sectors were affected during each week.

```
[LoadingsPM, specVarPM, TPM, stats, F] = factoran(stocks, 3,...
                                       'rotate','promax');
subplot(1,1,1);
plot3(F(:,1),F(:,2),F(:,3),'b.');
line([-4 4 NaN 0 0 NaN 0 0], [0 0 NaN -4 4 NaN 0 0],...
     [0 0 NaN 0 0 NaN -4 4], 'Color', 'black');
xlabel('Financial Sector');
ylabel('Retail Sector');
zlabel('Technology Sector');
grid on;
axis square;
view(-22.5, 8);
     4
     3
     2
     1
```



Oblique rotation often creates factors that are correlated. This plot shows some evidence of correlation between the first and third factors, and you can investigate further by computing the estimated factor correlation matrix.

Comparison of Factor Analysis and Principal Components Analysis

There is a good deal of overlap in terminology and goals between Principal Components Analysis (PCA) and Factor Analysis (FA). Much of the literature on the two methods does not distinguish between them, and some algorithms for fitting the FA model involve PCA. Both are dimension-reduction techniques, in the sense that they can be used to replace a large set of observed variables with a smaller set of new variables. However, the two methods are different in their goals and in their underlying models. Roughly speaking, you should use PCA when you simply need to summarize or approximate your data using fewer dimensions (to visualize it, for example), and you should use FA when you need an explanatory model for the correlations among your data.

Multivariate Analysis of Variance (MANOVA)

We reviewed the analysis of variance technique in "One-Way Analysis of Variance (ANOVA)" on page 4-4. With this technique we can take a set of grouped data and determine whether the mean of a variable differs significantly between groups. Often we have multiple variables, and we are interested in determining whether the entire set of means is different from one group to the next. There is a multivariate version of analysis of variance that can address that problem, as illustrated in the following example.

Example: Multivariate Analysis of Variance

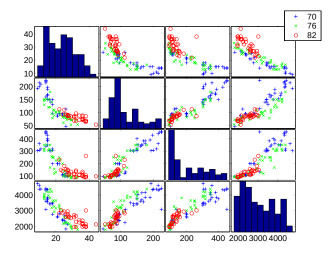
The carsmall data set has measurements on a variety of car models from the years 1970, 1976, and 1982. Suppose we are interested in whether the characteristics of the cars have changed over time.

First we load the data.

oad carsmall hos			
Name	Size	Bytes	Class
Acceleration	100x1	800	double array
Cylinders	100x1	800	double array
Displacement	100x1	800	double array
Horsepower	100x1	800	double array
MPG	100x1	800	double array
Model	100x36	7200	char array
Model_Year	100x1	800	double array
Origin	100x7	1400	char array
Weight	100x1	800	double array

Four of these variables (Acceleration, Displacement, Horsepower, and MPG) are continuous measurements on individual car models. The variable Model_Year indicates the year in which the car was made. We can create a grouped plot matrix of these variables using the gplotmatrix function.

```
x = [MPG Horsepower Displacement Weight];
gplotmatrix(x,[],Model_Year,[],'+xo')
```



(When the second argument of gplotmatrix is empty, the function graphs the columns of the x argument against each other, and places histograms along the diagonals. The empty fourth argument produces a graph with the default colors. The fifth argument controls the symbols used to distinguish between groups.)

It appears the cars do differ from year to year. The upper right plot, for example, is a graph of MPG versus Weight. The 1982 cars appear to have higher mileage than the older cars, and they appear to weigh less on average. But as a group, are the three years significantly different from one another? The manoval function can answer that question.

```
stats =
    W: [4x4 double]
    B: [4x4 double]
    T: [4x4 double]
    dfW: 90
    dfB: 2
    dfT: 92
    lambda: [2x1 double]
    chisqtf: [2x1 double]
    chisqdf: [2x1 double]
    eigenval: [4x1 double]
    eigenvec: [4x4 double]
    canon: [100x4 double]
    mdist: [100x1 double]
    gmdist: [3x3 double]
```

The manoval function produces three outputs:

- The first output, d, is an estimate of the dimension of the group means. If the means were all the same, the dimension would be 0, indicating that the means are at the same point. If the means differed but fell along a line, the dimension would be 1. In the example the dimension is 2, indicating that the group means fall in a plane but not along a line. This is the largest possible dimension for the means of three groups.
- The second output, p, is a vector of p-values for a sequence of tests. The first p-value tests whether the dimension is 0, the next whether the dimension is 1, and so on. In this case both p-values are small. That's why the estimated dimension is 2.
- The third output, stats, is a structure containing several fields, described in the following section.

The Fields of the stats Structure

The W, B, and T fields are matrix analogs to the within, between, and total sums of squares in ordinary one-way analysis of variance. The next three fields are the degrees of freedom for these matrices. Fields lambda, chisq, and chisqdf are the ingredients of the test for the dimensionality of the group means. (The p-values for these tests are the first output argument of manoval.)

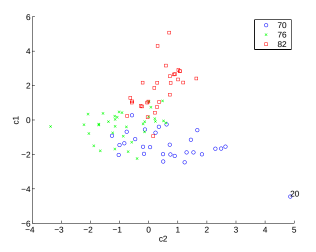
The next three fields are used to do a canonical analysis. Recall that in principal components analysis ("Principal Components Analysis" on page 7-3)

we look for the combination of the original variables that has the largest possible variation. In multivariate analysis of variance, we instead look for the linear combination of the original variables that has the largest separation between groups. It is the single variable that would give the most significant result in a univariate one-way analysis of variance. Having found that combination, we next look for the combination with the second highest separation, and so on.

The eigenvec field is a matrix that defines the coefficients of the linear combinations of the original variables. The eigenval field is a vector measuring the ratio of the between-group variance to the within-group variance for the corresponding linear combination. The canon field is a matrix of the canonical variable values. Each column is a linear combination of the mean-centered original variables, using coefficients from the eigenvec matrix.

A grouped scatter plot of the first two canonical variables shows more separation between groups then a grouped scatter plot of any pair of original variables. In this example it shows three clouds of points, overlapping but with distinct centers. One point in the bottom right sits apart from the others. By using the gname function, we can see that this is the 20th point.

```
c1 = stats.canon(:,1);
c2 = stats.canon(:,2);
gscatter(c2,c1,Model_Year,[],'oxs')
gname
```



Roughly speaking, the first canonical variable, c1, separates the 1982 cars (which have high values of c1) from the older cars. The second canonical variable, c2, reveals some separation between the 1970 and 1976 cars.

The final two fields of the stats structure are Mahalanobis distances. The mdist field measures the distance from each point to its group mean. Points with large values may be outliers. In this data set, the largest outlier is the one we saw in the scatter plot, the Buick Estate station wagon. (Note that we could have supplied the model name to the gname function above if we wanted to label the point with its model name rather than its row number.)

The gmdist field measures the distances between each pair of group means. The following commands examine the group means and their distances:

```
grpstats(x, Model Year)
ans =
  1.0e+003 *
    0.0177
               0.1489
                          0.2869
                                     3.4413
    0.0216
               0.1011
                          0.1978
                                     3.0787
    0.0317
               0.0815
                          0.1289
                                     2.4535
stats.gmdist
ans =
         0
               3.8277
                         11.1106
    3.8277
                    0
                          6.1374
   11.1106
               6.1374
                                0
```

As might be expected, the multivariate distance between the extreme years 1970 and 1982 (11.1) is larger than the difference between more closely spaced years (3.8 and 6.1). This is consistent with the scatter plots, where the points seem to follow a progression as the year changes from 1970 through 1976 to 1982. If we had more groups, we might have found it instructive to use the manovacluster function to draw a diagram that presents clusters of the groups, formed using the distances between their means.

Cluster Analysis

Cluster analysis, also called segmentation analysis or taxonomy analysis, is a way to way to create groups of objects, or *clusters*, in such a way that the profiles of objects in the same cluster are very similar and the profiles of objects in different clusters are quite distinct.

Cluster analysis can be performed on many different types of data sets. For example, a data set might contain a number of observations of subjects in a study where each observation contains a set of variables.

Many different fields of study, such as engineering, zoology, medicine, linguistics, anthropology, psychology, and marketing, have contributed to the development of clustering techniques and the application of such techniques. For example, cluster analysis can help in creating "balanced" treatment and control groups for a designed study. If you find that each cluster contains roughly equal numbers of treatment and control subjects, then statistical differences found between the groups can be attributed to the experiment and not to any initial difference between the groups.

This sections explores two kinds of clustering:

- "Hierarchical Clustering" on page 7-26
- "K-Means Clustering" on page 7-40

Hierarchical Clustering

Hierarchical clustering is a way to investigate grouping in your data, simultaneously over a variety of scales, by creating a cluster tree. The tree is not a single set of clusters, but rather a multi-level hierarchy, where clusters at one level are joined as clusters at the next higher level. This allows you to decide what level or scale of clustering is most appropriate in your application.

The following sections explore the hierarchical clustering features in the Statistics Toolbox:

- "Terminology and Basic Procedure" on page 7-27
- "Finding the Similarities Between Objects" on page 7-27
- "Defining the Links Between Objects" on page 7-30
- "Evaluating Cluster Formation" on page 7-32
- "Creating Clusters" on page 7-37

Terminology and Basic Procedure

To perform hierarchical cluster analysis on a data set using the Statistics Toolbox functions, follow this procedure:

- **1** Find the similarity or dissimilarity between every pair of objects in the data set. In this step, you calculate the *distance* between objects using the pdist function. The pdist function supports many different ways to compute this measurement. See "Finding the Similarities Between Objects" on page 7-27 for more information.
- 2 Group the objects into a binary, hierarchical cluster tree. In this step, you link together pairs of objects that are in close proximity using the linkage function. The linkage function uses the distance information generated in step 1 to determine the proximity of objects to each other. As objects are paired into binary clusters, the newly formed clusters are grouped into larger clusters until a hierarchical tree is formed. See "Defining the Links Between Objects" on page 7-30 for more information.
- **3** Determine where to divide the hierarchical tree into clusters. In this step, you divide the objects in the hierarchical tree into clusters using the cluster function. The cluster function can create clusters by detecting natural groupings in the hierarchical tree or by cutting off the hierarchical tree at an arbitrary point. See "Creating Clusters" on page 7-37 for more information.

The following sections provide more information about each of these steps.

Note The Statistics Toolbox includes a convenience function, clusterdata, which performs all these steps for you. You do not need to execute the pdist, linkage, or cluster functions separately. However, the clusterdata function does not give you access to the options each of the individual routines offers. For example, if you use the pdist function you can choose the distance calculation method, whereas if you use the clusterdata function you cannot.

Finding the Similarities Between Objects

You use the pdist function to calculate the distance between every pair of objects in a data set. For a data set made up of m objects, there are

 $m \cdot (m-1)/2$ pairs in the data set. The result of this computation is commonly known as a distance or dissimilarity matrix.

There are many ways to calculate this distance information. By default, the pdist function calculates the Euclidean distance between objects; however, you can specify one of several other options. See pdist for more information.

Note You can optionally normalize the values in the data set before calculating the distance information. In a real world data set, variables can be measured against different scales. For example, one variable can measure Intelligence Quotient (IQ) test scores and another variable can measure head circumference. These discrepancies can distort the proximity calculations. Using the zscore function, you can convert all the values in the data set to use the same proportional scale. See zscore for more information.

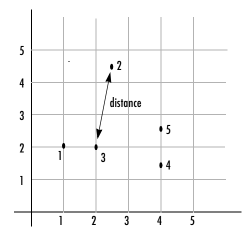
For example, consider a data set, X, made up of five objects where each object is a set of *x*,*y* coordinates.

- Object 1: 1, 2
- Object 2: 2.5, 4.5
- **Object 3**: 2, 2
- Object 4: 4, 1.5
- Object 5: 4, 2.5

You can define this data set as a matrix

 $X = [1 \ 2; 2.5 \ 4.5; 2 \ 2; 4 \ 1.5; 4 \ 2.5]$

and pass it to pdist. The pdist function calculates the distance between object 1 and object 2, object 1 and object 3, and so on until the distances between all the pairs have been calculated. The following figure plots these objects in a graph. The distance between object 2 and object 3 is shown to illustrate one interpretation of distance.



Returning Distance Information. The pdist function returns this distance information in a vector, Y, where each element contains the distance between a pair of objects.

```
Y = pdist(X)
Y =
Columns 1 through 7
2.9155 1.0000 3.0414 3.0414 2.5495 3.3541 2.5000
Columns 8 through 10
2.0616 2.0616 1.0000
```

To make it easier to see the relationship between the distance information generated by pdist and the objects in the original data set, you can reformat the distance vector into a matrix using the squareform function. In this matrix, element $i_{,j}$ corresponds to the distance between object i and object j in the original data set. In the following example, element 1,1 represents the distance between object 1 and itself (which is zero). Element 1,2 represents the distance between object 1 and object 2, and so on.

```
squareform(Y)
ans =
         0
               2.9155
                          1.0000
                                     3.0414
                                                3.0414
    2.9155
                    0
                          2.5495
                                     3.3541
                                                2,5000
    1.0000
               2.5495
                                     2.0616
                                                2.0616
                               0
```

3.0414	3.3541	2.0616	0	1.0000
3.0414	2.5000	2.0616	1.0000	0

Defining the Links Between Objects

Once the proximity between objects in the data set has been computed, you can determine which objects in the data set should be grouped together into clusters, using the linkage function. The linkage function takes the distance information generated by pdist and links pairs of objects that are close together into binary clusters (clusters made up of two objects). The linkage function then links these newly formed clusters to other objects to create bigger clusters until all the objects in the original data set are linked together in a hierarchical tree.

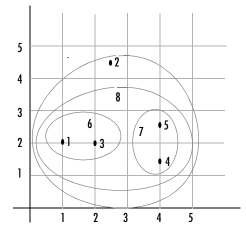
For example, given the distance vector Y generated by pdist from the sample data set of x and y coordinates, the linkage function generates a hierarchical cluster tree, returning the linkage information in a matrix, Z.

Ζ	=	linkage(Y)		
Ζ	=			
		1.0000	3.0000	1.0000
		4.0000	5.0000	1.0000
		6.0000	7.0000	2.0616
		8.0000	2.0000	2.5000

In this output, each row identifies a link. The first two columns identify the objects that have been linked, that is, object 1, object 2, and so on. The third column contains the distance between these objects. For the sample data set of x and y coordinates, the linkage function begins by grouping together objects 1 and 3, which have the closest proximity (distance value = 1.0000). The linkage function continues by grouping objects 4 and 5, which also have a distance value of 1.0000.

The third row indicates that the linkage function grouped together objects 6 and 7. If our original sample data set contained only five objects, what are objects 6 and 7? Object 6 is the newly formed binary cluster created by the grouping of objects 1 and 3. When the linkage function groups two objects together into a new cluster, it must assign the cluster a unique index value, starting with the value m+1, where m is the number of objects in the original data set. (Values 1 through m are already used by the original data set.) Object 7 is the index for the cluster formed by objects 4 and 5.

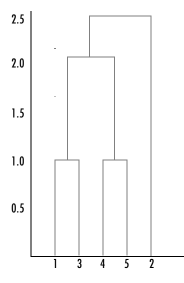
As the final cluster, the linkage function grouped object 8, the newly formed cluster made up of objects 6 and 7, with object 2 from the original data set. The following figure graphically illustrates the way linkage groups the objects into a hierarchy of clusters.



Plotting the Cluster Tree

The hierarchical, binary cluster tree created by the linkage function is most easily understood when viewed graphically. The Statistics Toolbox includes the dendrogram function that plots this hierarchical tree information as a graph, as in the following example.

```
dendrogram(Z)
```



In the figure, the numbers along the horizontal axis represent the indices of the objects in the original data set. The links between objects are represented as upside down U-shaped lines. The height of the U indicates the distance between the objects. For example, the link representing the cluster containing objects 1 and 3 has a height of 1. For more information about creating a dendrogram diagram, see the dendrogram function reference page.

Evaluating Cluster Formation

After linking the objects in a data set into a hierarchical cluster tree, you may want to verify that the tree represents significant similarity groupings. In addition, you may want more information about the links between the objects. The Statistics Toolbox provides functions to perform both these tasks, as described in the following sections:

- "Verifying the Cluster Tree" on page 7-32
- "Getting More Information About Cluster Links" on page 7-33

Verifying the Cluster Tree. One way to measure the validity of the cluster information generated by the linkage function is to compare it with the original proximity data generated by the pdist function. If the clustering is valid, the linking of objects in the cluster tree should have a strong correlation with the distances between objects in the distance vector. The cophenet

function compares these two sets of values and computes their correlation, returning a value called the *cophenetic correlation coefficient*. The closer the value of the cophenetic correlation coefficient is to 1, the better the clustering solution.

You can use the cophenetic correlation coefficient to compare the results of clustering the same data set using different distance calculation methods or clustering algorithms.

For example, you can use the cophenet function to evaluate the clusters created for the sample data set

where Z is the matrix output by the linkage function and Y is the distance vector output by the pdist function.

Execute pdist again on the same data set, this time specifying the City Block metric. After running the linkage function on this new pdist output, use the cophenet function to evaluate the clustering using a different distance metric.

The cophenetic correlation coefficient shows a stronger correlation when the City Block metric is used.

Getting More Information About Cluster Links. One way to determine the natural cluster divisions in a data set is to compare the height of each link in a cluster tree with the heights of neighboring links below it in the tree.

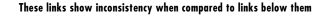
If a link is approximately the same height as neighboring links, it indicates that there are similarities between the objects joined at this level of the hierarchy. These links are said to exhibit a high level of consistency.

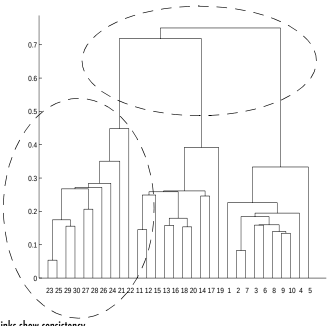
If the height of a link differs from neighboring links, it indicates that there are dissimilarities between the objects at this level in the cluster tree. This link is said to be inconsistent with the links around it. In cluster analysis, inconsistent links can indicate the border of a natural division in a data set. The cluster function uses a measure of inconsistency to determine where to divide a data set into clusters. (See "Creating Clusters" on page 7-37 for more information.)

The next section provides an example.

Example: Inconsistent Links. To illustrate, the following example creates a data set of random numbers with three deliberate natural groupings. In the dendrogram, note how the objects tend to collect into three groups. These three groups are then connected by three longer links. These longer links are inconsistent when compared with the links below them in the hierarchy.

```
rand('seed',3)
X = [rand(10,2)+1;rand(10,2)+2;rand(10,2)+3];
Y = pdist(X);
Z = linkage(Y);
dendrogram(Z);
```





These links show consistency

The relative consistency of each link in a hierarchical cluster tree can be quantified and expressed as the *inconsistency coefficient*. This value compares the height of a link in a cluster hierarchy with the average height of neighboring links. If the object is consistent with those around it, it will have a low inconsistency coefficient. If the object is inconsistent with those around it, it will have a higher inconsistency coefficient.

To generate a listing of the inconsistency coefficient for each link the cluster tree, use the inconsistent function. The inconsistent function compares each link in the cluster hierarchy with adjacent links two levels below it in the cluster hierarchy. This is called the *depth* of the comparison. Using the inconsistent function, you can specify other depths. The objects at the bottom of the cluster tree, called leaf nodes, that have no further objects below them, have an inconsistency coefficient of zero.

For example, returning to the sample data set of x and y coordinates, we can use the inconsistent function to calculate the inconsistency values for the links created by the linkage function, described in "Defining the Links Between Objects" on page 7-30.

Ι	=	inconsistent(Z)											
Ι	=												
		1.0000	0	1.0000	0								
		1.0000	0	1.0000	0								
		1.3539	0.8668	3.0000	0.8165								
		2.2808	0.3100	2.0000	0.7071								

The inconsistent function returns data about the links in an (m-1)-by-4 matrix where each column provides data about the links.

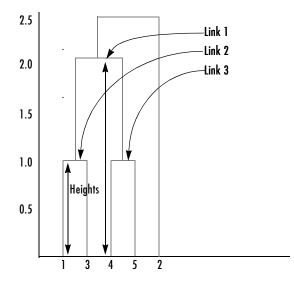
Column	Description
1	Mean of the heights of all the links included in the calculation
2	Standard deviation of all the links included in the calculation
3	Number of links included in the calculation
4	Inconsistency coefficient

In the sample output, the first row represents the link between objects 1 and 3. (This cluster is assigned the index 6 by the linkage function.) Because this a

leaf node, the inconsistency coefficient is zero. The second row represents the link between objects 4 and 5, also a leaf node. (This cluster is assigned the index 7 by the linkage function.)

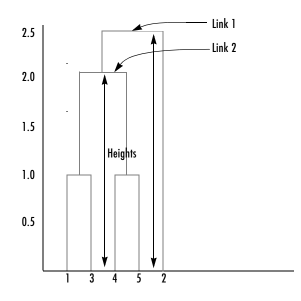
The third row evaluates the link that connects these two leaf nodes, objects 6 and 7. (This cluster is called object 8 in the linkage output). Column three indicates that three links are considered in the calculation: the link itself and the two links directly below it in the hierarchy. Column one represents the mean of the heights of these links. The inconsistent function uses the height information output by the linkage function to calculate the mean. Column two represents the standard deviation between the links. The last column contains the inconsistency value for these links, 0.8165.

The following figure illustrates the links and heights included in this calculation.



Row four in the output matrix describes the link between object 8 and object 2. Column three indicates that two links are included in this calculation: the link itself and the link directly below it in the hierarchy. The inconsistency coefficient for this link is 0.7071.

The following figure illustrates the links and heights included in this calculation.



Creating Clusters

After you create the hierarchical tree of binary clusters, you can divide the hierarchy into larger clusters using the cluster function. The cluster function lets you create clusters in two ways, as discussed in the following sections:

- "Finding the Natural Divisions in the Data Set" on page 7-37
- "Specifying Arbitrary Clusters" on page 7-38

Finding the Natural Divisions in the Data Set. In the hierarchical cluster tree, the data set may naturally align itself into clusters. This can be particularly evident in a dendrogram diagram where groups of objects are densely packed in certain areas and not in others. The inconsistency coefficient of the links in the cluster tree can identify these points where the similarities between objects change. (See "Evaluating Cluster Formation" on page 7-32 for more information about the inconsistency coefficient.) You can use this value to determine where the cluster function draws cluster boundaries.

For example, if you use the cluster function to group the sample data set into clusters, specifying an inconsistency coefficient threshold of 0.9 as the value of the cutoff argument, the cluster function groups all the objects in the sample

data set into one cluster. In this case, none of the links in the cluster hierarchy had an inconsistency coefficient greater than 0.9.

```
T = cluster(Z,0.9)
T =
1
1
1
1
1
```

The cluster function outputs a vector, T, that is the same size as the original data set. Each element in this vector contains the number of the cluster into which the corresponding object from the original data set was placed.

If you lower the inconsistency coefficient threshold to 0.8, the cluster function divides the sample data set into three separate clusters.

```
T = cluster(Z,0.8)
T =
1
3
1
2
2
```

This output indicates that objects 1 and 3 were placed in cluster 1, objects 4 and 5 were placed in cluster 2, and object 2 was placed in cluster 3.

Specifying Arbitrary Clusters. Instead of letting the cluster function create clusters determined by the natural divisions in the data set, you can specify the number of clusters you want created. In this case, the value of the cutoff argument specifies the point in the cluster hierarchy at which to create the clusters.

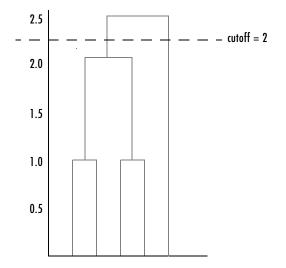
For example, you can specify that you want the cluster function to divide the sample data set into two clusters. In this case, the cluster function creates one cluster containing objects 1, 3, 4, and 5 and another cluster containing object 2.

```
T = cluster(Z,2)
T =
1
2
```

1

1

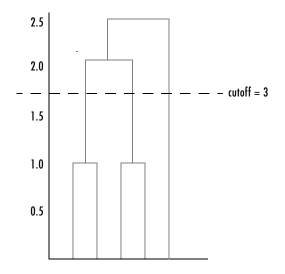
To help you visualize how the cluster function determines how to create these clusters, the following figure shows the dendrogram of the hierarchical cluster tree. When you specify a value of 2, the cluster function draws an imaginary horizontal line across the dendrogram that bisects two vertical lines. All the objects below the line belong to one of these two clusters.



If you specify a cutoff value of 3, the cluster function cuts off the hierarchy at a lower point, bisecting three lines.

```
T = cluster(Z,3)
T =
1
3
1
2
2
```

This time, objects 1 and 3 are grouped in a cluster, objects 4 and 5 are grouped in a cluster, and object 2 is placed into a cluster, as seen in the following figure.



K-Means Clustering

This section gives a description and an example of using the MATLAB function for K-means clustering, kmeans.

- "Overview of K-Means Clustering" on page 7-40
- "Example: Clustering Data in Four Dimensions" on page 7-41

Overview of K-Means Clustering

K-means clustering can best be described as a partitioning method. That is, the function kmeans partitions the observations in your data into K mutually exclusive clusters, and returns a vector of indices indicating to which of the k clusters it has assigned each observation. Unlike the hierarchical clustering methods used in linkage (see "Hierarchical Clustering" on page 7-26), kmeans does not create a tree structure to describe the groupings in your data, but rather creates a single level of clusters. Another difference is that K-means clustering uses the actual observations of objects or individuals in your data, and not just their proximities. These differences often mean that kmeans is more suitable for clustering large amounts of data.

kmeans treats each observation in your data as an object having a location in space. It finds a partition in which objects within each cluster are as close to

each other as possible, and as far from objects in other clusters as possible. You can choose from five different distance measures, depending on the kind of data you are clustering.

Each cluster in the partition is defined by its member objects and by its centroid, or center. The centroid for each cluster is the point to which the sum of distances from all objects in that cluster is minimized. kmeans computes cluster centroids differently for each distance measure, to minimize the sum with respect to the measure that you specify.

kmeans uses an iterative algorithm that minimizes the sum of distances from each object to its cluster centroid, over all clusters. This algorithm moves objects between clusters until the sum cannot be decreased further. The result is a set of clusters that are as compact and well-separated as possible. You can control the details of the minimization using several optional input parameters to kmeans, including ones for the initial values of the cluster centroids, and for the maximum number of iterations.

Example: Clustering Data in Four Dimensions

This example explores possible clustering in four-dimensional data by analyzing the results of partitioning the points into three, four, and five clusters.

Note Because each part of this example generates random numbers sequentially, i.e., without setting a new seed, you must perform all steps in sequence to duplicate the results shown. If you perform the steps out of sequence, the answers will be essentially the same, but the intermediate results, number of iterations, or ordering of the silhouette plots may differ. See "Random Numbers in Examples" on page 1-5 to set the correct seed.

Creating Clusters and Determining Separation. First, load some data.

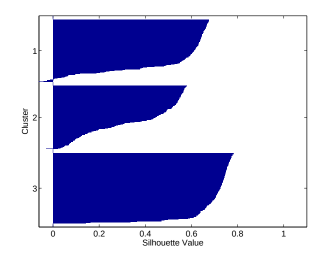
```
load kmeansdata;
size(X)
ans =
560 4
```

Even though these data are four-dimensional, and cannot be easily visualized, kmeans enables you to investigate if a group structure exists in them. Call kmeans with k, the desired number of clusters, equal to 3. For this example, specify the city block distance measure, and use the default starting method of initializing centroids from randomly selected data points.

```
idx3 = kmeans(X,3,'distance','city');
```

To get an idea of how well-separated the resulting clusters are, you can make a silhouette plot using the cluster indices output from kmeans. The silhouette plot displays a measure of how close each point in one cluster is to points in the neighboring clusters. This measure ranges from +1, indicating points that are very distant from neighboring clusters, through 0, indicating points that are not distinctly in one cluster or another, to -1, indicating points that are probably assigned to the wrong cluster. silhouette returns these values in its first output.

```
[silh3,h] = silhouette(X,idx3,'city');
xlabel('Silhouette Value')
ylabel('Cluster')
```



From the silhouette plot, you can see that most points in the third cluster have a large silhouette value, greater than 0.6, indicating that that cluster is somewhat separated from neighboring clusters. However, the second cluster contains many points with low silhouette values, and the first contains a few points with negative values, indicating that those two clusters are not well separated.

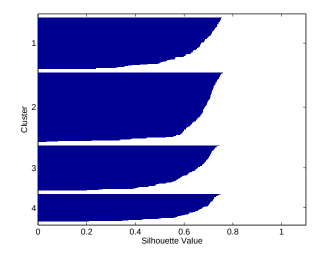
Determining the Correct Number of Clusters. Increase the number of clusters to see if kmeans can find a better grouping of the data. This time, use the optional 'display' parameter to print information about each iteration.

```
idx4 = kmeans(X,4, 'dist','city', 'display','iter');
  iter phase
                  num
                               sum
     1
            1
                  560
                           2897.56
     2
            1
                   53
                           2736.67
     3
            1
                   50
                           2476.78
     4
                           1779.68
            1
                  102
     5
            1
                    5
                            1771.1
     6
            2
                    0
                            1771.1
6 iterations, total sum of distances = 1771.1
```

Notice that the total sum of distances decreases at each iteration as kmeans reassigns points between clusters and recomputes cluster centroids. In this case, the second phase of the algorithm did not make any reassignments, indicating that the first phase reached a minimum after five iterations. In some problems, the first phase may not reach a minimum, but the second phase always will.

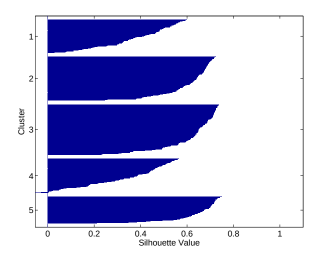
A silhouette plot for this solution indicates that these four clusters are better separated than the three in the previous solution.

```
[silh4,h] = silhouette(X,idx4,'city');
xlabel('Silhouette Value')
ylabel('Cluster')
```



A more quantitative way to compare the two solutions is to look at the average silhouette values for the two cases.

Finally, try clustering the data using five clusters.



This silhouette plot indicates that this is probably not the right number of clusters, since two of the clusters contain points with mostly low slhouette values. Without some knowledge of how many clusters are really in the data, it is a good idea to experiment with a range of values for k.

Avoiding Local Minima. Like many other types of numerical minimizations, the solution that kmeans reaches often depends on the starting points. It is possible for kmeans to reach a local minimum, where reassigning any one point to a new cluster would increase the total sum of point-to-centroid distances, but where a better solution does exist. However, you can use the optional 'replicates' parameter to overcome that problem.

For four clusters, specify five replicates, and use the 'display' parameter to print out the final sum of distances for each of the solutions.

The output shows that, even for this relatively simple problem, non-global minima do exist. Each of these five replicates began from a different randomly

selected set of initial centroids, and kmeans found two different local minima. However, the final solution that kmeans returns is the one with the lowest total sum of distances, over all replicates.

```
sum(sumdist)
ans =
1771.1
```

Classical Multidimensional Scaling

These two sections demonstrate how to use the function cmdscale to perform classical multidimensional scaling.

- "Overview" on page 7-47
- "Reconstructing a Map from Inter-City Distances" on page 7-49

Overview

One of the most important goals in visualizing data is to get a sense of how near or far points are from each other. Often, you can do this with a scatter plot. However, for some analyses, the data that you have may not be in the form of "points" at all, but rather in the form of pairwise similarities or dissimilarities between cases, observations, or subjects. Without any points, you cannot make a scatter plot.

Even if your data are in the form of points rather than pairwise distances, a scatter plot of those data may not be useful. For some kinds of data, the relevant way to measure how "near" two points are may not be their Euclidean distance. While scatter plots of the raw data make it easy to compare Euclidean distances, they are not always useful when comparing other kinds of interpoint distances, city block distance for example, or even more general dissimilarities. Also, with a large number of variables, it is very difficult to visualize distances unless the data can be represented in a small number of dimensions. Some sort of dimension reduction is usually necessary.

Multidimensional Scaling (MDS) is a set of methods that address all of these problems. MDS allows you to visualize how "near" points are to each other for many kinds of distance or dissimilarity measures, and can produce a representation of your data in a small number of dimensions. MDS does not require raw data, but only a matrix of pairwise distances or dissimilarities.

The function cmdscale performs classical (metric) multidimensional scaling, also known as Principal Coordinates Analysis. cmdscale takes as an input a matrix of interpoint distances, and creates a configuration of points. Ideally, those points are in two or three dimensions, and the Euclidean distances between them reproduce the original distance matrix. Thus, a scatter plot of the points created by cmdscale will provide a visual representation of the original distances.

A Simple Example

As a very simple example, you can reconstruct a set of points from only their interpoint distances. First, create some four dimensional points with a small component in their fourth coordinate, and reduce them to distances.

```
X = [ normrnd(0,1,10,3), normrnd(0,.1,10,1) ];
D = pdist(X,'euclidean');
```

Next, use cmdscale to find a configuration with those interpoint distances. cmdscale accepts distances as either a square matrix, or, as in this example, in the vector upper-triangular form produced by pdist.

[Y,eigvals] = cmdscale(D);

cmdscale produces two outputs. The first output, Y, is a matrix containing the reconstructed points. The second output, eigvals, is a vector containing the sorted eigenvalues of what is often referred to as the "scalar product matrix", which, in the simplest case, is equal to Y*Y'. The relative magnitudes of those eigenvalues indicate the relative contribution of the corresponding columns of Y in reproducing the original distance matrix D with the reconstructed points.

```
format short g
[eigvals eigvals/max(abs(eigvals))]
ans =
       12.623
                          1
       4.3699
                   0.34618
       1.9307
                   0.15295
     0.025884
                 0.0020505
  1.7192e-015
              1.3619e-016
  6.8727e-016
              5.4445e-017
  4.4367e-017 3.5147e-018
 -9.2731e-016 -7.3461e-017
  -1.327e-015 -1.0513e-016
 -1.9232e-015 -1.5236e-016
```

If eigvals contains only positive and zero (within roundoff error) eigenvalues, then the columns of Y corresponding to the positive eigenvalues provide an exact reconstruction of D, in the sense that their interpoint Euclidean distances, computed using pdist for example, are identical (within roundoff) to the values in D.

maxerr4 = max(abs(D - pdist(Y))) % exact reconstruction

maxerr4 = 2.6645e-015

If two or three of the eigenvalues in eigvals are much larger than the rest, then the distance matrix based on the corresponding columns of Y nearly reproduces the original distance matrix D. In this sense, those columns form a lower-dimensional representation that adequately describes the data. However it is not always possible to find a good low-dimensional reconstruction.

The reconstruction in three dimensions reproduces D very well, but the reconstruction in two dimensionss has errors that are the same order of magnitude as the largest values in D.

Often, eigvals contains some negative eigenvalues, indicating that the distances in D cannot be reproduced exactly. That is, there may not be any configuration of points whose interpoint Euclidean distances are given by D. If the largest negative eigenvalue is small in magnitude relative to the largest positive eigenvalues, then the configuration returned by cmdscale may still reproduce D well. The following example demonstrates this.

Reconstructing a Map from Inter-City Distances

Given only the distances between 10 US cities, cmdscale can construct a map of those cities. First, create the distance matrix and pass it to cmdscale. In this example, D is a full distance matrix: it is square, symmetric, has positive entries, and zeros on the diagonal.

```
cities =
{'Atl','Chi','Den','Hou','LA','Mia','NYC','SF','Sea','WDC'};
D = [
            587 1212
                       701 1936 604
                                      748 2139 2182
         0
                                                       543;
       587
                 920
                       940 1745 1188
                                      713 1858 1737
                                                       597;
              0
      1212
            920
                    0
                       879
                            831 1726 1631
                                            949 1021
                                                      1494;
       701
            940
                 879
                         0 1374
                                 968 1420 1645 1891
                                                      1220;
      1936 1745
                 831 1374
                              0 2339 2451
                                            347
                                                 959
                                                      2300;
       604 1188 1726 968 2339
                                   0 1092 2594 2734
                                                       923;
       748
            713 1631 1420 2451 1092
                                         0 2571
                                                2408
                                                       205;
      2139 1858 949 1645
                            347 2594 2571
                                              0
                                                 678
                                                      2442;
      2182 1737 1021 1891
                            959 2734 2408
                                            678
                                                   0
                                                      2329;
       543
            597 1494 1220 2300 923
                                      205 2442 2329
                                                         0];
[Y,eigvals] = cmdscale(D);
```

Next, look at the eigenvalues returned by cmdscale. Some of these are negative, indicating that the original distances are not Euclidean. This is because of the curvature of the earth.

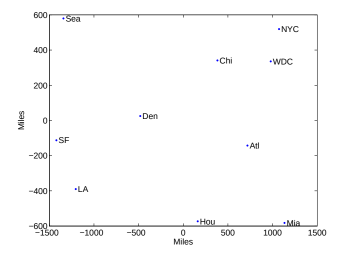
```
format short g
[eigvals eigvals/max(abs(eigvals))]
ans =
  9.5821e+006
                          1
  1.6868e+006
                   0.17604
       8157.3
                 0.0008513
       1432.9
                0.00014954
       508.67 5.3085e-005
       25.143
                2.624e-006
  5.3394e-010 5.5722e-017
       -897.7 -9.3685e-005
      -5467.6
                 -0.0005706
       -35479
                 -0.0037026
```

However, in this case, the two largest positive eigenvalues are much larger in magnitude than the remaining eigenvalues. So, despite the negative eigenvalues, the first two coordinates of Y are sufficient for a reasonable reproduction of D.

```
Dtriu = D(find(tril(ones(10),-1)))';
maxrelerr = max(abs(Dtriu - pdist(Y(:,1:2)))) ./ max(Dtriu)
maxrelerr =
    0.0075371
```

Here is a plot of the reconstructed city locations as a map. The orientation of the reconstruction is arbitrary: in this case, it happens to be close to, although not exactly, the correct orientation.

```
plot(Y(:,1),Y(:,2),'.');
text(Y(:,1)+25,Y(:,2),cities);
xlabel('Miles'); ylabel('Miles');
```





8

Statistical Plots

Introduction	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	8-2
Box Plots	•	•	•	•		•	•	•		•	•	•	•			•	•	8-3
Distribution Plots																		8-4
Normal Probability Pl	ots	;																8-4
Quantile-Quantile Plo	\mathbf{ts}																	8-6
Weibull Probability Pl																		
Empirical Cumulative																		
Scatter Plots																		8-10

Introduction

The Statistics Toolbox adds specialized plots to the extensive graphics capabilities of MATLAB.

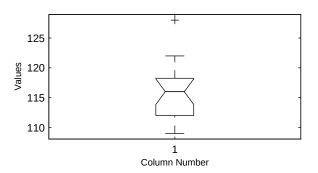
- *Box plots* are graphs for describing data samples. They are also useful for graphic comparisons of the means of many samples (see "One-Way Analysis of Variance (ANOVA)" on page 4-4).
- *Distribution plots* are graphs for visualizing the distribution of one or more samples. They include normal and Weibull probability plots, quantile-quantile plots, and empirical cumulative distribution plots.
- *Scatter plots* are graphs for visualizing the relationship between a pair of variables or several such pairs. Grouped versions of these plots use different plotting symbols to indicate group membership. The gname function can label points on these plots with a text label or an observation number.

The plot types are described further in the following sections:

- "Box Plots" on page 8-3
- "Distribution Plots" on page 8-4
- "Scatter Plots" on page 8-10

Box Plots

The graph shows an example of a notched box plot.



This plot has several graphic elements:

- The lower and upper lines of the "box" are the 25th and 75th percentiles of the sample. The distance between the top and bottom of the box is the interquartile range.
- The line in the middle of the box is the sample median. If the median is not centered in the box, that is an indication of skewness.
- The "whiskers" are lines extending above and below the box. They show the extent of the rest of the sample (unless there are outliers). Assuming no outliers, the maximum of the sample is the top of the upper whisker. The minimum of the sample is the bottom of the lower whisker. By default, an outlier is a value that is more than 1.5 times the interquartile range away from the top or bottom of the box.
- The plus sign at the top of the plot is an indication of an outlier in the data. This point may be the result of a data entry error, a poor measurement or a change in the system that generated the data.
- The notches in the box are a graphic confidence interval about the median of a sample. Box plots do not have notches by default.

A side-by-side comparison of two notched box plots is the graphical equivalent of a t-test. See "Hypothesis Tests" on page 6-1.

Distribution Plots

There are several types of plots for examining the distribution of one or more samples, as described in the following sections:

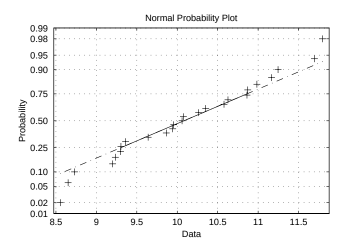
- "Normal Probability Plots" on page 8-4
- "Quantile-Quantile Plots" on page 8-6
- "Weibull Probability Plots" on page 8-7
- "Empirical Cumulative Distribution Function (CDF)" on page 8-8

Normal Probability Plots

A normal probability plot is a useful graph for assessing whether data comes from a normal distribution. Many statistical procedures make the assumption that the underlying distribution of the data is normal, so this plot can provide some assurance that the assumption of normality is not being violated, or provide an early warning of a problem with your assumptions.

This example shows a typical normal probability plot.

```
x = normrnd(10,1,25,1);
normplot(x)
```

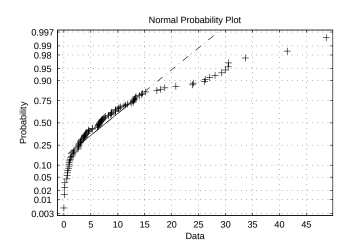


The plot has three graphical elements. The plus signs show the empirical probability versus the data value for each point in the sample. The solid line connects the 25th and 75th percentiles of the data and represents a robust linear fit (i.e., insensitive to the extremes of the sample). The dashed line extends the solid line to the ends of the sample.

The scale of the *y*-axis is not uniform. The *y*-axis values are probabilities and, as such, go from zero to one. The distance between the tick marks on the *y*-axis matches the distance between the quantiles of a normal distribution. The quantiles are close together near the median (probability = 0.5) and stretch out symmetrically moving away from the median. Compare the vertical distance from the bottom of the plot to the probability 0.25 with the distance from 0.25 to 0.50. Similarly, compare the distance from the top of the plot to the probability 0.75 with the distance from 0.75 to 0.50.

If all the data points fall near the line, the assumption of normality is reasonable. But, if the data is nonnormal, the plus signs may follow a curve, as in the example using exponential data below.

```
x = exprnd(10,100,1);
normplot(x)
```



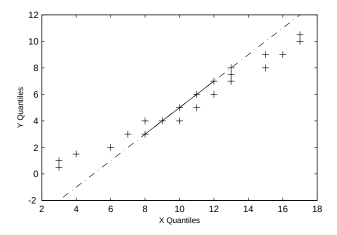
This plot is clear evidence that the underlying distribution is not normal.

Quantile-Quantile Plots

A quantile-quantile plot is useful for determining whether two samples come from the same distribution (whether normally distributed or not).

The example shows a quantile-quantile plot of two samples from a Poisson distribution.

```
x = poissrnd(10,50,1);
y = poissrnd(5,100,1);
qqplot(x,y);
```

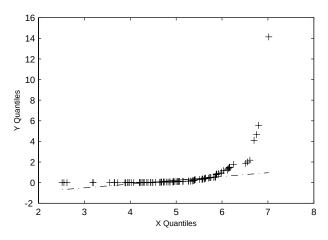


Even though the parameters and sample sizes are different, the straight line relationship shows that the two samples come from the same distribution.

Like the normal probability plot, the quantile-quantile plot has three graphical elements. The pluses are the quantiles of each sample. By default the number of pluses is the number of data values in the smaller sample. The solid line joins the 25th and 75th percentiles of the samples. The dashed line extends the solid line to the extent of the sample.

The example below shows what happens when the underlying distributions are not the same.

```
x = normrnd(5,1,100,1);
y = weibrnd(2,0.5,100,1);
qqplot(x,y);
```



These samples clearly are not from the same distribution.

It is incorrect to interpret a linear plot as a *guarantee* that the two samples come from the same distribution. But, for assessing the validity of a statistical procedure that depends on the two samples coming from the same distribution (e.g., ANOVA), a linear quantile-quantile plot should be sufficient.

Weibull Probability Plots

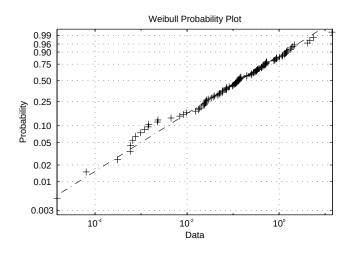
A Weibull probability plot is a useful graph for assessing whether data comes from a Weibull distribution. Many reliability analyses make the assumption that the underlying distribution of the lifetimes is Weibull, so this plot can provide some assurance that this assumption is not being violated, or provide an early warning of a problem with your assumptions.

The scale of the *y*-axis is not uniform. The *y*-axis values are probabilities and, as such, go from zero to one. The distance between the tick marks on the *y*-axis matches the distance between the quantiles of a Weibull distribution.

If the data points (pluses) fall near the line, the assumption that the data comes from a Weibull distribution is reasonable.

This example shows a typical Weibull probability plot.

```
y = weibrnd(2,0.5,100,1);
weibplot(y)
```

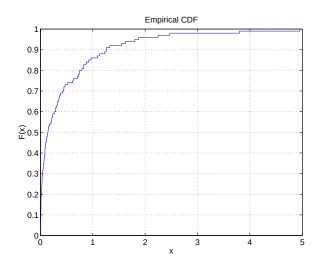


Empirical Cumulative Distribution Function (CDF)

If you are not willing to assume that your data follows a specific probability distribution, you can use the cdfplot function to graph an empirical estimate of the cumulative distribution function (cdf). This function computes the proportion of data points less than each x value, and plots the proportion as a function of x. The y-axis scale is linear, not a probability scale for a specific distribution.

This example shows the empirical cumulative distribution function for a Weibull sample.

y = weibrnd(2,0.5,100,1); cdfplot(y)



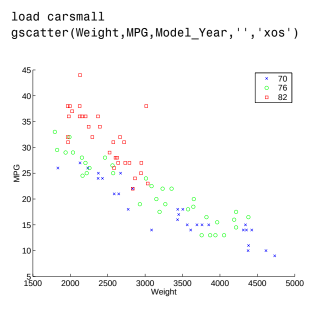
The plot shows a probability function that rises steeply near x=0 and levels off for larger values. Over 80% of the observations are less than 1, with the remaining values spread over the range [1 5].

Scatter Plots

A scatter plot is a simple plot of one variable against another. The MATLAB plot and scatter functions can produce scatter plots. The MATLAB plotmatrix function can produce a matrix of such plots showing the relationship between several pairs of variables.

The Statistics Toolbox adds functions that produce grouped versions of these plots. These are useful for determining whether the values of two variables or the relationship between those variables is the same in each group.

Suppose we want to examine the weight and mileage of cars from three different model years.

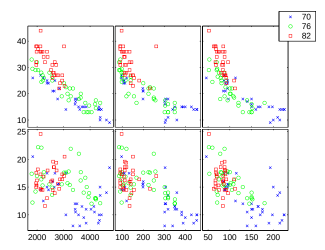


This shows that not only is there a strong relationship between the weight of a car and its mileage, but also that newer cars tend to be lighter and have better gas mileage than older cars.

(The default arguments for gscatter produce a scatter plot with the different groups shown with the same symbol but different colors. The last two arguments above request that all groups be shown in default colors and with different symbols.)

The carsmall data set contains other variables that describe different aspects of cars. We can examine several of them in a single display by creating a grouped plot matrix.

```
xvars = [Weight Displacement Horsepower];
yvars = [MPG Acceleration];
gplotmatrix(xvars,yvars,Model_Year,'','xos')
```



The upper right subplot displays MPG against Horsepower, and shows that over the years the horsepower of the cars has decreased but the gas mileage has improved.

The gplotmatrix function can also graph all pairs from a single list of variables, along with histograms for each variable. See "Multivariate Analysis of Variance (MANOVA)" on page 7-20.



9

Statistical Process Control

Introduction	•	•	•	•	•						•		•	•	•		•	•		•	9-2
Control Char	ts																				9-3
Xbar Charts .																					9-3
S Charts																					9-4
EWMA Charts	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	9-5
Capability Studies																					9-6

Introduction

Statistical Process Control (SPC) is an omnibus term for a number of methods for assessing and monitoring the quality of manufactured goods. These methods are simple, which makes them easy to implement even in a production environment. The following sections discuss some of the SPC features of the Statistics Toolbox:

- "Control Charts" on page 9-3
- "Capability Studies" on page 9-6

Control Charts

These graphs were popularized by Walter Shewhart in his work in the 1920s at Western Electric. A control chart is a plot of a measurements over time with statistical limits applied. Actually, *control* chart is a slight misnomer. The chart itself is actually a monitoring tool. The control activity may occur if the chart indicates that the process is changing in an undesirable systematic direction.

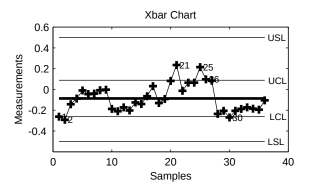
The Statistics Toolbox supports three common control charts, described in the following sections:

- "Xbar Charts" on page 9-3
- "S Charts" on page 9-4
- "EWMA Charts" on page 9-5

Xbar Charts

Xbar charts are a plot of the average of a sample of a process taken at regular intervals. Suppose we are manufacturing pistons to a tolerance of 0.5 thousandths of an inch. We measure the runout (deviation from circularity in thousandths of an inch) at four points on each piston.

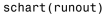
```
load parts
conf = 0.99;
spec = [-0.5 0.5];
xbarplot(runout,conf,spec)
```

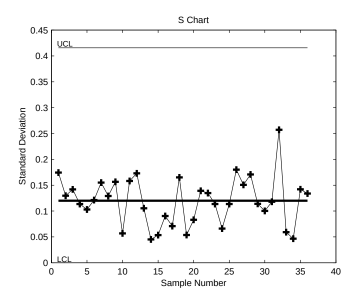


The lines at the bottom and the top of the plot show the process specifications. The central line is the average runout over all the pistons. The two lines flanking the center line are the 99% statistical control limits. By chance only one measurement in 100 should fall outside these lines. We can see that even in this small run of 36 parts, there are several points outside the boundaries (labeled by their observation numbers). This is an indication that the process mean is not in statistical control. This might not be of much concern in practice, since all the parts are well within specification.

S Charts

The S chart is a plot of the standard deviation of a process taken at regular intervals. The standard deviation is a measure of the variability of a process. So, the plot indicates whether there is any systematic change in the process variability. Continuing with the piston manufacturing example, we can look at the standard deviation of each set of four measurements of runout.





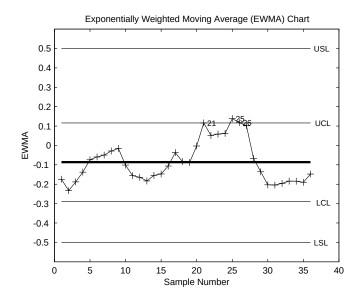
The average runout is about 0.1 thousandths of an inch. There is no indication of nonrandom variability.

EWMA Charts

The exponentially-weighted moving average (EWMA) chart is another chart for monitoring the process average. It operates on slightly different assumptions than the Xbar chart. The mathematical model behind the Xbar chart posits that the process mean is actually constant over time and any variation in individual measurements is due entirely to chance.

The EWMA model is a little looser. Here we assume that the mean may be varying in time. Here is an EWMA chart of our runout example. Compare this with the plot in "Xbar Charts" on page 9-3.

ewmaplot(runout,0.5,0.01,spec)



Capability Studies

Before going into full-scale production, many manufacturers run a pilot study to determine whether their process can actually build parts to the specifications demanded by the engineering drawing.

Using the data from these capability studies with a statistical model allows us to get a preliminary estimate of the percentage of parts that will fall outside the specifications.

```
[p,Cp,Cpk] = capable(mean(runout),spec)
p =
    1.3940e-09
Cp =
    2.3950
Cpk =
    1.9812
```

The result above shows that the probability (p = 1.3940e-09) of observing an unacceptable runout is extremely low. Cp and Cpk are two popular capability indices.

 ${\cal C}_p$ is the ratio of the range of the specifications to six times the estimate of the process standard deviation.

$$C_p = \frac{USL - LSL}{6\sigma}$$

For a process that has its average value on target, a C_p of 1 translates to a little more than one defect per thousand. Recently many industries have set a quality goal of one part per million. This would correspond to a $C_p = 1.6$. The higher the value of C_p , the more capable the process.

 C_{pk} is the ratio of difference between the process mean and the closer specification limit to three times the estimate of the process standard deviation.

$$C_{pk} = min\left(\frac{USL - \mu}{3\sigma}, \frac{\mu - LSL}{3\sigma}\right)$$

where the process mean is μ . For processes that do not maintain their average on target, C_{pk} , is a more descriptive index of process capability.

10

Design of Experiments

Introduction	• •	•	•	•	. 10-2
Full Factorial Designs		•	•	•	. 10-4
Fractional Factorial Designs		•		•	. 10-6
Response Surface Designs					
Central Composite Designs					. 10-8
Box-Behnken Designs		•	•	•	. 10-9
D-Optimal Designs					10-11
Generating D-Optimal Designs					10-11
Augmenting D-Optimal Designs					10-14
Designing Experiments with Uncontrolled Inputs					10-16
Controlling Candidate Points					
Including Categorical Factors					

Introduction

There is a world of difference between data and information. To extract information from data you have to make assumptions about the system that generated the data. Using these assumptions and physical theory you may be able to develop a mathematical model of the system.

Generally, even rigorously formulated models have some unknown constants. The goal of experimentation is to acquire data that allow us to estimate these constants.

But why do we need to experiment at all? We could instrument the system we want to study and just let it run. Sooner or later we would have all the data we could use.

In fact, this is a fairly common approach. There are three characteristics of historical data that pose problems for statistical modeling:

- Suppose we observe a change in the operating variables of a system followed by a change in the outputs of the system. That does *not* necessarily mean that the change in the system *caused* the change in the outputs.
- A common assumption in statistical modeling is that the observations are independent of each other. This is not the way a system in normal operation works.
- Controlling a system in operation often means changing system variables in tandem. But if two variables change together, it is impossible to separate their effects mathematically.

Designed experiments directly address these problems. The overwhelming advantage of a designed experiment is that you actively manipulate the system you are studying. With Design of Experiments (DOE) you may generate fewer data points than by using passive instrumentation, but the quality of the information you get will be higher.

The Statistics Toolbox provides several functions for generating experimental designs appropriate to various situations. These are discussed in the following sections:

- "Full Factorial Designs" on page 10-4
- "Fractional Factorial Designs" on page 10-6
- "Response Surface Designs" on page 10-8

• "D-Optimal Designs" on page 10-11

Full Factorial Designs

Suppose you want to determine whether the variability of a machining process is due to the difference in the lathes that cut the parts or the operators who run the lathes.

If the same operator always runs a given lathe then you cannot tell whether the machine or the operator is the cause of the variation in the output. By allowing every operator to run every lathe you can separate their effects.

This is a factorial approach. fullfact is the function that generates the design. Suppose we have four operators and three machines. What is the factorial design?

```
d = fullfact([4 3])
d =
      1
              1
      2
              1
      3
              1
      4
              1
      1
              2
      2
              2
      3
              2
              2
      4
      1
              3
      2
              3
      3
              3
       4
              3
```

Each row of d represents one operator/machine combination. Note that there are 4*3 = 12 rows.

One special subclass of factorial designs is when all the variables take only two values. Suppose you want to quickly determine the sensitivity of a process to high and low values of three variables.

```
d2 = ff2n(3)

d2 = 0 	 0 	 0
	 0 	 0 	 1
```

0	1	0
0	1	1
1	0	0
1	0	1
1	1	0
1	1	1

There are $2^3 = 8$ combinations to check.

Fractional Factorial Designs

One difficulty with factorial designs is that the number of combinations increases exponentially with the number of variables you want to manipulate.

For example, the sensitivity study discussed above might be impractical if there were seven variables to study instead of just three. A full factorial design would require $2^7 = 128$ runs!

If we assume that the variables do not act synergistically in the system, we can assess the sensitivity with far fewer runs. The theoretical minimum number is eight. A design known as the Plackett-Burman design uses a Hadamard matrix to define this minimal number of runs. To see the design (X) matrix for the Plackett-Burman design, we use the hadamard function.

X = hadamard(8)

X =								
	1	1	1	1	1	1	1	1
	1	- 1	1	- 1	1	- 1	1	- 1
	1	1	- 1	- 1	1	1	- 1	- 1
	1	- 1	- 1	1	1	- 1	- 1	1
	1	1	1	1	- 1	- 1	- 1	- 1
	1	- 1	1	- 1	- 1	1	- 1	1
	1	1	- 1	- 1	- 1	- 1	1	1
	1	- 1	- 1	1	- 1	1	1	- 1

The last seven columns are the actual variable settings (-1 for low, 1 for high.) The first column (all ones) allows us to measure the mean effect in the linear equation, $y = X\beta + \varepsilon$.

The Plackett-Burman design enables us to study the main (linear) effects of each variable with a small number of runs. It does this by using a fraction, in this case 8/128, of the runs required for a full factorial design. A drawback of this design is that if the effect of one variable does vary with the value of another variable, then the estimated effects will be biased (that is, they will tend to be off by a systematic amount).

At a cost of a somewhat larger design, we can find a fractional factorial that is much smaller than a full factorial, but that does allow estimation of main effects independent of interactions between pairs of variables. We can do this by specifying generators that control the confounding between variables. As an example, suppose we create a design with the first four variables varying independently as in a full factorial, but with the other three variables formed by multiplying different triplets of the first four. With this design the effects of the last three variables are confounded with three-way interactions among the first four variables. The estimated effect of any single variable, however, is not confounded with (is independent of) interaction effects between any pair of variables. Interaction effects are confounded with each other. Box, Hunter, and Hunter [5] present the properties of these designs and provide the generators needed to produce them.

The fracfact function can produce this fractional factorial design using the generator strings that Box, Hunter, and Hunter provide.

X = fracfact('a b c d abc bcd acd')

X =

- 1	- 1	- 1	- 1	- 1	- 1	- 1
- 1	- 1	- 1	1	- 1	1	1
- 1	- 1	1	- 1	1	1	1
- 1	- 1	1	1	1	- 1	- 1
- 1	1	- 1	- 1	1	1	- 1
- 1	1	- 1	1	1	- 1	1
- 1	1	1	- 1	- 1	- 1	1
- 1	1	1	1	- 1	1	- 1
1	- 1	- 1	- 1	1	- 1	1
1	- 1	- 1	1	1	1	- 1
1	- 1	1	- 1	- 1	1	- 1
1	- 1	1	1	- 1	- 1	1
1	1	- 1	- 1	- 1	1	1
1	1	- 1	1	- 1	- 1	- 1
1	1	1	- 1	1	- 1	- 1
1	1	1	1	1	1	1

Response Surface Designs

Sometimes simple linear and interaction models are not adequate. For example, suppose that the outputs are defects or yield, and the goal is to minimize defects and maximize yield. If these optimal points are in the interior of the region in which the experiment is to be conducted, we need a mathematical model that can represent curvature so that it has a local optimum. The simplest such model has the quadratic form

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \beta_{12} X_1 X_2 + \beta_{11} X_1^2 + \beta_{22} X_2^2$$

containing linear terms for all factors, squared terms for all factors, and products of all pairs of factors.

Designs for fitting these types of models are known as response surface designs. One such design is the full factorial design having three values for each input. Although the Statistics Toolbox is capable of generating this design, it is not really a satisfactory design in most cases because it has many more runs than are necessary to fit the model.

The two most common designs generally used in response surface modeling are central composite designs and Box-Behnken designs. In these designs the inputs take on three or five distinct values (levels), but not all combinations of these values appear in the design.

The functions described here produce specific response surface designs:

- "Central Composite Designs" on page 10-8
- "Box-Behnken Designs" on page 10-9

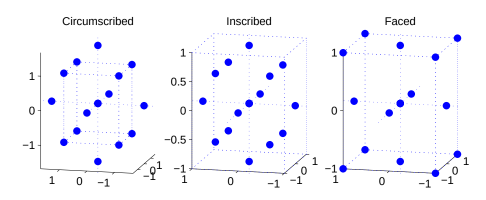
If these do not serve your purposes, consider creating a D-optimal design. For more information see "D-Optimal Designs" on page 10-11.

Central Composite Designs

Central composite designs are response surface designs that can fit a full quadratic model. To picture a central composite design, imagine you have several factors that can vary between low and high values. For convenience, suppose each factor varies from -1 to +1.

One central composite design consists of cube points at the corners of a unit cube that is the product of the intervals [-1,1], star points along the axes at or outside the cube, and center points at the origin.

Central composite designs are of three types. Circumscribed (CCC) designs are as described above. Inscribed (CCI) designs are as described above, but scaled so the star points take the values -1 and +1, and the cube points lie in the interior of the cube. Faced (CCF) designs have the star points on the faces of the cube. Faced designs have three levels per factor, in contrast with the other types that have five levels per factor. The following figure shows these three types of designs for three factors.



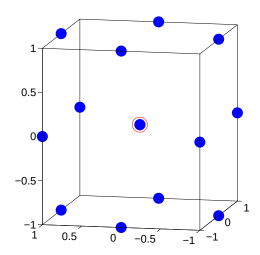
Box-Behnken Designs

Like central composite designs, Box-Behnken designs are response surface designs that can fit a full quadratic model. Unlike most central composite designs, Box-Behnken designs use just three levels of each factor. This makes them appealing when the factors are quantitative but the set of achievable values is small.

Central composite faced (CCF) designs also use just three factor levels. However, they are not rotatable as Box-Behnken designs are. On the other hand, Box-Behnken designs can be expected to have poorer prediction ability in the corners of the cube that encloses the design, because unlike CCF designs they do not include points at the corners of that cube.

The figure below shows a Box-Behnken design for three factors, with the circled point appearing at the origin and possibly repeated for several runs. A

repeated center point makes it possible to compute an estimate of the error term that does not depend on the fitted model. For this design all points except the center point appear at a distance $\sqrt{2}$ from the origin. That doesn't hold true for Box-Behnken designs with different numbers of factors.



D-Optimal Designs

The designs above pre-date the computer age, and some were in use by early in the 20th century. In the 1970s statisticians started to use the computer in experimental design by recasting the design of experiments (DOE) in terms of optimization. A D-optimal design is one that maximizes the determinant of Fisher's information matrix, X^TX . This matrix is proportional to the inverse of the covariance matrix of the parameters. So maximizing $det(X^TX)$ is equivalent to minimizing the determinant of the covariance of the parameters.

A D-optimal design minimizes the volume of the confidence ellipsoid of the regression estimates of the linear model parameters, β .

There are several functions in the Statistics Toolbox that generate D-optimal designs. These are cordexch, daugment, dcovary, and rowexch. The following sections explore D-optimal design in greater detail:

- "Generating D-Optimal Designs" on page 10-11
- "Augmenting D-Optimal Designs" on page 10-14
- "Designing Experiments with Uncontrolled Inputs" on page 10-16
- "Controlling Candidate Points" on page 10-16
- "Including Categorical Factors" on page 10-17

Generating D-Optimal Designs

The cordexch and rowexch functions provide two competing optimization algorithms for computing a D-optimal design given a model specification.

Both cordexch and rowexch are iterative algorithms. They operate by improving a starting design by making incremental changes to its elements. In the coordinate exchange algorithm, the increments are the individual elements of the design matrix. In row exchange, the elements are the rows of the design matrix. Atkinson and Donev [1] is a reference.

To generate a D-optimal design you must specify the number of inputs, the number of runs, and the order of the model you want to fit.

Both cordexch and rowexch take the following strings to specify the model:

- 'linear' or 'l' the default model with constant and first order terms
- 'interaction' or 'i' includes constant, linear, and cross product terms

- 'quadratic' or 'q' interactions plus squared terms
- 'purequadratic' or 'p' includes constant, linear and squared terms

Alternatively, you can use a matrix of integers to specify the terms. Details are in the help for the utility function x2fx.

For a simple example using the coordinate-exchange algorithm, consider the problem of quadratic modeling with two inputs. The model form is

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \epsilon_1 x_1^2 + \beta_{22} x_2^2 + \epsilon_2 x_2^2 + \epsilon_1 x_1^2 + \beta_{22} x_2^2 + \epsilon_2 x_2^2 + \epsilon_1 x_1^2 + \beta_{22} x_2^2 + \epsilon_2 x_2^2 + \epsilon_$$

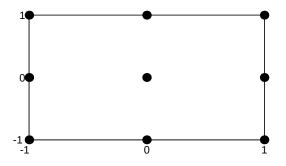
Suppose we want the D-optimal design for fitting this model with nine runs.

```
settings = cordexch(2,9,'q')
settings =
```

```
- 1
           1
 1
           1
 0
           1
 1
         - 1
- 1
         - 1
 0
         - 1
 1
           0
 0
           0
- 1
           0
```

We can plot the columns of settings against each other to get a better picture of the design.

```
h = plot(settings(:,1),settings(:,2),'.');
set(gca,'Xtick',[-1 0 1])
set(gca,'Ytick',[-1 0 1])
set(h,'Markersize',20)
```



For a simple example using the row-exchange algorithm, consider the interaction model with two inputs. The model form is

 $y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2 + \varepsilon$

Suppose we want the D-optimal design for fitting this model with four runs.

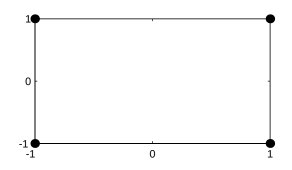
```
[settings, X] = rowexch(2,4,'i')
settings =
     - 1
             1
     - 1
            - 1
      1
            - 1
      1
             1
Х =
            - 1
      1
                     1
                           - 1
      1
            - 1
                    - 1
                            1
      1
             1
                    - 1
                           - 1
             1
      1
                     1
                             1
```

The settings matrix shows how to vary the inputs from run to run. The X matrix is the design matrix for fitting the above regression model. The first column of X is for fitting the constant term. The last column is the element-wise product of the second and third columns.

The associated plot is simple but elegant.

```
h = plot(settings(:,1),settings(:,2),'.');
set(gca,'Xtick',[-1 0 1])
```

set(gca,'Ytick',[-1 0 1])
set(h,'Markersize',20)



Augmenting D-Optimal Designs

In practice, experimentation is an iterative process. We often want to add runs to a completed experiment to learn more about our system. The function daugment allows you choose these extra runs optimally.

Suppose we have executed the eight-run design below for fitting a linear model to four input variables.

```
settings = cordexch(4,8)
settings =
       1
               - 1
                         1
                                 1
      - 1
              - 1
                         1
                                - 1
      - 1
                1
                         1
                                 1
       1
                1
                         1
                                - 1
      - 1
                1
                       - 1
                                 1
       1
              - 1
                       - 1
                                 1
      - 1
              - 1
                       - 1
                                - 1
                                - 1
       1
                1
                       - 1
```

This design is adequate to fit the linear model for four inputs, but cannot fit the six cross-product (interaction) terms. Suppose we are willing to do eight more runs to fit these extra terms. Here's how.

```
[augmented, X] = daugment(settings,8,'i');
```

augmen [.] augmen [.]										
1	- 1	1	1							
- 1	- 1	1	- 1							
- 1	1	1	1							
1	1	1	- 1							
- 1	1	- 1	1							
1	- 1	- 1	1							
- 1	- 1	- 1	- 1							
1	1	- 1	- 1							
- 1	- 1	- 1	1							
1	1	1	1							
- 1	- 1	1	1							
- 1	1	1	- 1							
1	- 1	1	- 1							
1	- 1	- 1	- 1							
- 1	1	- 1	- 1							
1	1	- 1	1							
info =	X'*X									
info =										
		_			-				_	_
16	0	0	0	0	0	0	0	0	0	0
0	16	0	0	0	0	0	0	0	0	0
0	0	16	0	0	0	0	0	0	0	0
0	0	0	16	0	0	0	0	0	0	0
0	0	0	0	16	0	0	0	0	0	0
0	0	0	0	0	16	0	0	0	0	0
0	0	0	0	0	0	16	0	0	0	0
0	0	0	0	0	0	0	16	0	0	0
0	0	0	0	0	0	0	0	16	0	0
0	0	0	0	0	0	0	0	0	16	0
0	0	0	0	0	0	0	0	0	0	16

The augmented design is orthogonal, since X'*X is a multiple of the identity matrix. In fact, this design is the same as a 2^4 factorial design.

Designing Experiments with Uncontrolled Inputs

Sometimes it is impossible to control every experimental input. But you may know the values of some inputs in advance. An example is the time each run takes place. If a process is experiencing linear drift, you may want to include the time of each test run as a variable in the model.

The function dcovary allows you to choose the settings for each run in order to maximize your information despite a linear drift in the process.

Suppose we want to execute an eight-run experiment with three factors that is optimal with respect to a linear drift in the response over time. First we create our drift input variable. Note, that drift is normalized to have mean zero. Its minimum is -1 and its maximum is 1.

```
drift = (linspace(-1,1,8))'
drift =
   -1.0000
   -0.7143
   -0.4286
   -0.1429
    0.1429
    0.4286
    0.7143
    1.0000
settings = dcovary(3,drift,'linear')
settings =
    1.0000
               1.0000
                         -1.0000
                                    -1.0000
   -1.0000
              -1.0000
                         -1.0000
                                    -0.7143
   -1.0000
                                    -0.4286
               1.0000
                          1.0000
    1.0000
              -1.0000
                          1.0000
                                    -0.1429
   -1.0000
               1.0000
                         -1.0000
                                     0.1429
    1.0000
               1.0000
                          1.0000
                                     0.4286
   -1.0000
              -1.0000
                          1.0000
                                     0.7143
    1.0000
              -1.0000
                         -1.0000
                                     1.0000
```

Controlling Candidate Points

The rowexch function generates a candidate set of possible design points, and then uses a D-optimal algorithm to select a design from those points. It does this by invoking the candgen and candexch functions. If you need to supply your own candidate set, or if you need to modify the one that the candgen function provides, then you may prefer to call these functions separately.

This example creates a design that represents proportions of a mixture, so the sum of the proportions cannot exceed 1.

```
% Generate a matrix of (x,y) values with x+y<=1
[x,y]=meshgrid(0:.1:1);
xy = [x(:) y(:)];
xy = xy(sum(xy,2) <= 1,:);
% Compute quadratic model terms for these points.
f = x2fx(xy, 'q');
% Generate a 10-point design and display it
r=candexch(f,10);
xy(r,:)
ans =
        0
                     0
                1.0000
        0
   1.0000
                     0
                0.5000
        0
   0.5000
                     0
        0
                1.0000
   1.0000
                     n
   0.5000
                0.5000
   0.5000
                     0
   0.5000
                0.5000
```

Including Categorical Factors

Another example where it is useful to call candexch directly is to generate a design that includes categorical factors. For these designs you create a candidate set containing "dummy variables" for the categorical factors. The dummyvar function is useful to create such a candidate set.

In this example we have three categorical factors, each taking three levels. We create a candidate set F containing all 27 combinations of these factor levels. Then we create a matrix C containing dummy variables for the factors, and we remove enough columns to make the resulting matrix full rank. (We remove

one column for each factor except the first factor.) Finally, we use the candexch function to generate a 9-run design. The resulting design has the property that for each pair of factors, each of the 9 possible combinations of levels appears exactly once.

Demos

Introduction	•			•	•		•	•	•	•	•	•			•	•	. 11-2
The disttool Demo	•									•	•					•	. 11-3
The polytool Demo																	. 11-5
Confidence Bounds .																	. 11-7
Overfitting																	
The aoctool Demo																	11-10
Example: aoctool with	n S	an	npl	e I	Da	ta	•	•	•	•	•	•			•	•	11-11
The randtool Demo	• •	•		•	•	•		•	•	•	•	•		•	•	•	11-18
The rsmdemo Demo	D																11-19
Part 1																	11-19
Part 2																	
The glmdemo Demo).	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	11-21
The robustdemo De	m	0								•						•	11-22

11

Introduction

The Statistics Toolbox has demonstration programs that create an interactive environment for exploring the probability distributions, random number generation, curve fitting, and design of experiments functions. Most of them provide a graphical user interface that can be used with your real data, not just with the sample data provided.

Demo	Purpose
aoctool	Interactive graphic prediction of anocova fits
disttool	Graphic interaction with probability distributions
glmdemo	Generalized linear models slide show
nlintool	Interactive fitting of nonlinear models
polytool	Interactive graphic prediction of polynomial fits
randtool	Interactive control of random number generation
robustdemo	Interactive comparison of robust and least squares fits
rsmdemo	Design of experiments and regression modeling
rstool	Exploring graphs of multidimensional polynomials
stepwise	Interactive stepwise regression

The available demos are listed below.

Most of these functions are described below. The nlintool, rstool, and stepwise demos are discussed in earlier sections:

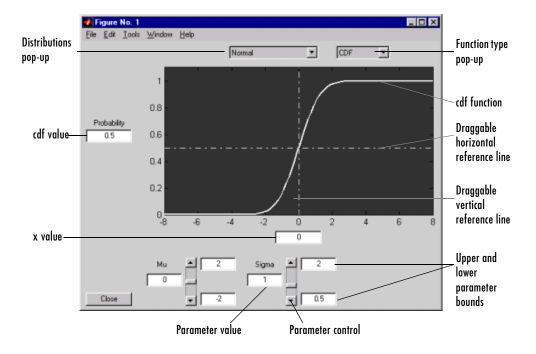
- nlintool: "An Interactive GUI for Nonlinear Fitting and Prediction" on page 5-7
- rstool: "Exploring Graphs of Multidimensional Polynomials" on page 4-22
- stepwise: "Example: Stepwise Regression" on page 4-24

The disttool Demo

disttool is a graphic environment for developing an intuitive understanding of probability distributions.

The disttool demo has the following features:

- A graph of the cdf (pdf) for the given parameters of a distribution.
- A pop-up menu for changing the distribution function.
- A pop-up menu for changing the function type (cdf <-> pdf).
- Sliders to change the parameter settings.
- Data entry boxes to choose specific parameter values.
- Data entry boxes to change the limits of the parameter sliders.
- Draggable horizontal and vertical reference lines to do interactive evaluation of the function at varying values.
- A data entry box to evaluate the function at a specific *x*-value.
- For cdf plots, a data entry box on the probability axis (*y*-axis) to find critical values corresponding to a specific probability.
- A Close button to end the demonstration.



The polytool Demo

The polytool demo is an interactive graphic environment for polynomial curve fitting and prediction.

The polytool demo has the following features:

- A graph of the data, the fitted polynomial, and global confidence bounds on a new predicted value.
- *y*-axis text to display the predicted *y*-value and its uncertainty at the current *x*-value.
- A data entry box to change the degree of the polynomial fit.
- A data entry box to evaluate the polynomial at a specific *x*-value.
- A draggable vertical reference line to do interactive evaluation of the polynomial at varying *x*-values.
- **Bounds** and **Method** menus to control the confidence bounds and choose between least squares or robust fitting.
- A Close button to end the demonstration.
- An **Export** list box to store fit results into variables.

You can use polytool to do curve fitting and prediction for any set of x-y data, but, for the sake of demonstration, the Statistics Toolbox provides a data set (polydata.mat) to teach some basic concepts.

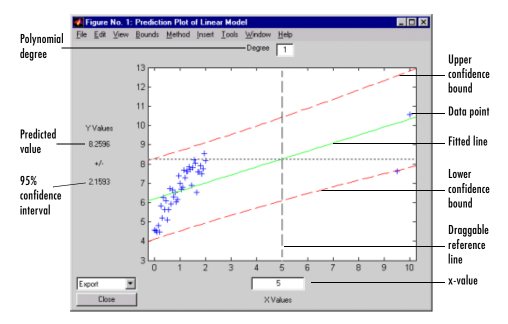
To start the demonstration, you must first load the data set.

load polydata who Your variables are: x x1 y y1

The variables x and y are observations made with error from a cubic polynomial. The variables x1 and y1 are data points from the "true" function without error.

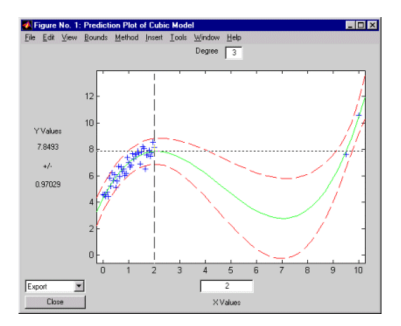
If you do not specify the degree of the polynomial, polytool does a linear fit to the data.

```
polytool(x,y)
```



The linear fit is not very good. The bulk of the data with *x*-values between zero and two has a steeper slope than the fitted line. The two points to the right are dragging down the estimate of the slope.

In the **Degree** box at the top, type 3 for a cubic model. Then, drag the vertical reference line to the *x*-value of 2 (or type 2 in the **X Values** text box).



This graph shows a much better fit to the data. The confidence bounds are closer together indicating that there is less uncertainty in prediction. The data at both ends of the plot tracks the fitted curve.

The following sections explore additional aspects of the tool:

- "Confidence Bounds" on page 11-7
- "Overfitting" on page 11-8

Confidence Bounds

By default, the confidence bounds are nonsimultaneous bounds for a new observation. What does this mean? Let p(x) be the true but unknown function we want to estimate. The graph contains the following three curves:

- f(x), our fitted function
- l(x), the lower confidence bounds
- u(x), the upper confidence bounds

Suppose we plan to take a new observation at the value x_{n+1} . Call it $y_{n+1}(x_{n+1})$. This new observation has its own error ε_{n+1} , so it satisfies the equation

$$y_{n+1}(x_{n+1}) = p(x_{n+1}) + \varepsilon_{n+1}$$

What are the likely values for this new observation? The confidence bounds provide the answer. The interval $[l_{n+1}, \, u_{n+1}]$ is a 95% confidence bound for $y_{n+1}(x_{n+1})$.

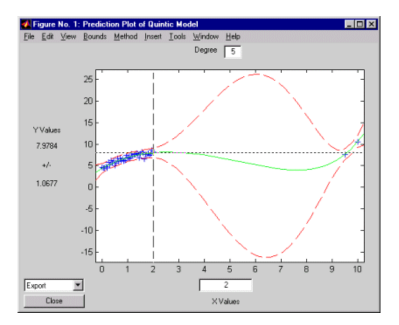
These are the default bounds, but the **Bounds** menu on the polytool figure window provides options for changing the meaning of these bounds. This menu has options that let you specify whether the bounds are to apply to the estimated function or to a new observation, and whether the bounds should be simultaneous or not. Using these options you can produce any of the following types of confidence bounds.

Simultaneous?	For Quantity	Yields Confidence Bounds for
Nonsimultaneous	Observation	$y_{n+1}(x_{n+1})$
Nonsimultaneous	Curve	$p(x_{n+1})$
Simultaneous	Observation	$y_{n+1}(x)$, globally for any x
Simultaneous	Curve	p(x), simultaneously for all x

Overfitting

If the cubic polynomial is a good fit, it is tempting to try a higher order polynomial to see if even more precise predictions are possible.

Since the true function is cubic, this amounts to overfitting the data. Use the data entry box for degree and type 5 for a quintic model.



As measured by the confidence bounds, the fit is precise near the data points. But, in the region between the data groups, the uncertainty of prediction rises dramatically.

This bulge in the confidence bounds happens because the data really does not contain enough information to estimate the higher order polynomial terms precisely, so even interpolation using polynomials can be risky in some cases.

The aoctool Demo

The aoctool demo is an interactive graphical environment for fitting and prediction with analysis of covariance (anocova) models. It is similar to the polytool demo.

Analysis of covariance is a technique for analyzing grouped data having a response (y, the variable to be predicted) and a predictor (x, the variable used to do the prediction). Using analysis of covariance, you can model y as a linear function of x, with the coefficients of the line possibly varying from group to group. The aoctool function fits the following models for the *i*th group:

l same mean	$y = \alpha + \varepsilon$
2 separate means	$y = (\alpha + \alpha_i) + \varepsilon$
3 same line	$y = \alpha + \beta x + \varepsilon$
4 parallel lines	$y = (\alpha + \alpha_i) + \beta x + \varepsilon$
5 separate lines	$y = (\alpha + \alpha_i) + (\beta + \beta_i)x + \varepsilon$

In the fourth model, for example, the intercept varies from one group to the next, but the slope is the same for each group. In the first model, there is a common intercept and no slope. In order to make the group coefficients well determined, we impose the constraints $\Sigma \alpha_i = \Sigma \beta_i = 0$.

The aoctool demo displays the results of the fit in three figure windows. One window displays estimates of the coefficients (α , α_i , β , β_i). A second displays an analysis of variance table that you can use to test whether a more complex model is significantly better than a simpler one. The third, main graphics window has the following features:

- A graph of the data with superimposed fitted lines and optional confidence bounds.
- *y*-axis text to display the predicted *y*-value and its uncertainty at the current *x*-value for the current group, if a group is currently selected.
- A data entry box to evaluate the fit at a specific *x*-value.
- A list box to evaluate the fit for a specific group or to display fitted lines for all groups.

- A draggable vertical reference line to do interactive evaluation of the fit at varying *x*-values.
- A Close button to end the demonstration.
- An **Export** list box to store fit results into variables.

The following section provides an illustrative example.

Example: aoctool with Sample Data

The Statistics Toolbox has a small data set named carsmall with information about cars. It is a good sample data set to use with aoctool. You can also use aoctool with your own data.

To start the demonstration, load the data set.

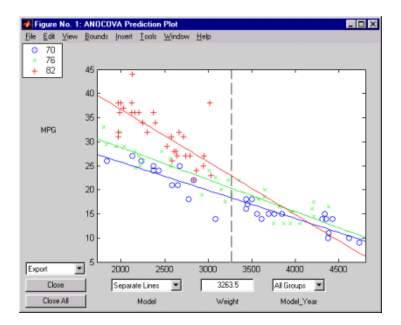
```
load carsmall
who
Your variables are:
Acceleration Horsepower Model_Year
Cylinders MPG Origin
Displacement Model Weight
```

Suppose we want to study the relationship between the weight of a car and its mileage, and whether this relationship has changed over the years.

Next, start up the tool.

```
[h,atab,ctab,stats] = aoctool(Weight,MPG,Model_Year);
Note: 6 observations with missing values have been removed.
```

The graphical output consists of the following main window, plus a table of coefficient estimates and an analysis of variance table.



The group of each data point is coded by its color and symbol, and the fit for each group has the same color as the data points.

🛃 Figure No. 3:	ANOCOVA Coeff	icients		
<u>File E</u> dit <u>T</u> ools	<u>W</u> indow <u>H</u> elp			
	Coeffi	cient Estima	ates	
Coefficient	Estimate	Std. Err.	Т	Prob> T 🛌
Intercept 70 76 82 Slope 70 76 82	45.9798 -8.5805 -3.8902 12.4707 -0.0078 0.002 0.0011 -0.0031	1.52085 1.96186 1.86864 2.5568 0.00056 0.00066 0.00065 0.001	30.23 -4.37 -2.08 4.88 -14 2.96 1.74 -3.1	0 0.0403 0 0.0039 0.0849 0.0826

The initial fit models the y variable, MPG, as a linear function of the x variable, Weight. Each group has a separate line. The coefficients of the three lines

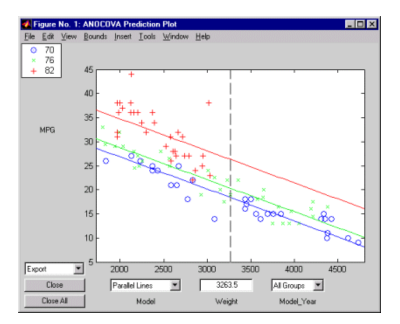
appear in the figure titled **ANOCOVA Coefficients**. You can see that the slopes are roughly -0.0078, with a small deviation for each group:

Model year 70:	$y = (45.9798 - 8.5805) + (-0.0078 + 0.002)x + \varepsilon$
Model year 76:	$y = (45.9798 - 3.8902) + (-0.0078 + 0.0011)x + \varepsilon$
Model year 82:	$y = (45.9798 + 12.4707) + (-0.0078 - 0.0031)x + \varepsilon$

Notice that the three fitted lines have slopes that are roughly similar. Could they really be the same? The Model_Year*Weight interaction expresses the difference in slopes, and the ANOVA table shows a test for the significance of this term. With an F statistic of 5.23 and a p-value of 0.0072, the slopes are significantly different.

🛃 Figure No. 2: ANOCOV	A Test R	esults			
<u>File E</u> dit <u>T</u> ools <u>W</u> indow	<u>H</u> elp				
		ANOVA Ta	able		
Source	d.f.	Sum Sq	Mean Sq	F	Prob>F 🛌
Model_Year Weight Model_Year*Weight Error	2 1 2 88	807.69 2050.2 81.22 683.74	403.84 2050.2 40.61 7.77	51.98 263.87 5.23	0 0 0.0072

To examine the fits when the slopes are constrained to be the same, return to the **ANOCOVA Prediction Plot** window and use the **Model** pop-up to select a **Parallel Lines** model. The window updates to show the graph below.



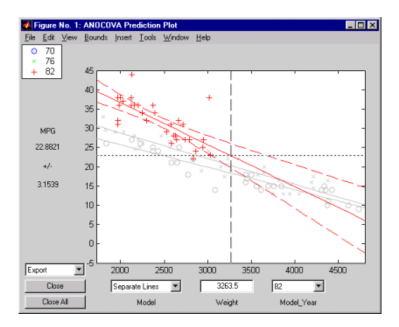
Though this fit looks reasonable, we know it is significantly worse than the **Separate Lines** model. Use the **Model** pop-up again to return to the original model.

The following sections focus on two other interesting aspects of aoctool:

- "Confidence Bounds" on page 11-14
- "Multiple Comparisons" on page 11-16

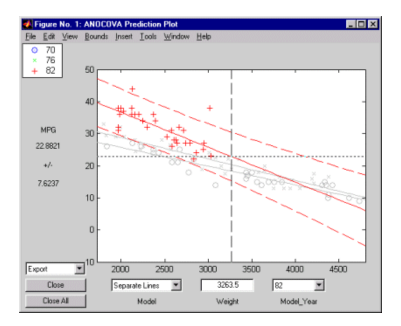
Confidence Bounds

Now we have estimates of the relationship between MPG and Weight for each Model_Year, but how accurate are they? We can superimpose confidence bounds on the fits by examining them one group at a time. In the **Model_Year** menu at the lower right of the figure, change the setting from **All Groups** to 82. The data and fits for the other groups are dimmed, and confidence bounds appear around the 82 fit.



The dashed lines form an envelope around the fitted line for model year 82. Under the assumption that the true relationship is linear, these bounds provide a 95% confidence region for the true line. Note that the fits for the other model years are well outside these confidence bounds for Weight values between 2000 and 3000.

Sometimes it is more valuable to be able to predict the response value for a new observation, not just estimate the average response value. Like the polytool function, the aoctool function has a **Bounds** menu to change the definition of the confidence bounds. Use that menu to change from **Line** to **Observation**. The resulting wider intervals reflect the uncertainty in the parameter estimates as well as the randomness of a new observation.



Also like the polytool function, the aoctool function has crosshairs you can use to manipulate the Weight and watch the estimate and confidence bounds along the *y*-axis update. These values appear only when a single group is selected, not when **All Groups** is selected.

Multiple Comparisons

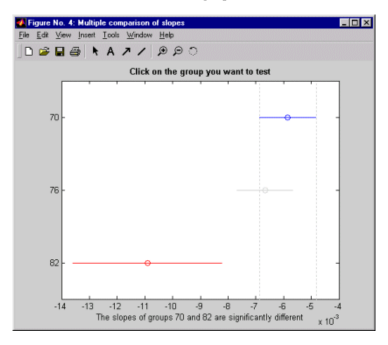
We can perform a multiple comparison test by using the stats output from aoctool as input to the multcompare function. The multcompare function can test either slopes, intercepts, or population marginal means (the heights of the four lines evaluated at the mean X value). In this example, we have already determined that the slopes are not all the same, but could it be that two are the same and only the other one is different? We can test that hypothesis.

```
multcompare(stats,0.05,'on','','s')
```

```
ans =
```

1.0000	2.0000	-0.0012	0.0008	0.0029
1.0000	3.0000	0.0013	0.0051	0.0088
2.0000	3.0000	0.0005	0.0042	0.0079

This matrix shows that the estimated difference between the intercepts of groups 1 and 2 (1970 and 1976) is 0.0008, and a confidence interval for the difference is [-0.0012, 0.0029]. There is no significant difference between the two. There are significant differences, however, between the intercept for 1982 and each of the other two. The graph shows the same information.



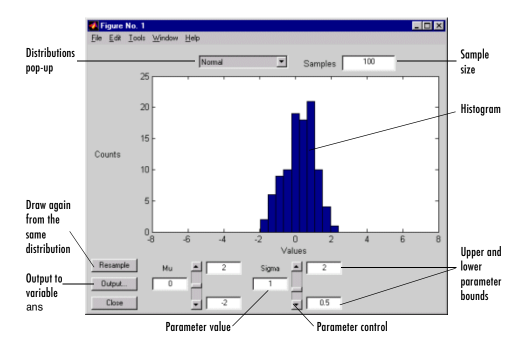
Note that the stats structure was created in the initial call to the aoctool function, so it is based on the initial model fit (typically a separate-lines model). If you change the model interactively and want to base your multiple comparisons on the new model, you need to run aoctool again to get another stats structure, this time specifying your new model as the initial model.

The randtool Demo

randtool is a graphic environment for generating random samples from various probability distributions and displaying the sample histogram.

The randtool demo has the following features:

- A histogram of the sample.
- A pop-up menu for changing the distribution function.
- Sliders to change the parameter settings.
- A data entry box to choose the sample size.
- Data entry boxes to choose specific parameter values.
- Data entry boxes to change the limits of the parameter sliders.
- An **Output** button to output the current sample to the variable ans.
- A **Resample** button to allow repetitive sampling with constant sample size and fixed parameters.
- A Close button to end the demonstration.



The rsmdemo Demo

The rsmdemo utility is an interactive graphic environment that demonstrates the design of experiments and surface fitting through the simulation of a chemical reaction. The goal of the demo is to find the levels of the reactants needed to maximize the reaction rate.

There are two parts to the demo:

- "Part 1" on page 11-19 Compare data gathered through trial and error with data from a designed experiment.
- "Part 2" on page 11-20 Compare response surface (polynomial) modeling with nonlinear modeling.

Part 1

Begin the demo by using the sliders in the **Reaction Simulator** window to control the partial pressures of three reactants: **Hydrogen**, **n-Pentane**, and **Isopentane**. Each time you click the **Run** button, the levels for the reactants and results of the run are entered in the **Trial and Error Data** window.

Based on the results of previous runs, you can change the levels of the reactants to increase the reaction rate. (The results are determined using an underlying model that takes into account the noise in the process, so even if you keep all of the levels the same, the results will vary from run to run.) You are allotted a budget of 13 runs. When you have completed the runs, you can use the **Plot** menu on the **Trial and Error Data** window to plot the relationships between the reactants and the reaction rate, or click the **Analyze** button. When you click **Analyze**, rsmdemo calls the rstool function, which you can then use to try to optimize the results.)

Next, perform another set of 13 runs, this time from a designed experiment. In the **Experimental Design Data** window, click the **Do Experiment** button. rsmdemo calls the cordexch function to generate a D-optimal design, and then, for each run, computes the reaction rate.

Now use the **Plot** menu on the **Experimental Design Data** window to plot the relationships between the levels of the reactants and the reaction rate, or click the **Response Surface** button to call rstool to find the optimal levels of the reactants.

Compare the analysis results for the two sets of data. It is likely (though not certain) that you'll find some or all of these differences:

- You can fit a full quadratic model with the data from the designed experiment, but the trial and error data may be insufficient for fitting a quadratic model or interactions model.
- Using the data from the designed experiment, you are more likely to be able to find levels for the reactants that result in the maximum reaction rate. Even if you find the best settings using the trial and error data, the confidence bounds are likely to be wider than those from the designed experiment.

Part 2

Now analyze the experimental design data with a polynomial model and a nonlinear model, and comparing the results. The true model for the process, which is used to generate the data, is actually a nonlinear model. However, within the range of the data, a quadratic model approximates the true model quite well.

To see the polynomial model, click the **Response Surface** button on the **Experimental Design Data** window. rsmdemo calls rstool, which fits a full quadratic model to the data. Drag the reference lines to change the levels of the reactants, and find the optimal reaction rate. Observe the width of the confidence intervals.

Now click the **Nonlinear Model** button on the **Experimental Design Data** window. rsmdemo calls nlintool, which fits a Hougen-Watson model to the data. As with the quadratic model, you can drag the reference lines to change the reactant levels. Observe the reaction rate and the confidence intervals.

Compare the analysis results for the two models. Even though the true model is nonlinear, you may find that the polynomial model provides a good fit. Because polynomial models are much easier to fit and work with than nonlinear models, a polynomial model is often preferable even when modeling a nonlinear process. Keep in mind, however, that such models are unlikely to be reliable for extrapolating outside the range of the data.

The glmdemo Demo

The glmdemo function presents a simple slide show describing generalized linear models. It presents examples of what functions and distributions are available with generalized linear models. It presents an example where traditional linear least squares fitting is not appropriate, and shows how to use the glmfit function to fit a logistic regression model and the glmval function to compute predictions from that model.

To run the glmdemo from the command line, type playshow glmdemo.

The robustdemo Demo

The robustdemo function presents a simple comparison of least squares and robust fits for a response and a single predictor. You can use robustdemo with your own data or with the sample data provided.

To begin using robustdemo with the built-in sample data, simply type the function name.

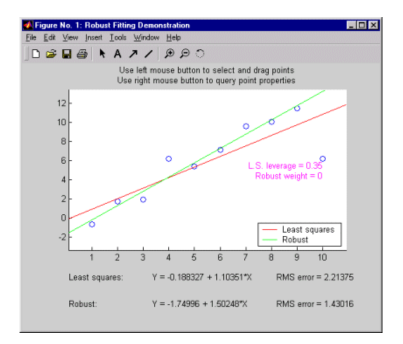
robustdemo

The resulting figure presents a scatter plot with two fitted lines. One line is the fit from an ordinary least squares regression. The other is from a robust regression. Along the bottom of the figure are the equations for the fitted line and the estimated error standard deviation for each fit.

The effect of any point on the least squares fit depends on the residual and leverage for that point. The residual is simply the vertical distance from the point to the line. The leverage is a measure of how far the point is from the center of the X data.

The effect of any point on the robust fit also depends on the weight assigned to the point. Points far from the line get lower weight.

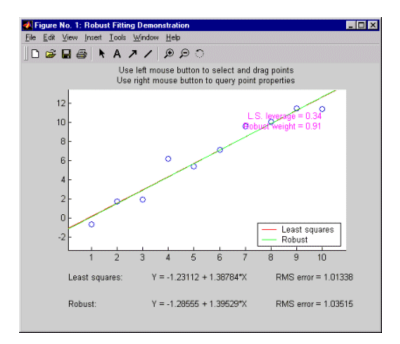
You can use the right mouse button to click on any point and see its least squares leverage and robust weight.



In this example, the rightmost point has a leverage value of 0.35. It is also far from the line, so it exerts a large influence on the least squares fit. It has a small weight, though, so it is effectively excluded from the robust fit.

Using the left mouse button, you can experiment to see how changes in the data affect the two fits. Select any point, and drag it to a new location while holding the left button down. When you release the point, both fits update.

Bringing the rightmost point closer to the line makes the two fitted lines nearly identical. Now, the point has nearly full weight in the robust fit.



12

Reference

This chapter contains detailed descriptions of all the Statistics Toolbox functions. It is divided into two sections:

- "Functions By Category" on page 12-3, a list of functions grouped by subject area.
- "Functions Alphabetical List" on page 12-18, detailed function descriptions in alphabetical order.

Functions – By Category

The Statistics Toolbox provides several categories of functions.

Probability Distributions	Specific functions for each supported distribution
Descriptive Statistics	Descriptive statistics for data samples
Statistical Plotting	Statistical plots
Statistical Process Control	Statistical Process Control
Linear Models	Fitting linear models to data
Nonlinear Regression	Fitting nonlinear regression models
Design of Experiments	Design of Experiments
Multivariate Statistics	Multivariate statistics
Decision Tree Techniques	Decision trees
Hypothesis Tests	Statistical tests of hypotheses
Distribution Testing	Tests for fitting distributions to data
Nonparametric Testing	Nonparametric testing
File I/O	Reading data from and writing data to operating-system files
Demonstrations	Demonstrations
Data	Data for examples

The following tables list the functions in each of these specific areas. The first seven tables contain probability distribution functions. The remaining tables describe the other categories of functions.

Probability Distributions

For each supported distribution, the Statistics Toolbox provides associated functions in each of these categories

Parameter Estimation

Cumulative Distribution Functions (cdf)

Probability Density Functions (pdf)

Inverse Cumulative Distribution Functions

Random Number Generators

Moments of Distribution Functions

Parameter Estimation

betafit	Parameter estimation for the beta distribution
betalike	Negative beta log-likelihood function
binofit	Parameter estimation for the binomial distribution
expfit	Parameter estimation for the exponential distribution
gamfit	Parameter estimation for the gamma distribution
gamlike	Negative gamma log-likelihood function
mle	Maximum likelihood estimation
nbinfit	Parameter estimates and confidence intervals for negative binomial data
normfit	Parameter estimation for the normal distribution
normlike	Negative normal log-likelihood function
poissfit	Parameter estimation for the Poisson distribution
raylfit	Rayleigh parameter estimation
unifit	Parameter estimation for the uniform distribution
weibfit	Weibull parameter estimation
weiblike	Weibull negative log-likelihood function

Cumulative Distribution Functions (cdf)

betacdf	Beta cdf
binocdf	Binomial cdf
cdf	Parameterized cdf routine
chi2cdf	Chi-square cdf
ecdf	Empirical (Kaplan-Meier) cdf
expcdf	Exponential cdf
fcdf	F cdf
gamcdf	Gamma cdf
geocdf	Geometric cdf
hygecdf	Hypergeometric cdf
logncdf	Lognormal cdf
nbincdf	Negative binomial cdf
ncfcdf	Noncentral F cdf
nctcdf	Noncentral t cdf
ncx2cdf	Noncentral Chi-square cdf
normcdf	Normal (Gaussian) cdf
poisscdf	Poisson cdf
raylcdf	Rayleigh cdf
tcdf	Student's t cdf
unidcdf	Discrete uniform cdf
unifcdf	Continuous uniform cdf
weibcdf	Weibull cdf

Probability Density Functions (pdf)

betapdf	Beta pdf
binopdf	Binomial pdf

chi2pdf	Chi-square pdf
exppdf	Exponential pdf
fpdf	F pdf
gampdf	Gamma pdf
geopdf	Geometric pdf
hygepdf	Hypergeometric pdf
lognpdf	Lognormal pdf
m∨npdf	Multivariate normal pdf
nbinpdf	Negative binomial pdf
ncfpdf	Noncentral F pdf
nctpdf	Noncentral t pdf
ncx2pdf	Noncentral Chi-square pdf
normpdf	Normal (Gaussian) pdf
pdf	Parameterized pdf routine
poisspdf	Poisson pdf
raylpdf	Rayleigh pdf
tpdf	Student's t pdf
unidpdf	Discrete uniform pdf
unifpdf	Continuous uniform pdf
weibpdf	Weibull pdf

Inverse Cumulative Distribution Functions

betainv	Beta critical values
binoinv	Binomial critical values
chi2inv	Chi-square critical values
expinv	Exponential critical values
finv	F critical values

gaminv	Gamma critical values
geoinv	Geometric critical values
hygeinv	Hypergeometric critical values
icdf	Parameterized inverse distribution routine
logninv	Lognormal critical values
nbininv	Negative binomial critical values
ncfinv	Noncentral F critical values
nctinv	Noncentral t critical values
ncx2inv	Noncentral Chi-square critical values
norminv	Normal (Gaussian) critical values
poissinv	Poisson critical values
raylinv	Rayleigh critical values
tinv	Student's t critical values
unidinv	Discrete uniform critical values
unifinv	Continuous uniform critical values
weibinv	Weibull critical values

Random Number Generators

betarnd	Beta random numbers
binornd	Binomial random numbers
chi2rnd	Chi-square random numbers
exprnd	Exponential random numbers
frnd	F random numbers
gamrnd	Gamma random numbers
geornd	Geometric random numbers
hygernd	Hypergeometric random numbers
iwishrnd	Inverse Wishart random matrix

lhsdesign	Latin hypercube sample
lhsnorm	Latin hypercube sample with normal distribution
lognrnd	Lognormal random numbers
mvnrnd	Multivariate normal random numbers
mvtrnd	Multivariate t random numbers
nbinrnd	Negative binomial random numbers
ncfrnd	Noncentral F random numbers
nctrnd	Noncentral t random numbers
ncx2rnd	Noncentral Chi-square random numbers
normrnd	Normal (Gaussian) random numbers
poissrnd	Poisson random numbers
random	Parameterized random number routine
raylrnd	Rayleigh random numbers
trnd	Student's t random numbers
unidrnd	Discrete uniform random numbers
unifrnd	Continuous uniform random numbers
weibrnd	Weibull random numbers
wishrnd	Wishart random matrix

Moments of Distribution Functions

betastat	Beta mean and variance
binostat	Binomial mean and variance
chi2stat	Chi-square mean and variance
expstat	Exponential mean and variance
fstat	F mean and variance
gamstat	Gamma mean and variance
geostat	Geometric mean and variance

Hypergeometric mean and variance
Lognormal mean and variance
Negative binomial mean and variance
Noncentral F mean and variance
Noncentral t mean and variance
Noncentral Chi-square mean and variance
Normal (Gaussian) mean and variance
Poisson mean and variance
Rayleigh mean and variance
Student's t mean and variance
Discrete uniform mean and variance
Continuous uniform mean and variance
Weibull mean and variance

Descriptive Statistics

bootstrp	Bootstrap statistics for any function
corrcoef	$Correlation \ coefficients \ (in \ MATLAB)$
COV	Covariance matrix (in MATLAB)
crosstab	Cross tabulation
geomean	Geometric mean
grpstats	Summary statistics by group
harmmean	Harmonic mean
iqr	Interquartile range
kurtosis	Sample kurtosis
mad	Mean absolute deviation
mean	Arithmetic average (in MATLAB)

median	50th percentile (in MATLAB)
moment	Central moments of all orders
nanmax	Maximum ignoring missing data
nanmean	Average ignoring missing data
nanmedian	Median ignoring missing data
nanmin	Minimum ignoring missing data
nanstd	Standard deviation ignoring missing data
nansum	Sum ignoring missing data
prctile	Empirical percentiles of a sample
range	Sample range
skewness	Sample skewness
std	Standard deviation (in MATLAB)
tabulate	Frequency table
trimmean	Trimmed mean
var	Variance

Statistical Plotting

boxplot	Bo	x plots
cdfplot	Plo	ot of empirical cumulative distribution function
errorbar	Er	ror bar plot
fsurfht	Int	ceractive contour plot of a function
gline	Int	ceractive line drawing
gname	Int	eractive point labeling
gplotmatr	ix Ma	atrix of scatter plots grouped by a common variable
gscatter	Sca	atter plot of two variables grouped by a third
lsline	Ad	d least-squares fit line to plotted data

normplot	Normal probability plots
pareto	Pareto charts
qqplot	Quantile-Quantile plots
rcoplot	Regression case order plot
refcurve	Reference polynomial
refline	Reference line
surfht	Interactive interpolating contour plot
weibplot	Weibull plotting

Statistical Process Control

capable	Quality capability indices
capaplot	Plot of process capability
ewmaplot	Exponentially weighted moving average plot
histfit	Histogram and normal density curve
normspec	Plot normal density between limits
schart	Time plot of standard deviation
xbarplot	Time plot of means

Linear Models

anova1	One-way Analysis of Variance (ANOVA)
anova2	Two-way Analysis of Variance
anovan	N-way analysis of variance
aoctool	Interactive tool for analysis of covariance
dummyvar	Dummy-variable coding
friedman	Friedman's test (nonparametric two-way anova)
glmfit	Generalized linear model fitting

glmval	Compute predictions for generalized linear model
kruskalwallis	Kruskal-Wallis test (nonparametric one-way anova)
leverage	Regression diagnostic
lscov	Regression given a covariance matrix (in MATLAB)
manova1	One-way multivariate analysis of variance
manovacluster	Draw clusters of group means for manoval
multcompare	Multiple comparisons of means and other estimates
polyconf	Polynomial prediction with confidence intervals
polyfit	Polynomial fitting (in MATLAB)
polyval	Polynomial prediction (in MATLAB)
rcoplot	Residuals case order plot
regress	Multiple linear regression
regstats	Regression diagnostics for linear models
ridge	Ridge regression
rstool	Response surface tool
robustfit	Robust regression model fitting
rstool	Multidimensional response surface visualization (RSM)
stepwise	Stepwise regression GUI
x2fx	Factor settings matrix (X) to design matrix (D)

Nonlinear Regression

nlinfit	Nonlinear least-squares fitting
nlintool	Prediction graph for nonlinear fits
nlparci	Confidence intervals on parameters
nlpredci	Confidence intervals for prediction
nnls	Nonnegative least squares (in MATLAB)

treefit	Fit a tree-based	l model for	classification	or regression.

- treeprune Produce a sequence of subtrees by pruning.
- treedisp Show classification or regression tree graphically.
- treetest Compute error rate for tree.
- treeval Compute fitted value for decision tree applied to data.

Design of Experiments

bbdesign	Box-Behnken design
candgen	Candidate set for D-optimal design
candexch	D-optimal design from candidate set using row exchanges
ccdesign	Central composite design
cordexch	D-optimal design using coordinate exchange
daugment	D-optimal augmentation of designs
dcovary	D-optimal design with fixed covariates
ff2n	Two-level full factorial designs
fracfact	Two-level fractional factorial design
fullfact	Mixed level full factorial designs
hadamard	Hadamard designs (in MATLAB)
rowexch	D-optimal design using row exchange

Multivariate Statistics

Cluster Analysis

cluster	Create clusters from linkage output
clusterdata	Create clusters from a dataset
cophenet	Calculate the cophenetic correlation coefficient
dendrogram	Plot a hierarchical tree in a dendrogram graph

inconsistent	Calculate the inconsistency values of objects in a cluster hierarchy tree
kmeans	K-means clustering
linkage	Link objects in a dataset into a hierarchical tree of binary clusters
pdist	Calculate the pairwise distance between objects in a dataset
silhouette	Silhouette plot for clustered data
squareform	Reformat output of pdist function from vector to square matrix

Dimension Reduction Techniques

factoran	Maximum Likelihood Common Factor Analysis
pcacov	PCA from covariance matrix
pcares	Residuals from PCA
princomp	PCA from raw data matrix

Other Multivariate Methods

barttest	Bartlett's test
canoncorr	Canonical correlation analysis
classify	Discriminant Analysis
cmdscale	Classical multidimensional scaling
mahal	Mahalanobis distance
manova1	One-way multivariate analysis of variance
manovacluster	Draw clusters of group means for manova1
procrustes	Procrustes Analysis
zscore	Normalize a dataset before calculating the distance

Decision Tree Techniques

treefit	Fit a tree-based model for classification or regression.
treeprune	Produce a sequence of subtrees by pruning.
treedisp	Show classification or regression tree graphically.
treetest	Compute error rate for tree.
treeval	Compute fitted value for decision tree applied to data.

Hypothesis Tests

ranksum	Wilcoxon rank sum test
signrank	Wilcoxon signed rank test
signtest	Sign test for paired samples
ttest	One sample t-test
ttest2	Two sample t-test
ztest	Z-test

Distribution Testing

jbtest	Jarque-Bera test of normality
kstest	Kolmogorov-Smirnov test for one sample
kstest2	Kolmogorov-Smirnov test for two samples
lillietest	Lilliefors test of normality

Nonparametric Testing

friedman	Friedman's test (nonparametric two-way anova)
kruskalwallis	Kruskal-Wallis test (nonparametric one-way anova)
ksdensity	Probability density estimate using a kernel smoothing method
ranksum	Wilcoxon rank sum test (independent samples)

signrank	Wilcoxon sign rank test (paired samples)
signtest	Sign test (paired samples)

File I/O

caseread	Read casenames from a file
casewrite	Write casenames from a string matrix to a file
tblread	Retrieve tabular data from the file system
tblwrite	Write data in tabular form to the file system
tdfread	Read in text and numeric data from tab-delimited file

Demonstrations

aoctool	Interactive tool for analysis of covariance
disttool	Interactive exploration of distribution functions
glmdemo	Generalized linear model slide show
randtool	Interactive random number generation
polytool	Interactive fitting of polynomial models
rsmdemo	Interactive process experimentation and analysis
robustdemo	Interactive tool to compare robust and least squares fits

Data

carbig.mat	Measurements on 406 car models
carsmall.mat	Measurements on 100 car models from 1970, 1976, and 1982
census.mat	U.S. Population 1790 to 1980
cities.mat	Names of U.S. metropolitan areas
discrim.mat	Classification data
gas.mat	Gasoline prices

hald.mat	Hald data
hogg.mat	Bacteria counts from milk shipments
lawdata.mat	GPA versus LSAT for 15 law schools
mileage.mat	Mileage data for three car models from two factories
moore.mat	Five factor – one response regression data
parts.mat	Dimensional runout on 36 circular parts
popcorn.mat	Data for popcorn example (anova2, friedman)
polydata.mat	Data for polytool demo
reaction.mat	Reaction kinetics data
sat.dat	ASCII data for tblread example

Functions – Alphabetical List

This section contains function reference pages listed alphabetically. The reference pages contain detailed descriptions of the Statistics Toolbox functions.

Purpose	One-way Analysis of Variance (ANOVA)				
Syntax					
	<pre>p = anova1(X) p = anova1(X,group) p = anova1(X,group,'displayopt') [p,table] = anova1() [p,table,stats] = anova1()</pre>				
Description	p = anova1(X) performs a balanced one-way ANOVA for comparing the means of two or more columns of data in the <i>m</i> -by- <i>n</i> matrix X, where each column represents an independent sample containing <i>m</i> mutually independent observations. The function returns the p-value for the null hypothesis that all samples in X are drawn from the same population (or from different populations with the same mean).				
	If the p-value is near zero, this casts doubt on the null hypothesis and suggests that at least one sample mean is significantly different than the other sample means. The choice of a critical p-value to determine whether the result is judged "statistically significant" is left to the researcher. It is common to declare a result significant if the p-value is less than 0.05 or 0.01.				
	The anova1 function displays two figures. The first figure is the standard ANOVA table, which divides the variability of the data in X into two parts:				
	• Variability due to the differences among the column means (variability <i>between</i> groups)				
	• Variability due to the differences between the data in each column and the column mean (variability <i>within</i> groups)				
	The ANOVA table has six columns:				
	• The first shows the source of the variability.				
	• The second shows the Sum of Squares (SS) due to each source.				
	• The third shows the degrees of freedom (df) associated with each source.				
	• The fourth shows the Mean Squares (MS) for each source, which is the ratio SS/df.				
	• The fifth shows the <i>F</i> statistic, which is the ratio of the MS's.				

• The sixth shows the p-value, which is derived from the cdf of *F*. As *F* increases, the p-value decreases.

The second figure displays box plots of each column of X. Large differences in the center lines of the box plots correspond to large values of F and correspondingly small p-values.

p = anova1(X,group) uses the values in group (a character array or cell array) as labels for the box plot of the samples in X, when X is a matrix. Each row of group contains the label for the data in the corresponding column of X, so group must have length equal to the number of columns in X.

When X is a vector, anoval performs a one-way ANOVA on the samples contained in X, as indexed by input group (a vector, character array, or cell array). Each element in group identifies the group (i.e., sample) to which the corresponding element in vector X belongs, so group must have the same length as X. The labels contained in group are also used to annotate the box plot. The vector-input form of anoval does not require equal numbers of observations in each sample, so it is appropriate for unbalanced data.

It is not necessary to label samples sequentially (1, 2, 3, ...). For example, if X contains measurements taken at three different temperatures, -27°, 65°, and 110°, you could use these numbers as the sample labels in group. If a row of group contains an empty cell or empty string, that row and the corresponding observation in X are disregarded. NaNs in either input are similarly ignored.

p = anova1(X,group, 'displayopt') enables the ANOVA table and box plot displays when 'displayopt' is 'on' (default) and suppresses the displays when 'displayopt' is 'off'.

[p,table] = anova1(...) returns the ANOVA table (including column and row labels) in cell array table. (Copy a text version of the ANOVA table to the clipboard by using the **Copy Text** item on the **Edit** menu.)

[p,table,stats] = anova1(...) returns a stats structure that you can use to perform a follow-up multiple comparison test. The anova1 test evaluates the hypothesis that the samples all have the same mean against the alternative that the means are not all the same. Sometimes it is preferable to perform a test to determine *which pairs* of means are significantly different, and which are not. Use the multcompare function to perform such tests by supplying the stats structure as input.

Assumptions

The ANOVA test makes the following assumptions about the data in X:

- All sample populations are normally distributed.
- All sample populations have equal variance.
- All observations are mutually independent.

The ANOVA test is known to be robust to modest violations of the first two assumptions.

Examples

Example 1

The five columns of X are the constants one through five plus a random normal disturbance with mean zero and standard deviation one.

X = meshgrid(1:5) X =

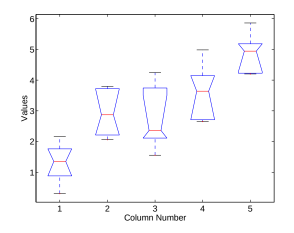
	1 1 1 1	2 2 2 2 2	3 3 3 3 3	4 4 4 4 4	5 5 5 5 5		
X =	X + noi	rmrnc	1(0,1,5	,5)			
X =							
p =	2.1650 1.6268 1.0751 1.3516 0.3035 anova1		3.6961 2.0591 3.7971 2.2641 2.8717	2.2 4.2 2.3	5538 2988 2460 3610 5774	3.6400 3.8644 2.6507 2.7296 4.9846	4.9551 4.2011 4.2348 5.8617 4.9438

p =

5.9952e-005

anova 1

ANOVA Table					
Source Columns Error Total	SS 32.93 14.62 47.55	df 4 20 24	MS 8.232 0.7312	F Prob>F 11.26 5.995e-005	



The very small p-value of 6e-5 indicates that differences between the column means are highly significant. The probability of this outcome under the null hypothesis (i.e., the probability that samples actually drawn from the same population would have means differing by the amounts seen in X) is less than 6 in 100,000. The test therefore strongly supports the alternate hypothesis, that one or more of the samples are drawn from populations with different means.

Example 2

The following example comes from a study of the material strength of structural beams in Hogg (1987). The vector strength measures the deflection of a beam in thousandths of an inch under 3,000 pounds of force. Stronger beams deflect less. The civil engineer performing the study wanted to determine whether the strength of steel beams was equal to the strength of two more expensive alloys. Steel is coded 'st' in the vector alloy. The other materials are coded 'all' and 'al2'.

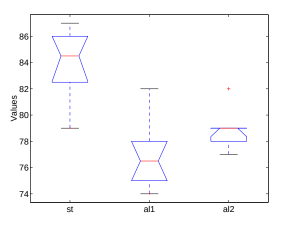
Though alloy is sorted in this example, you do not need to sort the grouping variable.

```
p = anova1(strength,alloy)
```

p =

1.5264e-004

ANOVA Table					
Source Groups Error Total	SS 184.8 102 286.8	df 2 17 19	MS 92.4 6	F 15.4	Prob>F 0.0001526



The p-value indicates that the three alloys are significantly different. The box plot confirms this graphically and shows that the steel beams deflect more than the more expensive alloys.

References	[1] Hogg, R. V., and J. Ledolter. Engineering Statistics. MacMillan Publishing
	Company, 1987.

See Also anova2, anovan, boxplot, ttest

Purpose	Two-way Analysis of Variance (ANOVA)
Syntax	p = anova2(X,reps)
	p = anova2(X,reps,' <i>displayopt</i> ')
	[p,table] = anova2()
	[p,table,stats] = anova2()
Description	anova2(X, reps) performs a balanced two-way ANOVA for comparing the means of two or more columns and two or more rows of the observations in The data in different columns represent changes in factor A. The data in

means of two or more columns and two or more rows of the observations in X. The data in different columns represent changes in factor A. The data in different rows represent changes in factor B. If there is more than one observation for each combination of factors, input reps indicates the number of replicates in each "cell," which much be constant. (For unbalanced designs, use anovan.)

The matrix below shows the format for a set-up where column factor A has two levels, row factor B has three levels, and there are two replications (reps=2). The subscripts indicate row, column, and replicate, respectively.

$$\begin{bmatrix} x & 111 & x_{121} \\ x_{112} & x_{122} \\ x_{211} & x_{221} \\ x_{212} & x_{222} \\ x_{311} & x_{321} \\ x_{312} & x_{322} \end{bmatrix}$$
 B = 3

When reps is 1 (default), anova2 returns two p-values in vector p:

- 1 The p-value for the null hypothesis, H_{0A} , that all samples from factor A (i.e., all column-samples in X) are drawn from the same population

When reps is greater than 1, anova2 returns a third p-value in vector p:

3 The p-value for the null hypothesis, H_{0AB} , that the effects due to factors A and B are *additive* (i.e., that there is no interaction between factors A and B)

If any p-value is near zero, this casts doubt on the associated null hypothesis. A sufficiently small p-value for H_{0A} suggests that at least one column-sample mean is significantly different that the other column-sample means; i.e., there is a main effect due to factor A. A sufficiently small p-value for H_{0B} suggests that at least one row-sample mean is significantly different than the other row-sample means; i.e., there is a main effect due to factor B. A sufficiently small p-value for H_{0AB} suggests that there is an interaction between factors A and B. The choice of a limit for the p-value to determine whether a result is "statistically significant" is left to the researcher. It is common to declare a result significant if the p-value is less than 0.05 or 0.01.

anova2 also displays a figure showing the standard ANOVA table, which divides the variability of the data in X into three or four parts depending on the value of reps:

- The variability due to the differences among the column means
- The variability due to the differences among the row means
- The variability due to the interaction between rows and columns (if reps is greater than its default value of one)
- The remaining variability not explained by any systematic source

The ANOVA table has five columns:

- The first shows the source of the variability.
- The second shows the Sum of Squares (SS) due to each source.
- The third shows the degrees of freedom (df) associated with each source.
- The fourth shows the Mean Squares (MS), which is the ratio SS/df.
- The fifth shows the F statistics, which is the ratio of the mean squares.

p = anova2(X,reps,'displayopt') enables the ANOVA table display when 'displayopt' is 'on' (default) and suppresses the display when 'displayopt' is 'off'. [p,table] = anova2(...) returns the ANOVA table (including column and row labels) in cell array table. (Copy a text version of the ANOVA table to the clipboard by using the Copy Text item on the Edit menu.)
 [p,table,stats] = anova2(...) returns a stats structure that you can use to perform a follow-up multiple comparison test.
 The anova2 test evaluates the hypothesis that the row, column, and interaction effects are all the same, against the alternative that they are not all the same. Sometimes it is preferable to perform a test to determine *which pairs* of effects are significantly different, and which are not. Use the multcompare function to perform such tests by supplying the stats structure as input.
 Examples
 The data below come from a study of popcorn brands and popper type (Hogg 1987). The columns of the matrix popcorn are brands (Gourmet, National, and Generic). The rows are popper type (Oil and Air.) The study popped a batch of

1987). The columns of the matrix popcorn are brands (Gourmet, National, and Generic). The rows are popper type (Oil and Air.) The study popped a batch of each brand three times with each popper. The values are the yield in cups of popped popcorn.

load popcorn					
popcorn					
popcorn =					
	5.5000	4.5000	3.5000		
	5.5000	4.5000	4.0000		
	6.0000	4.0000	3.0000		
	6.5000	5.0000	4.0000		
	7.0000	5.5000	5.0000		
	7.0000	5.0000	4.5000		
p = anova2(popcorn,3)					
р	=				
	0.0000	0.0001	0.7462		

ANOVA Table					
Source Columns Rows Interaction Error Total	SS 15.75 4.5 0.08333 1.667 22	df 2 1 2 12 17	MS 7.875 4.5 0.04167 0.1389	F 56.7 32.4 0.3	

The vector p shows the p-values for the three brands of popcorn, 0.0000, the two popper types, 0.0001, and the interaction between brand and popper type, 0.7462. These values indicate that both popcorn brand and popper type affect the yield of popcorn, but there is no evidence of a synergistic (interaction) effect of the two.

The conclusion is that you can get the greatest yield using the Gourmet brand and an Air popper (the three values popcorn(4:6,1)).

Reference [1] Hogg, R. V. and J. Ledolter. *Engineering Statistics*. MacMillan Publishing Company, 1987.

See Also anova1, anovan

Purpose	N-way Analysis of Variance (ANOVA)
Syntax	<pre>p = anovan(X,group) p = anovan(X,group, 'model') p = anovan(X,group, 'model',sstype) p = anovan(X,group, 'model',sstype,gnames) p = anovan(X,group, 'model',sstype,gnames, 'displayopt') [p,table] = anovan() [p,table,stats] = anovan() [p,table,stats,terms] = anovan()</pre>
Description	<pre>p = anovan(X,group) performs a balanced or unbalanced multi-way ANOVA for comparing the means of the observations in vector X with respect to N different factors. The factors and factor levels of the observations in X are</pre>

assigned by the cell array group. Each of the N cells in group contains a list of factor levels identifying the observations in X with respect to one of the N factors. The list within each cell can be a vector, character array, or cell array of strings, and must have the same number of elements as X.

As an example, consider the X and group inputs below.

```
X = [x1 x2 x3 x4 x5 x6 x7 x8];
group = {[1 2 1 2 1 2 1 2];...
['hi';'hi';'lo';'lo';'hi';'hi';'lo';'lo'];...
{'may' 'may' 'may' 'may' 'june' 'june' 'june' 'june'};
```

In this case, anovan(X, group) is a three-way ANOVA with two levels of each factor. Every observation in X is identified by a combination of factor levels in group. If the factors are A, B, and C, then observation x1 is associated with:

- Level 1 of factor A
- Level 'hi' of factor B
- Level 'may' of factor C

Similarly, observation x6 is associated with:

- Level 2 of factor A
- Level 'hi' of factor B
- \bullet Level 'june' of factor C

Output vector p contains p-values for the null hypotheses on the N main effects. Element p(1) contains the p-value for the null hypotheses, H_{0A} , that samples at all levels of factor A are drawn from the same population, element p(2) contains the p-value for the null hypotheses, H_{0B} , that samples at all levels of factor B are drawn from the same population, and so on.

If any p-value is near zero, this casts doubt on the associated null hypothesis. For example, a sufficiently small p-value for H_{0A} suggests that at least one A-sample mean is significantly different that the other A-sample means; i.e., there is a main effect due to factor A. The choice of a limit for the p-value to determine whether a result is "statistically significant" is left to the researcher. It is common to declare a result significant if the p-value is less than 0.05 or 0.01.

anovan also displays a figure showing the standard ANOVA table, which by default divides the variability of the data in X into:

- The variability due to differences between the levels of each factor accounted for in the model (one row for each factor)
- The remaining variability not explained by any systematic source

The ANOVA table has six columns:

- The first shows the source of the variability.
- The second shows the Sum of Squares (SS) due to each source.
- The third shows the degrees of freedom (df) associated with each source.
- The fourth shows the Mean Squares (MS), which is the ratio SS/df.
- The fifth shows the F statistics, which is the ratio of the mean squares.
- The sixth shows the p-values for the F statistics.

p = anovan(X,group, 'model') performs the ANOVA using the model specified by 'model', where 'model' can be 'linear', 'interaction', 'full', or an integer or vector. The default 'linear' model computes only the p-values for the null hypotheses on the N main effects. The 'interaction' model computes the p-values for null hypotheses on the N main effects and the $\binom{N}{2}$ two-factor interactions. The 'full' model computes the p-values for null hypotheses on the N main effects and interactions at all levels. For an integer value of 'model', k (k \leq N), anovan computes all interaction levels through the kth level. The values k=1 and k=2 are equivalent to the 'linear' and 'interaction' specifications, respectively, while the value k=N is equivalent to the 'full' specification.

For more precise control over the main and interaction terms that anovan computes, 'model' can specify a vector containing one element for each main or interaction term to include in the ANOVA model. Each vector element encodes the corresponding ANOVA term as the decimal equivalent of an N-bit number, where N is the number of factors. The table below illustrates the coding for a 3-factor ANOVA.

3-bit Code	Decimal Value	Corresponding ANOVA Terms
[0 0 1]	1	Main term A
[0 1 0]	2	Main term B
[1 0 0]	4	Main term C
[0 1 1]	3	Interaction term AB
[1 1 0]	6	Interaction term BC
[1 0 1]	5	Interaction term AC
[1 1 1]	7	Interaction term ABC

For example, if 'model' is the vector [2 4 6], then output vector p contains the p-values for the null hypotheses on the main effects B and C and the interaction effect BC, in that order. A simple way to generate the 'model' vector is to modify the terms output, which codes the terms in the current model using the format described above. If anovan returned [2 4 6] for terms, for example, and there was no significant result for interaction BC, you could recompute the ANOVA on just the main effects B and C by specifying [2 4] for 'model'.

p = anovan(X,group, 'model',sstype) computes the ANOVA using the type
of sum-of-squares specified by sstype, which can be 1, 2, or 3 to designate
Type 1, Type 2, or Type 3 sum-of-squares, respectively. The default is 3. The
value of sstype only influences computations on unbalanced data.

The sum of squares for any term is determined by comparing two models. The Type 1 sum of squares for a term is the reduction in residual sum of squares obtained by adding that term to a fit that already includes the terms listed before it. The Type 2 sum of squares is the reduction in residual sum of squares obtained by adding that term to a model consisting of all other terms that do not contain the term in question. The Type 3 sum of squares is the reduction in residual sum of squares obtained by adding that term to a model consisting of squares is the reduction in residual sum of squares obtained by adding that term to a model contain the terms in question. The Type 3 sum of squares is the reduction in residual sum of squares obtained by adding that term to a model containing all other terms, but with their effects constrained to obey the usual "sigma restrictions" that make models estimable.

Suppose we are fitting a model with two factors and their interaction, and that the terms appear in the order A, B, AB. Let $R(\cdot)$ represent the residual sum of squares for a model, so for example R(A,B,AB) is the residual sum of squares fitting the whole model, R(A) is the residual sum of squares fitting just the main effect of A, and R(1) is the residual sum of squares fitting just the mean. The three types of sums of squares are as follows:

Term	Type 1 SS	Type 2 SS	Type 3 SS
А	R(1)-R(A)	R(B)-R(A,B)	R(B,AB)-R(A,B,AB)
В	R(A)- $R(A,B)$	R(A)-R(A,B)	R(A,AB)-R(A,B,AB)
AB	R(A,B)- $R(A,B,AB)$	R(A,B)- $R(A,B,AB)$	R(A,B)- $R(A,B,AB)$

Table 12-1:

The models for Type 3 sum of squares have sigma restrictions imposed. This means, for example, that in fitting R(B,AB), the array of AB effects is constrained to sum to 0 over A for each value of B, and over B for each value of A.

p = anovan(X,group, 'model', sstype,gnames) uses the string values in character array gnames to label the N experimental factors in the ANOVA table. The array can be a string matrix with one row per observation, or a cell array of strings with one element per observation. When gnames is not specified, the default labels 'X1', 'X2', 'X3', ..., 'XN' are used.

p = anovan(X,group, 'model',sstype,gnames, 'displayopt') enables the ANOVA table display when 'displayopt' is 'on' (default) and suppresses the display when 'displayopt' is 'off'.

[p,table] = anovan(...) returns the ANOVA table (including factor labels) in cell array table. (Copy a text version of the ANOVA table to the clipboard by using the **Copy Text** item on the **Edit** menu.)

[p,table,stats] = anovan(...) returns a stats structure that you can use to perform a follow-up multiple comparison test.

The anovan test evaluates the hypothesis that the different levels of a factor (or more generally, a term) have the same effect, against the alternative that they do not all have the same effect. Sometimes it is preferable to perform a test to determine *which pairs* of levels are significantly different, and which are not. Use the multcompare function to perform such tests by supplying the stats structure as input.

[p,table,stats,terms] = anovan(...) returns the main and interaction terms used in the ANOVA computations. The terms are encoded in output vector terms using the same format described above for input 'model'. When 'model' itself is specified in this vector format, the vector returned in terms is identical.

Examples

In the previous section we used anova2 to analyze the effects of two factors on a response in a balanced design. For a design that is not balanced, we can use anovan instead.

The dataset carbig contains a number of measurements on 406 cars. Let's study how the mileage depends on where and when the cars were made.

```
load carbig
anovan(MPG,{org when},2,3,{'Origin';'Mfg date'})
ans =
0
0
0.30587
```

The p-value for the interaction term is not small, indicating little evidence that the effect of the car's year or manufacture (when) depends on where the car was made (org). The linear effects of those two factors, though, are significant.

Figure No. 1: N-Way Al	AVOVA				_ [
e <u>E</u> dit <u>V</u> iew <u>I</u> nsert <u>T</u> o	ols <u>W</u> indow <u>H</u>	<u>l</u> elp			
	Ana	lysis of	Variance		
Source	Sum Sq.	d.f.	Mean Sq.	F	Prob>F
Origin Mfg date Origin*Mfg date Error Total	5727.2 4710.3 120.5 9679.1 24252.6	2 2 4 389 397	2863.58 2355.15 30.12 24.88	115.09 94.65 1.21	0 0 0.3059
	Constrain	ed (Type II	I) sums of squares.		

- **Reference** [1] Hogg, R. V. and J. Ledolter. *Engineering Statistics*. MacMillan Publishing Company, 1987.
- See Also anova1, anova2, multcompare

Purpose	Interactive plot for fitting and predicting analysis of covariance models
Syntax	<pre>aoctool(x,y,g) aoctool(x,y,g,alpha) aoctool(x,y,g,alpha,xname,yname,gname) aoctool(x,y,g,alpha,xname,yname,gname,'displayopt') aoctool(x,y,g,alpha,xname,yname,gname,'displayopt','model') h = aoctool() [h,atab,ctab] = aoctool() [h,atab,ctab,stats] = aoctool()</pre>
Description	aoctool(x,y,g) fits a separate line to the column vectors, x and y, for each group defined by the values in the array g. These types of models are known as one-way analysis of covariance (ANOCOVA) models. The output consists of three figures:
	• An interactive graph of the data and prediction curves
	 An ANOVA table A table of parameter estimates
	You can use the figures to change models and to test different parts of the model. More information about interactive use of the aoctool function appears on "The aoctool Demo" on page 11-10.
	aoctool(x,y,g,alpha) determines the confidence levels of the prediction intervals. The confidence level is 100*(1-alpha)%. The default value of alpha is 0.05.
	aoctool(x,y,g,alpha,xname,yname,gname) specifies the name to use for the x, y, and g variables in the graph and tables. If you enter simple variable names for the x, y, and g arguments, the aoctool function uses those names. If you enter an expression for one of these arguments, you can specify a name to use in place of that expression by supplying these arguments. For example, if you enter m(:,2) as the x argument, you might choose to enter 'Col 2' as the xname argument.
	aoctool(x,y,g,alpha,xname,yname,gname,' <i>displayopt</i> ') enables the graph and table displays when ' <i>displayopt</i> ' is 'on' (default) and suppresses those displays when ' <i>displayopt</i> ' is 'off'.

aoctool(x,y,g,alpha,xname,yname,gname,'displayopt','model')
specifies the initial model to fit. The value of 'model' can be any of the
following:

- 'same mean' fit a single mean, ignoring grouping
- 'separate means' fit a separate mean to each group
- 'same line' fit a single line, ignoring grouping
- 'parallel lines' fit a separate line to each group, but constrain the lines to be parallel
- 'separate lines' fit a separate line to each group, with no constraints
- h = aoctool(...) returns a vector of handles to the line objects in the plot.

[h,atab,ctab] = aoctool(...) returns cell arrays containing the entries in ANOVA table (atab) and the table of coefficient estimates (ctab). (You can copy a text version of either table to the clipboard by using the **Copy Text** item on the **Edit** menu.)

[h,atab,ctab,stats] = aoctool(...) returns a stats structure that you can use to perform a follow-up multiple comparison test. The ANOVA table output includes tests of the hypotheses that the slopes or intercepts are all the same, against a general alternative that they are not all the same. Sometimes it is preferable to perform a test to determine which pairs of values are significantly different, and which are not. You can use the multcompare function to perform such tests by supplying the stats structure as input. You can test either the slopes, the intercepts, or population marginal means (the heights of the curves at the mean x value).

Example

This example illustrates how to fit different models non-interactively. First, we load the smaller car dataset and fit a separate-slopes model, then examine the coefficient estimates.

' 70'	[-8.58050531454973]
' 76'	[-3.89017396094922]
' 82'	[12.47067927549897]
'Slope'	[-0.00780212907455]
' 70'	[0.00195840368824]
' 76'	[0.00113831038418]
' 82'	[-0.00309671407243]

Roughly speaking, the lines relating MPG to Weight have an intercept close to 45.98 and a slope close to -0.0078. Each group's coefficients are offset from these values somewhat. For instance, the intercept for the cars made in 1970 is 45.98-8.58 = 37.40.

Next, we try a fit using parallel lines. (If we had examined the ANOVA table, we would have found that the parallel-lines fit is significantly worse than the separate-lines fit.)

```
[h,a,c,s] = aoctool(Weight,MPG,Model_Year,0.05,...
                       '','','','off','parallel lines');
c(:,1:2)
ans =
    'Term'
                    'Estimate'
    'Intercept'
                    [43.38984085130596]
       70'
                    [-3.27948192983761]
    I.
       76'
                    [-1.35036234809006]
    ı.
       82 '
                    [ 4.62984427792768]
    'Slope'
                    [-0.00664751826198]
```

Here we again have separate intercepts for each group, but this time the slopes are constrained to be the same.

See Also anova1, multcompare, polytool

barttest

Purpose	Bartlett's test for dimensionality
Syntax	ndim = barttest(x,alpha) [ndim,prob,chisquare] = barttest(x,alpha)
Description	<pre>ndim = barttest(x,alpha) returns the number of dimensions necessary to explain the nonrandom variation in the data matrix x, using the significance probability alpha. The dimension is determined by a series of hypothesis tests. The test for ndim=1 tests the hypothesis that the variances of the data values along each principal component are equal, the test for ndim=2 tests the hypothesis that the variances along the second through last components are equal, and so on.</pre>
	[ndim,prob,chisquare] = barttest(x,alpha) returns the number of dimensions, the significance values for the hypothesis tests, and the χ^2 values associated with the tests.
Example	<pre>x = mvnrnd([0 0],[1 0.99; 0.99 1],20); x(:,3:4) = mvnrnd([0 0],[1 0.99; 0.99 1],20); x(:,5:6) = mvnrnd([0 0],[1 0.99; 0.99 1],20); [ndim, prob] = barttest(x,0.05)</pre>
	ndim =
	3
	prob = 0 0 0.5081 0.6618
See Also	princomp, pcacov, pcares

Purpose	Generate Box-Behnken design
Syntax	D = bbdesign(nfactors) D = bbdesign(nfactors,'pname1',pvalue1,'pname2',pvalue2,) [D,blk] = bbdesign()
Description	D = bbdesign(nfactors) generates a Box-Behnken design for nfactors factors. The output matrix D is n-by-nfactors, where n is the number of points in the design. Each row lists the settings for all factors, scaled between -1 and 1.
	<pre>[D,blk] = bbdesign(nfactors) requests a blocked design. The output vector blk is a vector of block numbers. Blocks are groups of runs that are to be measured under similar conditions (for example, on the same day). Blocked designs minimize the effect of between-block differences on the parameter estimates.</pre>
	<pre>[] = bbdesign(nfactors, 'pname1', pvalue1, 'pname2', pvalue2,) allows you to specify additional parameters and their values. Valid parameters are:</pre>
	'center' Number of center points to include.
	'blocksize' Maximum number of points allowed in a block.
Remarks	Box and Behnken proposed designs when the number of factors was equal to 3-7, 9-12, or 16. This function produces those designs. For other values of nfactors, this function produces designs that are constructed in a similar way, even though they were not tabulated by Box and Behnken, and they may be too large to be practical.
See Also	ccdesign, cordexch, rowexch

betacdf

Purpose	Beta cumulative distribution function (cdf)
Syntax	<pre>p = betacdf(X,A,B)</pre>
Description	p = betacdf(X,A,B) computes the beta cdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must all be positive, and the values in X must lie on the interval [0,1].
	The beta cdf for a given value \boldsymbol{x} and given pair of parameters \boldsymbol{a} and \boldsymbol{b} is
	$p = F(x a, b) = \frac{1}{B(a, b)} \int_0^x t^{a-1} (1-t)^{b-1} dt$
	where $B(\cdot)$ is the Beta function. The result, p, is the probability that a single observation from a beta distribution with parameters a and b will fall in the interval [0,x].
Examples	<pre>x = 0.1:0.2:0.9; a = 2; b = 2; p = betacdf(x,a,b)</pre>
	p =
	0.0280 0.2160 0.5000 0.7840 0.9720
	a = [1 2 3]; p = betacdf(0.5,a,a)
	p =
	0.5000 0.5000 0.5000
• • •	

See Also betafit, betainv, betalike, betapdf, betarnd, betastat, cdf

Purpose Parameter estimates and confidence intervals for beta distributed data

Syntax phat = betafit(x)
[phat,pci] = betafit(x,alpha)

Description phat = betafit(x) computes the maximum likelihood estimates of the beta distribution parameters a and b from the data in vector x, where the beta cdf is given by

$$F(x|a,b) = \frac{1}{B(a,b)} \int_0^x t^{a-1} (1-t)^{b-1} dt$$

and $B(\cdot)$ is the Beta function. The elements of x must lie in the interval (0 1).

[phat,pci] = betafit(x,alpha) returns confidence intervals on the *a* and *b* parameters in the 2-by-2 matrix pci. The first column of the matrix contains the lower and upper confidence bounds for parameter *a*, and the second column contains the confidence bounds for parameter *b*. The optional input argument alpha is a value in the range $[0 \ 1]$ specifying the width of the confidence intervals. By default, alpha is 0.05, which corresponds to 95% confidence intervals.

Example

This example generates 100 beta distributed observations. The true a and b parameters are 4 and 3, respectively. Compare these to the values returned in p. Note that the columns of ci both bracket the true parameters.

Reference [1] Hahn, Gerald J., & Shapiro, Samuel, S. *Statistical Models in Engineering*. John Wiley & Sons, New York. 1994. p. 95.

betafit

See Also betalike, mle

Purpose Inverse of the beta cumulative distribution function

Syntax X = betainv(P,A,B)

Description X = betainv(P,A,B) computes the inverse of the beta cdf with parameters specified by A and B for the corresponding probabilities in P. Vector or matrix inputs for P, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must all be positive, and the values in P must lie on the interval [0 1].

The inverse beta cdf for a given probability p and a given pair of parameters a and b is

$$x = F^{-1}(p|a, b) = \{x: F(x|a, b) = p\}$$

where

$$p = F(x|a,b) = \frac{1}{B(a,b)} \int_0^x t^{a-1} (1-t)^{b-1} dt$$

and $B(\cdot)$ is the Beta function. Each element of output X is the value whose cumulative probability under the beta cdf defined by the corresponding parameters in A and B is specified by the corresponding value in P.

AlgorithmThe betainv function uses Newton's method with modifications to constrain
steps to the allowable range for x, i.e., [0 1].

Examples

p = [0.01 0.5 0.99]; x = betainv(p,10,5) x = 0.3726 0.6742 0.8981

According to this result, for a beta cdf with a=10 and b=5, a value less than or equal to 0.3726 occurs with probability 0.01. Similarly, values less than or equal to 0.6742 and 0.8981 occur with respective probabilities 0.5 and 0.99.

See Also betafit, icdf

betalike

Purpose	Negative beta log-likelihood function
Syntax	logL = betalike(params,data) [logL,avar] = betalike(params,data)
Description	logL = betalike(params,data) returns the negative of the beta log-likelihood function for the beta parameters a and b specified in vector params and the observations specified in column vector data. The length of logL is the length of data.
	<pre>[logL,avar] = betalike(params,data) also returns avar, which is the asymptotic variance-covariance matrix of the parameter estimates if the values in params are the maximum likelihood estimates. avar is the inverse of Fisher's information matrix. The diagonal elements of avar are the asymptotic variances of their respective parameters.</pre>
	betalike is a utility function for maximum likelihood estimation of the beta distribution. The likelihood assumes that all the elements in the data sample are mutually independent. Since betalike returns the negative beta log-likelihood function, minimizing betalike using fminsearch is the same as maximizing the likelihood.
Example	This example continues the betafit example where we calculated estimates of the beta parameters for some randomly generated beta distributed data.
	r = betarnd(4,3,100,1); [logl,avar] = betalike([3.9010 2.6193],r)
	logl =
	-33.0514
	avar =
	0.2856 0.1528 0.1528 0.1142
See Also	betafit, fminsearch, gamlike, mle, normlike, weiblike

betapdf

Purpose	Beta probability density function (pdf)
Syntax	Y = betapdf(X,A,B)
Description	Y = betapdf(X,A,B) computes the beta pdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions of the other inputs. The parameters in A and B must all be positive, and the values in X must lie on the interval [0 1].
	The beta probability density function for a given value x and given pair of parameters a and b is
	$y = f(x a,b) = \frac{1}{B(a,b)} x^{a-1} (1-x)^{b-1} I_{(0,1)}(x)$
	where $B(\cdot)$ is the Beta function. The result, y , is the probability that a single observation from a beta distribution with parameters a and b will have value x . The indicator function $I_{(0,1)}(x)$ ensures that only values of x in the range (0 1) have nonzero probability. The uniform distribution on (0 1) is a degenerate case of the beta pdf where $a = 1$ and $b = 1$.
	A <i>likelihood function</i> is the pdf viewed as a function of the parameters. Maximum likelihood estimators (MLEs) are the values of the parameters that maximize the likelihood function for a fixed value of x .
Examples	a = [0.5 1; 2 4]
	a =
	0.5000 1.0000 2.0000 4.0000
	y = betapdf(0.5,a,a)
	y = 0.6366 1.0000 1.5000 2.1875

See Also betacdf, betafit, betainv, betalike, betarnd, betastat, pdf

betarnd

Purpose	Random numbers from the beta distribution
Syntax	R = betarnd(A,B) R = betarnd(A,B,m) R = betarnd(A,B,m,n)
Description	R = betarnd(A,B) generates random numbers from the beta distribution with parameters specified by A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of R. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input. R = betarnd(A,B,m) generates a matrix of size m containing random numbers from the beta distribution with parameters A and B, where m is a 1-by-2 vector containing the row and column dimensions of R.
	R = betarnd(A,B,m,n) generates an m-by-n matrix containing random numbers from the beta distribution with parameters A and B.
Examples	a = [1 1;2 2]; b = [1 2;1 2];
	r = betarnd(a,b)
	r = 0.6987 0.6139 0.9102 0.8067
	r = betarnd(10,10,[1 5])
	r = 0.5974 0.4777 0.5538 0.5465 0.6327
	r = betarnd(4,2,2,3)
	r =
	0.3943 0.6101 0.5768 0.5990 0.2760 0.5474
See Also	betacdf, betafit, betainv, betalike, betapdf, betastat, rand, randtool

betastat

Purpose	Mean and variance for the beta distribution				
Syntax	[M,V] = betastat(A,B)				
Description	[M,V] = betastat(A,B) returns the mean and variance for the beta distribution with parameters specified by A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of M and V. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input.				
	The mean of the beta distribution with parameters a and b is $a/(a+b)$ and the variance is				
	$\frac{ab}{(a+b+1)(a+b)^2}$				
Examples	<pre>If parameters a and b are equal, the mean is 1/2. a = 1:6; [m,v] = betastat(a,a)</pre>				
	m = 0.5000 0.5000 0.5000 0.5000 0.5000 0.5000				
	v = 0.0833 0.0500 0.0357 0.0278 0.0227 0.0192				
See Also	betacdf, betafit, betainv, betalike, betapdf, betarnd				

binocdf

Purpose	Binomial cumulative distribution function (cdf)			
Syntax	Y = binocdf(X,N,P)			
Description	binocdf(X,N,P) computes a binomial cdf at each of the values in X using the corresponding parameters in N and P. Vector or matrix inputs for X, N, and P must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions of the other inputs. The values in N must all be positive integers, and the values in X and P must lie on the interval [0 1].			
	The binomial cdf for a given value \boldsymbol{x} and given pair of parameters \boldsymbol{n} and \boldsymbol{p} is			
	$y = F(x n,p) = \sum_{i=0}^{x} {n \choose i} p^{i} q^{(n-i)} I_{(0,1,,n)}(i)$			
	The result, y, is the probability of observing up to x successes in n independent trials, where the probability of success in any given trial is p. The indicator function $I_{(0,1, \ldots, n)}(i)$ ensures that x only adopts values of 0, 1,, n.			
Examples	If a baseball team plays 162 games in a season and has a 50-50 chance of winning any game, then the probability of that team winning more than 10 games in a season is:			
	1 - binocdf(100,162,0.5)			
	The result is 0.001 (i.e., 1-0.999). If a team wins 100 or more games in a season, this result suggests that it is likely that the team's true probability of winning any game is greater than 0.5.			
See Also	binofit, binoinv, binopdf, binornd, binostat, cdf			

Purpose	Parameter estimates and confidence intervals for binomial data		
Syntax	phat = binofit(x,n) [phat,pci] = binofit(x,n) [phat,pci] = binofit(x,n,alpha)		
Description	phat = binofit(x,n) returns a maximum likelihood estimate of the probability of success in a <i>given</i> binomial trial based on the number of successes, x, observed in n independent trials. A scalar value for x or n is expanded to the same size as the other input.		
	[phat,pci] = binofit(x,n) returns the probability estimate, phat, and the 95% confidence intervals, pci.		
	<pre>[phat,pci] = binofit(x,n,alpha) returns the 100(1-alpha)% confidence intervals. For example, alpha = 0.01 yields 99% confidence intervals.</pre>		
Example	First we generate a binomial sample of 100 elements, where the probability of success in a given trial is 0.6. Then, we estimate this probability from the outcomes in the sample.		
	r = binornd(100,0.6); [phat,pci] = binofit(r,100)		
	phat =		
	0.5800		
	pci =		
	0.4771 0.6780		
	The 95% confidence interval, pci, contains the true value, 0.6.		
Reference	[1] Johnson, N. L., S. Kotz, and A.W. Kemp, "Univariate Discrete Distributions, Second Edition," Wiley 1992. pp. 124–130.		
See Also	binocdf, binoinv, binopdf, binornd, binostat, mle		

binoinv

Purpose	Inverse of the binomial cumulative distribution function (cdf)		
Syntax	X = binoinv(Y,N,P)		
Description	X = binoinv(Y,N,P) returns the smallest integer X such that the binomial cdf evaluated at X is equal to or exceeds Y. You can think of Y as the probability of observing X successes in N independent trials where P is the probability of success in each trial. Each X is a positive integer less than or equal to N.		
	Vector or matrix inputs for Y, N, and P must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in N must be positive integers, and the values in both P and Y must lie on the interval [0 1].		
Examples	If a baseball team has a 50-50 chance of winning any game, what is a reasonable range of games this team might win over a season of 162 games? W assume that a surprising result is one that occurs by chance once in a decad binoinv([0.05 0.95],162,0.5)		
	ans =		
	71 91		
	This result means that in 90% of baseball seasons, a .500 team should win between 71 and 91 games.		
See Also	binocdf, binofit, binopdf, binornd, binostat, icdf		

Purpose	Binomial probability density function (pdf)				
Syntax	Y = binopdf(X,N,P)				
Description	Y = binopdf(X,N,P) computes the binomial pdf at each of the values in X using the corresponding parameters in N and P. Vector or matrix inputs for N, and P must all have the same size. A scalar input is expanded to a consta- matrix with the same dimensions of the other inputs.				
	The parameters in N must be positive integers, and the values in P must lie on the interval $\begin{bmatrix} 0 & 1 \end{bmatrix}$.				
	The binomial probability density function for a given value x and given pair of parameters n and p is				
	$y = f(x n,p) = {n \choose x} p^{x} q^{(n-x)} I_{(0, 1,, n)}(x)$				
	where $q = 1$ - p . The result, y , is the probability of observing x successes in n independent trials, where the probability of success in any <i>given</i> trial is p . The indicator function $I_{(0,1,,n)}(x)$ ensures that x only adopts values of 0, 1,, n .				
Examples	A Quality Assurance inspector tests 200 circuit boards a day. If 2% of the boards have defects, what is the probability that the inspector will find no defective boards on any given day?				
	binopdf(0,200,0.02) ans =				
	0.0176				
	What is the most likely number of defective boards the inspector will find?				
	y = binopdf([0:200],200,0.02); [x,i] = max(y);				
	i i = 5				
See Also	binocdf, binofit, binoinv, binornd, binostat, pdf				

binornd

Purpose	Random numbers from the binomial distribution			
Syntax	R = binornd(N,P) R = binornd(N,P,mm) R = binornd(N,P,mm,nn)			
Description	 R = binornd(N,P) generates random numbers from the binomial distribution with parameters specified by N and P. Vector or matrix inputs for N and P must have the same size, which is also the size of R. A scalar input for N or P is expanded to a constant matrix with the same dimensions as the other input. R = binornd(N,P,mm) generates a matrix of size mm containing random numbers from the binomial distribution with parameters N and P, where mm is a 1-by-2 vector containing the row and column dimensions of R. R = binornd(N,p,mm,nn) generates an mm-by-nn matrix containing random numbers from the binomial distribution with parameters N and P. 			
Algorithm	The binornd function uses the direct method using the definition of the binomial distribution as a sum of Bernoulli random variables.			
Examples	n = 10:10:60; $r1 = binornd(n,1./n)$ $r1 = 2 1 0 1 1 2$ $r2 = binornd(n,1./n,[1 6])$ $r2 = 0 1 2 1 3 1$ $r3 = binornd(n,1./n,1,6)$ $r3 = 0 1 1 1 0 3$			
See Also	binocdf, binofit, binoinv, binopdf, binostat, rand, randtool			

binostat

Purpose	Mean and variance for the binomial distribution				
Syntax	[M,V] = binostat(N,P)				
Description	[M,V] = binostat(N,P) returns the mean and variance for the binomial distribution with parameters specified by N and P. Vector or matrix inputs for N and P must have the same size, which is also the size of M and V. A scalar input for N or P is expanded to a constant matrix with the same dimensions as the other input.				
	The mean of the binomial distribution with parameters n and p is np . The variance is npq , where $q = 1$ - p .				
Examples	n = logspace(1,5,5) n = 10 100 1000 10000 100000				
	[m,v] = binostat(n,1./n)				
	m = 1 1 1 1 1				
	v = 0.9000 0.9900 0.9990 0.9999 1.0000				
	[m,v] = binostat(n,1/2)				
	m = 5 50 500 5000 50000				
	V =				
	1.0e+04 * 0.0003 0.0025 0.0250 0.2500 2.5000				
See Also	binocdf, binofit, binoinv, binopdf, binornd				

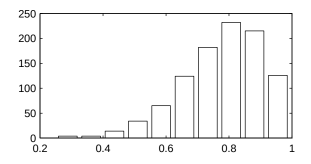
bootstrp

Purpose	Bootstrap statistics through resampling of data				
Syntax	bootstat = bootstrp(nboot,'bootfun',d1,d2,) [bootstat,bootsam] = bootstrp()				
Description	bootstat = bootstrp(nboot, 'bootfun', d1, d2,) draws nboot bootstrap samples from each of the input data sets, d1, d2, etc., and passes the bootstrap samples to function bootfun for analysis. nboot must be a positive integer, and each input data set must contain the same number of rows, n. Each bootstrap sample contains n rows chosen randomly (with replacement) from the corresponding input data set (d1, d2, etc.).				
	Each row of the output, bootstat, contains the results of applying bootfun to one set of bootstrap samples. If bootfun returns multiple outputs, only the first is stored in bootstat. If the first output from bootfun is a matrix, the matrix is reshaped to a row vector for storage in bootstat.				
	[bootstat,bootsam] = bootstrap() returns a matrix of bootstrap indices, bootsam. Each of the nboot columns in bootsam contains indices of the values that were drawn from the original data sets to constitute the corresponding bootstrap sample. For example, if d1, d2, etc., each contain 16 values, and nboot = 4, then bootsam is a 16-by-4 matrix. The first column contains the indices of the 16 values drawn from d1, d2, etc., for the first of the four bootstrap samples, the second column contains the indices for the second of the four bootstrap samples, and so on. (The bootstrap indices are the same for all input data sets.)				
Example	Correlate the LSAT scores and law-school GPA for 15 students. These 15 data points are resampled to create 1000 different data sets, and the correlation between the two variables is computed for each dataset.				
	load lawdata [bootstat,bootsam] = bootstrp(1000,'corrcoef',lsat,gpa);				
	<pre>bootstat(1:5,:)</pre>				
	ans =				
	1.0000 0.3021 0.3021 1.0000 1.0000 0.6869 0.6869 1.0000 1.0000 0.8346 0.8346 1.0000				

bootstrp

	1.0000		0.8711 0.8043		.8711 .8043	1.0000 1.0000
boot	bootsam(:,1:5)					
ans	=					
	4	7	5	12	8	
	1	11	10	8	4	
	11	9	12	4	2	
	11	14	15	5	15	
	15	13	6	6	2	
	6	8	4	3	8	
	8	2	15	8	6	
	13	10	11	14	5	
	1	7	12	14	14	
	1	11	10	1	8	
	8	14	2	14	7	
	11	12	10	8	15	
	1	4	14	8	1	
	6	1	5	5	12	
	2	12	7	15	12	

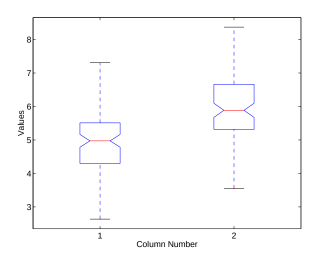
hist(bootstat(:,2))



The histogram shows the variation of the correlation coefficient across all the bootstrap samples. The sample minimum is positive, indicating that the relationship between LSAT score and GPA is not accidental.

boxplot

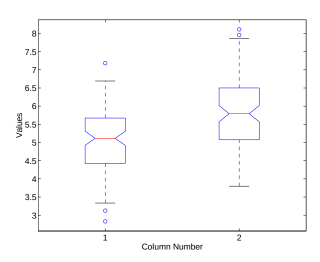
Purpose	Box plots of a data sample		
Syntax	<pre>boxplot(X) boxplot(X,notch) boxplot(X,notch,'sym') boxplot(X,notch,'sym',vert) boxplot(X,notch,'sym',vert,whis)</pre>		
Description	boxplot(X) produces a box and whisker plot for each column of X. The box has lines at the lower quartile, median, and upper quartile values. The whiskers are lines extending from each end of the box to show the extent of the rest of the data. Outliers are data with values beyond the ends of the whiskers. If there is no data outside the whisker, a dot is placed at the bottom whisker.		
	<pre>boxplot(X,notch) with notch = 1 produces a notched-box plot. Notches graph a robust estimate of the uncertainty about the means for box-to-box comparison. The default, notch = 0, produces a rectangular box plot.</pre>		
	boxplot(X,notch,'sym') where sym is a plotting symbol, affords control of the symbol for outliers. The default is '+'. See the MATLAB LineSpec property for information about the available marker symbols.		
	<pre>boxplot(X,notch,'sym',vert) with vert = 0 creates horizontal boxes rather than the default vertical boxes (vert = 1).</pre>		
	boxplot(X,notch, 'sym',vert,whis) enables you to specify the length of the "whiskers." whis defines the maximum length of the whiskers as a function of the inter-quartile range (default = 1.5). Each whisker extends to the most extreme data value within whis * IQR of the box. boxplot displays all data values beyond the whiskers using the plotting symbol, 'sym'. To display all data values outside the box using 'sym', set whis = 0.		
Examples	This example produces box plots for the sample data, and accepts the default $1.5 * IQR$ for the length of the whiskers.		
	<pre>x1 = normrnd(5,1,100,1); x2 = normrnd(6,1,100,1); x = [x1 x2]; boxplot(x,1)</pre>		



The difference between the means of the two columns of x is 1. We can detect this difference graphically by observing that the notches in the boxplot do not overlap.

The following figure shows the boxplot for same data with the length of the whiskers specified as 1.0. Points beyond the whiskers are displayed using 'o'.

boxplot(x,1,'o',[],1.0)



boxplot

See Also anova1, kruskalwallis

Purpose D-optimal design from candidate set using row exchanges

Syntax rlist = candexch(C,nrows) rlist = candexch(C,nrows,'param1',value1,'param2',value2,...)

Description rlist = candexch(C,nrows) uses a row-exchange algorithm to select a D-optimal design from the candidate set C. C is an n-by-p matrix containing the values of p model terms at each of n points. nrows is the desired number of rows in the design. rlist is a vector of length nrows listing the selected rows.

The candexch function selects a starting design X at random, and uses a row-exchange algorithm to iteratively replace rows of X by rows of C in an attempt to improve the determinant of X' * X.

rlist = candexch(C,nrows, 'param1',value1, 'param2',value2,...)
provides more control over the design generation through a set of
parameter/value pairs. Valid parameters are the following:

Parameter	Value
'display'	Either 'on' or 'off' to control display of iteration number The default is 'on'.
'init'	Initial design as an nrows-by-p matrix. The default is a random subset of the rows of C.
'maxiter'	Maximum number of iterations. The default is 10.

Note The rowexch function also generates D-optimal designs using a row-exchange algorithm, but it accepts a model type and automatically selects a candidate set that is appropriate for such a model.

Examples

Generate a D-optimal design when there is a restriction on the candidate set. In this case, the rowexch function isn't appropriate.

F = (fullfact([5 5 5])-1)/4; % Factor settings in unit cube. T = sum(F,2)<=1.51; % Find rows matching a restriction. F = F(T,:); % Take only those rows. C = [ones(size(F,1),1) F F.^2]; % Compute model terms including % a constant and all squared terms. R = candexch(C,12); % Find a D-optimal 12-point subset. X = F(R,:); % Get factor settings.

See Also candgen, cordexch, rowexch, x2fx

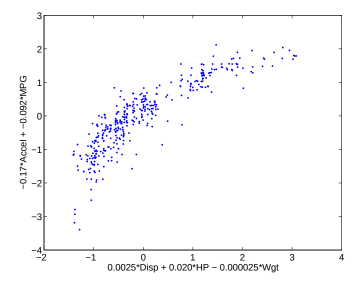
Purpose	Generate candidate set for D-optimal design		
Syntax	<pre>xcand = candgen(nfactors,'model') [xcand,fxcand] = candgen(nfactors,'model')</pre>		
Description	<pre>xcand = candgen(nfactors, 'model') generates a candidate set appropriate for a D-optimal design with nfactors factors and the model model. The output matrix xcand has nfactors columns, with each row representing the coordinates of a candidate point. model is one of:</pre>		
	'linear'	Constant and linear terms (the default)	
	'interaction'	Constant, linear, and cross product terms	
	'quadratic'	Interactions plus squared terms	
	'purequadratic'	Constant, linear, and squared terms	
	Alternatively, model can be a matrix of term definitions as accepted by the x2fx function. [xcand,fxcand] = candgen(nfactors,model) returns both the matrix of factor values xcand and the matrix of term values fxcand. You can input the latter to candexch to generate the D-optimal design. Note The rowexch function automatically generates a candidate set using candgen, and creates a D-optimal design from that candidate set using candexch. Call these functions separately if you want to modify the default candidate set.		
See Also	candexch, rowexch,	,x2fx	

canoncorr

Purpose	Canonical correlation analysis		
Syntax	<pre>[A,B] = canoncorr(X,Y) [A,B,r] = canoncorr(X,Y) [A,B,r,U,V] = canoncorr(X,Y) [A,B,r,U,V,stats] = canoncorr(X,Y)</pre>		
Description	[A,B] = canoncorr(X,Y) computes the sample canonical coefficients for the n-by-d1 and n-by-d2 data matrices X and Y. X and Y must have the same number of observations (rows) but can have different numbers of variables (columns). A and B are d1-by-d and d2-by-d matrices, where $d = min(rank(X), rank(Y))$. The jth columns of A and B contain the canonical coefficients, i.e., the linear combination of variables making up the jth canonical variable for X and Y, respectively. Columns of A and B are scaled to make the covariances of the canonical variables, or scores, the identity matrix (see U and V below). If X or Y is less than full rank, canoncorr gives a warning and returns zeros in the rows of A or B corresponding to dependent columns of X or Y.		
	[A,B,r] = canoncorr(X,Y) also returns a 1-by-d vector containing the sample canonical correlations. The jth element of r is the correlation between the jth columns of U and V (see below).		
	[A,B,r,U,V] = canoncorr(X,Y) also returns the canonical variables, known also as scores. U and V are n-by-d matrices computed as		
	U = (X - repmat(mean(X),N,1))*A V = (Y - repmat(mean(Y),N,1))*B		
	[A,B,r,U,V,stats] = canoncorr(X,Y) also returns a structure containing information relating to the sequence of d null hypotheses $H_0^{(k)}$, that the (k+1)st through dth correlations are all zero, for k = 0:(d-1). stats contains three fields, each a 1-by-d vector with elements corresponding to the values of k:		
	dfe	Error degrees of freedom, i.e., (d1-k)*(d2-k)	
	chisq	Bartlett's approximate chi-squared statistic for $H_0^{(k)}$	
	р	Right-tail significance level for $H_0^{(k)}$	

Examples

```
load carbig;
X = [Displacement Horsepower Weight Acceleration MPG];
nans = sum(isnan(X),2) > 0;
[A B r U V] = canoncorr(X(~nans,1:3), X(~nans,4:5));
plot(U(:,1),V(:,1),'.');
xlabel('0.0025*Disp + 0.020*HP - 0.000025*Wgt');
ylabel('-0.17*Accel + -0.092*MPG')
```



See Also manova1, princomp

References [1] Krzanowski, W.J., *Principles of Multivariate Analysis*, Oxford University Press, Oxford, 1988.

[2] Seber, G.A.F., Multivariate Observations, Wiley, New York, 1984.

capable

Purpose	Process capability indices		
Syntax	p = capable(data,specs) [p,Cp,Cpk] = capable(data,specs)		
Description	<pre>p = capable(data, specs) computes the probability that a sample, data, from some process falls outside the bounds specified in specs, a 2-element vector of the form [lower upper].</pre>		
	The assumptions are that the measured values in the vector data are normally distributed with constant mean and variance and that the measurements are statistically independent.		
	<pre>[p,Cp,Cpk] = capable(data,specs) additionally returns the capability indices Cp and Cpk.</pre>		
	C_p is the ratio of the range of the specifications to six times the estimate of the process standard deviation:		
	$C_p = \frac{USL - LSL}{6\sigma}$		
	For a process that has its average value on target, a C_p of 1 translates to a little more than one defect per thousand. Recently, many industries have set a quality goal of one part per million. This would correspond to $C_p = 1.6$. The higher the value of C_p , the more capable the process. C_{pk} is the ratio of difference between the process mean and the closer specification limit to three times the estimate of the process standard deviation:		
	$C_{pk} = min\left(\frac{USL-\mu}{3\sigma}, \frac{\mu-LSL}{3\sigma}\right)$		
	where the process mean is μ . For processes that do not maintain their average on target, C_{pk} is a more descriptive index of process capability.		
Example	Imagine a machined part with specifications requiring a dimension to be within three thousandths of an inch of nominal. Suppose that the machining process cuts too thick by one thousandth of an inch on average and also has a		

standard deviation of one thousandth of an inch. What are the capability indices of this process?

```
data = normrnd(1,1,30,1);
[p,Cp,Cpk] = capable(data,[-3 3]);
indices = [p Cp Cpk]
indices =
    0.0172    1.1144    0.7053
```

We expect 17 parts out of a thousand to be out-of-specification. Cpk is less than Cp because the process is not centered.

Reference [1] Montgomery, D., *"Introduction to Statistical Quality Control,"* John Wiley & Sons 1991. pp. 369–374.

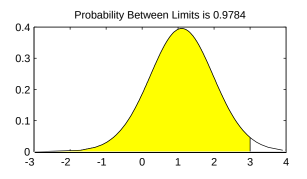
See Also capaplot, histfit

<u>capap</u>lot

Purpose	Process capability plot
Syntax	p = capaplot(data,specs) [p,h] = capaplot(data,specs)
Description	<pre>p = capaplot(data, specs) estimates the mean and variance of the observations in input vector data, and plots the pdf of the resulting T distribution. The observations in data are assumed to be normally distributed. The output, p, is the probability that a new observation from the estimated distribution will fall within the range specified by the two-element vector specs. The portion of the distribution between the lower and upper bounds specified in specs is shaded in the plot. [p,h] = capaplot(data, specs) additionally returns handles to the plot elements in h.</pre>
Example	<pre>Imagine a machined part with specifications requiring a dimension to be within 3 thousandths of an inch of nominal. Suppose that the machining process cuts too thick by one thousandth of an inch on average and also has a standard deviation of one thousandth of an inch. data = normrnd(1,1,30,1); p = capaplot(data,[-3 3]) p =</pre>

0.9784

The probability of a new observation being within specs is 97.84%.



See Also capable, histfit

caseread

Purpose	Read casenames from a file			
Syntax	names = caseread('filename') names = caseread			
Description	<pre>names = caseread('filename') reads the contents of filename and returns a string matrix of names. filename is the name of a file in the current directory, or the complete pathname of any file elsewhere. caseread treats each line as a separate case. names = caseread displays the Select File to Open dialog box for interactive selection of the input file.</pre>			
Example	Read the file months.dat created using the function casewrite on the next page. type months.dat January February March April May names = caseread('months.dat') names = January			
	February March April May			
See Also	tblread, gname, casewrite, tdfread			

casewrite

Purpose	Write casenames from a string matrix to a file
Syntax	casewrite(strmat,'filename') casewrite(strmat)
Description	<pre>casewrite(strmat, 'filename') writes the contents of string matrix strmat to filename. Each row of strmat represents one casename. filename is the name of a file in the current directory, or the complete pathname of any file elsewhere. casewrite writes each name to a separate line in filename. casewrite(strmat) displays the Select File to Write dialog box for interactive specification of the output file.</pre>
F	
Example	strmat = str2mat('January','February','March','April','May')
	strmat =
	January
	February
	March
	April
	May
	<pre>casewrite(strmat,'months.dat')</pre>
	type months.dat
	January
	February
	March
	April
	Мау
See Also	gname, caseread, tblwrite, tdfread

ccdesign

Purpose	Generate central composite design						
Syntax	<pre>D = ccdesign(nfactors) D = ccdesign(nfactors,'pname1',pvalue1,'pname2',pvalue2,) [D,blk] = ccdesign()</pre>						
Description	factors. The outp in the design. Ea of all factors for	D = ccdesign(nfactors) generates a central composite design for nfactors factors. The output matrix D is n-by-nfactors, where n is the number of points in the design. Each row represents one run of the design, and it has the settings of all factors for that run. Factor values are normalized so that the cube points take values between -1 and 1.					
	<pre>[D,blk] = ccdesign(nfactors) requests a blocked design. The output vector blk is a vector of block numbers. Blocks are groups of runs that are to be measured under similar conditions (for example, on the same day). Blocked designs minimize the effect of between-block differences on the parameter estimates.</pre>						
	<pre>[] = ccdesign(nfactors, 'pname1', pvalue1, 'pname2', pvalue2,) enables you to specify additional parameters and their values. Valid parameters are:</pre>						
	'center'	center' Number of center points:					
		Integer	Specific number of center points to include				
		'uniform'	Number of center points is selected to give uniform precision				
		'orthogonal'	Number of center points is selected to give an orthogonal design (default)				
	'fraction'	Fraction of full factorial for cube portion expressed as an exponent of 1/2. For example:					
		0	Whole design				
		1	1/2 fraction				
		2	1/4 fraction				
	'type'	Either 'inscribed', 'circumscribed', or 'faced'					
	'blocksize'	Maximum number of points allowed in a block.					

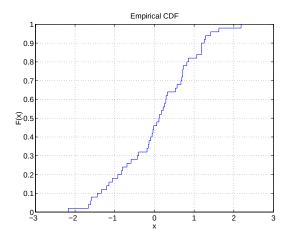
See Also bbdesign, cordexch, rowexch

Purpose	Computes a chosen cumulative distribution function (cdf)						
Syntax	P = cdf(' <i>name</i> ',X,A1,A2,A3)						
Description	P = cdf('name',X,A1,A2,A3) returns a matrix of probabilities, where name is a string containing the name of the distribution, X is a matrix of values, and A, A2, and A3 are matrices of distribution parameters. Depending on the distribution, some of these parameters may not be necessary.						
	Vector or matrix inputs for X, A1, A2, and A3 must have the same size, which is also the size of P . A scalar input for X, A1, A2, or A3 is expanded to a constant matrix with the same dimensions as the other inputs.						
	cdf is a utility routine allowing you to access all the cdfs in the Statistics Toolbox by using the name of the distribution as a parameter. See "Overview of the Distributions" on page 2-11 for the list of available distributions.						
Examples	p = cdf('Normal',-2:2,0,1) p =						
	0.0228 0.1587 0.5000 0.8413 0.9772						
	p = cdf('Poisson',0:5,1:6) p =						
	0.3679 0.4060 0.4232 0.4335 0.4405 0.4457						
See Also	betacdf, binocdf, chi2cdf, expcdf, fcdf, gamcdf, geocdf, hygecdf, icdf, logncdf, mle, nbincdf, ncfcdf, nctcdf, ncx2cdf, normcdf, pdf, poisscdf, random, raylcdf, tcdf, unidcdf, unifcdf, weibcdf						

Purpose	Plot of empirical cumulative distribution function						
Syntax	cdfplot(X) h = cdfplot(X) [h,stats] = cdf						
Description	cdfplot(X) displays a plot of the empirical cumulative distribution function (cdf) for the data in the vector X. The empirical cdf $F(x)$ is defined as the proportion of X values less than or equal to x .						
	the distribution of	This plot, like those produced by hist and normplot, is useful for examining the distribution of a sample of data. You can overlay a theoretical cdf on the same plot to compare the empirical distribution of the sample to the theoretical distribution.					
	The kstest, kstest2, and lillietest functions compute test statistics that are derived from the empirical cdf. You may find the empirical cdf plot produced by cdfplot useful in helping you to understand the output from those functions.						
	H = cdfplot(X) returns a handle to the cdf curve.						
	<pre>[h,stats] = cdfplot(X) also returns a stats structure with the following fields.</pre>						
	Field	Contents					
	stats.min	Minimum value					
	stats.max	Maximum value					
	stats.mean Sample mean						
	stats.median (50th percentile)						
	stats.std Sample standard deviation						
Examples	Generate a norm	al sample and an empirical cdf plot of the data.					

x = normrnd(0,1,50,1); cdfplot(x)

cdfplot





ecdf, hist, kstest, kstest2, lillietest, normplot

Purpose	Chi-square (χ^2) cumulative distribution function (cdf)					
Syntax	P = chi2cdf(X,V)					
Description	$P = chi2cdf(X,V)$ computes the χ^2 cdf at each of the values in X using the corresponding parameters in V. Vector or matrix inputs for X and V must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The degrees of freedom parameters in V must be positive integers, and the values in X must lie on the interval [0 1].					
	The χ^2 cdf for a given value x and degrees-of-freedom v is					
	$p = F(x v) = \int_0^x \frac{t^{(v-2)/2} e^{-t/2}}{2^{v/2} \Gamma(v/2)} dt$					
	where $\Gamma(\cdot)$ is the Gamma function. The result, p, is the probability that a single observation from a χ^2 distribution with v degrees of freedom will fall in the interval [0 x].					
	The χ^2 density function with v degrees-of-freedom is the same as the gamma density function with parameters v/2 and 2.					
Examples	probability = chi2cdf(5,1:5) probability =					
	0.9747 0.9179 0.8282 0.7127 0.5841					
	probability = chi2cdf(1:5,1:5) probability =					
	0.6827 0.6321 0.6084 0.5940 0.5841					
See Also	cdf, chi2inv, chi2pdf, chi2rnd, chi2stat					

chi2inv

Purpose	Inverse of the chi-square (χ^2) cumulative distribution function (cdf)				
Syntax	X = chi2inv(P,V)				
Description	$X = chi2inv(P,V)$ computes the inverse of the χ^2 cdf with parameters specified by V for the corresponding probabilities in P. Vector or matrix inputs for P and V must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.				
	The degrees of freedom parameters in V must be positive integers, and the values in P must lie in the interval [0 1].				
	The inverse χ^2 cdf for a given probability p and v degrees of freedom is				
	$x = F^{-1}(p v) = \{x:F(x v) = p\}$				
	where				
	$p = F(x v) = \int_0^x \frac{t^{(v-2)/2} e^{-t/2}}{2^{v/2} \Gamma(v/2)} dt$				
	and $\Gamma(\ \cdot\)$ is the Gamma function. Each element of output X is the value whose cumulative probability under the χ^2 cdf defined by the corresponding degrees of freedom parameter in V is specified by the corresponding value in P.				
Examples	Find a value that exceeds 95% of the samples from a χ^2 distribution with 10 degrees of freedom.				
	<pre>x = chi2inv(0.95,10) x =</pre>				
	18.3070				
	You would observe values greater than 18.3 only 5% of the time by chance.				
See Also	chi2cdf, chi2pdf, chi2rnd, chi2stat, icdf				

Purpose	Chi-square (χ^2) probability density function (pdf)
---------	--

Syntax Y = chi2pdf(X,V)

Description Y = chi2pdf(X,V) computes the χ^2 pdf at each of the values in X using the corresponding parameters in V. Vector or matrix inputs for X and V must have the same size, which is also the size of output Y. A scalar input is expanded to a constant matrix with the same dimensions as the other input.

The degrees of freedom parameters in V must be positive integers, and the values in X must lie on the interval $[0 \ 1]$.

The χ^2 pdf for a given value *x* and v degrees of freedom is

$$y = f(x|v) = \frac{x^{(v-2)/2}e^{-x/2}}{2^{v/2}\Gamma(v/2)}$$

where $\Gamma(\cdot)$ is the Gamma function. The result, *y*, is the probability that a single observation from a χ^2 distribution with v degrees of freedom will have value *x*.

If x is standard normal, then x^2 is distributed χ^2 with one degree of freedom. If $x_1, x_2, ..., x_n$ are *n* independent standard normal observations, then the sum of the squares of the *x*'s is distributed χ^2 with *n* degrees of freedom (and is equivalent to the gamma density function with parameters v/2 and 2).

```
Examples

nu = 1:6;

x = nu;

y = chi2pdf(x,nu)

y =

0.2420 0.1839 0.1542 0.1353 0.1220 0.1120
```

The mean of the χ^2 distribution is the value of the degrees of freedom parameter, nu. The above example shows that the probability density of the mean falls as nu increases.

See Also chi2cdf, chi2inv, chi2rnd, chi2stat, pdf

chi2rnd

Purpose	Random numbers from the chi-square (χ^2) distribution						
Syntax	<pre>R = chi2rnd(V) R = chi2rnd(V,m) R = chi2rnd(V,m,n)</pre>						
Description	R = chi2rnd(V) generates random numbers from the χ^2 distribution with degrees of freedom parameters specified by V. R is the same size as V.						
	R = chi2rnd(V,m) generates a matrix of size m containing random numbers from the χ^2 distribution with degrees of freedom parameter V, where m is a 1-by-2 vector containing the row and column dimensions of R.						
	R = chi2rnd(V,m,n) generates an m-by-n matrix containing random numbers from the χ^2 distribution with degrees of freedom parameter V.						
Examples	Note that the first and third commands are the same, but are different from the second command.						
	r = chi2rnd(1:6) r =						
	0.0037 3.0377 7.8142 0.9021 3.2019 9.0729						
	r = chi2rnd(6,[1 6]) r =						
	6.5249 2.6226 12.2497 3.0388 6.3133 5.0388						
	r = chi2rnd(1:6,1,6) r =						
	0.7638 6.0955 0.8273 3.2506 1.5469 10.9197						
See Also	chi2cdf, chi2inv, chi2pdf, chi2stat						

Purpose	Mean and variance for the chi-square (χ^2) distribution										
Syntax	[M,V] =	[M,V] = chi2stat(NU)									
Description	with dea The mea	$[M,V]$ = chi2stat(NU) returns the mean and variance for the χ^2 distribution with degrees of freedom parameters specified by NU. The mean of the χ^2 distribution is v, the degrees of freedom parameter, and the variance is $2v$.									
Example	nu = [m,v	1:10; nu'*n] = ch	u; i2stat	(nu)							
	m =										
	1	2	3	4	5	6	7	8	9	10	
	2	4	6	8	10	12	14	16	18	20	
	3	6	9	12	15	18	21	24	27	30	
	4	8	12	16	20	24	28	32	36	40	
	5	10 12	15	20	25	30	35	40	45	50	
	6 7	12	18 21	24 28	30 35	36 42	42 49	48 56	54 63	60 70	
	8	14	21	20 32	35 40	42 48	49 56	56 64	03 72	80	
	9	18	27	36	45	-0 54	63	72	81	90	
	10	20	30	40	50	60	70	80	90	100	
	v =										
			0	0	10	10		10	4.0		
	2 4	4 8	6 12	8 16	10 20	12 24	14 28	16 32	18 36	20 40	
	4 6	0 12	12	24	20 30	24 36	20 42	32 48	50 54	40 60	
	8	16	24	32	40	48	56	40 64	72	80	
	10	20	30	40	50	60	70	80	90	100	
	12	24	36	48	60	72	84	96	108	120	
	14	28	42	56	70	84	98	112	126	140	
	16	32	48	64	80	96	112	128	144	160	
	18	36	54	72	90	108	126	144	162	180	
	20	40	60	80	100	120	140	160	180	200	

See Also

chi2cdf, chi2inv, chi2pdf, chi2rnd

classify

Purpose	Discriminant Analysis					
Syntax	<pre>class = classify(sample,training,group) [class,err] = classify() [] = classify(,'type') [] = classify(,'type',prior)</pre>					
Description	<pre>class = classify(sample,training,group) classifies each row of the data in sample into one of the groups in training. sample and training must be matrices with the same number of columns. group is a grouping variable for training. Its unique values define groups, and each element defines which group the corresponding row of training belongs to. group can be a numeric vector, a string array, or a cell array of strings. training and group must have the same number of rows. classify treats NaNs or empty strings in group as missing values, and ignores the corresponding rows of training. class indicates which group each row of sample has been assigned to, and is of the same type as group. [class,err] = classify() also returns an estimate of the misclassification error rate. classify returns the apparent error rate, i.e., the percentage of observations in the training that are misclassified. [] = classify(,'type') allows you to specify the type of discriminant function, as one of:</pre>					
	'linear' Fits a multivariate normal density to each group, with a pooled estimate of covariance (default).					
	'quadratic'	Fits MVN densities with covariance estimates stratified by group.				
	'mahalanobis'	Uses Mahalanobis distances with stratified covariance estimates.				
	[] = classify(, ' <i>type</i> ', prior) enables you to specify prior probabilities for the groups in one of three ways. prior can be:					

• A numeric vector of the same length as the number of unique values in group. If group is numeric, the order of prior must correspond to the sorted

	values in group, or, if group contains strings, to the order of first occurrence of the values in group.A 1-by-1 structure with fields:				
	prob A numeric vector				
	group	group Of the same type as group, and containing unique values indicating which groups the elements of prob correspond to.			
	As a structure, prior can contain groups that do not appear in group. This can be useful if training is a subset a larger training set.				
		ing value 'empirical', indicating that classify should estimate the prior probabilities from the group relative frequencies in training.			
	prior defaults to a numeric vector of equal probabilities, i.e., a uniform distribution. prior is not used for discrimination by Mahalanobis distance, except for error rate calculation.				
Examples	sample traini g = gr class	<pre>discrim e = ratings(idx,:); ing = ratings(1:200,:); roup(1:200); = classify(sample,training,g); 5 = class(1:5)</pre>			
	first5	5 =			
See Also	mahal				
References		owski, W.J., <i>Principles of Multivariate Analysis</i> , Oxford University ford, 1988.			
	[2] Seber, G.A.F., Multivariate Observations, Wiley, New York, 1984				

cluster

Purpose	Construct clusters from linkage output			
Syntax	<pre>T = cluster(Z,'cutoff',c) T = cluster(Z,'maxclust',n) T = cluster(,'criterion','crit') T = cluster(,'depth',d)</pre>			
Description	T = cluster(Z, 'cutoff', c) constructs clusters from the hierarchical cluster tree, Z, generated by the linkage function. Z is a matrix of size (<i>m</i> -1)-by-3, where <i>m</i> is the number of observations in the original data. c is a threshold for cutting Z into clusters. Clusters are formed when inconsistent values are less than c. See the inconsistent function for more information. The output T is a vector of size <i>m</i> that contains the cluster number for each observation in the original data.			
	T = cluster(Z, 'maxclust', n) specifies n as the maximum number of clusters to form from the hierarchical tree in Z.			
	T = cluster(,'criterion',' <i>crit</i> ') uses the specified criterion for forming clusters, where crit is either 'inconsistent' or 'distance'.			
	<pre>T = cluster(, 'depth',d) evaluates inconsistent values to a depth of d in the tree. The default is d = 2. An inconsistency coefficient computation compares a link between two objects in the cluster tree with neighboring links up to the specified depth. See the inconsistent function for more information.</pre>			
Example	The example uses the pdist function to calculate the distance between items in a matrix of random numbers and then uses the linkage function to compute the hierarchical cluster tree based on the matrix. The example passes the output of the linkage function to the cluster function. The 'maxclust' value 3 indicates that you want to group the items into three clusters. The find function lists all the items grouped into cluster 1.			
	<pre>rand('seed', 0); X = [rand(10,3); rand(10,3)+1; rand(10,3)+2]; Y = pdist(X); Z = linkage(Y); T = cluster(Z,'maxclust',3); find(T==1)</pre>			

ans	=
	11
	12
	13
	14
	15
	16
	17
	18
	19
	20
cluste	rdata, cophenet, inconsistent, linkage, pdist

See Also

clusterdata

Purpose	Construct clusters from data					
Syntax		ata(X, cutoff) ata(X,'param1',val1,'param2',val2,)				
Description	T = clusterdata(X, cutoff) uses the pdist, linkage, and cluster functions to construct clusters from data X. X is an <i>m</i> -by- <i>n</i> matrix, treated as <i>m</i> observations of <i>n</i> variables. cutoff is a threshold for cutting the hierarchical tree generated by linkage into clusters. When 0 < cutoff < 2, clusterdata forms clusters when inconsistent values are greater than cutoff (see the inconsistent function). When cutoff is an integer and cutoff >= 2, then clusterdata inteprets cutoff as the maximum number of clusters to keep in the hierarchical tree generated by linkage. The output T is a vector of size <i>m</i> containing a cluster number for each observation.					
	T = clusterd	ata(X,cutoff) is the same as				
	Y = pdist(X,'euclid'); Z = linkage(Y,'single'); T = cluster(Z,'cutoff',cutoff);					
	T = clusterdata(X, 'param1', val1, 'param2', val2,) provides more control over the clustering through a set of parameter/value pairs. Valid parameters are:					
	'distance'	Any of the distance metric names allowed by pdist (follow the 'minkowski' option by the value of the exponent p).				
	'linkage'	Any of the linkage methods allowed by the linkage function				
	'cutoff'	Cutoff for inconsistent or distance measure				
	'maxclust' Maximum number of clusters to form					
	'criterion' Either 'inconsistent' or 'distance'					
	'depth' Depth for computing inconsistent values					
Example	clusterdatat	first creates a sample dataset of random numbers. It then uses to compute the distances between items in the dataset and create cluster tree from the dataset. Finally, the clusterdata function				

groups the items in the dataset into three clusters. The example uses the find function to list all the items in cluster 2.

```
See Also
```

cluster, inconsistent, kmeans, linkage, pdist

cmdscale

Purpose	Classical multidimensional scaling				
Syntax	Y = cmdscale(D) [Y,e] = cmdscale(D)				
Description	Y = cmdscale(D) takes an n-by-n distance matrix D, and returns an n-by-p configuration matrix Y. Rows of Y are the coordinates of n points in p-dimensional space for some $p < n$. When D is a Euclidean distance matrix, the distances between those points are given by D. p is the dimension of the smallest space in which the n points whose interpoint distances are given by D can be embedded.				
	[Y,e] = cmdscale(D) also returns the eigenvalues of $Y*Y'$. When D is Euclidean, the first p elements of e are positive, the rest zero. If the first k elements of e are much larger than the remaining $(n-k)$, then you can use the first k columns of Y as k-dimensional points whose interpoint distances approximate D. This can provide a useful dimension reduction for visualization, e.g., for k = 2.				
	D need not be a Euclidean distance matrix. If it is non-Euclidean or a more general dissimilarity matrix, then some elements of e are negative, and cmdscale choses p as the number of positive eigenvalues. In this case, the reduction to p or fewer dimensions provides a reasonable approximation to D only if the negative elements of e are small in magnitude.				
	You can specify D as either a full dissimilarity matrix, or in upper triangle vector form such as is output by pdist. A full dissimilarity matrix must be real and symmetric, and have zeros along the diagonal and positive elements everywhere else. A dissimilarity matrix in upper triangle form must have real, positive entries. You can also specify D as a full similarity matrix, with ones along the diagonal and all other elements less than one. cmdscale tranforms a similarity matrix to a dissimilarity matrix in such a way that distances between the points returned in Y equal or approximate sqrt(1-D). To use a different transformation, you must transform the similarities prior to calling cmdscale.				
Examples	Generate some points in 4-dimensional space, but "close" to 3-dimensional space, then reduce them to distances only.				

```
X = [normrnd(0,1,10,3) normrnd(0,.1,10,1)];
D = pdist(X,'euclidean');
```

Find a configuration with those inter-point distances.

```
[Y,e] = cmdscale(D);
% Four, but fourth one small
dim = sum(e > eps^{(3/4)})
% Poor reconstruction
maxerr2 = max(abs(pdist(X) - pdist(Y(:,1:2))))
% Good reconstruction
maxerr3 = max(abs(pdist(X) - pdist(Y(:,1:3))))
% Exact reconstruction
maxerr4 = max(abs(pdist(X) - pdist(Y)))
% D is now non-Euclidean
D = pdist(X, 'cityblock');
[Y,e] = cmdscale(D);
% One is large negative
min(e)
% Poor reconstruction
maxerr = max(abs(pdist(X) - pdist(Y)))
```

See Also pdist, procrustes

References [1] Seber, G.A.F., *Multivariate Observations*, Wiley, 1984

combnk

Purpose	Enumeration of all combinations of n objects k at a time			
Syntax	C = combnk(v,k)			
Description	C = combnk(v,k) returns all combinations of the <i>n</i> elements in v taken k at a time.			
	C = combnk(v,k) produces a matrix C with k columns and $n! / k!(n-k)!$ rows, where each row contains k of the elements in the vector v.			
	It is not practical to use this function if v has more than about 15 elements.			
Example	Combinations of characters from a string.			
	C = combnk('tendril',4); last5 = C(31:35,:)			
	last5 =			
	tedr tenl teni tenr tend			
	Combinations of elements from a numeric vector.			
	c = combnk(1:4,2)			
	$ \begin{array}{c} c \\ & 3 \\ & 2 \\ & 2 \\ & 4 \\ & 2 \\ & 3 \\ & 1 \\ & 1 \\ & 1 \\ & 3 \\ & 1 \end{array} $			

cophenet

Purpose	Cophenetic correlation	coefficient
---------	------------------------	-------------

Syntax

c = cophenet(Z,Y)

Description

c = cophenet(Z,Y) computes the cophenetic correlation coefficient which compares the distance information in Z, generated by linkage, and the distance information in Y, generated by pdist. Z is a matrix of size (*m*-1)-by-3, with distance information in the third column. Y is a vector of size $m \cdot (m-1)/2$.

For example, given a group of objects $\{1, 2, ..., m\}$ with distances Y, the function linkage produces a hierarchical cluster tree. The cophenet function measures the distortion of this classification, indicating how readily the data fits into the structure suggested by the classification.

The output value, c, is the cophenetic correlation coefficient. The magnitude of this value should be very close to 1 for a high-quality solution. This measure can be used to compare alternative cluster solutions obtained using different algorithms.

The cophenetic correlation between Z(:,3) and Y is defined as

$$c = \frac{\sum_{i < j} (Y_{ij} - y)(Z_{ij} - z)}{\sqrt{\sum_{i < j} (Y_{ij} - y)^2 \sum_{i < j} (Z_{ij} - z)^2}}$$

where:

- Y_{ij} is the distance between objects i and j in Y.
- Z_{ij} is the distance between objects i and j in Z(:,3).
- *y* and *z* are the average of Y and Z(:,3), respectively.

```
Example
```

cophenet

See Also cluster, dendrogram, inconsistent, linkage, pdist, squareform

Purpose	D-optimal design of experiments – coordinate exchange algorithm					
Syntax	<pre>settings = cordexch(nfactors,nruns) [settings,X] = cordexch(nfactors,nruns) [settings,X] = cordexch(nfactors,nruns,'model') [settings,X] = cordexch(,'param1',value1,'param2',value2,)</pre>					
Description	settings = cordexch(nfactors,nruns) generates the factor settings matrix, settings, for a D-optimal design using a linear additive model with a constant term. settings has nruns rows and nfactors columns.					
	[settings,X design matrix] = cordexch(nfactors,nruns) also generates the associated x X.				
] = cordexch(nfactors,nruns,'model') produces a design for ified regression model. The input, 'model', can be one of these				
	'linear'	Includes constant and linear terms (the default)				
	'interactio	n' Includes constant, linear, and cross-product terms.				
	'quadratic'	Includes interactions and squared terms.				
	'purequadra	tic' Includes constant, linear and squared terms.				
	Alternatively model can be a matrix of term definitions as accepted by the x2fx function.					
	[settings,X] = cordexch(,'param1',value1,'param2',value2,) provides more control over the design generation through a set of parameter/value pairs. Valid parameters are:					
	'display'	Either 'on' or 'off' to control display of iteration counter. The default is 'on'.				
	'init'	Initial design as an nruns-by-nfactors matrix. The default is a randomly selected set of points.				
	'maxiter'	Maximum number of iterations. The default is 10.				

Example The D-optimal design for two factors in nine runs using a quadratic model is the 3² factorial as shown below:

```
settings = cordexch(2,9,'quadratic')
settings =
     - 1
             1
      1
             1
      0
             1
      1
            - 1
     - 1
            - 1
      0
            - 1
             0
      1
             0
      0
     - 1
             0
```

- **Algorithm** The cordexch function searches for a D-optimal design using a coordinate exchange algorithm. It creates a starting design, and then iterates by changing each coordinate of each design point in an attempt to reduce the variance of the coefficients that would be estimated using this design.
- See Also bbdesign, candexch, candgen, ccdesign, daugment, dcovary, rowexch, x2fx

Purpose	Correlation coefficients with tests and confidence bounds			
Syntax	<pre>R = corr(X) R = corr(x,y) [R,P] = corr() [R,P,RL0,RUP] = corr() [] = corr(, 'param1',val1, 'param2',val2,)</pre>			
Description	R = corr(X) calculates a matrix R of correlation coefficients for an array X Each row of X is an observation and each column is a variable. NaN (not a number) values in X are treated as missing.			
	If C is the covariance matrix, C = $cov(X)$, then $corr(X)$ is the matrix whose (i, j)th element is			
	$rac{C(i,j))}{\sqrt{C(i,i)C(j,j)}}$			
	<pre>R = corr(x,y) where x and y are column vectors, is the same as R = corr([x y]).</pre>			
	 [R,P] = corr() also returns P, a matrix of p-values for testing the hypothesis of no correlation. Each p-value is the probability of getting a correlation as large as the observed value by random chance, when the true correlation is zero. If P(i,j) is small, say less than 0.05, then the correlation R(i,j) is significant. See "Algorithm" for details. [R,P,RLO,RUP] = corr() also returns matrices RLO and RUP, of the same size as R, containing lower and upper bounds for a 95% confidence interval for each coefficient. 			
	[] = corr(, 'param1', val1, 'param2', val2,) specifies additional parameters and their values. Valid parameters are the following:			
	'alpha' A number between 0 and 1 that specifies a confidence level of 100*(1-alpha)%. Default is 0.05 for 95% confidence intervals.			

	'rows'	'all'	Use all rows.	
		'complete'	Use rows with no missing values (default).	
		'pairwise'	Compute R(i,j) using rows with no missing values in column i or j.	
Examples		lo,rup] = cor	<pre>% load sample data r([x1 x2 x3 x4 x5],'alpha',0.10);); % find significant correlations % display their indices</pre>	
Algorithm	having 1 a confidence 0.5*log((These bou	he p-value is computed by transforming the correlation to create an F statistic aving 1 and n-2 degrees of freedom, where n is the number of rows of X. The onfidence bounds are based on an asymptotic normal distribution for .5*log((1+R)/(1-R)), with an approximate variance equal to 1/(n-3). hese bounds are accurate for large samples when X has a multivariate normal istribution.		
See Also	corrcoef,	cov, nanmean, i	nanstd, std	

corrcoef

Purpose	Correlation coefficients
Syntax	R = corrcoef(X)
	R = corrcoef(x,y)
	[R,P]=corrcoef()
	[R,P,RLO,RUP]=corrcoef()
	[]=corrcoef(,'param1',val1,'param2',val2,)

Description R = corrcoef(X) returns a matrix R of correlation coefficients calculated from an input matrix X whose rows are observations and whose columns are variables. The (i,j)th element of the matrix R is related to the covariance matrix C = cov(X) by

$$R(i,j) = \frac{C(i,j)}{\sqrt{C(i,i)C(j,j)}}$$

corrcoef(X) is the zeroth lag of the covariance function, that is, the zeroth lag of xcov(x, 'coeff') packed into a square array.

R = corrcoef(x,y) where x and y are column vectors is the same as corrcoef([x y]).

[R,P]=corrcoef(...) also returns P, a matrix of p-values for testing the hypothesis of no correlation. Each p-value is the probability of getting a correlation as large as the observed value by random chance, when the true correlation is zero. If P(i,j) is small, say less than 0.05, then the correlation R(i,j) is significant.

[R, P, RLO, RUP]=corrcoef(...) also returns matrices RLO and RUP, of the same size as R, containing lower and upper bounds for a 95% confidence interval for each coefficient.

[...]=corrcoef(..., 'param1', val1, 'param2', val2,...) specifies additional parameters and their values. Valid parameters are the following.

	'alpha'	A number between 0 and 1 to specify a confidence level of 100*(1 - alpha)%. Default is 0.05 for 95% confidence intervals. Either 'all' (default) to use all rows, 'complete' to use rows with no NaN values, or 'pairwise' to compute R(i,j) using rows with no NaN values in either column i or j.					
	'rows'						
	having n-2 d confidence b 0.5*log((1 These bound	the is computed by transforming the correlation to create a t statistic 2 degrees of freedom, where n is the number of rows of X. The e bounds are based on an asymptotic normal distribution of $(1+R)/(1-R)$, with an approximate variance equal to $1/(n-3)$. nds are accurate for large samples when X has a multivariate normal on. The 'pairwise' option can produce an R matrix that is not effinite.					
	The corrcoe	ef function is p	art of the star	ndard MATLAB language.			
Examples	Generate ra columns.	enerate random data having correlation between column 4 and the other olumns.					
	x(:,4) = [r,p] =	<pre>x = randn(30,4); % Uncorrelated data x(:,4) = sum(x,2); % Introduce correlation. [r,p] = corrcoef(x) % Compute sample correlation and p-values. [i,j] = find(p<0.05); % Find significant correlations. [i,j] % Display their (row,col) indices.</pre>					
	r =						
	1.00		0.1929	0.3457			
	-0.35		-0.1429	0.4461			
	0.19 0.34		1.0000 0.5183	0.5183 1.0000			
	p = 1.00	00 0.0531	0.3072	0.0613			
	0.05		0.3072	0.0135			
	0.30		1.0000	0.0033			
	0.06		0.0033	1.0000			
	ans = 4						

4	3
2	4
3	4

See Also o

cov, mean, std, var xcorr, xcov in the Signal Processing Toolbox

τον

Purpose	Covariance matrix.
Syntax	C = cov(X) C = cov(x,y)
Description	C = cov(X) computes the covariance matrix. For a single vector, $cov(x)$ returns a scalar containing the variance. For matrices, where each row is an observation, and each column a variable, $cov(X)$ is the covariance matrix.
	The variance function, $var(X)$ is the same as $diag(cov(X))$.
	The standard deviation function, $std(X)$ is equivalent to $sqrt(diag(cov(X)))$.
	$cov(x,y)$, where x and y are column vectors of equal length, gives the same result as $cov([x \ y])$.
	The cov function is part of the standard MATLAB language.
Algorithm	The algorithm for cov is
	<pre>[n,p] = size(X); X = X - ones(n,1) * mean(X); Y = X'*X/(n-1);</pre>
See Also	corrcoef, mean, std, var
	xcov, xcorr (Signal Processing Toolbox)

Purpose	Cross-tabulation of several vectors
Syntax	<pre>table = crosstab(col1,col2) table = crosstab(col1,col2,col3,) [table,chi2,p] = crosstab(col1,col2) [table,chi2,p,label] = crosstab(col1,col2)</pre>
Description	table = crosstab(col1,col2) takes two vectors of positive integers and returns a matrix, table, of cross-tabulations. The <i>ij</i> th element of table contains the count of all instances where $col1 = i$ and $col2 = j$.
	Alternatively, col1 and col2 can be vectors containing noninteger values, character arrays, or cell arrays of strings. crosstab implicitly assigns a positive integer group number to each distinct value in col1 and col2, and creates a cross-tabulation using those numbers.
	table = crosstab(col1,col2,col3,) returns table as an <i>n</i> -dimensional array, where <i>n</i> is the number of arguments you supply. The value of table(i,j,k,) is the count of all instances where col1 = <i>i</i> , col2 = <i>j</i> , col3 = k , and so on.
	<pre>[table,chi2,p] = crosstab(col1,col2) also returns the chi-square statistic, chi2, for testing the independence of the rows and columns of table. The scalar p is the significance level of the test. Values of p near zero cast doubt on the assumption of independence of the rows and columns of table.</pre>
	<pre>[table,chi2,p,label] = crosstab(col1,col2) also returns a cell array label that has one column for each input argument. The value in label(i,j) is the value of colj that defines group i in the jth dimension.</pre>
Example	Example 1 We generate 2 columns of 50 discrete uniform random numbers. The first column has numbers from 1 to 3. The second has only the numbers 1 and 2. The two columns are independent so we would be surprised if p were near zero. r1 = unidrnd(3,50,1);
	r2 = unidrnd(2,50,1); [table,chi2,p] = crosstab(r1,r2)
	table =

```
10 5
8 8
6 13
chi2 =
4.1723
p =
0.1242
```

The result, 0.1242, is not a surprise. A very small value of p would make us suspect the "randomness" of the random number generator.

Example 2

We have data collected on several cars over a period of time. How many four-cylinder cars were made in the USA during the late part of this period?

```
[t,c,p,1] = crosstab(cyl4,when,org);
l
l =
    'Other' 'Early' 'USA'
    'Four' 'Mid' 'Europe'
    [] 'Late' 'Japan'
t(2,3,1)
ans =
    38
```

See Also

tabulate

Purpose	D-optimal augmentation of an experimental design				
Syntax	<pre>settings = daugment(startdes,nruns) [settings,X] = daugment(startdes,nruns) [settings,X] = daugment(startdes,nruns,'model') [settings, X] = daugment(,'param1',value1,'param2',value2,)</pre>				
Description	<pre>settings = daugment(startdes,nruns) adds nruns runs to an experimental design using the coordinate exchange D-optimal algorithm. startdes is a matrix of factor settings in the original design. The output matrix settings is the matrix of factor settings for the design.</pre>				
	[settings,X] = daugment(startdes,nruns) also generates the associated design matrix, X.				
	<pre>[settings,X] = daugment(startdes,nruns,'model') also controls the order of the regression model. The input, 'model', can be one of these:</pre>				
	'linear'	linear' Includes constant and linear terms (the default)			
	<pre>'interaction' Includes constant, linear, and cross-product terms. 'quadratic' Includes interactions and squared terms.</pre>				
	'purequadratic'	Includes constant, linear and squared terms.			
	Alternatively model can be a matrix of term definitions as accepted by the x2fx function.				
	provides more con	daugment(,'param1',value1,'param2',value2,) trol over the design generation through a set of pairs. Valid parameters are the following::			
	Parameter Value				
		Either 'on' or 'off' to control display of iteration counter. The default is 'on'.			

daugment

Parameter	Value			
'init'	Initial design as an nruns-by-nfactors matrix. The default is a randomly selected set of points.			
'maxiter'	Maximum number of iterations. The default is 10.			
We add 5 runs to a 2^2 factorial design to allow us to fit a quadratic model.				
	= [-1 -1; 1 -1; -1 1; 1 1]; = daugment(startdes,5,'quadratic')			
settings =	=			
- 1	- 1			
1	-1			
- 1	1			
1	1			
1	0			
- 1	0			
0	1			
0	0			
0	-1			
	<pre>'maxiter' We add 5 runs startdes = settings = settings = -1 1 -1 1 1 -1 0 0 0</pre>			

See Also cordexch, x2fx

Purpose	D-optimal design with specified fixed covariates				
Syntax	<pre>settings = dcovary(nfactors,covariates) [settings,X] = dcovary(nfactors,covariates) [settings,X] = dcovary(nfactors,covariates,'model') [settings,X] = dcovary(,'param1',value1,'param2',value2,)</pre>				
Description	<pre>settings = dcovary(nfactors, covariates) uses a coordinate exchange algorithm to generate a D-optimal design for nfactors factors, subject to the constraint that it also include the fixed covariate values in the input matrix covariates. The number of runs in the design is taken to be the number of rows in the covariates matrix. The output matrix settings is the matrix of factor settings for the design, including the fixed covariates. [settings,X] = dcovary(nfactors, covariates) also generates the associated design matrix, X. [settings,X] = dcovary(nfactors, covariates, 'model') also controls the order of the regression model. The input, 'model', can be one of these:</pre>				
	'linear' Includes constant and linear terms (the default)				
	'interaction' Includes constant, linear, and cross-product terms.				
	'quadratic' Includes interactions and squared terms.				
	'purequadratic'	Includes constant, linear and squared terms.			
	Alternatively 'model' can be a matrix of term definitions as accept				

Alternatively 'model' can be a matrix of term definitions as accepted by the x2fx function. The model is applied to the fixed covariates as well as the regular factors. If you want to treat the fixed covariates specially, for example by including linear terms for them but quadratic terms for the regular factors, you can do this by creating the proper 'model' matrix.

[settings, X] = dcovary(...,'param1',value1,'param2',value2,...)
provides more control over the design generation through a set of
parameter/value pairs. Valid parameters are:

	'display'	Either 'on' or 'off' to control display of iteration counter. The default is 'on'.	
	'init'	Initial design as an nruns-by-nfactors matrix. The default is a randomly selected set of points.	
	'maxiter'	Maximum number of iterations. The default is 10.	
Example	<pre>Example 1. Generate a design for three factors in 2 blocks of 4 runs. blk = [-1 -1 -1 -1 1 1 1 1]'; dsgn = dcovary(3,blk)</pre>		
	dsgn = -1	1 1 -1	

- 1	1	1	- 1
1	- 1	- 1	- 1
- 1	1	- 1	- 1
1	- 1	1	- 1
1	1	- 1	1
1	1	1	1
- 1	- 1	1	1
- 1	- 1	- 1	1

Example 2. Suppose we want to block an eight run experiment into 4 blocks of size 2 to fit a linear model on two factors.

covariate	es = d	ummyvar	([1 1	2233	3 4 4]);
settings	= dco	vary(2,	covar	iates(:,	1:3),'linear')
settings	=				
1	1	1	0	0	
- 1	- 1	1	0	0	
- 1	1	0	1	0	
1	- 1	0	1	0	
1	1	0	0	1	
- 1	- 1	0	0	1	
- 1	1	0	0	0	
1	- 1	0	0	0	

The first two columns of the output matrix contain the settings for the two factors. The last three columns are *dummy variable* codings for the four blocks.

Algorithm	The dcovary function creates a starting design that includes the fixed covariate values, and then iterates by changing the non-fixed cordinates of each design
	point in an attempt to reduce the variance of the coefficients that would be estimated using this design.

See Also cordexch, daugment, rowexch, x2fx

dendrogram

Purpose	Plot dendrogram graphs				
Syntax	<pre>H = dendrogram(Z) H = dendrogram(Z,p) [H,T] = dendrogram() [H,T,perm] = dendrogram() [] = dendrogram(,'colorthreshold',t) [] = dendrogram(,'orientation','orient')</pre>				
Description	H = dendrogram(Z) generates a dendrogram plot of the hierarchical, binary cluster tree, Z. Z is an (m-1)-by-3 matrix, generated by the linkage function, where m is the number of objects in the original dataset.				
	A dendrogram consists of many U-shaped lines connecting objects in a hierarchical tree. Except for the Ward linkage (see linkage), the height of each U represents the distance between the two objects being connected. The output, H, is a vector of line handles.				
	H = dendrogram(Z,p) generates a dendrogram with only the top p nodes. By default, dendrogram uses 30 as the value of p. When there are more than 30 initial nodes, a dendrogram may look crowded. To display every node, set $p = 0$.				
	[H,T] = dendrogram() generates a dendrogram and returns T, a vector of length m that contains the leaf node number for each object in the original dataset. T is useful when p is less than the total number of objects, so some leaf nodes in the display correspond to multiple objects. For example, to find out which objects are contained in leaf node k of the dendrogram, use find(T==k). When there are fewer than p objects in the original data, all objects are displayed in the dendrogram. In this case, T is the identity map, i.e., T = (1:m)', where each node contains only itself.				
	When there are fewer than p objects in the original data, all objects are displayed in the dendrogram. In this case, T is the identity map, i.e., $T = (1:m)'$, where each node contains only itself.				
	[H,T,perm] = dendrogram() generates a dendrogram and returns the permutation vector of the node labels of the leaves of the dendrogram. perm is				

ordered from left to right on a horizontal dendrogram and bottom to top for a vertical dendrogram.

[...] = dendrogram(..., 'colorthreshold',t) assigns a unique color to each group of nodes in the dendrogram where the linkage is less than the threshold t. t is a value in the interval [0,max(Z(:,3))]. Setting t to the string 'default' is the same as t = .7(max(Z(:,3))). 0 is the same as not specifying 'colorthreshold'. The value max(Z(:,3)) treats the entire tree as one group and colors it all one color.

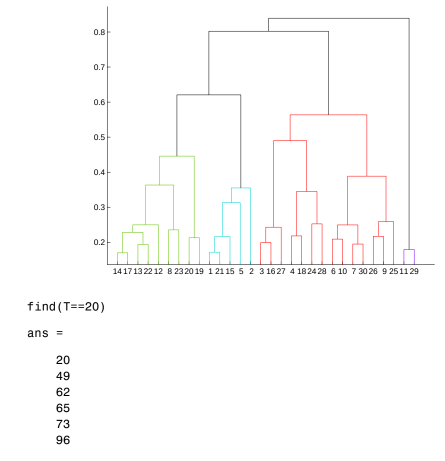
[...] = dendrogram(..., 'orientation', '*orient*') orients the dendrogram within the figure window. The options for '*orient*' are

'top'	Top to bottom (default)
'bottom'	Bottom to top
'left'	Left to right
'right'	Right to left

Example

rand('seed',12); X= rand(100,2); Y= pdist(X,'cityblock'); Z= linkage(Y,'average'); [H,T] = dendrogram(Z,'colorthreshold','default');

dendrogram



This output indicates that leaf node 20 in the dendrogram contains the original data points 20, 49, 62, 65, 73, and 96.

See Also cluster, clusterdata, cophenet, inconsistent, linkage, pdist, silhouette, squareform

Purpose	Interactive graph of cdf (or pdf) for many probability distributions				
Syntax	disttool				
Description	The disttool command displays a graphic user interface for exploring the effects of changing parameters on the plot of a cdf or pdf. Clicking and draggin a vertical line on the plot allows you to interactively evaluate the function ov its entire domain.				
	Evaluate the plotted function by typing a value in the <i>x</i> -axis edit box or dragging the vertical reference line on the plot. For cdfs, you can evaluate the inverse function by typing a value in the <i>y</i> -axis edit box or dragging the horizontal reference line on the plot. The shape of the pointer changes from an arrow to a crosshair when it is over the vertical or horizontal line to indicate that the reference line is draggable.				
	To change the distribution function, choose an option from the menu of functions at the top left of the figure. To change from cdfs to pdfs, choose an option from the menu at the top right of the figure.				
	To change the parameter settings, move the sliders or type a value in the edit box under the name of the parameter. To change the limits of a parameter, type a value in the edit box at the top or bottom of the parameter slider.				
	To close the tool, press the Close button.				
See Also	randtool				

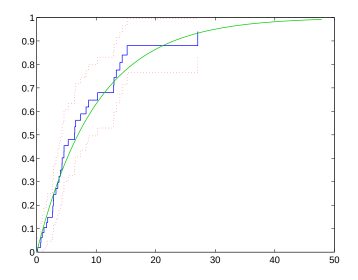
dummyvar

Purpose	Matrix of 0-1 "dummy" variables					
Syntax	D = du	mmyvar	(group)		
Description	D = dummyvar(group) generates a matrix, D, of 0-1 columns. D has one column for each unique value in each column of the matrix group. Each column of group contains positive integers that indicate the group membership of an individual row.					
Example	<pre>Suppose we are studying the effects of two machines and three operators on a process. The first column of group would have the values 1 or 2 depending on which machine was used. The second column of group would have the values 1, 2, or 3 depending on which operator ran the machine. group = [1 1;1 2;1 3;2 1;2 2;2 3]; D = dummyvar(group)</pre>					
	D =					
		1	0	1	0	0
		1	0	0	1	0
		1	0	0	0	1
		0	1	1	0	0
		0	1	0	1	0
		0	1	0	0	1
See Also	pinv,r	egress				

Purpose	Empirical (Kaplan-Meier) cumulative distribution function				
Syntax	<pre>[f,x] = ecdf(y) [f,x,flo,fup] = ecdf(y) [] = ecdf(y,'param1',value1,'param2',value2,)</pre>				
Description	<pre>[f,x] = ecdf(y) calculates the Kaplan-Meier estimate of the cumulative distribution function (cdf), also known as the empirical cdf. y is a vector of data values. f is a vector of values of the empirical cdf evaluated at x.</pre>				
	the cdf. These k	= ecdf(y) also returns lower and upper confidence bounds for bounds are calculated using Greenwood's formula, and are not onfidence bounds.			
		y, 'param1', value1, 'param2', value2,) specifies meter name-value pairs chosen from the following:			
	'censoring' Boolean vector of the same size as x. Elements are 1 for observations that are right-censored and 0 for observation that are observed exactly. Default is all observations observed exactly.				
	'frequency'	Vector of the same size as x containing non-negative integer counts. The jth element of this vector gives the number of times the jth element of x was observed. Default is 1 observation per element of x.			
	'alpha'	Value between 0 and 1 for a confidence level of 100*(1-alpha)%. Default is alpha=0.05 for 95% confidence.			
	'function'	Type of function returned as the f output argument, chosen from 'cdf' (default), 'survivor', or 'cumulative hazard'.			
Examples		m failure times and random censoring times, and compare the the known true cdf.			
	d = exprnd(t = min(y,d	<pre>10,50,1); % Random failure times are exponential(10) 20,50,1); % Drop-out times are exponential(20)); % Observe the minimum of these times (y>d); % Observe also whether the subject failed</pre>			

```
% Calculate and plot the empirical cdf and confidence bounds
[f,x,flo,fup] = ecdf(t,'censoring',censored);
stairs(x,f);
hold on;
stairs(x,flo,'r:'); stairs(x,fup,'r:');
```

```
% Superimpose a plot of the known true cdf
xx = 0:.1:max(t); yy = 1-exp(-xx/10); plot(xx,yy,'g-')
hold off;
```



See Also



References [1] Cox, D.R. and D. Oakes, *Analysis of Survival Data*, Chapman & Hall, London, 1984.

errorbar

Purpose	Plot error bars along a curve				
Syntax	errorbar(X,Y,L,U,symbol) errorbar(X,Y,L) errorbar(Y,L)				
Description	<pre>errorbar(X,Y,L,U,Symbol) plots X versus Y with error bars specified by L and U. X, Y, L, and U must be the same length. If X, Y, L, and U are matrices, then each column produces a separate line. The error bars are each drawn a distance of U(i) above and L(i) below the points in (X,Y). symbol is a string that controls the line type, plotting symbol, and color of the error bars. errorbar(X,Y,L) plots X versus Y with symmetric error bars about Y.</pre>				
	errorbar(Y,L) plots Y with error bars [Y-L Y+L].				
	The errorbar function is a part of the standard MATLAB language.				
Example	<pre>lambda = (0.1:0.2:0.5); r = poissrnd(lambda(ones(50,1),:)); [p,pci] = poissfit(r,0.001); L = p - pci(1,:) U = pci(2,:) - p errorbar(1:3,p,L,U,'+')</pre>				
	L = 0.1200 0.1600 0.2600				
	U = 0.2000 0.2200 0.3400				
	0.8				
	$\begin{array}{c} 0.4 \\ 0.2 \\ 0 \\ 0.5 \end{array} \begin{array}{c} 1 \\ 1.5 \end{array} \begin{array}{c} 2 \\ 2.5 \end{array} \begin{array}{c} 1 \\ 2.5 \end{array} \begin{array}{c} 1 \\ 1.5 \end{array} \begin{array}{c} 2 \\ 2.5 \end{array} \begin{array}{c} 3 \\ 3.5 \end{array}$				

ewmaplot

Purpose	Exponentially Weighted Moving Average (EWMA) chart for Statistical Process Control (SPC)
Syntax	ewmaplot(data) ewmaplot(data,lambda) ewmaplot(data,lambda,alpha) ewmaplot(data,lambda,alpha,specs) h = ewmaplot()
Description	ewmaplot(data) produces an EWMA chart of the grouped responses in data. The rows of data contain replicate observations taken at a given time. The rows should be in time order.
	ewmaplot(data,lambda) produces an EWMA chart of the grouped responses in data, and specifies how much the current prediction is influenced by past observations. Higher values of lambda give more weight to past observations. By default, lambda = 0.4; lambda must be between 0 and 1.
	ewmaplot(data,lambda,alpha) produces an EWMA chart of the grouped responses in data, and specifies the significance level of the upper and lower plotted confidence limits. alpha is 0.0027 by default. This value produces three-sigma limits:
	norminv(1-0.0027/2)
	ans = 3
	To get k -sigma limits, use the expression 2*(1-normcdf(k)). For example, the correct alpha value for 2-sigma limits is 0.0455, as shown below.
	k = 2; 2*(1-normcdf(k))
	ans = 0.0455

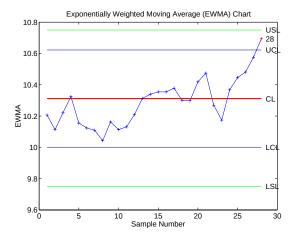
ewmaplot(data,lambda,alpha,specs) produces an EWMA chart of the grouped responses in data, and specifies a two-element vector, specs, for the lower and upper specification limits of the response.

h = ewmaplot(...) returns a vector of handles to the plotted lines.

Example

Consider a process with a slowly drifting mean. An EWMA chart is preferable to an x-bar chart for monitoring this kind of process. The simulation below demonstrates an EWMA chart for a slow linear drift.

t = (1:28)'; r = normrnd(10+0.02*t(:,ones(4,1)),0.5); ewmaplot(r,0.4,0.01,[9.75 10.75])



The EWMA value for group 28 is higher than would be expected purely by chance. If we had been monitoring this process continuously, we would have detected the drift when group 28 was collected, and we would have had an opportunity to investigate its cause.

- **Reference** [1] Montgomery, D., *Introduction to Statistical Quality Control*, John Wiley & Sons 1991. p. 299.
- See Also xbarplot, schart

expcdf

Purpose	Exponential cumulative distribution function (cdf)				
Syntax	P = expcdf(X,MU)				
Description	P = expcdf(X,MU) computes the exponential cdf at each of the values in X using the corresponding parameters in MU. Vector or matrix inputs for X and MU must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in MU must be positive.				
	The exponential cdf is				
	$p = F(x \mu) = \int_0^x \frac{1}{\mu} e^{-\frac{t}{\mu}} dt = 1 - e^{-\frac{x}{\mu}}$				
	The result, p , is the probability that a single observation from an exponential distribution will fall in the interval $[0 x]$.				
Examples	The median of the exponential distribution is $\mu * \log(2)$. Demonstrate this fact.				
	<pre>mu = 10:10:60; p = expcdf(log(2)*mu,mu)</pre>				
	p = 0.5000 0.5000 0.5000 0.5000 0.5000 0.5000				
	What is the probability that an exponential random variable will be less than or equal to the mean, μ ?				
	<pre>mu = 1:6; x = mu; p = expcdf(x,mu)</pre>				
	p = 0.6321 0.6321 0.6321 0.6321 0.6321 0.6321				
See Also	cdf, expfit, expinv, exppdf, exprnd, expstat				

Purpose	Parameter estimates and confidence intervals for exponential data		
Syntax	<pre>muhat = expfit(x) [muhat,muci] = expfit(x) [muhat,muci] = expfit(x,alpha)</pre>		
Description	muhat = expfit(x) returns the estimate of the parameter, μ , of the exponential distribution given data x.		
	[muhat,muci] = expfit(x) also returns the 95% confidence interval in muci.		
	[muhat,muci] = expfit(x,alpha) gives 100(1-alpha)% confidence intervals. For example, alpha = 0.01 yields 99% confidence intervals.		
Example	We generate 100 independent samples of exponential data with $\mu = 3$. muhat is an estimate of true_mu and muci is a 99% confidence interval around muhat. Notice that muci contains true_mu.		
	true_mu = 3; [muhat,muci] = expfit(r,0.01)		
	muhat =		
	2.8835		
	muci =		
	2.1949 3.6803		
See Also	expcdf, expinv, exppdf, exprnd, expstat, betafit, binofit, gamfit, normfit, poissfit, unifit, weibfit		

expinv

Purpose	Inverse of the exponential cumulative distribution function (cdf)		
Syntax	X = expinv(P,MU)		
Description	X = expinv(P,MU) computes the inverse of the exponential cdf with parameters specified by MU for the corresponding probabilities in P. Vector or matrix inputs for P and MU must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in MU must be positive and the values in P must lie on the interval [0 1].		
	The inverse of the exponential cdf is		
	$x = F(p \mu) = -\mu \ln(1-p)$		
	The result, x , is the value such that an observation from an exponential distribution with parameter μ will fall in the range $[0 x]$ with probability p .		
Examples	Let the lifetime of light bulbs be exponentially distributed with μ = 700 hours. What is the median lifetime of a bulb?		
	expinv(0.50,700)		
	ans =		
	485.2030		
	So, suppose you buy a box of "700 hour" light bulbs. If 700 hours is the mean life of the bulbs, then half them will burn out in less than 500 hours.		
See Also	expcdf, expfit, exppdf, exprnd, expstat, icdf		

Purpose	Exponential probability density function (pdf)					
Syntax	Y = exppdf(X,MU)					
Description	exppdf(X,MU) computes the exponential pdf at each of the values in X using the corresponding parameters in MU. Vector or matrix inputs for X and MU must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in MU must be positive.					
	The exponential pdf is					
	$y = f(x \mu) = \frac{1}{\mu}e^{-\frac{x}{\mu}}$					
	The exponential pdf is the gamma pdf with its first parameter equal to 1.					
	The exponential distribution is appropriate for modeling waiting times when the probability of waiting an additional period of time is independent of how long you've already waited. For example, the probability that a light bulb will burn out in its next minute of use is relatively independent of how many minutes it has already burned.					
Examples	y = exppdf(5,1:5)					
	y = 0.0067 0.0410 0.0630 0.0716 0.0736					
	y = exppdf(1:5,1:5)					
	y = 0.3679 0.1839 0.1226 0.0920 0.0736					
See Also	expcdf, expfit, expinv, exprnd, expstat, pdf					

exprnd

Purpose	Random numbers from the exponential distribution				
Syntax	R = exprnd(MU) R = exprnd(MU,m) R = exprnd(MU,m,n)				
Description	R = exprnd(MU) generates exponential random numbers with mean MU. The size of R is the size of MU.				
	R = exprnd(MU,m) generates exponential random numbers with mean MU, where m is a 1-by-2 vector that contains the row and column dimensions of R.				
	R = exprnd(MU,m,n) generates exponential random numbers with mean MU, where scalars m and n are the row and column dimensions of R.				
Examples	n1 = exprnd(5:10)				
	n1 = 7.5943 18.3400 2.7113 3.0936 0.6078 9.5841				
	n2 = exprnd(5:10,[1 6])				
	n2 = 3.2752 1.1110 23.5530 23.4303 5.7190 3.9876				
	n3 = exprnd(5,2,3)				
	n3 = 24.3339 13.5271 1.8788 4.7932 4.3675 2.6468				
See Also	expcdf, expfit, expinv, exppdf, expstat				

Purpose	Mean and variance for the exponential distribution					
Syntax	[M,V] = ex	pstat(MU)				
Description	$[M,V] = expstat(MU)$ returns the mean and variance for the exponential distribution with parameters MU. The mean of the exponential distribution is μ , and the variance is μ^2 .					
Examples	[m,v] =	expstat([1 10 100 10	00])		
	m =		10	100	1000	
		1	10	100	1000	
	V =	1	100	10000	1000000	
See Also	expcdf, expfit, expinv, exppdf, exprnd					

Purpose	Maximum Likelihood Common Factor Analysis
Syntax	<pre>lambda = factoran(X,m) [lambda,psi] = factoran(X,m) [lambda,psi,T] = factoran(X,m) [lambda,psi,T,stats] = factoran(X,m) [lambda,psi,T,stats,F] = factoran(X,m) [] = factoran(,'param1',value1,'param2',value2,)</pre>
Definition	factoran computes the maximum likelihood estimate (MLE) of the factor loadings matrix Λ in the Factor Analysis model $x = \mu + \Lambda f + e$
	where x is a vector of observed variables, μ is a constant vector of means, Λ is a constant d-by-m matrix of factor loadings, <i>f</i> is a vector of independent, standardized common factors, and <i>e</i> is a vector of independent specific factors. <i>x</i> , μ , and <i>e</i> are of length d. <i>f</i> is of length m.
	Alternatively, the Factor Analysis model can be specified as
	$cov(x) = \Lambda \Lambda^T + \Psi$
	where $\Psi = cov(e)$ is a d-by-d diagonal matrix of specific variances.
Description	<pre>lambda = factoran(X,m) returns the maximum likelihood estimate, lambda, of the factor loadings matrix, in a common factor analysis model with m common factors. X is an n-by-d matrix where each row is an observation of d variables. The (i,j)th element of the d-by-m matrix lambda is the coefficient, or loading, of the jth factor for the ith variable. By default, the estimated factor loadings are rotated using the 'varimax' option of the 'rotate' parameter.</pre>
	[lambda,psi] = factoran(X,m) also returns maximum likelihood estimates of the specific variances as a column vector psi of length d.
	<pre>[lambda,psi,T] = factoran(X,m) also returns the m-by-m factor loadings rotation matrix T.</pre>

[lambda,psi,T,stats] = factoran(X,m) also returns a structure stats containing information relating to the null hypothesis, H₀, that the number of common factors is m. stats includes the fields:

loglike	Maximized log-likelihood value
dfe	Error degrees of freedom = $((d-m)^2 - (d+m))/2$
chisq	$\label{eq:approximate} Approximate \ chi-squared \ statistic \ for \ the \ null \ hypothesis$
р	Right-tail significance level for the null hypothesis

factoran does not compute the chisq and p fields unless dfe is positive, and all the specific valiance estimates in psi are positive (see "Heywood Case" below). If X is a covariance matrix, then you must also specify the 'nobs' parameter if you want factoran to compute the chisq and p fields.

[lambda,psi,T,stats,F] = factoran(X,m) also returns, in F, predictions of the common factors, known as factor scores. F is an n-by-m matrix where each row is a prediction of m common factors. If X is a covariance matrix, factoran cannot compute F. factoran rotates F using the same criterion as for lambda.

 $[\ldots] = factoran(\ldots, 'param1', value1, 'param2', value2, \ldots)$ enables you to specify optional parameter name/value pairs to control the model fit and the outputs. These are the valid parameters. The most commonly used parameters are listed first.

Parameter	Value	Value		
'xtype'	The type of input in the matrix X. 'xtype' can be one of:			
	'data'	Raw data (default)		
	'covariance'	Positive definite covariance or correlation matrix		
'scores'	-	Method for predicting factor scores. 'scores' is ignored if X is not raw data.		

Parameter (Continued)	Value			
	'wls' 'Bartlett'	Synonyms for a weighted least squares estimate that treats F as fixed (default)		
	'regression' 'Thomson'	Synonyms for a minimum mean squared error prediction that is equivalent to a ridge regression		
'start'	Starting point for the specific variances psi in the maximum likelihood optimization. May be specified as:			
	'random'	Chooses d uniformly distributed values on the interval [0,1].		
	'Rsquared'	Chooses the starting vector as a scale factor times diag(inv(corrcoef(X))) (default). E.g., see Jöreskog [2].		
	Positive integer	Performs the given number of maximum likelihood fits, each initialized as with 'random'. factoran returns the fit with the highest likelihood.		
	Matrix	Performs one maximum likelihood fit for each column of the specified matrix. The ith optimization is initialized with the values from the ith column. The matrix must have d rows.		
'rotate'	Method used to	rotate factor loadings and scores.		
	'none'	Performs no rotation		

Parameter (Continued)	Value			
	'orthomax'	Orthogonal rotation that maximizes a criterion based on the variance of the loadings. Use the 'coeffom', 'normalize', 'reltol', and 'maxit' parameters to control the details of the rotation.		
	'varimax'	Special case of the orthomax rotation (default). Use the 'normalize', 'reltol', and 'maxit' parameters to control the details of the rotation.		
	'procrustes'	 Performs either an oblique (the default) or an orthogonal rotation to best match a specified target matrix in the least squares sense. Use the 'typeprocr' parameter to choose the type of rotation. Use 'targetprocr' to specify the target matrix. 		
	'promaxpm'	Performs an oblique procrustes rotation to a target matrix determined by factoran as a function of an orthomax solution. Use the 'powerpm' parameter to specify the exponent for creating the target matrix. Because 'promaxpm' uses 'orthomax' internally, you can also specify the parameters that apply to 'orthomax'.		

Parameter (Continued)	Value			
	Function	<pre>Function handle to rotation function of the form [B,T] = myrotation(A,)</pre>		
		where A is a d-by-m matrix of unrotated factor loadings. B is a d-by-m matrix of rotated loadings, and T is the corresponding m-by-m rotation matrix.		
		Use the factoran parameter 'userargs' to pass additional arguments to this rotation function. See Example 4.		
'coeffom'	Coefficient, often denoted as γ , defining the specific 'orthomax' criterion. Must be between 0 and 1. The value 0 corresponds to quartimax, and 1 corresponds to varimax. Default is 1.			
'normalize'	Flag indicating whether the loading matrix should be row-normalized (1) or left unnormalized (0) for 'orthomax' or 'varimax' rotation. Default is 1.			
'reltol'	Relative convergence tolerance for 'orthomax' or 'varimax' rotation. Default is sqrt(eps).			
'maxit'	Iteration limit for 'orthomax' or 'varimax' rotation. Default is 250.			
'targetprocr'	Target factor loading matrix for 'procrustes' rotation. Required for 'procrustes' rotation. No default value.			
'typeprocr'		Type of 'procrustes' rotation. Can be 'oblique' (default) or 'orthogonal'.		
'powerpm'	Exponent for creating the target matrix in the 'promaxpm' rotation. Must be >= 1. Default is 4.			

Parameter (Continued)	Value				
'userargs'	Denotes the beginning of additional input values for a user-defined rotation function. factoran appends all subsequent values, in order and without processing, to the rotation function argument list, following the unrotated factor loadings matrix A. See Example 4.				
'nobs'	number of obser estimation. This the null hypothe	If X is a covariance or correlation matrix, indicates the number of observations that were used in its estimation. This allows calculation of significance for the null hypothesis even when the original data are not available. There is no default. 'nobs' is ignored if X is raw data.			
'delta'	A lower bound for the specific variances psi during the maximum likelihood optimization. Default is 0.005.				
'optimopts'	likelihood optim	ning control options for the maximum ization. Create this structure with the on optimset. factoran uses the ts:			
	'Display'	'notify'			
	'MaxFunEvals '	100*d			
	'MaxIter'	400			
	'TolFun'	1e-8			
	'TolX'	1e-8			

Remarks **Observed Data Variables.** The variables in the observed data matrix X must be

linearly independent, i.e., cov(X) must have full rank, for maximum likelihood estimation to succeed. factoran reduces both raw data and a covariance matrix to a correlation matrix before performing the fit.

factoran standardizes the observed data X to zero mean and unit variance before estimating the loadings lambda. This does not affect the model fit,

because MLEs in this model are invariant to scale. However, lambda and psi are returned in terms of the standardized variables, i.e., lambda*lambda'+diag(psi) is an estimate of the correlation matrix of the original data X (although not after an oblique rotation). See Examples 1 and 3.

Heywood Case. If elements of psi are equal to the value of the 'delta' parameter (i.e, they are essentially zero), the fit is known as a Heywood case, and interpretation of the resulting estimates is problematic. In particular, there may be multiple local maxima of the likelihood, each with different estimates of the loadings and the specific variances. Heywood cases can indicate overfitting (i.e., m is too large), but can also be the result of underfitting.

Rotation of Factor Loadings and Scores. Unless you explicitly specify no rotation using the 'rotate' parameter, factoran rotates the estimated factor loadings, lambda, and the factor scores, F. The output matrix T is used to rotate the loadings, i.e., lambda = lambda0*T, where lambda0 is the initial (unrotated) MLE of the loadings. T is an orthogonal matrix for orthogonal rotations, and the identity matrix for no rotation. The inverse of T is known as the primary axis rotation matrix, while T itself is related to the reference axis rotation matrix. For orthogonal rotations, the two are identical.

factor an computes factor scores that have been rotated by inv(T'), i.e., F = F0 * inv(T'), where F0 contains the unrotated predictions. The estimated covariance of F is inv(T'*T), which, for orthogonal or no rotation, is the identity matrix. Rotation of factor loadings and scores is an attempt to create a more easily interpretable structure in the loadings matrix after maximum likelihood estimation.

Examples Example 1. Load the carbig data, and fit the default model with two factors.

load carbig
X = [Acceleration Displacement Horsepower MPG Weight];
X = X(all(~isnan(X),2),:);

```
[Lambda,Psi,T,stats,F] = factoran(X,2,'scores','regression')
inv(T'*T) % Estimated correlation matrix of F, == eye(2)
Lambda*Lambda' + diag(Psi) % Estimated correlation matrix of X
Lambda*inv(T) % Unrotate the loadings
F*T' % Unrotate the factor scores
```

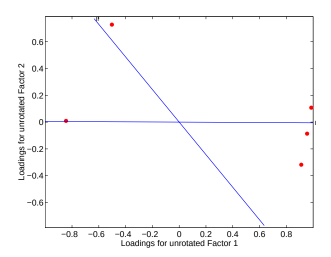
Example 2. Although the estimates are the same, the use of a covariance matrix rather than raw data doesn't let you request scores or significance level.

```
[Lambda,Psi,T] = factoran(cov(X),2,'xtype','cov')
[Lambda,Psi,T] = factoran(corrcoef(X),2,'xtype','cov')
```

Example 3. Use promax rotation.

Plot the rotated variables against the oblique axes.

```
invT = inv(T)
Lambda0 = Lambda*invT
plot(Lambda0(:,1),Lambda0(:,2), 'ro');
line([-invT(1,1) invT(1,1) NAN -invT(2,1) invT(2,1)], ...
     [-invT(1,2) invT(1,2) NAN -invT(2,2) invT(2,2)]);
text(invT(:,1), invT(:,2),[' I '; ' II']);
xlabel('Loadings for unrotated Factor 1')
ylabel('Loadings for unrotated Factor 2')
```



	<pre>Example 4. Syntax for passing additional arguments to a user-defined rotation function. [Lambda,Psi,T] = factoran(X,2,'rotate',@myrotation,'userargs',1,'two')</pre>			
See Also	optimset, princomp, procrustes			
References	[1] Harman, H.H., <i>Modern Factor Analysis</i> , 3rd Ed., University of Chicago Press, Chicago, 1976.			
	[2] Jöreskog, K.G., "Some Contributions to Maximum Likelihood Factor Analysis", <i>Psychometrika</i> , Vol.32, pp.443-482, 1967.			
	[3] Lawley, D.N. and A.E. Maxwell, <i>Factor Analysis as a Statistical Method</i> , 2nd Ed., American Elsevier Pub. Co., New York, 1971.			

Purpose F cumulative distribution function (cdf)

Syntax P = fcdf(X,V1,V2)

Description P = fcdf(X,V1,V2) computes the F cdf at each of the values in X using the corresponding parameters in V1 and V2. Vector or matrix inputs for X, V1, and V2 must all be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in V1 and V2 must be positive integers.

The F cdf is

$$p = F(x|v_1, v_2) = \int_0^x \frac{\Gamma\left[\frac{(v_1 + v_2)}{2}\right]}{\Gamma\left(\frac{v_1}{2}\right)\Gamma\left(\frac{v_2}{2}\right)} \left(\frac{v_1}{v_2}\right)^{\frac{v_1}{2}} \frac{t^{\frac{v_1 - 2}{2}}}{\left[1 + \left(\frac{v_1}{v_2}\right)t\right]^{\frac{v_1 + v_2}{2}}} dt$$

The result, p, is the probability that a single observation from an F distribution with parameters v_1 and v_2 will fall in the interval [0 x].

Examples This example illustrates an important and useful mathematical identity for the F distribution.

nu1 = 1:5;nu2 = 6:10;x = 2:6;F1 = fcdf(x, nu1, nu2)F1 = 0.7930 0.8854 0.9481 0.9788 0.9919 F2 = 1 - fcdf(1./x, nu2, nu1)F2 = 0.7930 0.8854 0.9481 0.9788 0.9919

See Also cdf, finv, fpdf, frnd, fstat

ff2n

Purpose	Two-level full-factorial designs			
Syntax	X = ff2n(n)			
Description	$X = ff2n(n)$ creates a two-level full-factorial design, X, where n is the desired number of columns of X. The number of rows in X is 2^n .			
Example	X = ff2n(3)			
	X = 0 0 0			
	0 0 1 0 1 0			
	0 1 1			
	1 0 0			
	1 0 1			
	1 1 0			
	1 1 1			
	\boldsymbol{X} is the binary representation of th	e numbers from 0 to 2^n -1.		
See Also	fracfact, fullfact			

Purpose Inverse of the F cumulative distribution function (cdf)

Syntax X = finv(P,V1,V2)

Description X = finv(P,V1,V2) computes the inverse of the F cdf with numerator degrees of freedom V1 and denominator degrees of freedom V2 for the corresponding probabilities in P. Vector or matrix inputs for P, V1, and V2 must all be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.

The parameters in V1 and V2 must all be positive integers, and the values in P must lie on the interval [0 1].

The F inverse function is defined in terms of the F cdf as

$$x = F^{-1}(p|v_1,v_2) = \{x: F(x|v_1,v_2) = p\}$$

where

$$p = F(x|v_1, v_2) = \int_0^x \frac{\Gamma\left[\frac{(v_1 + v_2)}{2}\right]}{\Gamma\left(\frac{v_1}{2}\right)\Gamma\left(\frac{v_2}{2}\right)} \left(\frac{v_1}{v_2}\right)^{\frac{v_1}{2}} \frac{t^{\frac{v_1 - 2}{2}}}{\left[1 + \left(\frac{v_1}{v_2}\right)t\right]^{\frac{v_1 + v_2}{2}}} dt$$

Examples Find a value that should exceed 95% of the samples from an F distribution with 5 degrees of freedom in the numerator and 10 degrees of freedom in the denominator.

You would observe values greater than 3.3258 only 5% of the time by chance.

See Also fcdf, fpdf, frnd, fstat, icdf

Purpose	F probability density function	(pdf)
---------	--------------------------------	-------

Syntax Y = fpdf(X, V1, V2)

Description Y = fpdf(X,V1,V2) computes the F pdf at each of the values in X using the corresponding parameters in V1 and V2. Vector or matrix inputs for X, V1, and V2 must all be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in V1 and V2 must all be positive integers, and the values in X must lie on the interval $[0 \infty)$.

The probability density function for the F distribution is

$$y = f(x|v_1, v_2) = \frac{\Gamma\left[\frac{(v_1 + v_2)}{2}\right]}{\Gamma\left(\frac{v_1}{2}\right)\Gamma\left(\frac{v_2}{2}\right)} \left(\frac{v_1}{v_2}\right)^{\frac{v_1}{2}} \frac{x^{\frac{v_1 - 2}{2}}}{\left[1 + \left(\frac{v_1}{v_2}\right)x\right]^{\frac{v_1 + v_2}{2}}}$$

Examples	y = fpdf(1:6,2,2)					
	y = 0.2500	0.1111	0.0625	0.0400	0.0278	0.0204
	z = fpdf(3,5:10,5:10)					
	z = 0.0689	0.0659	0.0620	0.0577	0.0532	0.0487
See Also	fcdf,finv,frnd,f	stat, pdf				

Purpose	Generate fractional factorial design from generators					
Syntax	x = fracfact('gen') [x,conf] = fracfact('gen')					
Description	x = fracfact('gen') generates a fractional factorial design as specified by the generator string gen, and returns a matrix x of design points. The input string gen is a generator string consisting of "words" separated by spaces. Each word describes how a column of the output design should be formed from columns of a full factorial. Typically gen will include single-letter words for the first few factors, plus additional multiple-letter words describing how the remaining factors are confounded with the first few.					
	The output matrix x is a fraction of a two-level full-factorial design. Suppose there are m words in gen, and that each word is formed from a subset of the first n letters of the alphabet. The output matrix x has 2^n rows and m columns. Let F represent the two-level full-factorial design as produced by ff2n(n). The values in column j of x are computed by multiplying together the columns of F corresponding to letters that appear in the j th word of the generator string.					
	[x,conf] = fracfact('gen') also returns a cell array, conf, that describes the confounding pattern among the main effects and all two-factor interactions.					
Examples	 Example 1 We want to run an experiment to study the effects of four factors on a response, but we can only afford eight runs. (A run is a single repetition of the experiment at a specified combination of factor values.) Our goal is to determine which factors affect the response. There may be interactions between some pairs of factors. A total of sixteen runs would be required to test all factor combinations. However, if we are willing to assume there are no three-factor interactions, we can estimate the main factor effects in just eight runs. 					
	[x,conf] = fracfact('a b c abc')					
	x = -1 -1 -1 -1 -1 -1 1 1					

fracfact

- 1	1	- 1	1	
- 1	1	1	- 1	
1	- 1	- 1	1	
1	- 1	1	- 1	
1	1	- 1	- 1	
1	1	1	1	
conf =				
'Ter	·m '	'Gene	rator'	'Confounding'
'X1 '		'a'		'X1'
'X2'		'b'		'X2'
'X3 '		'c'		'X3'
'X4 '		'abc'		'X4 '
'X1*	X2 '	'ab'		'X1*X2 + X3*X4'
'X1*	X3 '	'ac'		'X1*X3 + X2*X4'
'X1*	X4 '	'bc'		'X1*X4 + X2*X3'
'X2*	X3 '	'bc'		'X1*X4 + X2*X3'
'X2*	X4 '	'ac'		'X1*X3 + X2*X4'
'X3*	X4 '	'ab'		'X1*X2 + X3*X4'

The first three columns of the x matrix form a full-factorial design. The final column is formed by multiplying the other three. The confounding pattern shows that the main effects for all four factors are estimable, but the two-factor interactions are not. For example, the X1*X2 and X3*X4 interactions are confounded, so it is not possible to estimate their effects separately.

After conducting the experiment, we may find out that the 'ab' effect is significant. In order to determine whether this effect comes from X1*X2 or X3*X4 we would have to run the remaining eight runs. We can obtain those runs by reversing the sign of the final generator.

```
fracfact('a b c -abc')
ans =
     - 1
             - 1
                    - 1
                             1
     - 1
            - 1
                    1
                            - 1
     - 1
             1
                    -1
                            - 1
             1
                    1
     - 1
                             1
      1
            -1
                    -1
                            - 1
      1
            - 1
                    1
                             1
      1
             1
                    -1
                             1
```

1 1 1 -1

Example 2

Suppose now we need to study the effects of eight factors. A full factorial would require 256 runs. By clever choice of generators, we can find a sixteen-run design that can estimate those eight effects with no confounding from two-factor interactions.

```
[x,c] = fracfact('a b c d abc acd abd bcd');
c(1:10,:)
```

```
ans =
```

'Term'	'Generator'	'Confounding'
'X1'	'a'	'X1'
'X2'	'b'	'X2'
' X3 '	'C'	' X3 '
' X4 '	'd'	' X4 '
' X5 '	'abc'	' X5 '
' X6 '	'acd'	'X6'
' X7 '	'abd'	' X7 '
' X8 '	'bcd'	'X8'
'X1*X2'	'ab'	'X1*X2 + X3*X5 + X4*X7 + X6*X8'

This confounding pattern shows that the main effects are not confounded with two-factor interactions. The final row shown reveals that a group of four two-factor interactions is confounded. Other choices of generators would not have the same desirable property.

```
[x,c] = fracfact('a b c d ab cd ad bc');
c(1:10,:)
```

ans =

'Term'	'Generator'	'Confounding'
'X1'	'a'	'X1 + X2*X5 + X4*X7'
'X2'	'b'	'X2 + X1*X5 + X3*X8'
' X3 '	' C '	'X3 + X2*X8 + X4*X6'
' X4 '	' d '	'X4 + X1*X7 + X3*X6'
' X5 '	'ab'	'X5 + X1*X2'
'X6'	'cd'	'X6 + X3*X4'
' X7 '	'ad'	'X7 + X1*X4'

fracfact

'X8 '	'bc'	'X8 + X2*X3'
'X1*X2'	'ab'	'X5 + X1*X2'
Here all the main	effects ar	e confounded with one or more two-factor
interactions.		

References [1] Box, G. A. F., W. G. Hunter, and J. S. Hunter (1978), *Statistics for Experimenters*, Wiley, New York.

See Also ff2n, fullfact, hadamard

Purpose	Friedman's nonparametric two-way Analysis of Variance (ANOVA)
Syntax	<pre>p = friedman(X,reps) p = friedman(X,reps,'displayopt') [p,table] = friedman() [p,table,stats] = friedman()</pre>
Description	<pre>p = friedman(X,reps) performs the nonparametric Friedman's test to compare the means of the columns of X. Friedman's test is similar to classical two-way ANOVA, but it tests only for column effects after adjusting for possible row effects. It does not test for row effects or interaction effects. Friedman's test is appropriate when columns represent treatments that are under study, and rows represent nuisance effects (blocks) that need to be taken into account but are not of any interest.</pre>
	The different columns represent changes in factor A. The different rows represent changes in the blocking factor B. If there is more than one observation for each combination of factors, input reps indicates the number of replicates in each "cell," which must be constant.
	The matrix below illustrates the format for a set-up where column factor A has three levels, row factor B has two levels, and there are two replicates (reps=2).

The matrix below illustrates the format for a set-up where column factor A has three levels, row factor B has two levels, and there are two replicates (reps=2). The subscripts indicate row, column, and replicate, respectively.

 $\begin{bmatrix} x_{111} & x_{121} & x_{131} \\ x_{112} & x_{122} & x_{132} \\ x_{211} & x_{221} & x_{231} \\ x_{212} & x_{222} & x_{232} \end{bmatrix}$

Friedman's test assumes a model of the form

 $x_{ijk} = \mu + \alpha_i + \beta_j + \varepsilon_{ijk}$

where μ is an overall location parameter, α_i represents the column effect, β_j represents the row effect, and ε_{ijk} represents the error. This test ranks the data within each level of B, and tests for a difference across levels of A. The p that friedman returns is the p-value for the null hypothesis that $\alpha_i = 0$. If the p-value is near zero, this casts doubt on the null hypothesis. A sufficiently

friedman

small p-value suggests that at least one column-sample mean is significantly different than the other column-sample means; i.e., there is a main effect due to factor A. The choice of a limit for the p-value to determine whether a result is "statistically significant" is left to the researcher. It is common to declare a result significant if the p-value is less than 0.05 or 0.01.

friedman also displays a figure showing an ANOVA table, which divides the variability of the ranks into two or three parts:

- The variability due to the differences among the column means
- The variability due to the interaction between rows and columns (if reps is greater than its default value of 1)
- The remaining variability not explained by any systematic source

The ANOVA table has six columns:

- The first shows the source of the variability.
- The second shows the Sum of Squares (SS) due to each source.
- The third shows the degrees of freedom (df) associated with each source.
- The fourth shows the Mean Squares (MS), which is the ratio SS/df.
- The fifth shows Friedman's chi-square statistic.
- The sixth shows the p-value for the chi-square statistic.

p = friedman(X,reps,'displayopt') enables the ANOVA table display when 'displayopt' is 'on' (default) and suppresses the display when 'displayopt' is 'off'.

[p,table] = friedman(...) returns the ANOVA table (including column and row labels) in cell array table. (You can copy a text version of the ANOVA table to the clipboard by selecting **Copy Text** from the **Edit** menu.

[p,table,stats] = friedman(...) returns a stats structure that you can use to perform a follow-up multiple comparison test. The friedman test evaluates the hypothesis that the column effects are all the same against the alternative that they are not all the same. Sometimes it is preferable to perform a test to determine which pairs of column effects are significantly different, and which are not. You can use the multcompare function to perform such tests by supplying the stats structure as input.

Examples

Let's repeat the example from the anova2 function, this time applying Friedman's test. Recall that the data below come from a study of popcorn brands and popper type (Hogg 1987). The columns of the matrix popcorn are brands (Gourmet, National, and Generic). The rows are popper type (Oil and Air). The study popped a batch of each brand three times with each popper. The values are the yield in cups of popped popcorn.

load popcorn popcorn				
popcorn =				
5.5000	4.5000	3.5000		
5.5000	4.5000	4.0000		
6.0000	4.0000	3.0000		
6.5000	5.0000	4.0000		
7.0000	5.5000	5.0000		
7.0000	5.0000	4.5000		
p = friedman	(popcorn,3)			
p =				
0.0010				

🛃 Figure No. 2:	Friedman's Test					_ 🗆 🗵
<u>F</u> ile <u>E</u> dit <u>V</u> iew	<u>I</u> nsert <u>T</u> ools <u>W</u> ir	ndow	<u>H</u> elp			
	Fi	riedr	nan's ANOV	/A Table		
Source	SS	df	MS	Chi-sq	Prob>Chi-sq	A
Columns Interaction Error Total	99.75 0.0833 16.1667 116	2 2 12 17	0.0417	13.76	0.001	1
1	Test for	colum	n effects after row a	effects are remo	ved	

The small p-value of 0.001 indicates the popcorn brand affects the yield of popcorn. This is consistent with the results from anova2.

We could also test popper type by permuting the popcorn array as described on "Friedman's Test" on page 4-34 and repeating the test.

References [1] Hogg, R. V. and J. Ledolter. *Engineering Statistics*. MacMillan Publishing Company, 1987.

friedman

Hollander, M. and D. A. Wolfe. *Nonparametric Statistical Methods*. Wiley, 1973.

See Also anova2, multcompare

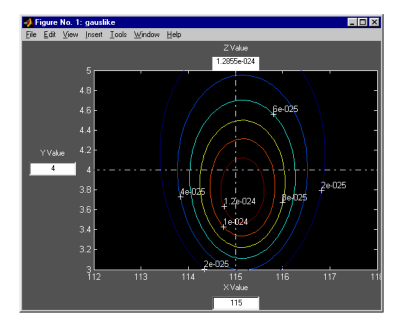
Purpose	Random numbers from the F distribution
Syntax	R = frnd(V1,V2) R = frnd(V1,V2,m) R = frnd(V1,V2,m,n)
Description	R = frnd(V1,V2) generates random numbers from the F distribution with numerator degrees of freedom V1 and denominator degrees of freedom V2. Vector or matrix inputs for V1 and V2 must have the same size, which is also the size of R. A scalar input for V1 or V2 is expanded to a constant matrix with the same dimensions as the other input.
	R = frnd(V1,V2,m) generates random numbers from the F distribution with parameters V1 and V2, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = frnd(V1,V2,m,n) generates random numbers from the F distribution with parameters V1 and V2, where scalars m and n are the row and column dimensions of R.
Examples	n1 = frnd(1:6,1:6)
	n1 = 0.0022 0.3121 3.0528 0.3189 0.2715 0.9539
	n2 = frnd(2,2,[2 3])
	n2 = 0.3186 0.9727 3.0268 0.2052 148.5816 0.2191
	n3 = frnd([1 2 3;4 5 6],1,2,3)
	n3 = 0.6233 0.2322 31.5458 2.5848 0.2121 4.4955
See Also	fcdf, finv, fpdf, fstat

fstat

Purpose	Mean and variance for the F distribution			
Syntax	[M,V] = fstat(V1,V2)			
Description	[M,V] = fstat(V1,V2) returns the mean and variance for the F distribution with parameters specified by V1 and V2. Vector or matrix inputs for V1 and V2 must have the same size, which is also the size of M and V. A scalar input for V1 or V2 is expanded to a constant matrix with the same dimensions as the other input.			
	The mean of the F distribution for values of v_2 greater than 2 is			
	$\frac{\nu_2}{\nu_2-2}$			
	The variance of the F distribution for values of v_2 greater than 4 is			
	$\frac{2{v_2}^2(v_1+v_2-2)}{v_1(v_2-2)^2(v_2-4)}$			
	The mean of the F distribution is undefined if v_2 is less than 3. The variance is			
	undefined for v_2 less than 5.			
Examples	fstat returns NaN when the mean and variance are undefined.			
	[m,v] = fstat(1:5,1:5)			
	m = NaN NaN 3.0000 2.0000 1.6667			
	v = NaN NaN NaN NaN 8.8889			
See Also	fcdf, finv, frnd, frnd			

Purpose	Interactive contour plot of a function
Syntax	fsurfht('fun',xlims,ylims) fsurfht('fun',xlims,ylims,p1,p2,p3,p4,p5)
Description	fsurfht('fun',xlims,ylims) is an interactive contour plot of the function specified by the text variable fun. The <i>x</i> -axis limits are specified by xlims in the form [xmin xmax], and the <i>y</i> -axis limits are specified by ylims in the form [ymin ymax].
	fsurfht('fun',xlims,ylims,p1,p2,p3,p4,p5) allows for five optional parameters that you can supply to the function fun.
	The intersection of the vertical and horizontal reference lines on the plot defines the current <i>x</i> -value and <i>y</i> -value. You can drag these reference lines and watch the calculated <i>z</i> -values (at the top of the plot) update simultaneously. Alternatively, you can type the <i>x</i> -value and <i>y</i> -value into editable text fields on the <i>x</i> -axis and <i>y</i> -axis.
Example	Plot the Gaussian likelihood function for the gas.mat data.
	load gas
	Create a function containing the following commands, and name it gauslike.m.
	<pre>function z = gauslike(mu,sigma,p1) n = length(p1); z = ones(size(mu)); for i = 1:n z = z .* (normpdf(p1(i),mu,sigma)); end</pre>
	The gauslike function calls normpdf, treating the data sample as fixed and the parameters μ and σ as variables. Assume that the gas prices are normally

distributed, and plot the likelihood surface of the sample.
fsurfht('gauslike',[112 118],[3 5],price1)



The sample mean is the *x*-value at the maximum, but the sample standard deviation is *not* the *y*-value at the maximum.

```
mumax = mean(price1)
mumax =
    115.1500
sigmamax = std(price1)*sqrt(19/20)
sigmamax =
    3.7719
```

Purpose	Full-factorial experimental design		
Syntax	<pre>design = fullfact(levels)</pre>		
Description	design = fullfact(levels) give the factor settings for a full factorial design. Each element in the vector levels specifies the number of unique values in the corresponding column of design.		
	For example, if the first element of levels is 3, then the first column of design contains only integers from 1 to 3.		
Example	If levels = [2 4], fullfact generates an eight-run design with two levels in the first column and four in the second column.		
	<pre>d = fullfact([2 4])</pre>		
	d =		
	1 1		
	2 1		
	1 2		
	2 2		
	1 3		
	2 3		
	1 4		
	2 4		
See Alee			

See Also

ff2n, dcovary, daugment, cordexch

gamcdf

Purpose	Gamma	cumulative	distribution	function (cdf)
---------	-------	------------	--------------	------------	------

Syntax P = gamcdf(X,A,B)

Description gamcdf(X,A,B) computes the gamma cdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must be positive.

The gamma cdf is

$$p = F(x|a, b) = \frac{1}{b^{a}\Gamma(a)}\int_{0}^{x} t^{a-1}e^{-\frac{t}{b}}dt$$

The result, p, is the probability that a single observation from a gamma distribution with parameters a and b will fall in the interval [0 x].

gammainc is the gamma distribution with b fixed at 1.

Examples

The mean of the gamma distribution is the product of the parameters, *ab*. In this example, the mean approaches the median as it increases (i.e., the distribution becomes more symmetric).

See Also cdf, gamfit, gaminv, gamlike, gampdf, gamrnd, gamstat

Purpose	Parameter estimates and confidence intervals for gamma distributed data
Syntax	phat = gamfit(x) [phat,pci] = gamfit(x) [phat,pci] = gamfit(x,alpha)
Description	phat = $gamfit(x)$ returns the maximum likelihood estimates (MLEs) for the parameters of the gamma distribution given the data in vector x.
	<pre>[phat,pci] = gamfit(x) returns MLEs and 95% percent confidence intervals. The first row of pci is the lower bound of the confidence intervals; the last row is the upper bound.</pre>
	<pre>[phat,pci] = gamfit(x,alpha) returns 100(1-alpha)% confidence intervals. For example, alpha = 0.01 yields 99% confidence intervals.</pre>
Example	Note that the 95% confidence intervals in the example below bracket the true parameter values of 2 and 4.
	a = 2; b = 4; r = gamrnd(a,b,100,1); [p,ci] = gamfit(r)
	p = 2.1990 3.7426
	ci = 1.6840 2.8298 2.7141 4.6554
Reference	[1] Hahn, G. J. and S.S. Shapiro. <i>Statistical Models in Engineering</i> . John Wiley & Sons, New York. 1994. p. 88.
See Also	gamcdf, gaminv, gamlike, gampdf, gamrnd, gamstat, betafit, binofit, expfit, normfit, poissfit, unifit, weibfit

gaminv

Purpose	Inverse of the gamma cumulative distribution function (cdf)			
Syntax	X = gaminv(P,A,B)			
Description	X = gaminv(P,A,B) computes the inverse of the gamma cdf with parameters A and B for the corresponding probabilities in P. Vector or matrix inputs for P, A, and B must all be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must all be positive, and the values in P must lie on the interval [0 1].			
	The gamma inverse function in terms of the gamma cdf is			
	$x = F^{-1}(p a,b) = \{x:F(x a,b) = p\}$			
	where			
	$p = F(x a, b) = \frac{1}{b^{a} \Gamma(a)} \int_{0}^{x} t^{a-1} e^{-\frac{t}{b}} dt$			
Algorithm	There is no known analytical solution to the integral equation above. gaminv uses an iterative approach (Newton's method) to converge on the solution.			
Examples	This example shows the relationship between the gamma cdf and its inverse function.			
	a = 1:5; b = 6:10; x = gaminv(gamcdf(1:5,a,b),a,b)			
	x = 1.0000 2.0000 3.0000 4.0000 5.0000			
See Also	gamcdf, gamfit, gamlike, gampdf, gamrnd, gamstat, icdf			

Purpose	Negative gamma log-likelihood function
Syntax	logL = gamlike(params,data) [logL,avar] = gamlike(params,data)
Description	<pre>logL = gamlike(params,data) returns the negative of the gamma log-likelihood function for the parameters, params, given data. The length of output vector logL is the length of vector data. [logL,avar] = gamlike(params,data) also returns avar, which is the asymptotic variance-covariance matrix of the parameter estimates when the values in params are the maximum likelihood estimates. avar is the inverse of Fisher's information matrix. The diagonal elements of avar are the asymptotic variances of their respective parameters.</pre>
	gamlike is a utility function for maximum likelihood estimation of the gamma distribution. Since gamlike returns the negative gamma log-likelihood function, minimizing gamlike using fminsearch is the same as maximizing the likelihood.
Example	This example continues the example for gamfit.
	a = 2; b = 3; r = gamrnd(a,b,100,1); [logL,info] = gamlike([2.1990 2.8069],r)
	logL =
	267.5585
	info =
	0.0690 -0.0790 -0.0790 0.1220
See Also	betalike, gamcdf, gamfit, gaminv, gampdf, gamrnd, gamstat, mle, normlike, weiblike

gampdf

PurposeGamma probability density function (pdf)

Syntax Y = gampdf(X,A,B)

Description gampdf(X,A,B) computes the gamma pdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must all be positive, and the values in X must lie on the interval $[0 \infty)$.

The gamma pdf is

$$y = f(x|a,b) = \frac{1}{b^{a}\Gamma(a)}x^{a-1}e^{-\frac{x}{b}}$$

The gamma probability density function is useful in reliability models of lifetimes. The gamma distribution is more flexible than the exponential distribution in that the probability of a product surviving an additional period may depend on its current age. The exponential and χ^2 functions are special cases of the gamma function.

Examples The exponential distribution is a special case of the gamma distribution.

mu = 1:5; y = gampdf(1	,1,mu)			
y = 0.3679	0.3033	0.2388	0.1947	0.1637
y1 = exppdf(1,mu)			
y1 = 0.3679	0.3033	0.2388	0.1947	0.1637

See Also gamcdf, gamfit, gaminv, gamlike, gamrnd, gamstat, pdf

gamrnd

Purpose	Random numbers from the gamma distribution					
Syntax	R = gamrnd(A,B) R = gamrnd(A,B,m) R = gamrnd(A,B,m,n)					
Description	R = gamrnd(A,B) generates gamma random numbers with parameters A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of R. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input.					
	R = gamrnd(A,B,m) generates gamma random numbers with parameters A and B, where m is a 1-by-2 vector that contains the row and column dimensions of R.					
	R = gamrnd(A,B,m,n) generates gamma random numbers with parameters A and B, where scalars m and n are the row and column dimensions of R.					
Examples	n1 = gamrnd(1:5,6:10)					
	n1 = 9.1132 12.8431 24.8025 38.5960 106.4164					
	n2 = gamrnd(5,10,[1 5])					
	n2 = 30.9486 33.5667 33.6837 55.2014 46.8265					
	n3 = gamrnd(2:6,3,1,5)					
	n3 = 12.8715 11.3068 3.0982 15.6012 21.6739					
See Also	gamcdf, gamfit, gaminv, gamlike, gampdf, gamstat					

gamstat

Purpose	Mean and variance for the gamma distribution			
Syntax	[M,V] = gamstat(A,B)			
Description	[M,V] = gamstat(A,B) returns the mean and variance for the gamma distribution with parameters specified by A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of M and V. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input.			
	The mean of the gamma distribution with parameters a and b is ab . The variance is ab^2 .			
Examples	[m,v] = gamstat(1:5,1:5) m = 1 4 9 16 25 v = 1 8 27 64 125			
	[m,v] = gamstat(1:5,1./(1:5))			
	m = 1 1 1 1 1			
	v = 1.0000 0.5000 0.3333 0.2500 0.2000			
See Also	gamcdf, gamfit, gaminv, gamlike, gampdf, gamrnd			

Purpose	Geometric cumulative distribution function (cdf)		
Syntax	Y = geocdf(X,P)		
Description	geocdf(X,P) computes the geometric cdf at each of the values in X using the corresponding probabilities in P. Vector or matrix inputs for X and P must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in P must lie on the interval [0 1].		
	The geometric cdf is		
	$y = F(x p) = \sum_{i=0}^{floor(x)} pq^{i}$		
	where $q = 1-p$.		
	The result, y , is the probability of observing up to x trials before a success, when the probability of success in any given trial is p .		
Examples	Suppose you toss a fair coin repeatedly. If the coin lands face up (heads), that is a success. What is the probability of observing three or fewer tails before getting a heads?		
	p = geocdf(3, 0.5)		
	p = 0.9375		
See Also	cdf, geoinv, geopdf, geornd, geostat		

geoinv

Purpose	Inverse of the geometric cumulative distribution function (cdf)
Syntax	X = geoinv(Y,P)
Description	X = geoinv(Y,P) returns the smallest positive integer X such that the geometric cdf evaluated at X is equal to or exceeds Y. You can think of Y as the probability of observing X successes in a row in independent trials where P is the probability of success in each trial.
	Vector or matrix inputs for P and Y must have the same size, which is also the size of X. A scalar input for P and Y is expanded to a constant matrix with the same dimensions as the other input. The values in P and Y must lie on the interval $[0\ 1]$.
Examples	The probability of correctly guessing the result of 10 coin tosses in a row is less than 0.001 (unless the coin is not fair).
	<pre>psychic = geoinv(0.999,0.5)</pre>
	psychic =
	9
	The example below shows the inverse method for generating random numbers from the geometric distribution.
	<pre>rndgeo = geoinv(rand(2,5),0.5)</pre>
	rndgeo =
	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
See Also	geocdf, geopdf, geornd, geostat, icdf

Purpose	Geometric mean of a sample
---------	----------------------------

Syntax m = geomean(X)

Description geomean calculates the geometric mean of a sample. For vectors, geomean(x) is the geometric mean of the elements in x. For matrices, geomean(X) is a row vector containing the geometric means of each column.

The geometric mean is

$$m = \left[\prod_{i=1}^{n} x_i\right]^{\frac{1}{n}}$$

Examples The sample average is greater than or equal to the geometric mean.

x = exprnd(1, 10, 6);geometric = geomean(x) geometric = 0.7466 0.6061 0.6038 0.2569 0.7539 0.3478 average = mean(x)average = 1.3509 1.1583 0.9741 0.5319 1.0088 0.8122

See Also mean, median, harmmean, trimmean

geopdf

Purpose	Geometric probability density function (pdf)		
Syntax	Y = geopdf(X,P)		
Description	geocdf(X,P) computes the geometric pdf at each of the values in X using the corresponding probabilities in P. Vector or matrix inputs for X and P must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in P must lie on the interval [0 1].		
	The geometric pdf is		
	$y = f(x p) = pq^{x}I_{(0, 1, K)}(x)$		
	where $q = 1 - p$.		
Examples	Suppose you toss a fair coin repeatedly. If the coin lands face up (heads), that is a success. What is the probability of observing exactly three tails before getting a heads?		
	p = geopdf(3,0.5)		
	p = 0.0625		
See Also	geocdf, geoinv, geornd, geostat, pdf		

Purpose	Random numbers from the geometric distribution				
Syntax	R = geornd(P) R = geornd(P,m) R = geornd(P,m,n)				
Description	The geometric distribution is useful when you want to model the number of successive failures preceding a success, where the probability of success in any given trial is the constant P.				
	R = geornd(P) generates geometric random numbers with probability parameter P. The size of R is the size of P.				
	R = geornd(P,m) generates geometric random numbers with probability parameter P, where m is a 1-by-2 vector that contains the row and column dimensions of R.				
	R = geornd(P,m,n) generates geometric random numbers with probability parameter P, where scalars m and n are the row and column dimensions of R.				
	The parameters in P must lie on the interval [0 1].				
Examples	r1 = geornd(1 ./ 2.^(1:6))				
	r1 =				
	2 10 2 5 2 60				
	r2 = geornd(0.01,[1 5])				
	r2 = 65 18 334 291 63				
	r3 = geornd(0.5,1,6)				
	r3 = 0 7 1 3 1 0				
See Also	geocdf, geoinv, geopdf, geostat				

geostat

Purpose	Mean and variance for the geometric distribution					
Syntax	[M,V] = ge	[M,V] = geostat(P)				
Description	[M,V] = geostat(P) returns the mean and variance for the geometric distribution with parameters specified by P.					
	The mean of The variance	<u> </u>	tric distribu	ition with pa	arameter <i>p</i> i	is q/p , where $q = 1-p$.
Examples	[m,v] =	geostat(1	./(1:6))			
	m = 0 v =	1.0000	2.0000	3.0000	4.0000	5.0000
	0	2.0000	6.0000	12.0000	20.0000	30.0000
See Also	geocdf, geo	inv,geopdf	,geornd			

Purpose	Interactively draw a line in a figure
Syntax	gline(fig) h = gline(fig) gline
Description	gline (fig) allows you to draw a line segment in the figure fig by clicking the pointer at the two end-points. A rubber band line tracks the pointer movement.
	h = gline(fig) returns the handle to the line in h .
	gline with no input arguments draws in the current figure.
See Also	refline, gname

glmdemo

Purpose	Demo of generalized linear models
Syntax	glmdemo
Description	glmdemo begins a slide show demonstration of generalized linear models. The slides indicate when generalized linear models are useful, how to fit generalized linear models using the glmfit function, and how to make predictions using the glmval function.
	Note To run this demo from the command line, type playshow glmdemo.
See Also	glmfit,glmval

Purpose	Generalized linear model fitting
Syntax	<pre>b = glmfit(X,Y,'distr') b = glmfit(X,Y,'distr','link','estdisp',offset,pwts,'const') [b,dev,stats] = glmfit()</pre>
Description	b = glmfit(x,y, 'distr') fits the generalized linear model for response Y, predictor variable matrix X, and distribution 'distr'. The following distributions are available: 'binomial', 'gamma', 'inverse gaussian', 'lognormal', 'normal' (the default), and 'poisson'. In most cases Y is a vector of response measurements, but for the binomial distribution Y is a two-column array having the measured number of counts in the first column and the number of trials (the binomial N parameter) in the second column. X is a matrix having the same number of rows as Y and containing the values of the predictor variables for each observation. The output b is a vector of coefficient estimates. This syntax uses the canonical link (see below) to relate the distribution parameter to the predictors.

b = glmfit(x,y, 'distr', 'link', 'estdisp', offset, pwts, 'const')provides additional control over the fit. The 'link' argument specifies the relationship between the distribution parameter (μ) and the fitted linear combination of predictor variables (*xb*). In most cases 'link' is one of the following:

'link'	Meaning	Default (Canonical) Link
'identity'	$\mu = xb$	'normal'
'log'	$\log(\mu) = xb$	'poisson'
'logit'	$\log(\mu / (1-\mu)) = xb$	'binomial'
'probit'	$\operatorname{norminv}(\mu) = xb$	
'comploglog'	$\log(-\log(1-\mu)) = xb$	
'logloglink'	$\log(-\log(\mu)) = xb$	
'reciprocal'	$1/\mu = xb$	'gamma'
p (a number)	$\mu^p = xb$	'inverse gaussian' (with $p=-2$)

Alternatively, you can write functions to define your own custom link. You specify the link argument as a three-element cell array containing functions that define the link function, its derivative, and its inverse. For example, suppose you want to define a reciprocal square root link using inline functions. You could define the variable mylinks to use as your '*link*' argument by writing:

```
FL = inline('x.^-.5')
FD = inline('-.5*x.^-1.5')
FI = inline('x.^-2')
mylinks = {FL FI FD}
```

Alternatively, you could define functions named FL, FD, and FI in their own M-files, and then specify mylinks in the form

```
mylinks = {@FL @FD @FI}
```

The 'estdisp' argument can be 'on' to estimate a dispersion parameter for the binomial or Poisson distribution, or 'off' (the default) to use the theoretical value of 1.0 for those distributions. The glmfit function always estimates dispersion parameters for other distributions.

The offset and pwts parameters can be vectors of the same length as Y, or can be omitted (or specified as an empty vector). The offset vector is a special predictor variable whose coefficient is known to be 1.0. As an example, suppose that you are modeling the number of defects on various surfaces, and you want to construct a model in which the expected number of defects is proportional to the surface area. You might use the number of defects as your response, along with the Poisson distribution, the log link function, and the log surface area as an offset.

The pwts argument is a vector of prior weights. As an example, if the response value Y(i) is the average of f(i) measurements, you could use f as a vector of prior weights.

The 'const' argument can be 'on' (the default) to estimate a constant term, or 'off' to omit the constant term. If you want the constant term, use this argument rather than specifying a column of ones in the X matrix.

[b, dev, stats] = glmfit(...) returns the additional outputs dev and stats. dev is the deviance at the solution vector. The deviance is a generalization of the residual sum of squares. It is possible to perform an analysis of deviance to compare several models, each a subset of the other, and to test whether the model with more terms is significantly better than the model with fewer terms.

stats is a structure with the following fields:

- stats.dfe = degrees of freedom for error
- stats.s = theoretical or estimated dispersion parameter
- stats.sfit = estimated dispersion parameter
- stats.estdisp = 1 if dispersion is estimated, 0 if fixed
- stats.beta = vector of coefficient estimates (same as b)
- stats.se = vector of standard errors of the coefficient estimates b
- stats.coeffcorr = correlation matrix for b
- stats.t = t statistics for b
- stats.p = p-values for b
- stats.resid = vector of residuals
- stats.residp = vector of Pearson residuals
- stats.residd = vector of deviance residuals
- stats.resida = vector of Anscombe residuals

If you estimate a dispersion parameter for the binomial or Poisson distribution, then stats.s is set equal to stats.sfit. Also, the elements of stats.se differ by the factor stats.s from their theoretical values.

Example We have data on cars weighing between 2100 and 4300 pounds. For each car weight we have the total number of cars of that weight, and the number that can be considered to get "poor mileage" according to some test. For example, 8 out of 21 cars weighing 3100 pounds get poor mileage according to a measurement of the miles they can travel on a gallon of gasoline.

```
w = (2100:200:4300)';
poor = [1 2 0 3 8 8 14 17 19 15 17 21]';
total = [48 42 31 34 31 21 23 23 21 16 17 21]';
```

We can compare several fits to these data. First, let's try fitting logit and probit models:

```
[bl,dl,sl] = glmfit(w,[poor total],'binomial');
[bp,dp,sp] = glmfit(w,[poor total],'binomial','probit');
```

```
dl
dl =
6.4842
dp
dp =
7.5693
```

The deviance for the logit model is smaller than for the probit model. Although this is not a formal test, it leads us to prefer the logit model.

We can do a formal test comparing two logit models. We already fit one model using w as a linear predictor. Let's fit another logit model using both linear and squared terms in w. If there is no true effect for the squared term, the difference in their deviances should be small compared with a chi-square distribution having one degree of freedom.

```
[b2,d2,s2] = glmfit([w w.^2],[poor total],'binomial')
dl-d2
ans =
            0.7027
chi2cdf(dl-d2,1)
ans =
            0.5981
```

A difference of 0.7072 is not at all unusual for a chi-square distribution with one degree of freedom, so the quadratic model does not give a significantly better fit than the simpler linear model.

The following are the coefficient estimates, their standard errors, t-statistics, and p-values for the linear model:

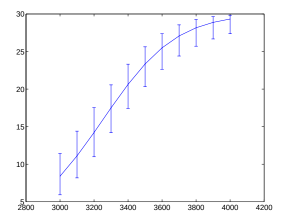
```
[b sl.se sl.t sl.p]
ans =
    -13.3801    1.3940   -9.5986    0.0000
    0.0042    0.0004    9.4474    0.0000
```

	This shows that we cannot simplify the model any further. Both the intercept and slope coefficients are significantly different from 0, as indicated by p-values that are 0.0000 to four decimal places.
See Also	glmval, glmdemo, nlinfit, regress, regstats
References	[1] Dobson, A. J. An Introduction to Generalized Linear Models. 1990, CRC Press.
	[2] MuCullagh, P. and J. A. Nelder. <i>Generalized Linear Models</i> . 2nd edition, 1990, Chapman and Hall.

glmval

Purpose	Compute predictions for generalized linear model
Syntax	<pre>yfit = glmval(b,X,'link') [yfit,dlo,dhi] = glmval(b,X,'link',stats,clev) [yfit,dlo,dhi] = glmval(b,X,'link',stats,clev,N,offset,'const')</pre>
Description	yfit = glmval(b,X, 'link') computes the predicted distribution parameters for observations with predictor values X using the coefficient vector b and link function 'link'. Typically, b is a vector of coefficient estimates computed by the glmfit function. The value of 'link' must be the same as that used in glmfit. The result yfit is the value of the inverse of the link function at the linear combination X*b.
	<pre>[yfit,dlo,dhi] = glmval(b,X,'link',stats,clev) returns confidence bounds for the predicted values when you supply the stats structure returned from glmfit, and optionally specify a confidence level as the clev argument. (The default confidence level is 0.95 for 95% confidence.) The interval [yfit-dlo, yfit+dhi] is a confidence bound for the true parameter value at the specified X values.</pre>
	<pre>[yhat,dlo,dhi] = glmval(beta,X,'link',stats,clev,N,offset,'const') specifies three additional arguments that may be needed if you used certain arguments to glmfit. If you fit a binomial distribution using glmfit, specify N as the value of the binomial N parameter for the predictions. If you included an offset variable, specify offset as the new value of this variable. Use the same 'const' value('on' or 'off') that you used with glmfit.</pre>
Example	Let's model the number of cars with poor gasoline mileage using the binomial distribution. First we use the binomial distribution with the default logit link to model the probability of having poor mileage as a function of the weight and squared weight of the cars. Then we compute a vector wnew of new car weights at which we want to make predictions. Next we compute the expected number of cars, out of a total of 30 cars of each weight, that would have poor mileage. Finally we graph the predicted values and 95% confidence bounds as a function of weight.
	<pre>w = [2100 2300 2500 2700 2900 3100 3300 3500 3700 3900 4100 4300]'; poor = [1 2 0 3 8 8 14 17 19 15 17 21]'; total = [48 42 31 34 31 21 23 23 21 16 17 21]';</pre>

```
[b2,d2,s2] = glmfit([w w.^2],[poor total],'binomial')
wnew = (3000:100:4000)';
[yfit,dlo,dhi] = glmval(b2,[wnew wnew.^2],'logit',s2,0.95,30)
errorbar(wnew,yfit,dlo,dhi);
```

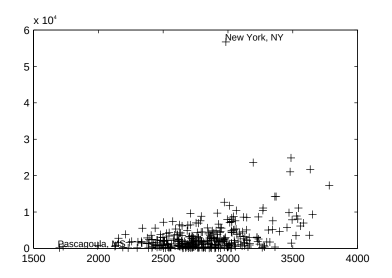


See Also

glmfit,glmdemo

gname

Purpose	Label plotted points with their case names or case number
Syntax	gname('cases') gname h = gname('cases',line_handle)
Description	<pre>gname('cases') displays a figure window, displays cross-hairs, and waits for a mouse button or keyboard key to be pressed. Position the cross-hair with the mouse and click once near each point to label that point. Input 'cases' is a string matrix with each row the case name of a data point. You can also click and drag a selection rectangle to label all points within the rectangle. When you are done, press the Enter or Escape key. gname with no arguments labels each case with its case number. h = gname('cases',line_handle) returns a vector of handles to the text objects on the plot. Use the scalar line_handle to identify the correct line if there is more than one line object on the plot.</pre>
	You can use gname to label plots created by the plot, scatter, gscatter, plotmatrix, and gplotmatrix functions.
Example	Let's use the city ratings data sets to find out which cities are the best and worst for education and the arts. We create a graph, call the gname function, and click on the points at the extreme left and at the top.
	<pre>load cities education = ratings(:,6); arts = ratings(:,7); plot(education,arts,'+') gname(names)</pre>



See Also

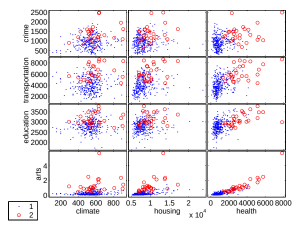
gplotmatrix, gscatter, gtext, plot, plotmatrix, scatter

gplotmatrix

Purpose	Plot matrix of scatter plots by group
Syntax	<pre>gplotmatrix(x,y,g) gplotmatrix(x,y,g,'clr','sym',siz) gplotmatrix(x,y,g,'clr','sym',siz,'doleg') gplotmatrix(x,y,g,'clr','sym',siz,'doleg','dispopt') gplotmatrix(x,y,g,'clr','sym',siz,'doleg','dispopt','xnam','ynam') [h,ax,bigax] = gplotmatrix()</pre>
Description	gplotmatrix(x,y,g) creates a matrix of scatter plots. Each individual set of axes in the resulting figure contains a scatter plot of a column of x against a column of y. All plots are grouped by the grouping variable g.
	x and y are matrices with the same number of rows. If x has p columns and y has q columns, the figure contains a p-by-q matrix of scatter plots. If you omit y or specify it as the empty matrix, [], gplotmatrix creates a square matrix of scatter plots of columns of x against each other.
	g is a grouping variable that can be a vector, string array, or cell array of strings. g must have the same number of rows as x and y. Points with the same value of g are placed in the same group, and appear on the graph with the same marker and color. Alternatively, g can be a cell array containing several grouping variables (such as {G1 G2 G3}); in that case, observations are in the same group if they have common values of all grouping variables.
	<pre>gplotmatrix(x,y,g,'clr','sym',siz) specifies the color, marker type, and size for each group. clr is a string array of colors recognized by the plot function. The default is 'clr' = 'bgrcmyk'. 'sym' is a string array of symbols recognized by the plot command, with the default value '.'. siz is a vector of sizes, with the default determined by the 'defaultlinemarkersize' property. If you do not specify enough values for all groups, gplotmatrix cycles through the specified values as needed.</pre>
	<pre>gplotmatrix(x,y,g,'clr','sym',siz,'doleg') controls whether a legend is displayed on the graph ('doleg' = 'on', the default) or not ('doleg' = 'off').</pre>
	gplotmatrix(x,y,g,' <i>clr</i> ',' <i>sym</i> ',siz,' <i>doleg</i> ',' <i>dispopt</i> ') controls what appears along the diagonal of a plot matrix of x versus x. Allowable values are

	'none' to leave the diagonals blank, 'hist' (the default) to plot histograms, or 'variable' to write the variable names.
	gplotmatrix(x,y,g,' <i>clr</i> ',' <i>sym</i> ',siz,' <i>doleg</i> ',' <i>dispopt</i> ','xnam','ynam') specifies the names of the columns in the x and y arrays. These names are used to label the <i>x</i> - and <i>y</i> -axes. 'xnam' and 'ynam' must be character arrays with one row for each column of x and y, respectively.
	<pre>[h,ax,bigax] = gplotmatrix() returns three arrays of handles. h is an array of handles to the lines on the graphs. ax is a matrix of handles to the axes of the individual plots. bigax is a handle to big (invisible) axes framing the entire plot matrix. These are left as the current axes, so a subsequent title, xlabel, or ylabel command will produce labels that are centered with respect to the entire plot matrix.</pre>
Example	Load the cities data. The ratings array has ratings of the cities in nine categories (category names are in the array categories). group is a code whose value is 2 for the largest cities. We can make scatter plots of the first three categories against the other four, grouped by the city size code.
	load discrim gplotmatrix(ratings(:,1:3),ratings(:,4:7),group)
	The output figure (not shown) has an array of graphs with each city group represented by a different color. The graphs are a little easier to read if we specify colors and plotting symbols, label the axes with the rating categories, and move the legend off the graphs.
	gplotmatrix(ratings(:,1:3),ratings(:,4:7),group, 'br','.o',[],'on','',categories(1:3,:), categories(4:7,:))

gplotmatrix



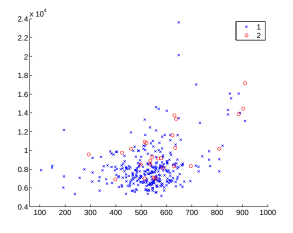
See Also

grpstats, gscatter, plotmatrix

_				
Purpose	Summary statistics by group			
Syntax	means = grpstats(X,group) [means,sem,counts,name] = grpstats(X,group) grpstats(x,group,alpha)			
Description	<pre>means = grpstats(X,group) returns the means of each column of X by group, where X is a matrix of observations. group is an array that defines the grouping such that two elements of X are in the same group if their corresponding group values are the same. The grouping variable group can be a vector, string array, or cell array of strings. It can also be a cell array containing several grouping variables (such as {G1 G2 G3}); in that case observations are in the same group if they have common values of all grouping variables.</pre>			
	[means,sem,counts,name] = grpstats(x,group,alpha) supplies the standard error of the mean in sem, the number of elements in each group in counts, and the name of each group in name. name is useful to identify and label the groups when the input group values are not simple group numbers.			
	grpstats(x,group,alpha) plots 100(1-alpha)% confidence intervals around each mean.			
Example	We assign 100 observations to one of four groups. For each observation we measure five quantities with <i>true means</i> from 1 to 5. grpstats allows us to compute the means for each group.			
	group = unidrnd(4,100,1); true_mean = 1:5; true_mean = true_mean(ones(100,1),:); x = normrnd(true_mean,1); means = grpstats(x,group)			
	means =			
	0.79472.09082.89693.67494.65550.93771.76003.02853.94844.81691.05492.02552.87934.07995.37400.71071.92642.82323.88154.9689			
See Also	tabulate, crosstab			

gscatter

Purpose	Scatter plot by group
Syntax	gscatter(x,y,g) gscatter(x,y,g,'clr','sym',siz) gscatter(x,y,g,'clr','sym',siz,'doleg') gscatter(x,y,g,'clr','sym',siz,'doleg','xnam','ynam') h = gscatter()
Description	gscatter(x,y,g) creates a scatter plot of x and y, grouped by g, where x and y are vectors with the same size and g can be a vector, string array, or cell array of strings. Points with the same value of g are placed in the same group, and appear on the graph with the same marker and color. Alternatively, g can be a cell array containing several grouping variables (such as {G1 G2 G3}); in that case, observations are in the same group if they have common values of all grouping variables.
	gscatter(x,y,g,'clr','sym',siz) specifies the color, marker type, and size for each group. 'clr' is a string array of colors recognized by the plot function. The default is 'clr' = 'bgrcmyk'.'sym' is a string array of symbols recognized by the plot command, with the default value '.'. siz is a vector of sizes, with the default determined by the 'defaultlinemarkersize' property. If you do not specify enough values for all groups, gscatter cycles through the specified values as needed.
	<pre>gscatter(x,y,g,'clr','sym',siz,'doleg') controls whether a legend is displayed on the graph ('doleg' = 'on', the default) or not ('doleg' = 'off').</pre>
	gscatter(x,y,g, 'clr', 'sym', siz, 'doleg', 'xnam', 'ynam') specifies the name to use for the x-axis and y-axis labels. If the x and y inputs are simple variable names and xnam and ynam are omitted, gscatter labels the axes with the variable names.
	h = gscatter() returns an array of handles to the lines on the graph.
Example	Load the cities data and look at the relationship between the ratings for climate (first column) and housing (second column) grouped by city size. We'll also specify the colors and plotting symbols. load discrim



gscatter(ratings(:,1),ratings(:,2),group,'br','xo')



gplotmatrix, grpstats, scatter

harmmean

Purpose	Harmonic mean of a sample of data	a
---------	-----------------------------------	---

Syntax m = harmmean(X)

Description m = harmmean(X) calculates the harmonic mean of a sample. For vectors, harmmean(x) is the harmonic mean of the elements in x. For matrices, harmmean(X) is a row vector containing the harmonic means of each column.

The harmonic mean is

$$m = \frac{n}{\sum_{i=1}^{n} \frac{1}{x_i}}$$

Examples The sample average is greater than or equal to the harmonic mean.

	exprnd(1, monic = ha					
har	monic =					
	0.3382	0.3200	0.3710	0.0540	0.4936	0.0907
ave	erage = mea	an(x)				
ave	erage =					
	1.3509	1.1583	0.9741	0.5319	1.0088	0.8122

See Also mean, median, geomean, trimmean

Purpose	Plot histograms
Syntax	hist(y) hist(y,nb) hist(y,x) [n,x] = hist(y,)
Description	hist(y) draws a 10-bin histogram for the data in vector y. The bins are equally spaced between the minimum and maximum values in y.
	hist(y,nb) draws a histogram with nb bins.
	hist(y,x) draws a histogram using the bins in the vector x.
	[n,x] = hist(y,) do not draw graphs, but return vectors n and x containing the frequency counts and the bin locations such that bar(x,n) plots the histogram. This is useful in situations where more control is needed over the appearance of a graph, for example, to combine a histogram into a more elaborate plot statement.
	The hist function is a part of the standard MATLAB language.
Examples	Generate bell-curve histograms from Gaussian data.
	<pre>x = -2.9:0.1:2.9; y = normrnd(0,1,1000,1); hist(y,x)</pre>
	-3 -2 -1 0 1 2 3

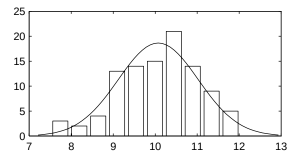
histfit

Purpose	Histogram with superimposed normal density
Syntax	histfit(data) histfit(data,nbins) h = histfit(data,nbins)
Description	histfit(data,nbins) plots a histogram of the values in the vector data using nbins bars in the histogram. With nbins is omitted, its value is set to the square root of the number of elements in data.

h = histfit(data,nbins) returns a vector of handles to the plotted lines, where h(1) is the handle to the histogram, h(2) is the handle to the density curve.

Example

r = normrnd(10,1,100,1);
histfit(r)





hist, normfit

Purpose	Hougen-Watson model for reaction kinetics	
Syntax	yhat = hougen(beta,x)	
Description	yhat = hougen(beta,x) returns the predicted values of the reaction rate, yhat, as a function of the vector of parameters, beta, and the matrix of data, X. beta must have 5 elements and X must have three columns.	
	hougen is a utility function for rsmdemo.	
	The model form is:	
	$\hat{y} = \frac{\beta_1 x_2 - x_3 / \beta_5}{1 + \beta_2 x_1 + \beta_3 x_2 + \beta_4 x_3}$	
Reference	[1] Bates, D., and D. Watts. <i>Nonlinear Regression Analysis and Its Applications</i> . Wiley 1988. p. 271–272.	
See Also	rsmdemo	

hygecdf

Purpose	Hypergeometric cumulative distribution function (cdf)			
Syntax	P = hygecdf(X,M,K,N)			
Description	hygecdf(X,M,K,N) computes the hypergeometric cdf at each of the values in X using the corresponding parameters in M, K, and N. Vector or matrix inputs for X, M, K, and N must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.			
	The hypergeometric cdf is			
	$p = F(x M, K, N) = \sum_{i=0}^{x} \frac{\binom{K}{i}\binom{M-K}{N-i}}{\binom{M}{N}}$			
	The result, p , is the probability of drawing up to x of a possible K items in N drawings without replacement from a group of M objects.			
Examples	Suppose you have a lot of 100 floppy disks and you know that 20 of them are defective. What is the probability of drawing zero to two defective floppies if you select 10 at random?			
	p = hygecdf(2,100,20,10)			
	p = 0.6812			
See Also	cdf, hygeinv, hygepdf, hygernd, hygestat			

Purpose	Inverse of the hypergeometric cumulative distribution function (cdf)		
Syntax	X = hygeinv(P,M,K,N)		
Description	hygeinv(P,M,K,N) returns the smallest integer X such that the hypergeometric cdf evaluated at X equals or exceeds P. You can think of P as the probability of observing X defective items in N drawings without replacement from a group of M items where K are defective.		
Examples	Suppose you are the Quality Assurance manager for a floppy disk manufacturer. The production line turns out floppy disks in batches of 1,000. You want to sample 50 disks from each batch to see if they have defects. You want to accept 99% of the batches if there are no more than 10 defective disks in the batch. What is the maximum number of defective disks should you allow in your sample of 50?		
	x = hygeinv(0.99,1000,10,50)		
	x = 3		
	What is the median number of defective floppy disks in samples of 50 disks from batches with 10 defective disks?		
	x = hygeinv(0.50,1000,10,50)		
	x = 0		
See Also	hygecdf, hygepdf, hygernd, hygestat, icdf		

hygepdf

Purpose	Hypergeometric probability density function (pdf)
Syntax	Y = hygepdf(X,M,K,N)
Description	Y = hygecdf(X,M,K,N) computes the hypergeometric pdf at each of the values in X using the corresponding parameters in M, K, and N. Vector or matrix inputs for X, M, K, and N must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.
	The parameters in M, K, and N must all be positive integers, with $N \le M$. The values in X must be less than or equal to all the parameter values.
	The hypergeometric pdf is
	$y = f(x M, K, N) = \frac{\binom{K}{x}\binom{M-K}{N-x}}{\binom{M}{N}}$
	The result, y , is the probability of drawing exactly x of a possible K items in n drawings without replacement from a group of M objects.
Examples	Suppose you have a lot of 100 floppy disks and you know that 20 of them are defective. What is the probability of drawing 0 through 5 defective floppy disks if you select 10 at random?
	p = hygepdf(0:5,100,20,10)
	p = 0.0951 0.2679 0.3182 0.2092 0.0841 0.0215
See Also	hygecdf, hygeinv, hygernd, hygestat, pdf

hygernd

Purpose	Random numbers from the hypergeometric distribution
Syntax	R = hygernd(M,K,N) R = hygernd(M,K,N,mm) R = hygernd(M,K,N,mm,nn)
Description	 R = hygernd(M,K,N) generates hypergeometric random numbers with parameters M, K, and N. Vector or matrix inputs for M, K, and N must have the same size, which is also the size of R. A scalar input for M, K, or N is expanded to a constant matrix with the same dimensions as the other inputs. R = hygernd(M,K,N,mm) generates hypergeometric random numbers with parameters M, K, and N, where mm is a 1-by-2 vector that contains the row and
	column dimensions of R. R = hygernd(M,K,N,mm,nn) generates hypergeometric random numbers with
	parameters $M,K,{\rm and}N,{\rm where}{\rm scalars}mm$ and nn are the row and column dimensions of $R.$
Examples	<pre>numbers = hygernd(1000,40,50)</pre>
	numbers =
	1
See Also	hygecdf, hygeinv, hygepdf, hygestat

hygestat

Purpose	Mean and variance for the hypergeometric distribution				
Syntax	[MN,V] = hygestat(M,K,N)				
Description	[MN,V] = hygestat(M,K,N) returns the mean and variance for the hypergeometric distribution with parameters specified by M, K, and N. Vector or matrix inputs for M, K, and N must have the same size, which is also the size of MN and V. A scalar input for M, K, or N is expanded to a constant matrix with the same dimensions as the other inputs.				
	The mean of the hypergeometric distribution with parameters M, K , and N is NK/M , and the variance is				
	$Nrac{K}{M}rac{M-K}{M}rac{M-N}{M-1}$				
Examples	The hypergeometric distribution approaches the binomial distribution, where $p = K / M$ as M goes to infinity.				
	[m,v] = hygestat(10.^(1:4),10.^(0:3),9)				
	m =				
	0.9000 0.9000 0.9000 0.9000				
	v = 0.0900 0.7445 0.8035 0.8094				
	[m,v] = binostat(9,0.1)				
	m = 0.9000				
	v = 0.8100				
See Also	hygecdf, hygeinv, hygepdf, hygernd				

Purpose	Inverse of a specified cumulative distribution function (icdf)			
Syntax	X = icdf('name',P,A1,A2,A3)			
Description	X = icdf('name', P,A1,A2,A3) returns a matrix of critical values, X, where 'name' is a string containing the name of the distribution. P is a matrix of probabilities, and A, B, and C are matrices of distribution parameters. Depending on the distribution some of the parameters may not be necessary.			
	Vector or matrix inputs for P, A1, A2, and A3 must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.			
	icdf is a utility routine allowing you to access all the inverse cdfs in the Statistics Toolbox using the name of the distribution as a parameter. See "Overview of the Distributions" on page 2-11 for the list of available distributions.			
Examples	x = icdf('Normal',0.1:0.2:0.9,0,1)			
	x = -1.2816 -0.5244 0 0.5244 1.2816			
	<pre>x = icdf('Poisson',0.1:0.2:0.9,1:5)</pre>			
	x = 1 1 3 5 8			
See Also	betainv, binoinv, cdf, chi2inv, expinv, finv, gaminv, geoinv, hygeinv, logninv, nbininv, ncfinv, nctinv, ncx2inv, norminv, pdf, poissinv, random, raylinv, tinv, unidinv, unifinv, weibinv			

inconsistent

Purpose	Calculate the inconsistency coefficient of a cluster tree			
Syntax	Y = inconsistent(Z) Y = inconsistent(Z,d)			
Description	Y = inconsistent(Z) computes the inconsistency coefficient for each link of the hierarchical cluster tree Z, where Z is an $(m-1)$ -by-3 matrix generated by the linkage function. The inconsistency coefficient characterizes each link in a cluster tree by comparing its length with the average length of other links at the same level of the hierarchy. The higher the value of this coefficient, the less similar the objects connected by the link.			

Y = inconsistent(Z,d) computes the inconsistency coefficient for each link in the hierarchical cluster tree Z to depth d, where d is an integer denoting the number of levels of the cluster tree that are included in the calculation. By default, d=2.

The output, Y, is an (m-1)-by-4 matrix formatted as follows.

Column	Description
1	Mean of the lengths of all the links included in the calculation.
2	Standard deviation of all the links included in the calculation.
3	Number of links included in the calculation.
4	Inconsistency coefficient.

For each link, k, the inconsistency coefficient is calculated as:

Y(k, 4) = (z(k, 3) - Y(k, 1))/Y(k, 2)

For leaf nodes, nodes that have no further nodes under them, the inconsistency coefficient is set to 0.

```
Example rand('seed',12);
X = rand(10,2);
Y = pdist(X);
```

<pre>Z = linkage(Y,'centroid'); W = inconsistent(Z,3)</pre>				
W =				
0.04	23	0 1.0000) 0	
0.14	06	0 1.0000	0 0	
0.11	63 0.104	7 2.0000	0.7071	
0.21	01	0 1.0000) 0	
0.20	54 0.088	6 3.0000	0.6792	
0.17	42 0.176	2 3.0000	0.6568	
0.23	36 0.131	7 4.0000	0.6408	
0.30	81 0.210	9 5.0000	0.7989	
0.46	10 0.372	8 4.0000	0.8004	

See Also

cluster, cophenet, clusterdata, dendrogram, linkage, pdist, squareform

Purpose	Interquartile range (IQR) of a sample			
Syntax	y = iqr(X)			
Description	y = iqr(X) computes the difference between the 75th and the 25th percentiles of the sample in X. The IQR is a robust estimate of the spread of the data, since changes in the upper and lower 25% of the data do not affect it.			
	If there are outliers in the data, then the IQR is more representative than the standard deviation as an estimate of the spread of the body of the data. The IQR is less efficient than the standard deviation as an estimate of the spread when the data is all from the normal distribution.			
	Multiply the IQR by 0.7413 to estimate σ (the second parameter of the normal distribution.)			
Examples	This Monte Carlo simulation shows the relative efficiency of the IQR to the sample standard deviation for normal data.			
	x = normrnd(0,1,100,100); s = std(x); s_IQR = 0.7413 * iqr(x); efficiency = (norm(s - 1)./norm(s_IQR - 1)).^2			
	efficiency =			
	0.3297			
See Also	std, mad, range			

iwishrnd

Purpose	Generate inverse Wishart random matrix			
Syntax	W=iwishrnd(SIGMA,df) W=iwishrnd(SIGMA,df,DI) [W,DI]=iwishrnd(SIGMA,df)			
Description	W=iwishrnd(SIGMA,df) generates a random matrix W whose inverse has the Wishart distribution with covariance matrix inv(SIGMA) and with df degrees of freedom.			
	W=iwishrnd(SIGMA,df,DI) expects DI to be the Cholesky factor of the inverse of SIGMA. If you call iwishrnd multiple times using the same value of SIGMA, it's more efficient to supply DI instead of computing it each time.			
	[W,DI]=iwishrnd(SIGMA,df) returns DI so you can provide it as input in future calls to iwishrnd.			
See Also	wishrnd			

jbtest

Purpose	Jarque-Bera test for goodness-of-fit to a normal distribution				
Syntax	H = jbtest(X) H = jbtest(X,alpha) [H,P,JBSTAT,CV] = jbtest(X,alpha)				
Description	H = jbtest(X) performs the Jarque-Bera test on the input data vector X and returns H, the result of the hypothesis test. The result is H=1 if we can reject the hypothesis that X has a normal distribution, or H=0 if we cannot reject that hypothesis. We reject the hypothesis if the test is significant at the 5% level.				
	The Jarque-Bera test evaluates the hypothesis in the test is significant at the 5% level. The Jarque-Bera test evaluates the hypothesis that X has a normal distribution with unspecified mean and variance, against the alternative that X does not have a normal distribution. The test is based on the sample skewness and kurtosis of X. For a true normal distribution, the sample skewness should be near 0 and the sample kurtosis should be near 3. The Jarque-Bera test determines whether the sample skewness and kurtosis are unusually different than their expected values, as measured by a chi-square statistic.				
	The Jarque-Bera test is an asymptotic test, and should not be used with small samples. You may want to use lillietest in place of jbtest for small samples.				
	H = jbtest(X,alpha) performs the Jarque-Bera test at the 100*alpha% level rather than the 5% level, where alpha must be between 0 and 1.				
	[H,P,JBSTAT,CV] = jbtest(X,alpha) returns three additional outputs. P is the p-value of the test, JBSTAT is the value of the test statistic, and CV is the critical value for determining whether to reject the null hypothesis.				
Example	We can use jbtest to determine if car weights follow a normal distribution.				
	load carsmall [h,p,j] = jbtest(Weight)				
	h = 1				
	p = 0.026718				
	j =				

7.2448

With a p-value of 2.67%, we reject the hypothesis that the distribution is normal. With a log transformation, the distribution becomes closer to normal but is still significantly different at the 5% level.

See lillietest for a different test of the same hypothesis.

Reference [1] Judge, G. G., R. C. Hill, W. E. Griffiths, H. Lutkepohl, and T.-C. Lee. *Introduction to the Theory and Practice of Econometrics*. New York, Wiley.

See Also hist, kstest2, lillietest

Purpose	K-means clustering			
Syntax	<pre>IDX = kmeans(X,k) [IDX,C] = kmeans(X,k) [IDX,C,sumd] = kmeans(X,k) [IDX,C,sumd,D] = kmeans(X,k) [] = kmeans(,'param1',val1,'param2',val2,)</pre>			
Description	clusters. This ite within-cluster su correspond to poi n-by-1 vector IDX	K = kmeans(X, k) partitions the points in the n-by-p data matrix X into k sters. This iterative partitioning minimizes the sum, over all clusters, of the hin-cluster sums of point-to-cluster-centroid distances. Rows of X respond to points, columns correspond to variables. kmeans returns an y-1 vector IDX containing the cluster indices of each point. By default, ans uses squared Euclidean distances.		
	[IDX,C] = kmeans(X,k) returns the k cluster centroid locations in the k-by-p matrix C.			
	[IDX,C,sumd] = kmeans(X,k) returns the within-cluster sums of point-to-centroid distances in the 1-by-k vector sumd.			
	[IDX,C,sumd,D] = kmeans(X,k) returns distances from each point to every centroid in the n-by-k matrix D.			
	<pre>[] = kmeans(,'param1',val1,'param2',val2,) enables you to specify optional parameter name-value pairs to control the iterative algorithm used by kmeans. Valid parameters are the following. Parameter Value</pre>			
	'distance'	Distance measure, in p-dimensional space, that kmeans minimizes with respect to. kmeans computes centroid clusters differently for the different supported distance measures: 'sqEuclidean' Squared Euclidean distance (default). Each centroid is the mean of the points in that cluster.		

Parameter (Continued)	Value		
	'cityblock'	Sum of absolute differences, i.e., L1. Each centroid is the component-wise median of the points in that cluster.	
	'cosine'	One minus the cosine of the included angle between points (treated as vectors). Each centroid is the mean of the points in that cluster, after normalizing those points to unit Euclidean length.	
	'correlation'	One minus the sample correlation between points (treated as sequences of values). Each centroid is the component-wise mean of the points in that cluster, after centering and normalizing those points to zero mean and unit standard deviation.	
	'Hamming'	Percentage of bits that differ (only suitable for binary data). Each centroid is the component-wise median of points in that cluster.	
'start'	Method used to choose the initial cluster centroid positions, sometimes known as "seeds". Valid starting values are:		
	'sample'	Select k observations from X at random (default).	
	'uniform'	Select k points uniformly at random from the range of X. Not valid with Hamming distance.	
	'cluster'	Perform a preliminary clustering phase on a random 10% subsample of X. This preliminary phase is itself initialized using 'sample'.	

Parameter (Continued)	Value		
	Matrix	<pre>k-by-p matrix of centroid starting locations. In this case, you can pass in [] for k, and kmeans infers k from the first dimension of the matrix. You can also supply a 3-dimensional array, implying a value for the 'replicates' parameter from the array's third dimension.</pre>	
'replicates'	Number of times to repeat the clustering, each with a new set of initial cluster centroid positions. kmeans returns the solution with the lowest value for sumd. You can supply 'replicates' implicitly by supplying a 3-dimensional array as the value for the 'start' parameter.		
'maxiter'	Maximum numb	per of iterations. Default is 100.	
'emptyaction'	Action to take if a cluster loses all its member observations. Can be one of:		
	'error'	Treat an empty cluster as an error. (default)	
	'drop'	Remove any clusters that become empty. kmeans sets the corresponding return values in C and D to NaN.	
	'singleton'	Create a new cluster consisting of the one point furthest from its centroid.	
'display'	Controls display of output.		
	'off'	Display no output.	
	'iter'	Display information about each iteration during minimization, including the iteration number, the optimization phase (see "Algorithm"), the number of points moved, and the total sum of distances.	

Parameter (Continued)		Value		
		'final'	Display a summary of each replication.	
		'notify'	Display only warning and error messages. (default)	
Algorithm	 kmeans uses a two-phase iterative algorithm to minimize the sum of point-to-centroid distances, summed over all k clusters: The first phase uses what the literature often describes as "batch" undetes 			
	• The first phase uses what the literature often describes as "batch" updates, where each iteration consists of reassigning points to their nearest cluster centroid, all at once, followed by recalculation of cluster centroids. You can think of this phase as providing a fast but potentially only approximate solution as a starting point for the second phase.			

• The second phase uses what the literature often describes as "on-line" updates, where points are individually reassigned if doing so will reduce the sum of distances, and cluster centroids are recomputed after each reassignment. Each iteration during this second phase consists of one pass though all the points.

kmeans can converge to a local optimum, in this case, a partition of points in which moving any single point to a different cluster increases the total sum of distances. This problem can only be solved by a clever (or lucky, or exhaustive) choice of starting points.

- See Also clusterdata, linkage, silhouette
- **References** [1] Seber, G.A.F., *Multivariate Observations*, Wiley, New York, 1984.

 [2] Spath, H., Cluster Dissection and Analysis: Theory, FORTRAN Programs, Examples, translated by J. Goldschmidt, Halsted Press, New York, 1985, 226 pp.

kruskalwallis

Purpose	Kruskal-Wallis nonparametric one-way Analysis of Variance (ANOVA)								
Syntax	p = kruskalwa p = kruskalwa p = kruskalwa [p,table] = k [p,table,stat	llis(X,q llis(X,q ruskalwa	group,'a allis(.)					
Description	<pre>p = kruskalwallis(X) performs a Kruskal-Wallis test for comparing the means of columns of the <i>m</i>-by-<i>n</i> matrix X, where each column represents an independent sample containing <i>m</i> mutually independent observations. The Kruskal-Wallis test is a nonparametric version of the classical one-way ANOVA. The function returns the p-value for the null hypothesis that all samples in X are drawn from the same population (or from different populations with the same mean).</pre>								
	If the p-value is near zero, this casts doubt on the null hypothesis and suggests that at least one sample mean is significantly different than the other sample means. The choice of a critical p-value to determine whether the result is judged "statistically significant" is left to the researcher. It is common to declare a result significant if the p-value is less than 0.05 or 0.01.						er sample sult is		
	The kruskalwallis function displays two figures. The first figure is a standard ANOVA table, calculated using the ranks of the data rather than their numeric values. Ranks are found by ordering the data from smallest to largest across all groups, and taking the numeric index of this ordering. The rank for a tied observation is equal to the average rank of all observations tied with it. For example, the following table shows the ranks for a small sample.								
	X value	1.4	2.7	1.6	1.6	3.3	0.9	1.1	
	Rank	3	6	4.5	4.5	7	1	2	
	The entries in t freedom, and ot					-			

The entries in the ANOVA table are the usual sums of squares, degrees of freedom, and other quantities calculated on the ranks. The usual F statistic is replaced by a chi-square statistic. The p-value measures the significance of the chi-square statistic.

The second figure displays box plots of each column of X (not the ranks of X).

p = kruskalwallis(X,group) uses the values in group (a character array or cell array) as labels for the box plot of the samples in X, when X is a matrix. Each row of group contains the label for the data in the corresponding column of X, so group must have length equal to the number of columns in X.

When X is a vector, kruskalwallis performs a Kruskal-Wallis test on the samples contained in X, as indexed by input group (a vector, character array, or cell array). Each element in group identifies the group (i.e., sample) to which the corresponding element in vector X belongs, so group must have the same length as X. The labels contained in group are also used to annotate the box plot.

It is not necessary to label samples sequentially (1, 2, 3, ...). For example, if X contains measurements taken at three different temperatures, -27°, 65°, and 110°, you could use these numbers as the sample labels in group. If a row of group contains an empty cell or empty string, that row and the corresponding observation in X are disregarded. NaNs in either input are similarly ignored.

p = kruskalwallis(X,group, 'displayopt') enables the table and box plot displays when 'displayopt' is 'on' (default) and suppresses the displays when 'displayopt' is 'off'.

[p,table] = kruskalwallis(...) returns the ANOVA table (including column and row labels) in cell array table. (You can copy a text version of the ANOVA table to the clipboard by using the **Copy Text** item on the **Edit** menu.)

[p,table,stats] = kruskalwallis(...) returns a stats structure that you can use to perform a follow-up multiple comparison test. The kruskalwallis test evaluates the hypothesis that all samples have the same mean, against the alternative that the means are not all the same. Sometimes it is preferable to perform a test to determine *which* pairs of means are significantly different, and which are not. You can use the multcompare function to perform such tests by supplying the stats structure as input.

Assumptions

The Kruskal-Wallis test makes the following assumptions about the data in X:

- All sample populations have the same continuous distribution, apart from a possibly different location.
- All observations are mutually independent.

kruskalwallis

The classical one-way ANOVA test replaces the first assumption with the stronger assumption that the populations have normal distributions.

Example Let's revisit the same material strength study that we used with the anova1 function, to see if the nonparametric Kruskal-Wallis procedure leads to the same conclusion. Recall we are studying the strength of beams made from three alloys:

strength = [82 86 79 83 84 85 86 87 74 82 78 75 76 77 79 ... 79 77 78 82 79]; alloy = {'st','st','st','st','st','st','st',... 'al1','al1','al1','al1','al1','al1',... 'al2','al2','al2','al2','al2','al2'};

This time we try both classical and Kruskal-Wallis anova, omitting displays:

```
anova1(strength,alloy,'off')
ans =
    1.5264e-004
kruskalwallis(strength,alloy,'off')
ans =
    0.0018
```

Both tests find that the three alloys are significantly different, though the result is less significant according to the Kruskal-Wallis test. It is typical that when a dataset has a reasonable fit to the normal distribution, the classical ANOVA test will be more sensitive to differences between groups.

To understand when a nonparametric test may be more appropriate, let's see how the tests behave when the distribution is not normal. We can simulate this by replacing one of the values by an extreme value (an outlier).

```
strength(20)=120;
anova1(strength,alloy,'off')
ans =
        0.2501
kruskalwallis(strength,alloy,'off')
```

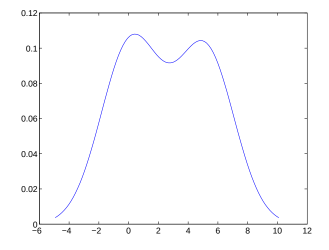
ans =
0.0060Now the classical ANOVA test does not find a significant difference, but the
nonparametric procedure does. This illustrates one of the properties of
nonparametric procedures – they are often not severely affected by changes in
a small portion of the data.Reference[1] Hollander, M., and D. A. Wolfe, Nonparametric Statistical Methods, Wiley,
1973.See Alsoanova1, boxplot, multcompare

ksdensity

Compute density estimate using a kernel smoothing method				
<pre>[f,xi] = ksdensity(x) f = ksdensity(x,xi) [f,xi,u] = ksdensity() [] = ksdensity(,'param1',val1,'param2',val2,)</pre>				
<pre>[f,xi] = ksdensity(x) computes a probability density estimate of the sample in the vector x. f is the vector of density values evaluated at the points in xi. The estimate is based on a normal kernel function, using a window parameter('width') that is a function of the number of points in x. The density is evaluated at 100 equally-spaced points covering the range of the data in x.</pre>				
<pre>f = ksdensity(x,xi) specifies the vector xi of values where the density estimate is to be evaluated.</pre>				
<pre>[f,xi,u] = ksdensity() also returns the width of the kernel smoothing window.</pre>				
[] = ksdensity(, 'param1', val1, 'param2', val2,) specifies parameter name/value pairs to control the density estimation. Valid parameters and their possible values are:				
'kernel'	The type of kernel smoother to use. Choose the value as 'normal' (default), 'box', 'triangle', or 'epanechinikov'.			
	Alternatively, you can specify some other function, as a function handle or as a string, e.g., @normpdf or 'normpdf'. The function must take a single argument that is an array of distances between data values and places where the density is evaluated. It must return an array of the same size containing corresponding values of the kernel function.			
'npoints'	The number of equally-spaced points in xi. Default is 100.			
'width'	The bandwidth of the kernel smoothing window. The default is optimal for estimating normal densities, but you may want to choose a smaller value to reveal features such as multiple modes.			
	<pre>[f,xi] = ksden f = ksdensity [f,xi,u] = ksden [f,xi] = ksden sample in the va- in xi. The estim parameter ('wid is evaluated at f = ksdensity estimate is to be [f,xi,u] = ksden parameter nam parameters and 'kernel' 'npoints'</pre>			

Examples This example generates a mixture of two normal distributions, and plots the estimated density.

```
x = [randn(30,1); 5+randn(30,1)];
[f,xi] = ksdensity(x);
plot(xi,f);
```



See Also hist, @(function handle)

References [1] Bowman, A.W. and A. Azzalini, *Applied Smoothing Techniques for Data Analysis*, Oxford University Press, 1997.

kstest

Purpose	Kolmogorov-Smirnov test of the distribution of one sample
Syntax	<pre>H = kstest(X) H = kstest(X,cdf) H = kstest(X,cdf,alpha,tail) [H,P,KSSTAT,CV] = kstest(X,cdf,alpha,tail)</pre>
Description	H = kstest(X) performs a Kolmogorov-Smirnov test to compare the values in the data vector X with a standard normal distribution (that is, a normal distribution having mean 0 and variance 1). The null hypothesis for the Kolmogorov-Smirnov test is that X has a standard normal distribution. The alternative hypothesis that X does not have that distribution. The result H is 1 if we can reject the hypothesis that X has a standard normal distribution, or 0 if we cannot reject that hypothesis. We reject the hypothesis if the test is significant at the 5% level.
	For each potential value x , the Kolmogorov-Smirnov test compares the proportion of values less than x with the expected number predicted by the standard normal distribution. The kstest function uses the maximum difference over all x values is its test statistic. Mathematically, this can be written as
	$\max(F(x) - G(x))$
	where $F(x)$ is the proportion of X values less than or equal to x and $G(x)$ is the standard normal cumulative distribution function evaluated at x .
	H = kstest(X, cdf) compares the distribution of X to the hypothesized distribution defined by the two-column matrix cdf. Column one contains a set of possible x values, and column two contains the corresponding hypothesized cumulative distribution function values $G(x)$. If possible, you should define cdf so that column one contains the values in X. If there are values in X not found in column one of cdf, kstest will approximate $G(X)$ by interpolation. All values in X must lie in the interval between the smallest and largest values in the first column of cdf. If the second argument is empty (cdf = []), kstest uses the standard normal distribution as if there were no second argument.
	The Kolmogorov-Smirnov test requires that cdf be predetermined. It is not

The Kolmogorov-Smirnov test requires that cdf be predetermined. It is not accurate if cdf is estimated from the data. To test X against a normal distribution without specifying the parameters, use lillietest instead.

 $\begin{array}{l} \mathsf{H} = \mathsf{kstest}(\mathsf{X},\mathsf{cdf},\mathsf{alpha},\mathit{tail}) \text{ specifies the significance level alpha and a } \\ \mathsf{code} \ \mathit{tail} \ \mathsf{for} \ \mathsf{the type of alternative hypothesis. If} \ \mathit{tail} = \mathsf{0} \ (\mathsf{the default}), \\ \mathsf{kstest} \ \mathsf{performs} \ \mathsf{a} \ \mathsf{two-sided} \ \mathsf{test} \ \mathsf{with the general alternative} \ F \neq G \ \mathsf{. If} \\ \mathit{tail} = \mathsf{-1}, \ \mathsf{the alternative is that} \ F < G \ \mathsf{. If} \ \mathit{tail} = \mathsf{1}, \ \mathsf{the alternative is} \ F > G \ \mathsf{.} \\ \mathsf{The form of the test statistic depends on the value of \ \mathit{tail} \ \mathsf{as follows}. \end{array}$

```
tail = 0: \max(|F(x) - G(x)|)
tail = -1: \max(G(x) - F(x))
tail = 1: \max(F(x) - G(x))
```

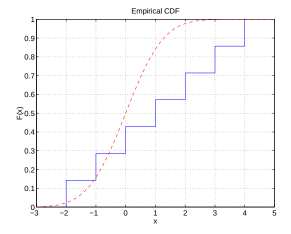
[H,P,KSSTAT,CV] = kstest(X,cdf,alpha,tail) also returns the observed p-value P, the observed Kolmogorov-Smirnov statistic KSSTAT, and the cutoff value CV for determining if KSSTAT is significant. If the return value of CV is NaN, then kstest determined the significance calculating a p-value according to an asymptotic formula rather than by comparing KSSTAT to a critical value.

Examples Example 1. Let's generate some evenly spaced numbers and perform a Kolmogorov-Smirnov test to see how well they fit to a normal distribution:

```
x = -2:1:4
x =
                  0
                       1
                               2
                                      3
    -2
          - 1
                                            4
[h,p,k,c] = kstest(x,[],0.05,0)
h =
     0
p =
      0.13632
k =
      0.41277
c =
      0.48342
```

We cannot reject the null hypothesis that the values come from a standard normal distribution. Although intuitively it seems that these evenly-spaced integers could not follow a normal distribution, this example illustrates the difficulty in testing normality in small samples. To understand the test, it is helpful to generate an empirical cumulative distribution plot and overlay the theoretical normal distribution.

```
xx = -3:.1:5;
cdfplot(x)
hold on
plot(xx,normcdf(xx),'r--')
```



The Kolmogorov-Smirnov test statistic is the maximum difference between these curves. It appears that this maximum of 0.41277 occurs as we approach x = 1.0 from below. We can see that the empirical curve has the value 3/7 here, and we can easily verify that the difference between the curves is 0.41277.

```
normcdf(1) - 3/7
ans =
0.41277
```

We can also perform a one-sided test. By setting tail = -1 we indicate that our alternative is F < G, so the test statistic counts only points where this inequality is true.

0.068181 k = 0.41277

The test statistic is the same as before because in fact F < G at x = 1.0. However, the p-value is smaller for the one-sided test. If we carry out the other one-sided test, we see that the test statistic changes, and is the difference between the two curves near x = -1.0.

Example 2. Now let's generate random numbers from a Weibull distribution, and test against that Weibull distribution and an exponential distribution.

See Also kstest2, lillietest

kstest2

Purpose	Kolmogorov-Smirnov test to compare the distribution of two samples
Syntax	H = kstest2(X1,X2) H = kstest2(X1,X2,alpha, <i>tail</i>) [H,P,KSSTAT] = kstest(X,cdf,alpha, <i>tail</i>)
Description	H = kstest2(X1, X2) performs a two-sample Kolmogorov-Smirnov test to compare the distributions of values in the two data vectors X1 and X2 of length n1 and n2, respectively. The null hypothesis for this test is that X1 and X2 have the same continuous distribution. The alternative hypothesis is that they have different continuous distributions. The result H is 1 if we can reject the hypothesis that the distributions are the same, or 0 if we cannot reject that hypothesis. We reject the hypothesis if the test is significant at the 5% level.
	For each potential value x , the Kolmogorov-Smirnov test compares the proportion of X1 values less than x with proportion of X2 values less than x . The kstest2 function uses the maximum difference over all x values is its test statistic. Mathematically, this can be written as
	$\max(F1(x) - F2(x))$
	where $F1(x)$ is the proportion of X1 values less than or equal to x and $F2(x)$ is the proportion of X2 values less than or equal to x . Missing observations, indicated by NaNs are ignored.
	$ \begin{array}{l} {\sf H} \ = \ {\sf kstest2(X1,X2,alpha,tail)} \ {\sf specifies the significance level alpha and a code tail for the type of alternative hypothesis. If tail = 0 (the default), {\sf kstest2} \ {\sf performs a two-sided test with the general alternative } F1 \neq F2 \ . If tail = -1, the alternative is that F1 < F2 \ . If tail = 1, the alternative is {\sf F1} > F2 \ . The form of the test statistic depends on the value of tail as follows: } \end{array} $
	$tail = 0: \max(F1(x) - F2(x))$
	$tail = -1: \max(F2(x) - F1(x))$
	$tail = 1: \max(F1(x) - F2(x))$
	[H,P,KSSTAT] = kstest2() also returns the observed p-value P, and the Kolmogorov-Smirnov test statistic KSSTAT defined above for the test type

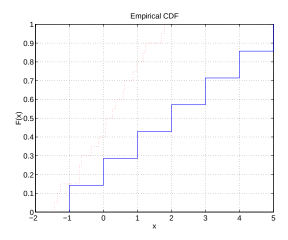
The asymptotic p-value becomes very accurate for large sample sizes, and is believed to be reasonably accurate for sample sizes n1 and n2 such that $(n1*n2)/(n1 + n2) \ge 4$.

Examples

Let's compare the distributions of a small evenly-spaced sample and a larger normal sample:

The difference between their distributions is significant at the 5% level (p = 4%). To visualize the difference, we can overlay plots of the two empirical cumulative distribution functions. The Kolmogorov-Smirnov statistic is the maximum difference between these functions. After changing the color and line style of one of the two curves, we can see that the maximum difference appears to be near x = 1.9. We can also verify that the difference equals the k value that kstest2 reports:

```
cdfplot(x)
hold on
cdfplot(y)
h = findobj(gca,'type','line');
set(h(1),'linestyle',':','color','r')
1 - 3/7
ans =
0.5714
```



See Also

kstest, lillietest

kurtosis

Purpose	Sample kurtosis	
Syntax	k = kurtosis(X) k = kurtosis(X, <i>flag</i>)	
Description	<pre>k = kurtosis(X) returns the sample kurtosis of X. For vectors, kurtosis(x) is the kurtosis of the elements in the vector x. For matrices kurtosis(X) returns the sample kurtosis for each column of X.</pre>	
	Kurtosis is a measure of how outlier-prone a distribution is. The kurtosis of the normal distribution is 3. Distributions that are more outlier-prone than the normal distribution have kurtosis greater than 3; distributions that are less outlier-prone have kurtosis less than 3.	
	The kurtosis of a distribution is defined as	
	$k = \frac{E(x-\mu)^4}{\sigma^4}$	
	where μ is the mean of x , σ is the standard deviation of x , and $E(t)$ represents the expected value of the quantity t .	
Note Some definitions of kurtosis subtract 3 from the computed value, so that the normal distribution has kurtosis of 0. The kurtosis function does use this convention.		
	k = kurtosis(X, flag) specifies whether to correct for bias $(flag = 0)$ or not $(flag = 1, the default)$. When X represents a sample from a population, the kurtosis of X is biased, that is, it will tend to differ from the population kurtosis by a systematic amount that depends on the size of the sample. You can set $flag = 0$ to correct for this systematic bias.	
Example	X = randn([5 4])	
-	X =	

X =			
1.1650	1.6961	-1.4462	-0.3600
0.6268	0.0591	-0.7012	-0.1356
0.0751	1.7971	1.2460	-1.3493

0.3516 0.2641 -0.6390 -1.2704 -0.6965 0.8717 0.5774 0.9846 k = kurtosis(X) k = 2.1658 1.2967 1.6378 1.9589

See Also

mean, moment, skewness, std, var

leverage

Purpose	Leverage values for a regression	
Syntax	h = leverage(data) h = leverage(data,' <i>model</i> ')	
Description	<pre>h = leverage(data) finds the leverage of each row (point) in the matrix data for a linear additive regression model.</pre>	
	<pre>h = leverage(data, 'model') finds the leverage on a regression, using a specified model type, where 'model' can be one of these strings:</pre>	
	 'linear' – includes constant and linear terms 	
	• 'interaction' – includes constant, linear, and cross product terms	
	 'quadratic' – includes interactions and squared terms 	
	ullet 'purequadratic' – includes constant, linear, and squared terms	
	Leverage is a measure of the influence of a given observation on a regression due to its location in the space of the inputs.	
Example	One rule of thumb is to compare the leverage to $2p/n$ where <i>n</i> is the number of observations and <i>p</i> is the number of parameters in the model. For the Hald dataset this value is 0.7692.	
	load hald h = max(leverage(ingredients,'linear'))	
	h = 0.7004	
	Since 0.7004 < 0.7692, there are no high leverage points using this rule.	
Algorithm	<pre>[Q,R] = qr(x2fx(data,'model'));</pre>	
	leverage = (sum(Q'.*Q'))'	
Reference	[1] Goodall, C. R. (1993). <i>Computation using the QR decomposition</i> . Handbook in Statistics, Volume 9. Statistical Computing (C. R. Rao, ed.). Amsterdam, NL Elsevier/North-Holland.	
See Also	regstats	

Ihsdesign

Purpose	Generate a latin hypercube sample		
Syntax	X = lhsdesign(n,p) ,'smooth','off') ,'criterion','c') ,'iterations',k)	
Description	X = 1hsdesign(n,p) generates a latin hypercube sample X containing n values on each of p variables. For each column, the n values are randomly distributed with one from each interval (0,1/n), (1/n,2/n),, (1-1/n,1), and they are randomly permuted.		
	X = lhsdesign(, 'smooth', 'off') produces points at the midpoints of the above intervals: 0.5/n, 1.5/n,, 1-0.5/n. The default is 'on'.		
	X = lhsdesign(, criterion', c') iteratively generates latin hypercube samples to find the best one according to the criterion 'c', which can be:		
	'none'	No iteration	
	'maximin'	Maximize minimum distance between points	
	'correlation'	Reduce correlation	
	X = lhsdesign(, 'iterations', k) iterates up to k times in an attempt improve the design according to the specified criterion. Default is $K = 5$. Latin hypercube designs are useful when you need a sample that is random b that is guaranteed to be relatively uniformly distributed over each dimension		
See Also	lhsnorm, unifrn	d	

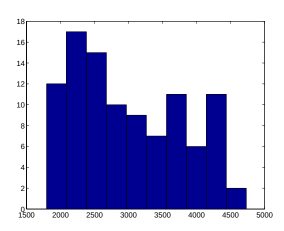
Purpose	Generate a latin hypercube sample with a normal distribution
Syntax	X = lhsnorm(mu,SIGMA,n) X = lhsnorm(mu,SIGMA,n,' <i>onoff</i> ')
Description	X = lhsnorm(mu,SIGMA,n) generates a latin hypercube sample X of size n from the multivariate normal distribution with mean vector mu and covariance matrix SIGMA. X is similar to a random sample from the multivariate normal distribution, but the marginal distribution of each column is adjusted so that its sample marginal distribution is close to its theoretical normal distribution. X = lhsnorm(mu,SIGMA,n,'onoff') controls the amount of smoothing in the sample. If 'onoff' is 'off', each column has points equally spaced on the probability scale. In other words, each column is a permutation of the values G(0.5/n), $G(1.5/n)$,, $G(1-0.5/n)$ where G is the inverse normal cumulative distribution for that column's marginal distribution. If 'onoff' is 'on' (the default), each column has points uniformly distributed on the probability scale. For example, in place of 0.5/n we use a value having a uniform distribution on the interval $(0/n, 1/n)$.
See Also	lhsdesign, mvnrnd

lillietest

Purpose	Lilliefors test for goodness of fit to a normal distribution	
Syntax	H = lillietest(X) H = lillietest(X,alpha) [H,P,LSTAT,CV] = lillietest(X,alpha)	
Description	H = lillietest(X) performs the Lilliefors test on the input data vector X and returns H, the result of the hypothesis test. The result H is 1 if we can reject the hypothesis that X has a normal distribution, or 0 if we cannot reject that hypothesis. We reject the hypothesis if the test is significant at the 5% level.	
	The Lilliefors test evaluates the hypothesis that X has a normal distribution with unspecified mean and variance, against the alternative that X does not have a normal distribution. This test compares the empirical distribution of X with a normal distribution having the same mean and variance as X. It is similar to the Kolmogorov-Smirnov test, but it adjusts for the fact that the parameters of the normal distribution are estimated from X rather than specified in advance.	
	H = lillietest(X, alpha) performs the Lilliefors test at the 100*alpha% level rather than the 5% level. alpha must be between 0.01 and 0.2.	
	[H,P,LSTAT,CV] = lillietest(X,alpha) returns three additional outputs. P is the p-value of the test, obtained by linear interpolation in a set of table created by Lilliefors. LSTAT is the value of the test statistic. CV is the critical value for determining whether to reject the null hypothesis. If the value of LSTAT is outside the range of the Lilliefors table, P is returned as NaN but H indicates whether to reject the hypothesis.	
Example	Do car weights follow a normal distribution? Not exactly, because weights are always positive, and a normal distribution allows both positive and negative values. However, perhaps the normal distribution is a reasonable approximation.	
	load carsmall [h p l c] = lillietest(Weight); [h p l c]	
	ans = 1.0000 0.0232 0.1032 0.0886	

The Lilliefors test statistic of 0.10317 is larger than the cutoff value of 0.0886 for a 5% level test, so we reject the hypothesis of normality. In fact, the p-value of this test is approximately 0.02.

To visualize the distribution, we can make a histogram. This graph shows that the distribution is skewed to the right – from the peak near 2250, the frequencies drop off abruptly to the left but more gradually to the right.



hist(Weight)

Sometimes it is possible to transform a variable to make its distribution more nearly normal. A log transformation, in particular, tends to compensate for skewness to the right.

Now the p-value is approximately 0.13, so we do not reject the hypothesis.

Reference [1] Conover, W. J. (1980). *Practical Nonparametric Statistics*. New York, Wiley.

See Also hist, jbtest, kstest2

linkage

Purpose	Create hierarchical cluster tree	
Syntax	Z = linkage(Y) Z = linkage(Y,'/	method')
Description	 Z = linkage(Y) creates a hierarchical cluster tree, using the Single Linkage algorithm. The input matrix, Y, is a distance vector of length ((m - 1) · m/2) -by-1, where m is the number of objects in the original dataset. You can generate such a vector with the pdist function. Y can also be a more general dissimilarity matrix conforming to the output format of pdist. Z = linkage(Y, 'method') computes a hierarchical cluster tree using the algorithm specified by 'method', where 'method' can be any of the following character strings that identify ways to create the cluster hierarchy. Their definitions are explained in "Mathematical Definitions" on page 12-219. 	
	'single'	Shortest distance (default)
	'complete'	Largest distance
	'average'	Average distance
	'centroid'	Centroid distance. The output Z is meaningful only if Y contains Euclidean distances.
	'ward'	Incremental sum of squares

The output, Z, is an (m-1)-by-3 matrix containing cluster tree information. The leaf nodes in the cluster hierarchy are the objects in the original dataset, numbered from 1 to m. They are the singleton clusters from which all higher clusters are built. Each newly formed cluster, corresponding to row i in Z, is assigned the index m+i, where m is the total number of initial leaves.

Columns 1 and 2, Z(i, 1:2), contain the indices of the objects that were linked in pairs to form a new cluster. This new cluster is assigned the index value m+i. There are m-1 higher clusters that correspond to the interior nodes of the hierarchical cluster tree.

Column 3, Z(i,3), contains the corresponding linkage distances between the objects paired in the clusters at each row *i*.

For example, consider a case with 30 initial nodes. If the tenth cluster formed by the linkage function combines object 5 and object 7 and their distance is 1.5, then row 10 of Z will contain the values (5, 7, 1.5). This newly formed cluster will have the index 10+30=40. If cluster 40 shows up in a later row, that means this newly formed cluster is being combined again into some bigger cluster.

Mathematical Definitions

The 'method' argument is a character string that specifies the algorithm used to generate the hierarchical cluster tree information. These linkage algorithms are based on various measurements of proximity between two groups of objects. If n_r is the number of objects in cluster r and n_s is the number of objects in cluster s, and x_{ri} is the *i*th object in cluster r, the definitions of these various measurements are as follows:

• *Single linkage*, also called *nearest neighbor*, uses the smallest distance between objects in the two groups.

$$d(r,s) = min(dist(x_{ri}, x_{si})), i \in (i, ..., n_r), j \in (1, ..., n_s)$$

• *Complete linkage*, also called *furthest neighbor*, uses the largest distance between objects in the two groups.

 $d(r,s) = max(dist(x_{ri}, x_{si})), i \in (1, ..., n_r), j \in (1, ..., n_s)$

• *Average linkage* uses the average distance between all pairs of objects in cluster *r* and cluster *s*.

$$d(r,s) = \frac{1}{n_r n_s} \sum_{i=1}^{n_r} \sum_{j=1}^{n_s} dist(x_{ri}, x_{sj})$$

• Centroid linkage uses the distance between the centroids of the two groups. $d(r,s)\,=\,d(\bar{x}_r,\bar{x}_s)$

where

$$\bar{x}_r = \frac{1}{n_r} \sum_{i=1}^{n_r} x_{ri}$$

and \bar{x}_s is defined similarly.

The centroid method can produce a cluster tree that is not monotonic. This occurs when the distance from the union of two clusters, $r \cup s$, to a third cluster is less than the distance from either r or s to that third cluster. In this case, sections of the dendrogram change direction. This is an indication that you should use another method.

• *Ward linkage* uses the incremental sum of squares; that is, the increase in the total within-group sum of squares as a result of joining groups *r* and *s*. It is given by

$$d(r,s) = n_r n_s d_{rs}^2 / (n_r + n_s)$$

where d_{rs}^2 is the distance between cluster *r* and cluster *s* defined in the Centroid linkage. The within-group sum of squares of a cluster is defined as the sum of the squares of the distance between all objects in the cluster and the centroid of the cluster.

Example	X = [3 1.7; 1 1; 2 3; 2 2.5; 1.2 1; 1.1 1.5; 3 1]; Y = pdist(X); Z = linkage(Y)		
	Z =		
	2.0000 5.0000 0.2000		
	3.0000 4.0000 0.5000		
	8.0000 6.0000 0.5099		
	1.0000 7.0000 0.7000		
	11.0000 9.0000 1.2806		
	12.0000 10.0000 1.3454		
See Also	cluster, clusterdata, cophenet, dendrogram, inconsistent, kmeans, pdist,		

silhouette, squareform

Purpose Lognormal cumulative distribution function

Syntax P = logncdf(X,MU,SIGMA)

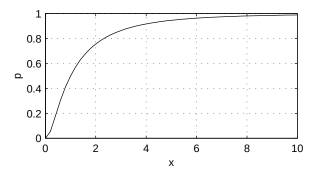
Description P = logncdf(X,MU,SIGMA) computes the lognormal cdf at each of the values in X using the corresponding means in MU and standard deviations in SIGMA. Vector or matrix inputs for X, MU, and SIGMA must have the same size, which is also the size of P. A scalar input for X, MU, or SIGMA is expanded to a constant matrix with the same dimensions as the other inputs.

The lognormal cdf is

$$p = F(x|\mu, \sigma) = \frac{1}{\sigma\sqrt{2\pi}} \int_0^x \frac{\frac{-(\ln(t) - \mu)^2}{2\sigma^2}}{t} dt$$

Example

x = (0:0.2:10); y = logncdf(x,0,1); plot(x,y); grid; xlabel('x'); ylabel('p');



Reference [1] Evans, M., N. Hastings, and B. Peacock, *Statistical Distributions, Second Edition, John Wiley and Sons, 1993.* p. 102–105.

See Also cdf, logninv, lognpdf, lognrnd, lognstat

logninv

Purpose Inverse of the lognormal cumulative distribution function (cdf)

Syntax X = logninv(P,MU,SIGMA)

Description X = logninv(P,MU,SIGMA) computes the inverse lognormal cdf with mean MU and standard deviation SIGMA, at the corresponding probabilities in P. Vector or matrix inputs for P, MU, and SIGMA must have the same size, which is also the size of X. A scalar input for P, MU, or SIGMA is expanded to a constant matrix with the same dimensions as the other inputs.

We define the lognormal inverse function in terms of the lognormal cdf as

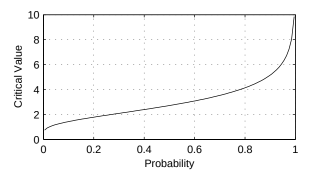
$$x = F^{-1}(p|\mu, \sigma) = \{x: F(x|\mu, \sigma) = p\}$$

where

$$p = F(x|\mu, \sigma) = \frac{1}{\sigma\sqrt{2\pi}} \int_0^x \frac{\frac{-(\ln(t) - \mu)^2}{2\sigma^2}}{t} dt$$

Example

p = (0.005:0.01:0.995); crit = logninv(p,1,0.5); plot(p,crit) xlabel('Probability');ylabel('Critical Value'); grid



Reference [1] Evans, M., N. Hastings, and B. Peacock, *Statistical Distributions, Second Edition*, John Wiley and Sons, 1993. p. 102–105.

See Also icdf, logncdf, lognpdf, lognrnd, lognstat

lognpdf

Purpose L	ognormal probability	v density function (pdf)
-----------	----------------------	--------------------------

Syntax Y = lognpdf(X,MU,SIGMA)

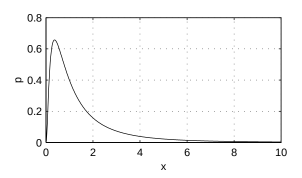
Description Y = logncdf(X,MU,SIGMA) computes the lognormal cdf at each of the values in X using the corresponding means in MU and standard deviations in SIGMA. Vector or matrix inputs for X, MU, and SIGMA must have the same size, which is also the size of Y. A scalar input for X, MU, or SIGMA is expanded to a constant matrix with the same dimensions as the other inputs

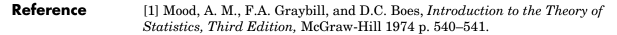
The lognormal pdf is

$$y = f(x|\mu, \sigma) = \frac{1}{x\sigma\sqrt{2\pi}}e^{\frac{-(\ln(x)-\mu)^2}{2\sigma^2}}$$

Example

x = (0:0.02:10); y = lognpdf(x,0,1); plot(x,y); grid; xlabel('x'); ylabel('p')





See Also logncdf, logninv, lognrnd, lognstat, pdf

lognrnd

Purpose	Random matrices from the lognormal distribution
Syntax	<pre>R = lognrnd(MU,SIGMA) R = lognrnd(MU,SIGMA,m) R = lognrnd(MU,SIGMA,m,n)</pre>
Description	 R = lognrnd(MU,SIGMA) generates lognormal random numbers with parameters MU and SIGMA. Vector or matrix inputs for MU and SIGMA must have the same size, which is also the size of R. A scalar input for MU or SIGMA is expanded to a constant matrix with the same dimensions as the other input. R = lognrnd(MU,SIGMA,m) generates lognormal random numbers with parameters MU and SIGMA, where m is a 1-by-2 vector that contains the row and column dimensions of R. R = lognrnd(MU,SIGMA,m,n) generates lognormal random numbers with parameters MU and SIGMA, where m is a 1-by-2 vector that contains the row and column dimensions of R.
Example	dimensions of R. $(0, 1, 4, 2)$
Example	r = lognrnd(0,1,4,3)
	$r = \\ 3.2058 & 0.4983 & 1.3022 \\ 1.8717 & 5.4529 & 2.3909 \\ 1.0780 & 1.0608 & 0.2355 \\ 1.4213 & 6.0320 & 0.4960 \\ \end{cases}$
Reference	[1] Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition,</i> John Wiley and Sons, 1993. p. 102–105.
See Also	random, logncdf, logninv, lognpdf, lognstat

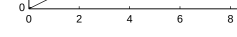
lognstat

Purpose	Mean and variance for the lognormal distribution	
Syntax	[M,V] = lognstat(MU,SIGMA)	
Description	[M,V] = lognstat(MU,SIGMA) returns the mean and variance of the lognormal distribution with parameters MU and SIGMA. Vector or matrix inputs for MU and SIGMA must have the same size, which is also the size of M and V. A scalar input for MU or SIGMA is expanded to a constant matrix with the same dimensions as the other input.	
	The mean of the lognormal distribution with parameters μ and σ is	
	$e^{\left(\mu+rac{oldsymbol{\sigma}^2}{2} ight)}$	
	and the variance is	
	$e^{(2\mu+2\sigma^2)}-e^{(2\mu+\sigma^2)}$	
Example	[m,v]= lognstat(0,1)	
	m = 1.6487	
	v = 4.6708	
Reference	[1] Mood, A. M., F.A. Graybill, and D.C. Boes, <i>Introduction to the Theory of Statistics, Third Edition,</i> McGraw-Hill 1974 p. 540–541.	
See Also	logncdf, logninv, lognrnd, lognrnd	

Isline

Purpose	Least squares fit line(s)	
Syntax	lsline h = lsline	
Description	lsline superimposes the least squares line on each line object in the current axes (except LineStyles '-','','').	
	h = lsline returns the handles to the line objects.	
Example	y = [2 3.4 5.6 8 11 12.3 13.8 16 18.8 19.9]'; plot(y,'+'); lsline;	
	5	

10



See Also

polyfit, polyval

mad

Purpose	Mean absolute deviation (MAD) of a sample of data
Syntax	y = mad(X)
Description	y = mad(X) computes the average of the absolute differences between a set of data and the sample mean of that data. For vectors, $mad(x)$ returns the mean absolute deviation of the elements of x. For matrices, $mad(X)$ returns the MAD of each column of X.
	The MAD is less efficient than the standard deviation as an estimate of the spread when the data is all from the normal distribution.
	Multiply the MAD by 1.3 to estimate σ (the second parameter of the normal distribution).
Examples	This example shows a Monte Carlo simulation of the relative efficiency of the MAD to the sample standard deviation for normal data.
	<pre>x = normrnd(0,1,100,100); s = std(x); s_MAD = 1.3 * mad(x); efficiency = (norm(s - 1)./norm(s_MAD - 1)).^2</pre>
	efficiency =
	0.5972
See Also	std, range

mahal

Purpose	Mahalanobis distance
Syntax	<pre>d = mahal(Y,X)</pre>
Description	mahal(Y,X) computes the Mahalanobis distance of each point (row) of the matrix Y from the sample in the matrix X.
	The number of columns of Y must equal the number of columns in X, but the number of rows may differ. The number of rows in X must exceed the number of columns.
	The Mahalanobis distance is a multivariate measure of the separation of a data set from a point in space. It is the criterion minimized in linear discriminant analysis.
Example	The Mahalanobis distance of a matrix \mbox{r} when applied to itself is a way to find outliers.
	r = mvnrnd([0 0],[1 0.9;0.9 1],100);
	r = [r;10 10]; d = mahal(r,r);
	last6 = $d(96:101)$
	last6 =
	1.1036
	2.2353
	2.0219
	0.3876
	1.5571
	52.7381
	The last element is clearly an outlier.

See Also classify

manova 1

Purpose	One-way Multivariate Analysis of Variance (MANOVA)
Syntax	<pre>d = manova1(X,group) d = manova1(X,group,alpha) [d,p] = manova1() [d,p,stats] = anova1()</pre>
Description	d = manoval(X, group) performs a one-way Multivariate Analysis of Variance (MANOVA) for comparing the multivariate means of the columns of X, grouped by group. X is an <i>m</i> -by- <i>n</i> matrix of data values, and each row is a vector of measurements on <i>n</i> variables for a single observation. group is a grouping variable defined as a vector, string array, or cell array of strings. Two observations are in the same group if they have the same value in the group array. The observations in each group represent a sample from a population.
	The function returns d, an estimate of the dimension of the space containing the group means. manoval tests the null hypothesis that the means of each group are the same <i>n</i> -dimensional multivariate vector, and that any difference observed in the sample X is due to random chance. If $d = 0$, there is no evidence to reject that hypothesis. If $d = 1$, then you can reject the null hypothesis at the 5% level, but you cannot reject the hypothesis that the multivariate means lie on the same line. Similarly, if $d = 2$ the multivariate means may lie on the same plane in <i>n</i> -dimensional space, but not on the same line.
	d = manova1(X,group,alpha) gives control of the significance level, alpha. The return value d will be the smallest dimension having p > alpha, where p is a p-value for testing whether the means lie in a space of that dimension.
	[d,p] = manova1() also returns a p, a vector of p-values for testing whether the means lie in a space of dimension 0, 1, and so on. The largest possible dimension is either the dimension of the space, or one less than the number of groups. There is one element of p for each dimension up to, but not including, the largest.
	If the <i>i</i> th p-value is near zero, this casts doubt on the hypothesis that the group means lie on a space of <i>i</i> -1 dimensions. The choice of a critical p-value to determine whether the result is judged "statistically significant" is left to the researcher and is specified by the value of the input argument alpha. It is common to declare a result significant if the p-value is less than 0.05 or 0.01.

[d,p,stats] = anova1(...) also returns stats, a structure containing additional MANOVA results. The structure contains the following fields.

Field	Contents
W	Within-groups sum of squares and cross-products matrix
В	Between-groups sum of squares and cross-products matrix
Т	Total sum of squares and cross-products matrix
dfW	Degrees of freedom for W
dfB	Degrees of freedom for B
dfT	Degrees of freedom for T
lambda	Vector of values of Wilk's lambda test statistic for testing whether the means have dimension 0, 1, etc.
chisq	Transformation of lambda to an approximate chi-square distribution
chisqdf	Degrees of freedom for chisq
eigenval	Eigenvalues of $W^{-1}B$
eigenvec	Eigenvectors of $W^{-1}B$; these are the coefficients for the canonical variables C, and they are scaled so the within-group variance of the canonical variables is 1
canon	Canonical variables C, equal to XC*eigenvec, where XC is X with columns centered by subtracting their means
mdist	A vector of Mahalanobis distances from each point to the mean of its group
gmdist	A matrix of Mahalanobis distances between each pair of group means

The canonical variables C are linear combinations of the original variables, chosen to maximize the separation between groups. Specifically, C(:, 1) is the linear combination of the X columns that has the maximum separation between groups. This means that among all possible linear combinations, it is the one with the most significant F statistic in a one-way analysis of variance.

 $C(:\,,2)$ has the maximum separation subject to it being orthogonal to $C(:\,,1)\,,$ and so on.

You may find it useful to use the outputs from manoval along with other functions to supplement your analysis. For example, you may want to start with a grouped scatter plot matrix of the original variables using gplotmatrix. You can use gscatter to visualize the group separation using the first two canonical variables. You can use manovacluster to graph a dendrogram showing the clusters among the group means.

Assumptions

The MANOVA test makes the following assumptions about the data in X:

- The populations for each group are normally distributed.
- The variance-covariance matrix is the same for each population.
- All observations are mutually independent.

Example

We can use manoval to determine whether there are differences in the averages of four car characteristics, among groups defined by the country where the cars were made.

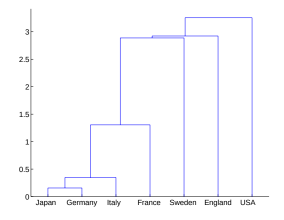
There are four dimensions in the input matrix, so the group means must lie in a four-dimensional space. manoval shows that we cannot reject the hypothesis that the means lie in a three-dimensional subspace.

References [1] Krzanowski, W. J. *Principles of Multivariate Analysis*. Oxford University Press, 1988.

See Also anova1, canoncorr, gscatter, gplotmatrix, manovacluster

manovacluster

Purpose	Plot dendrogram showing group mean clusters after MANOVA			
Syntax	manovacluster(stats) manovacluster(stats,' <i>method</i> ') H = manovacluster(stats)			
Description	manovacluster(stats) generates a dendrogram plot of the group means after a multivariate analysis of variance (MANOVA). stats is the output stats structure from manova1. The clusters are computed by applying the single linkage method to the matrix of Mahalanobis distances between group means.			
	-	r more information on the graphical output from this rogram is most useful when the number of groups is large.		
	linkage. 'method'	ats, 'method') uses the specified method in place of single can be any of the following character strings that identify cluster hierarchy. See linkage for further explanation.		
	'single'	Shortest distance (default)		
	'complete'	Largest distance		
	'average'	Average distance		
	'centroid'	Centroid distance		
	'ward'	Incremental sum of squares		
	H = manovacluster(stats, ' <i>method</i> ') returns a vector of handles to the lining the figure.			
Example	•	arger car dataset to determine which countries produce cars lar characteristics.		
		leration Weight Displacement]; manova1(X,Origin); (stats)		



See Also

cluster, dendrogram, linkage, manova1

mean

Purpose	Average or mean value of vectors and matrices		
Syntax	m = mean(X)		
Description	m = mean(X) calculates the sample average		
	$\overline{x_j} = \frac{1}{n} \sum_{i=1}^n x_{ij}$		
	For vectors, mean(x) is the mean value of the elements in vector x. For matrices, mean(X) is a row vector containing the mean value of each column.		
	The mean function is part of the standard MATLAB language.		
Example	These commands generate five samples of 100 normal random numbers with mean, zero, and standard deviation, one. The sample averages in xbar are much less variable (0.00 ± 0.10) .		
	x = normrnd(0,1,100,5); xbar = mean(x)		
	xbar =		
	0.0727 0.0264 0.0351 0.0424 0.0752		
See Also	median, std, cov, corrcoef, var		

median

Purpose	Median value of vectors and matrices
Syntax	<pre>m = median(X)</pre>
Description	m = median(X) calculates the median value, which is the 50th percentile of a sample. The median is a robust estimate of the center of a sample of data, since outliers have little effect on it.
	For vectors, median(x) is the median value of the elements in vector x. For matrices, median(X) is a row vector containing the median value of each column. Since median is implemented using sort, it can be costly for large matrices.
	The median function is part of the standard MATLAB language.
Examples	<pre>xodd = 1:5; modd = median(xodd)</pre>
	modd =
	3
	meven = median(xeven)
	meven =
	2.5000
	This example shows robustness of the median to outliers.
	<pre>xoutlier = [x 10000]; moutlier = median(xoutlier)</pre>
	moutlier =
	3
See Also	mean, std, cov, corrcoef

mle

Purpose	Maximum likelihood estimation
Syntax	phat = mle(' <i>dist</i> ',data) [phat,pci] = mle(' <i>dist</i> ',data) [phat,pci] = mle(' <i>dist</i> ',data,alpha) [phat,pci] = mle(' <i>dist</i> ',data,alpha,p1)
Description	phat = $mle('dist', data)$ returns the maximum likelihood estimates (MLEs) for the distribution specified in 'dist' using the sample in the vector, data. See "Overview of the Distributions" on page 2-11 for the list of available distributions.
	<pre>[phat,pci] = mle('dist',data) returns the MLEs and 95% percent confidence intervals.</pre>
	<pre>[phat,pci] = mle('dist',data,alpha) returns the MLEs and 100(1-alpha)% confidence intervals given the data and the specified alpha.</pre>
	<pre>[phat,pci] = mle('dist',data,alpha,p1) is used for the binomial distribution only, where p1 is the number of trials.</pre>
Example	rv = binornd(20,0.75)
	rv = 16
	<pre>[p,pci] = mle('binomial',rv,0.05,20)</pre>
	p = 0.8000
	pci = 0.5634 0.9427
See Also	betafit, binofit, expfit, gamfit, nbinfit, normfit, normlike, poissfit,

weibfit

moment

Purpose	Central moment	of all orders	5			
Syntax	<pre>m = moment(X,or</pre>	der)				
Description	<pre>m = moment(X,order) returns the central moment of X specified by the positive integer order. For vectors, moment(x,order) returns the central moment of the specified order for the elements of x. For matrices, moment(X,order) returns central moment of the specified order for each column.</pre>				entral	
	the variance com	Note that the central first moment is zero, and the second central moment is the variance computed using a divisor of n rather than n -1, where n is the length of the vector x or the number of rows in the matrix X.				
	The central mom	ent of order	k of a distr	ibution is d	efined as	
	$m_n = E(x-\mu)$	$)^k$				
	where $E(x)$ is the	expected v	alue of <i>x</i> .			
Example	X = randn([6	5])				
	X =					
	1.1650	0.0591	1.2460	-1.2704	-0.0562	
	0.6268	1.7971	-0.6390	0.9846	0.5135	
	0.0751		0.5774	-0.0449	0.3967	
	0.3516	0.8717	-0.3600	-0.7989	0.7562	
	-0.6965	-1.4462	-0.1356	-0.7652	0.4005	
	1.6961	-0.7012	-1.3493	0.8617	-1.3414	
	<pre>m = moment(X</pre>	,3)				
	m =					
	-0.0282	0.0571	0.1253	0.1460	-0.4486	
See Also	kurtosis,mean,s	skewness, s	td,var			

multcompare

Purpose	Multiple comparison test of means or other estimates
Syntax	<pre>c = multcompare(stats) c = multcompare(stats,alpha) c = multcompare(stats,alpha,'displayopt') c = multcompare(stats,alpha,'displayopt','ctype') c = multcompare(stats,alpha,'displayopt','ctype','estimate') c = multcompare(stats,alpha,'displayopt','ctype','estimate',dim) [c,m] = multcompare() [c,m,h] = multcompare()</pre>
Description	c = multcompare(stats) performs a multiple comparison test using the information in the stats structure, and returns a matrix c of pairwise comparison results. It also displays an interactive figure presenting a graphical representation of the test.
	In a one-way analysis of variance, you compare the means of several groups to test the hypothesis that they are all the same, against the general alternative that they are not all the same. Sometimes this alternative may be too general. You may need information about which pairs of means are significantly different, and which are not. A test that can provide such information is called a "multiple comparison procedure."
	When you perform a simple t-test of one group mean against another, you specify a significance level that determines the cutoff value of the t statistic. For example, you can specify the value $alpha = 0.05$ to insure that when there is no real difference, you will incorrectly find a significant difference no more than 5% of the time. When there are many group means, there are also many pairs to compare. If you applied an ordinary t-test in this situation, the $alpha$ value would apply to each comparison, so the chance of incorrectly finding a significant difference would increase with the number of comparisons. Multiple comparison procedures are designed to provide an upper bound on the probability that <i>any</i> comparison will be incorrectly found significant.
	The output c contains the results of the test in the form of a five-column matrix. Each row of the matrix represents one test, and there is one row for each pair of groups. The entries in the row indicate the means being compared, the estimated difference in means, and a confidence interval for the difference.
	For example, suppose one row contains the following entries.

2.0000 5.0000 1.9442 8.2206 14.4971

These numbers indicate that the mean of group 2 minus the mean of group 5 is estimated to be 8.2206, and a 95% confidence interval for the true mean is [1.9442, 14.4971].

In this example the confidence interval does not contain 0.0, so the difference is significant at the 0.05 level. If the confidence interval did contain 0.0, the difference would not be significant at the 0.05 level.

The multcompare function also displays a graph with each group mean represented by a symbol and an interval around the symbol. Two means are significantly different if their intervals are disjoint, and are not significantly different if their intervals overlap. You can use the mouse to select any group, and the graph will highlight any other groups that are significantly different from it.

c = multcompare(stats, alpha) determines the confidence levels of the intervals in the c matrix and in the figure. The confidence level is 100*(1-alpha)%. The default value of alpha is 0.05.

c = multcompare(stats, alpha, 'displayopt') enables the graph display when 'displayopt' is 'on' (default) and suppresses the display when 'displayopt' is 'off'.

<pre>c = multcompare(stats,alpha,'displayopt','ctype') specifies the critical</pre>
value to use for the multiple comparison, which can be any of the following.

ctype	Meaning
'hsd',or 'tukey-kramer'	Use Tukey's honestly significant difference criterion. This is the default, and it is based on the Studentized range distribution. It is optimal for balanced one-way ANOVA and similar procedures with equal sample sizes. It has been proven to be conservative for one-way ANOVA with different sample sizes. According to the unproven Tukey-Kramer conjecture, it is also accurate for problems where the quantities being compared are correlated, as in analysis of covariance with unbalanced covariate values.
'lsd'	Use Tukey's least significant difference procedure. This procedure is a simple t-test. It is reasonable if the preliminary test (say, the one-way ANOVA F statistic) shows a significant difference. If it is used unconditionally, it provides no protection against multiple comparisons.
'bonferroni'	Use critical values from the t distribution, after a Bonferroni adjustment to compensate for multiple comparisons. This procedure is conservative, but usually less so than the Scheffé procedure.
'dunn-sidak'	Use critical values from the t distribution, after an adjustment for multiple comparisons that was proposed by Dunn and proved accurate by Šidák. This procedure is similar to, but less conservative than, the Bonferroni procedure.
'scheffe'	Use critical values from Scheffé's S procedure, derived from the F distribution. This procedure provides a simultaneous confidence level for comparisons of all linear combinations of the means, and it is conservative for comparisons of simple differences of pairs.

c = multcompare(stats,alpha,'displayopt','ctype','estimate')
specifies the estimate to be compared. The allowable values of estimate depend
on the function that was the source of the stats structure, according to the
following table.

Source	Allowable Values of Estimate
'anova1'	Ignored. Always compare the group means.
'anova2'	Either 'column' (the default) or 'row' to compare column or row means.
'anovan'	Ignored. Always compare the population marginal means as specified by the dim argument.
'aoctool'	Either 'slope', 'intercept', or 'pmm' to compare slopes, intercepts, or population marginal means. If the analysis of covariance model did not include separate slopes, then 'slope' is not allowed. If it did not include separate intercepts, then no comparisons are possible.
'friedman'	Ignored. Always compare average column ranks.
'kruskalwallis'	Ignored. Always compare average group ranks.

c = multcompare(stats, alpha, 'displayopt', 'ctype', 'estimate', dim) specifies the population marginal means to be compared. This argument is used only if the input stats structure was created by the anovan function. For n-way ANOVA with n factors, you can specify dim as a scalar or a vector of integers between 1 and n. The default value is 1.

For example, if dim = 1, the estimates that are compared are the means for each value of the first grouping variable, adjusted by removing effects of the other grouping variables as if the design were balanced. If dim = $[1 \ 3]$, population marginal means are computed for each combination of the first and third grouping variables, removing effects of the second grouping variable. If you fit a singular model, some cell means may not be estimable and any population marginal means that depend on those cell means will have the value NaN.

Population marginal means are described by Milliken and Johnson (1992) and by Searle, Speed, and Milliken (1980). The idea behind population marginal means is to remove any effect of an unbalanced design by fixing the values of the factors specified by dim, and averaging out the effects of other factors as if each factor combination occurred the same number of times. The definition of population marginal means does not depend on the number of observations at each factor combination. For designed experiments where the number of observations at each factor combination has no meaning, population marginal means can be easier to interpret than simple means ignoring other factors. For surveys and other studies where the number of observations at each combination does have meaning, population marginal means may be harder to interpret.

[c,m] = multcompare(...) returns an additional matrix m. The first column
of m contains the estimated values of the means (or whatever statistics are
being compared) for each group, and the second column contains their standard
errors.

[c,m,h] = multcompare(...) returns a handle h to the comparison graph. Note that the title of this graph contains instructions for interacting with the graph, and the *x*-axis label contains information about which means are significantly different from the selected mean. If you plan to use this graph for presentation, you may want to omit the title and the *x*-axis label. You can remove them using interactive features of the graph window, or you can use the following commands.

```
title('')
xlabel('')
```

Example

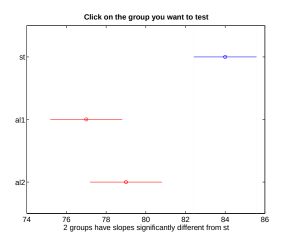
Let's revisit the anoval example testing the material strength in structural beams. From the anoval output we found significant evidence that the three types of beams are not equivalent in strength. Now we can determine where those differences lie. First we create the data arrays and we perform one-way ANOVA.

```
strength = [82 86 79 83 84 85 86 87 74 82 78 75 76 77 79 ...
79 77 78 82 79];
alloy = {'st','st','st','st','st','st','st',...
'al1','al1','al1','al1','al1','al1',...
'al2','al2','al2','al2','al2','al2'};
```

```
[p,a,s] = anova1(strength,alloy);
```

Among the outputs is a structure that we can use as input to multcompare.

multcompare(s)				
ans =				
1.0000	2.0000	3.6064	7.0000	10.3936
1.0000	3.0000	1.6064	5.0000	8.3936
2.0000	3.0000	-5.6280	-2.0000	1.6280



The third row of the output matrix shows that the differences in strength between the two alloys is not significant. A 95% confidence interval for the difference is [-5.6, 1.6], so we cannot reject the hypothesis that the true difference is zero.

The first two rows show that both comparisons involving the first group (steel) have confidence intervals that do not include zero. In other words, those differences are significant. The graph shows the same information.

See Also anova1, anova2, anovan, aoctool, friedman, kruskalwallis

References [1] Hochberg, Y., and A. C. Tamhane, *Multiple Comparison Procedures*, 1987, Wiley.

[2] Milliken, G. A., and D. E. Johnson, *Analysis of Messy Data, Volume 1: Designed Experiments*, 1992, Chapman & Hall.

[3] Searle, S. R., F. M. Speed, and G. A. Milliken, "Population marginal means in the linear model: an alternative to least squares means," *American Statistician*, 1980, pp. 216-221.

Purpose	Multivariate normal probability density function (pdf)
Syntax	<pre>y = mvnpdf(X) y = mvnpdf(X,MU) y = mvnpdf(X,MU,SIGMA)</pre>
Description	y = mvnpdf(X) returns the n-by-1 vector y, containing the probability density of the multivariate normal distribution with zero mean and identity covariance matrix, evaluated at each row of the n-by-d matrix X. Rows of X correspond to observations and columns correspond to variables or coordinates.
	y = mvnpdf(X,MU) returns the density of the multivariate normal distribution with mean MU and identity covariance matrix, evaluated at each row of X. MU is a 1-by-d vector, or an n-by-d matrix. If MU is a matrix, the density is evaluated for each row of X with the corresponding row of MU. MU can also be a scalar value, which mvnpdf replicates to match the size of X.
	<pre>y = mvnpdf(X,MU,SIGMA) returns the density of the multivariate normal distribution with mean MU and covariance SIGMA, evaluated at each row of X. SIGMA is a d-by-d matrix, or an d-by-d-by-n array, in which case the density is evaluated for each row of X with the corresponding page of SIGMA, i.e., mvnpdf computes y(i) using X(i,:) and SIGMA(:,:,i). Specify [] for MU to use its default value when you want to specify only SIGMA.</pre>
	If X is a 1-by-d vector, mvnpdf replicates it to match the leading dimension of MU or the trailing dimension of SIGMA.
Example	<pre>mu = [1 -1]; Sigma = [.9 .4; .4 .3]; X = mvnrnd(mu,Sigma,10); p = mvnpdf(X,mu,Sigma);</pre>
See Also	mvnrnd, normpdf

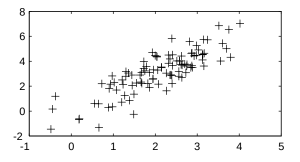
mvnrnd

Purpose	Random matrices from the multivariate normal distribution
Syntax	R = mvnrnd(MU,SIGMA) R = mvnrnd(MU,SIGMA,cases)
Description	R = mvnrnd(MU, SIGMA) returns an n-by-d matrix R of random vectors chosen from the multivariate normal distribution with mean MU, and covariance SIGMA. MU is an n-by-d matrix, and mvnrnd generates each row of R using the corresponding row of MU. SIGMA is a d-by-d symmetric positive semi-definite matrix, or a d-by-d-by-n array. If SIGMA is an array, mvnrnd generates each row of R using the corresponding page of SIGMA, i.e., mvnrnd computes $R(i,:)$ using MU(i,:) and SIGMA(:,:,i). If MU is a 1-by-d vector, mvnrnd replicates it to match the trailing dimension of SIGMA.

r = mvnrnd(MU, SIGMA, cases) returns a cases-by-d matrix R of random vectors chosen from the multivariate normal distribution with a common 1-by-d mean vector MU, and a common d-by-d covariance matrix SIGMA.

Example

mu = [2 3]; sigma = [1 1.5; 1.5 3]; r = mvnrnd(mu,sigma,100); plot(r(:,1),r(:,2),'+')





lhsnorm, mvnpdf, normrnd

Purpose Random matrices from the multivariate t distribution

r = mvtrnd(C, df, cases)

Syntax

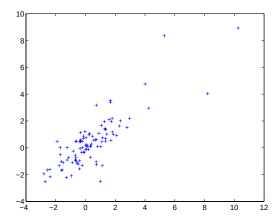
Description r = mvtrnd(C,df,cases) returns a matrix of random numbers chosen from the multivariate t distribution, where C is a correlation matrix. df is the degrees of freedom and is either a scalar or is a vector with cases elements. If p is the number of columns in C, then the output r has cases rows and p columns.

Let t represent a row of r. Then the distribution of t is that of a vector having a multivariate normal distribution with mean 0, variance 1, and covariance matrix C, divided by an independent chi-square random value having df degrees of freedom. The rows of r are independent.

C must be a square, symmetric and positive definite matrix. If its diagonal elements are not all 1 (that is, if C is a covariance matrix rather than a correlation matrix), mvtrnd computes the equivalent correlation matrix before generating the random numbers.

Example

sigma = [1 0.8;0.8 1]; r = mvtrnd(sigma,3,100); plot(r(:,1),r(:,2),'+')



See Also

mvnrnd, trnd

nanmax

Purpose	Maximum ignoring NaNs
Syntax	<pre>m = nanmax(a) [m,ndx] = nanmax(a) m = nanmax(a,b)</pre>
Description	<pre>m = nanmax(a) returns the maximum with NaNs treated as missing. For vectors, nanmax(a) is the largest non-NaN element in a. For matrices, nanmax(A) is a row vector containing the maximum non-NaN element from each column.</pre>
	[m,ndx] = nanmax(a) also returns the indices of the maximum values in vector ndx.
	<pre>m = nanmax(a,b) returns the larger of a or b, which must match in size.</pre>
Example	m = magic(3); m([1 6 8]) = [NaN NaN NaN]
	m =
	NaN 1 6
	3 5 NaN
	4 NaN 2
	[nmax,maxidx] = nanmax(m)
	nmax =
	4 5 6
	maxidx =
	3 2 1
Can Alen	

See Also nanmin, nanmean, nanmedian, nanstd, nansum

nanmean

Purpose	Mean ignoring NaNs
Syntax	y = nanmean(X)
Description	y = nanmean(X) is the average computed by treating NaNs as missing values. For vectors, nanmean(x) is the mean of the non-NaN elements of x. For matrices, nanmean(X) is a row vector containing the mean of the non-NaN elements in each column.
Example	m = magic(3); m([1 6 8]) = [NaN NaN NaN] m =
	NaN 1 6 3 5 NaN 4 NaN 2
	nmean = nanmean(m)
	nmean =
	3.5000 3.0000 4.0000
See Also	nanmin, nanmax, nanmedian, nanstd, nansum

nanmedian

Purpose	Median ignoring NaNs
Syntax	y = nanmedian(X)
Description	y = nanmedian(X) is the median computed by treating NaNs as missing values.
	For vectors, nanmedian(x) is the median of the non-NaN elements of x. For matrices, nanmedian(X) is a row vector containing the median of the non-NaN elements in each column of X.
Example	m = magic(4); m([1 6 9 11]) = [NaN NaN NaN NaN]
	m =
	NaN 2 NaN 13
	5 NaN 10 8
	9 7 NaN 12
	4 14 15 1
	nmedian = nanmedian(m)
	nmedian =
	5.0000 7.0000 12.5000 10.0000
See Also	nanmin, nanmax, nanmean, nanstd, nansum

nanmin

Purpose	Minimum ignoring NaNs
Syntax	m = nanmin(a) [m,ndx] = nanmin(a) m = nanmin(a,b)
Description	<pre>m = nanmin(a) is the minimum computed by treating NaNs as missing values. For vectors, nanmin(a) is the smallest non-NaN element in a. For matrices, nanmin(A) is a row vector containing the minimum non-NaN element from each column.</pre>
	[m,ndx] = nanmin(a) also returns the indices of the minimum values in vector ndx.
	<pre>m = nanmin(a,b) returns the smaller of a or b, which must match in size.</pre>
Example	m = magic(3); m([1 6 8]) = [NaN NaN NaN]
	m =
	NaN 1 6
	3 5 NaN 4 NaN 2
	[nmin,minidx] = nanmin(m)
	nmin =
	3 1 2
	minidx =
	2 1 3
Can Alan	

See Also nanmax, nanmean, nanmedian, nanstd, nansum

nanstd

Purpose	Standard deviation ignoring NaNs
Syntax	y = nanstd(X)
Description	y = nanstd(X) is the standard deviation computed by treating NaNs as missing values.
	For vectors, $nanstd(x)$ is the standard deviation of the non-NaN elements of x. For matrices, $nanstd(X)$ is a row vector containing the standard deviations of the non-NaN elements in each column of X.
Example	<pre>m = magic(3); m([1 6 8]) = [NaN NaN]</pre>
	m = NaN 1 6 3 5 NaN 4 NaN 2
	<pre>nstd = nanstd(m)</pre>
	nstd =
	0.7071 2.8284 2.8284
See Also	nanmax, nanmin, nanmean, nanmedian, nansum

nansum

Purpose	Sum ignoring NaNs
Syntax	y = nansum(X)
Description	y = nansum(X) is the sum computed by treating NaNs as missing values. For vectors, $nansum(x)$ is the sum of the non-NaN elements of x. For matrices, $nansum(X)$ is a row vector containing the sum of the non-NaN elements in each column of X.
Example	<pre>m = magic(3); m([1 6 8]) = [NaN NaN NaN] m =</pre>
	7 6 8
See Also	nanmax, nanmin, nanmean, nanmedian, nanstd

nbincdf

Syntax Y = nbincdf(X,R,P)

Description Y = nbincdf(X,R,P) computes the negative binomial cdf at each of the values in X using the corresponding parameters in R and P. Vector or matrix inputs for X, R, and P must have the same size, which is also the size of Y. A scalar input for X, R, or P is expanded to a constant matrix with the same dimensions as the other inputs.

The negative binomial cdf is

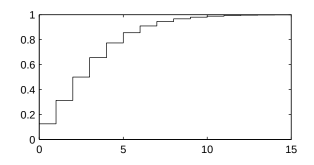
$$y = F(x|r,p) = \sum_{i=0}^{x} {r+i-1 \choose i} p^{r} q^{i} I_{(0,1,...)}(i)$$

The simplest motivation for the negative binomial is the case of successive random trials, each having a constant probability P of success. The number of *extra* trials you must perform in order to observe a given number R of successes has a negative binomial distribution. However, consistent with a more general interpretation of the negative binomial, nbincdf allows R to be any positive value, including nonintegers. When R is noninteger, the binomial coefficient in the definition of the cdf is replaced by the equivalent expression

$$\frac{\Gamma(r+i)}{\Gamma(r)\Gamma(i+1)}$$

Example

x = (0:15); p = nbincdf(x,3,0.5); stairs(x,p)



See Also

cdf, nbinfit, nbininv, nbinpdf, nbinrnd, nbinstat

nbinfit

Purpose	Parameter estimates and confidence intervals for negative binomial data
Syntax	parmhat = nbinfit(x) [parmhat,parmci] = nbinfit(x,alpha) [] = nbinfit(,options)
Description	parmhat = $nbinfit(x)$ returns the maximum likelihood estimates (MLEs) of the parameters of the negative binomial distribution given the data in the vector x.
	<pre>[parmhat,parmci] = nbinfit(x,alpha) returns MLEs and 100*(1-alpha) percent confidence intervals. By default, alpha = 0.05, which corresponds to 95% confidence intervals.</pre>
	[] = nbinfit(,options) specifies control parameters for the numerical optimization used to compute MLEs. Create this argument with the MATLAB optimset function. The default is optimset('Display', 'notify').
	Note The variance of a negative binomial distribution is greater than its mean. If the sample variance of the data x is less than its sample mean, nbinfit cannot compute MLEs. You should use the poissfit function instead.
See Also	nbincdf, nbininv, nbinpdf, nbinrnd, nbinstat, mle, optimset

Purpose	Inverse of the negative binomial cumulative distribution function (cdf)
Syntax	X = nbininv(Y,R,P)
Description	X = nbininv(Y,R,P) returns the inverse of the negative binomial cdf with parameters R and P at the corresponding probabilities in P. Since the binomial distribution is discrete, nbininv returns the least integer X such that the negative binomial cdf evaluated at X equals or exceeds Y. Vector or matrix inputs for Y, R, and P must have the same size, which is also the size of X. A scalar input for Y, R, or P is expanded to a constant matrix with the same dimensions as the other inputs.
	The simplest motivation for the negative binomial is the case of successive random trials, each having a constant probability P of success. The number of <i>extra</i> trials you must perform in order to observe a given number R of successes has a negative binomial distribution. However, consistent with a more general interpretation of the negative binomial, nbininv allows R to be any positive value, including nonintegers.
Example	How many times would you need to flip a fair coin to have a 99% probability of having observed 10 heads?
	flips = nbininv(0.99,10,0.5) + 10
	flips =
	33
	Note that you have to flip at least 10 times to get 10 heads. That is why the second term on the right side of the equals sign is a 10.
See Also	icdf, nbincdf, nbinfit, nbinpdf, nbinrnd, nbinstat

nbinpdf

Purpose	Negative binomial probability density function
---------	--

Syntax Y = nbinpdf(X,R,P)

Description Y = nbinpdf(X,R,P) returns the negative binomial pdf at each of the values in X using the corresponding parameters in R and P. Vector or matrix inputs for X, R, and P must have the same size, which is also the size of Y. A scalar input for X, R, or P is expanded to a constant matrix with the same dimensions as the other inputs. Note that the density function is zero unless the values in X are integers.

The negative binomial pdf is

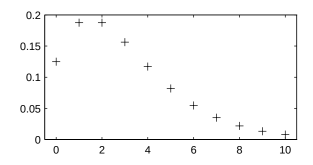
$$y = f(x|r,p) = {r+x-1 \choose x} p^r q^x I_{(0, 1, ...)}(x)$$

The simplest motivation for the negative binomial is the case of successive random trials, each having a constant probability P of success. The number of *extra* trials you must perform in order to observe a given number R of successes has a negative binomial distribution. However, consistent with a more general interpretation of the negative binomial, nbinpdf allows R to be any positive value, including nonintegers. When R is noninteger, the binomial coefficient in the definition of the pdf is replaced by the equivalent expression

$$\frac{\Gamma(r+x)}{\Gamma(r)\Gamma(x+1)}$$

Example

x = (0:10); y = nbinpdf(x,3,0.5); plot(x,y,'+') set(gca,'Xlim',[-0.5,10.5])



See Also

nbincdf, nbinfit, nbininv, nbinrnd, nbinstat, pdf

nbinrnd

Purpose	Random matrices from a negative binomial distribution
Syntax	RND = nbinrnd(R,P) RND = nbinrnd(R,P,m) RND = nbinrnd(R,P,m,n)
Description	RND = nbinrnd(R,P) is a matrix of random numbers chosen from a negative binomial distribution with parameters R and P. Vector or matrix inputs for R and P must have the same size, which is also the size of RND. A scalar input for R or P is expanded to a constant matrix with the same dimensions as the other input.
	RND = nbinrnd(R,P,m) generates random numbers with parameters R and P, where m is a 1-by-2 vector that contains the row and column dimensions of RND.
	RND = nbinrnd(R,P,m,n) generates random numbers with parameters R and P, where scalars m and n are the row and column dimensions of RND.
	The simplest motivation for the negative binomial is the case of successive random trials, each having a constant probability P of success. The number of <i>extra</i> trials you must perform in order to observe a given number R of successes has a negative binomial distribution. However, consistent with a more general interpretation of the negative binomial, nbinrnd allows R to be any positive value, including nonintegers.
Example	Suppose you want to simulate a process that has a defect probability of 0.01. How many units might Quality Assurance inspect before finding three defective items?
	r = nbinrnd(3,0.01,1,6) + 3
	r =
	496 142 420 396 851 178
See Also	nbincdf, nbinfit, nbininv, nbinpdf, nbinstat

Mean and variance of the negative binomial distribution
[M,V] = nbinstat(R,P)
[M,V] = nbinstat(R,P) returns the mean and variance of the negative binomial distribution with parameters R and P. Vector or matrix inputs for R and P must have the same size, which is also the size of M and V. A scalar input for R or P is expanded to a constant matrix with the same dimensions as the other input.
The mean of the negative binomial distribution with parameters r and p is rq/p , where $q = 1-p$. The variance is rq/p^2 .
The simplest motivation for the negative binomial is the case of successive random trials, each having a constant probability P of success. The number of <i>extra</i> trials you must perform in order to observe a given number R of successes has a negative binomial distribution. However, consistent with a more general interpretation of the negative binomial, nbinstat allows R to be any positive value, including nonintegers.
<pre>p = 0.1:0.2:0.9; r = 1:5; [R,P] = meshgrid(r,p); [M,V] = nbinstat(R,P)</pre>
M =
$\begin{array}{cccccccccccccccccccccccccccccccccccc$

nbinstat

See Also nbincdf, nbinfit, nbininv, nbinpdf, nbinrnd

Purpose Noncentral F cumulative distribution function (cdf)

Syntax P = ncfcdf(X,NU1,NU2,DELTA)

Example

Description P = ncfcdf(X,NU1,NU2,DELTA) computes the noncentral F cdf at each of the values in X using the corresponding numerator degrees of freedom in NU1, denominator degrees of freedom in NU2, and positive noncentrality parameters in DELTA. Vector or matrix inputs for X, NU1, NU2, and DELTA must have the same size, which is also the size of P. A scalar input for X, NU1, NU2, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.

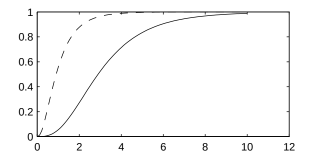
The noncentral F cdf is

$$F(x|v_1, v_2, \delta) = \sum_{j=0}^{\infty} \left(\frac{\left(\frac{1}{2}\delta\right)^j}{j!} e^{-\frac{\delta}{2}} \right) I\left(\frac{v_1 \cdot x}{v_2 + v_1 \cdot x} \middle| \frac{v_1}{2} + j, \frac{v_2}{2} \right)$$

where I(x | a, b) is the incomplete beta function with parameters *a* and *b*.

Compare the noncentral F cdf with $\delta = 10$ to the F cdf with the same number of numerator and denominator degrees of freedom (5 and 20 respectively).

x = (0.01:0.1:10.01)'; p1 = ncfcdf(x,5,20,10); p = fcdf(x,5,20); plot(x,p,'--',x,p1,'-')



References	[1] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2,</i> John Wiley and Sons, 1970. pp. 189–200.
See Also	cdf, ncfpdf, ncfinv, ncfrnd, ncfstat

Purpose	Inverse of the noncentral F cumulative distribution function (cdf)
Syntax	<pre>X = ncfinv(P,NU1,NU2,DELTA)</pre>
Description	X = ncfinv(P,NU1,NU2,DELTA) returns the inverse of the noncentral F cdf with numerator degrees of freedom NU1, denominator degrees of freedom NU2, and positive noncentrality parameter DELTA for the corresponding probabilities in P. Vector or matrix inputs for P, NU1, NU2, and DELTA must have the same size, which is also the size of X. A scalar input for P, NU1, NU2, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
Example	One hypothesis test for comparing two sample variances is to take their ratio and compare it to an F distribution. If the numerator and denominator degrees of freedom are 5 and 20 respectively, then you reject the hypothesis that the first variance is equal to the second variance if their ratio is less than that computed below.
	critical = finv(0.95,5,20)
	critical =
	2.7109
	Suppose the truth is that the first variance is twice as big as the second variance. How likely is it that you would detect this difference?
	<pre>prob = 1 - ncfcdf(critical,5,20,2)</pre>
	prob =
	0.1297
References	[1] Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition, John Wiley and Sons, 1993. p. 102–105.</i>
	[2] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2</i> , John Wiley and Sons, 1970. pp. 189–200.
See Also	icdf, ncfcdf, ncfpdf, ncfrnd, ncfstat

ncfpdf

Purpose	Noncentral F probability density function
Syntax	Y = ncfpdf(X,NU1,NU2,DELTA)
Description	Y = ncfpdf(X, NU1, NU2, DELTA) computes the noncentral F pdf at each of the values in X using the corresponding numerator degrees of freedom in NU1, denominator degrees of freedom in NU2, and positive noncentrality parameters in DELTA. Vector or matrix inputs for X, NU1, NU2, and DELTA must have the same size, which is also the size of Y. A scalar input for P, NU1, NU2, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
	The F distribution is a special case of the noncentral F where $\delta = 0$. As δ increases, the distribution flattens like the plot in the example.
Example	<pre>Compare the noncentral F pdf with δ = 10 to the F pdf with the same number of numerator and denominator degrees of freedom (5 and 20 respectively). x = (0.01:0.1:10.01)'; p1 = ncfpdf(x,5,20,10); p = fpdf(x,5,20); plot(x,p,'',x,p1,'-')</pre>
	$\begin{array}{c} 0.8 \\ 0.6 \\ 0.4 \\ 0.2 \\ 0 \\ 0 \\ 2 \\ 4 \\ 6 \\ 8 \\ 10 \\ 12 \\ 10 \\ 10 \\ 10 \\ 10 \\ 10 \\ 10$
References	[1] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2,</i> John Wiley and Sons, 1970. pp. 189–200.

See Also ncfcdf, ncfinv, ncfrnd, ncfstat, pdf

Purpose	Random matrices from the noncentral F distribution
Syntax	R = ncfrnd(NU1,NU2,DELTA) R = ncfrnd(NU1,NU2,DELTA,m) R = ncfrnd(NU1,NU2,DELTA,m,n)
Description	R = ncfrnd(NU1,NU2,DELTA) returns a matrix of random numbers chosen from the noncentral F distribution with parameters NU1, NU2 and DELTA. Vector or matrix inputs for NU1, NU2, and DELTA must have the same size, which is also the size of R. A scalar input for NU1, NU2, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
	R = ncfrnd(NU1,NU2,DELTA,m) returns a matrix of random numbers with parameters NU1, NU2, and DELTA, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = ncfrnd(NU1,NU2,DELTA,m,n) generates random numbers with parameters NU1, NU2, and DELTA, where scalars m and n are the row and column dimensions of R.
Example	Compute six random numbers from a noncentral F distribution with 10 numerator degrees of freedom, 100 denominator degrees of freedom and a noncentrality parameter, δ , of 4.0. Compare this to the F distribution with the same degrees of freedom.
	r = ncfrnd(10,100,4,1,6)
	r = 2.5995 0.8824 0.8220 1.4485 1.4415 1.4864
	r1 = frnd(10,100,1,6)
	r1 = 0.9826 0.5911 1.0967 0.9681 2.0096 0.6598
References	[1] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2,</i> John Wiley and Sons, 1970. pp. 189–200.
See Also	ncfcdf, ncfinv, ncfpdf, ncfstat

ncfstat

Purpose	Mean and variance of the noncentral F distribution	
---------	--	--

Syntax [M,V] = ncfstat(NU1,NU2,DELTA)

Description [M,V] = ncfstat(NU1,NU2,DELTA) returns the mean and variance of the noncentral F pdf with NU1 and NU2 degrees of freedom and noncentrality parameter DELTA. Vector or matrix inputs for NU1, NU2, and DELTA must have the same size, which is also the size of M and V. A scalar input for NU1, NU2, or DELTA is expanded to a constant matrix with the same dimensions as the other input.

The mean of the noncentral F distribution with parameters v_1 , v_2 , and δ is

$$\frac{\nu_2(\delta+\nu_1)}{\nu_1(\nu_2-2)}$$

where $v_2 > 2$.

The variance is

$$2 \left(\frac{\nu_2}{\nu_1}\right)^2 \left[\frac{\left(\delta + \nu_1\right)^2 + (2\delta + \nu_1)(\nu_2 - 2)}{\left(\nu_2 - 2\right)^2(\nu_2 - 4)}\right]$$

where $v_2 > 4$.

Example

[m,v]= ncfstat(10,100,4)

m = 1.4286

v = 3.9200

References[1] Evans, M., N. Hastings, and B. Peacock, Statistical Distributions, Second
Edition, John Wiley and Sons, 1993. p. 73–74.

[2] Johnson, N., and S. Kotz, *Distributions in Statistics: Continuous Univariate Distributions-2*, John Wiley and Sons, 1970. pp. 189–200.

See Also ncfcdf, ncfinv, ncfpdf, ncfrnd

Purpose	Noncentral T cumulative distribution function
Syntax	P = nctcdf(X,NU,DELTA)
Description	P = nctcdf(X,NU,DELTA) computes the noncentral T cdf at each of the values in X using the corresponding degrees of freedom in NU and noncentrality parameters in DELTA. Vector or matrix inputs for X, NU, and DELTA must have the same size, which is also the size of P. A scalar input for X, NU, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
Example	Compare the noncentral T cdf with DELTA = 1 to the T cdf with the same number of degrees of freedom (10) .
	<pre>x = (-5:0.1:5)'; p1 = nctcdf(x,10,1); p = tcdf(x,10); plot(x,p,'',x,p1,'-')</pre>
	1
	0.4
	0.2

References [1] Evans, M., N. Hastings, and B. Peacock, *Statistical Distributions, Second Edition*, John Wiley and Sons, 1993. p. 147–148.

0

[2] Johnson, N., and S. Kotz, *Distributions in Statistics: Continuous Univariate Distributions-2*, John Wiley and Sons, 1970. pp. 201–219.

5

See Also cdf, nctcdf, nctinv, nctpdf, nctrnd, nctstat

0l -5

nctinv

Purpose	Inverse of the noncentral T cumulative distribution
Syntax	X = nctinv(P,NU,DELTA)
Description	X = nctinv(P,NU,DELTA) returns the inverse of the noncentral T cdf with NU degrees of freedom and noncentrality parameter DELTA for the corresponding probabilities in P. Vector or matrix inputs for P, NU, and DELTA must have the same size, which is also the size of X. A scalar input for P, NU, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
Example	<pre>x = nctinv([0.1 0.2],10,1) x = -0.2914 0.1618</pre>
References	 Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition, John Wiley and Sons, 1993. p. 147–148.</i> Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate</i>
	<i>Distributions-2,</i> John Wiley and Sons, 1970. pp. 201–219.
See Also	icdf, nctcdf, nctpdf, nctrnd, nctstat

Purpose	Noncentral T probability density function (pdf)
Syntax	Y = nctpdf(X,V,DELTA)
Description	Y = nctpdf(X,V,DELTA) computes the noncentral T pdf at each of the values in X using the corresponding degrees of freedom in V and noncentrality parameters in DELTA. Vector or matrix inputs for X, V, and DELTA must have the same size, which is also the size of Y. A scalar input for X, V, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
Example	Compare the noncentral T pdf with DELTA = 1 to the T pdf with the same number of degrees of freedom (10). x = (-5:0.1:5)'; $p1 = nctpdf(x,10,1);$ $p = tpdf(x,10);$ $plot(x,p,'',x,p1,'-')$ 0.4 0.4 0.7 0.4 0.7 0.7 0.7 0.7 0.7 0.7 0.7 0.7 0.7 0.7
References	[1] Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition, John Wiley and Sons, 1993.</i> p. 147–148.

[2] Johnson, N., and S. Kotz, *Distributions in Statistics: Continuous Univariate Distributions-2*, John Wiley and Sons, 1970. pp. 201–219.

See Also nctcdf, nctinv, nctrnd, nctstat, pdf

nctrnd

Purpose	Random matrices from noncentral T distribution
Syntax	R = nctrnd(V,DELTA) R = nctrnd(V,DELTA,m) R = nctrnd(V,DELTA,m,n)
Description	R = nctrnd(V, DELTA) returns a matrix of random numbers chosen from the noncentral T distribution with parameters V and DELTA. Vector or matrix inputs for V and DELTA must have the same size, which is also the size of R. A scalar input for V or DELTA is expanded to a constant matrix with the same dimensions as the other input.
	R = nctrnd(V,DELTA,m) returns a matrix of random numbers with parameters V and DELTA, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = nctrnd(V, DELTA, m, n) generates random numbers with parameters V and DELTA, where scalars m and n are the row and column dimensions of R.
Example	nctrnd(10,1,5,1)
	ans =
	1.6576 1.0617
	1.4491
	0.2930 3.6297
References	[1] Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition</i> , John Wiley and Sons, 1993. p. 147–148.
	[2] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2,</i> John Wiley and Sons, 1970. pp. 201–219.
See Also	nctcdf, nctinv, nctpdf, nctstat

Purpose Mean and variance for the noncentral t distribution

Syntax [M,V] = nctstat(NU,DELTA)

Description [M,V] = nctstat(NU, DELTA) returns the mean and variance of the noncentral t pdf with NU degrees of freedom and noncentrality parameter DELTA. Vector or matrix inputs for NU and DELTA must have the same size, which is also the size of M and V. A scalar input for NU or DELTA is expanded to a constant matrix with the same dimensions as the other input.

The mean of the noncentral t distribution with parameters ν and δ is

$$\frac{\delta(\nu/2)^{1/2}\Gamma((\nu-1)/2)}{\Gamma(\nu/2)}$$

where v > 1.

The variance is

$$\frac{\nu}{(\nu-2)}(1+\delta^2) - \frac{\nu}{2} \delta^2 \bigg[\frac{\Gamma((\nu-1)/2)}{\Gamma(\nu/2)} \bigg]^2$$

where v > 2.

Example

```
[m,v] = nctstat(10,1)
```

```
m =
1.0837
```

v = 1.3255

References[1] Evans, M., N. Hastings, and B. Peacock, Statistical Distributions, Second
Edition, John Wiley and Sons, 1993. p. 147–148.[2] Johnson, N., and S. Kotz, Distributions in Statistics: Continuous Univariate
Distributions-2, John Wiley and Sons, 1970. pp. 201–219.

See Also nctcdf, nctinv, nctpdf, nctrnd

ncx2cdf

 Purpose
 Noncentral chi-square cumulative distribution function (cdf)

Syntax P = ncx2cdf(X,V,DELTA)

Description P = ncx2cdf(X,V,DELTA) computes the noncentral chi-square cdf at each of the values in X using the corresponding degrees of freedom in V and positive noncentrality parameters in DELTA. Vector or matrix inputs for X, V, and DELTA must have the same size, which is also the size of P. A scalar input for X, V, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.

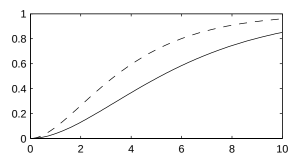
Some texts refer to this distribution as the generalized Rayleigh, Rayleigh-Rice, or Rice distribution.

The noncentral chi-square cdf is

$$F(x|\nu, \delta) = \sum_{j=0}^{\infty} \left(\frac{\left(\frac{1}{2}\delta\right)^{j}}{j!} e^{-\frac{\delta}{2}} \right) Pr[\chi_{\nu+2j}^{2} \le x]$$

Example

x = (0:0.1:10)'; p1 = ncx2cdf(x,4,2); p = chi2cdf(x,4); plot(x,p,'--',x,p1,'-')



References [1] Johnson, N., and S. Kotz, *Distributions in Statistics: Continuous Univariate Distributions-2,* John Wiley and Sons, 1970. pp. 130–148.

See Also cdf, ncx2inv, ncx2pdf, ncx2rnd, ncx2stat

ncx2inv

Purpose	Inverse of the noncentral chi-square cdf
Syntax	X = ncx2inv(P,V,DELTA)
Description	X = ncx2inv(P,V,DELTA) returns the inverse of the noncentral chi-square cdf with parameters V and DELTA at the corresponding probabilities in P. Vector or matrix inputs for P, V, and DELTA must have the same size, which is also the size of X. A scalar input for P, V, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
Algorithm	ncx2inv uses Newton's method to converge to the solution.
Example	ncx2inv([0.01 0.05 0.1],4,2)
	ans =
	0.4858 1.1498 1.7066
References	Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition,</i> John Wiley and Sons, 1993. p. 50–52.
	Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2</i> , John Wiley and Sons, 1970. pp. 130–148.
See Also	icdf, ncx2cdf, ncx2pdf, ncx2rnd, ncx2stat

Purpose	Noncentral chi-square probability density function (pdf)
Syntax	Y = ncx2pdf(X,V,DELTA)
Description	Y = ncx2pdf(X,V,DELTA) computes the noncentral chi-square pdf at each of the values in X using the corresponding degrees of freedom in V and positive noncentrality parameters in DELTA. Vector or matrix inputs for X, V, and DELTA must have the same size, which is also the size of Y. A scalar input for X, V, or DELTA is expanded to a constant matrix with the same dimensions as the other inputs.
	Some texts refer to this distribution as the generalized Rayleigh, Rayleigh-Rice, or Rice distribution.
Example	As the noncentrality parameter δ increases, the distribution flattens as shown in the plot. x = (0:0.1:10)'; p1 = ncx2pdf(x,4,2); p = chi2pdf(x,4); plot(x,p,'',x,p1,'-')
	$0 \frac{1}{0} \frac{1}{2} \frac{1}{4} \frac{1}{6} \frac{1}{8} \frac{1}{10}$
References	[1] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2</i> , John Wiley and Sons, 1970. pp. 130–148.

See Also ncx2cdf, ncx2inv, ncx2rnd, ncx2stat, pdf

ncx2rnd

Purpose	Random matrices from the noncentral chi-square distribution
Syntax	<pre>R = ncx2rnd(V,DELTA) R = ncx2rnd(V,DELTA,m) R = ncx2rnd(V,DELTA,m,n)</pre>
Description	R = ncx2rnd(V, DELTA) returns a matrix of random numbers chosen from the non-central chi-square distribution with parameters V and DELTA. Vector or matrix inputs for V and DELTA must have the same size, which is also the size of R. A scalar input for V or DELTA is expanded to a constant matrix with the same dimensions as the other input.
	R = ncx2rnd(V,DELTA,m) returns a matrix of random numbers with parameters V and DELTA, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = ncx2rnd(V, DELTA, m, n) generates random numbers with parameters V and DELTA, where scalars m and n are the row and column dimensions of R.
Example	ncx2rnd(4,2,6,3)
	ans =
	6.8552 5.9650 11.2961 5.2631 4.2640 5.9495
	9.1939 6.7162 3.8315
	10.3100 4.4828 7.1653
	2.1142 1.9826 4.6400
	3.8852 5.3999 0.9282
References	[1] Evans, M., N. Hastings, and B. Peacock, <i>Statistical Distributions, Second Edition,</i> John Wiley and Sons, 1993. p. 50–52.
	[2] Johnson, N., and S. Kotz, <i>Distributions in Statistics: Continuous Univariate Distributions-2,</i> John Wiley and Sons, 1970. pp. 130–148.
See Also	ncx2cdf, ncx2inv, ncx2pdf, ncx2stat

Purpose	Mean and variance for the noncentral chi-square distribution
Syntax	[M,V] = ncx2stat(NU,DELTA)
Description	[M,V] = ncx2stat(NU, DELTA) returns the mean and variance of the noncentral chi-square pdf with NU degrees of freedom and noncentrality parameter DELTA. Vector or matrix inputs for NU and DELTA must have the same size, which is also the size of M and V. A scalar input for NU or DELTA is expanded to a constant matrix with the same dimensions as the other input.
	The mean of the noncentral chi-square distribution with parameters ν and δ is $\nu+\delta$, and the variance is $2(\nu+2\delta)$.
Example	<pre>[m,v] = ncx2stat(4,2) m =</pre>
References	 Evans, M., N. Hastings, and B. Peacock, Statistical Distributions, Second Edition, John Wiley and Sons, 1993. p. 50–52. Johnson, N., and S. Kotz, Distributions in Statistics: Continuous Univariate Distributions-2, John Wiley and Sons, 1970. pp. 130–148.
See Also	ncx2cdf, ncx2inv, ncx2pdf, ncx2rnd

nlinfit

Purpose	Nonlinear least-squares data fitting by the Gauss-Newton method
Syntax	beta = nlinfit(X,y,FUN,beta0) [beta,r,J] = nlinfit(X,y,FUN,beta0)
Description	<pre>beta = nlinfit(X,y,FUN,beta0) estimates the coefficients of a nonlinear function using least squares. y is a vector of response (dependent variable) values. Typically, X is a design matrix of predictor (independent variable) values, with one row for each value in y. However, X can be any array that FUN can accept. FUN is a function of the form</pre>
	yhat = myfun(beta,X)
	where beta is a coefficient vector, and X is the design matrix. FUN returns a vector yhat of fitted y values. beta0 is a vector containing initial values for the coefficients.
	[beta,r,J] = nlinfit(X,y,FUN,beta0) returns the fitted coefficients, beta, the residuals, r, and the Jacobian, J. You can use these outputs with nlpredci to produce error estimates on predictions, and with nlparci to produce error estimates on the estimated coefficients.
	Note nlintool provides a GUI for performing nonlinear fits and computing confidence intervals.
Example	Find the coefficients that best fit the data in reaction.mat. The chemistry behind this data set deals with reaction kinetics as a function of the partial pressure of three chemical reactants: hydrogen, n-pentane, and isopentane.
	The hougen function uses the Hougen-Watson model for reaction kinetics to return the predicted values of the reaction rate.
	load reaction betafit = nlinfit(reactants,rate,@hougen,beta)
	betafit =
	1.2526 0.0628

0.0400 0.1124 1.1914

See Also

hougen, nlintool, nlparci, nlpredci

nlintool

Purpose	Fits a nonlinear equation to data and displays an interactive graph
Syntax	nlintool(x,y,FUN,betaO) nlintool(x,y,FUN,betaO,alpha) nlintool(x,y,FUN,betaO,alpha,'xname','yname')
Description	nlintool(x,y,FUN,beta0) is a prediction plot that provides a nonlinear curve fit to (x,y) data. It plots a 95% global confidence interval for predictions as two red curves. beta0 is a vector containing initial guesses for the parameters.
	nlintool(x,y,FUN,beta0,alpha) plots a 100(1-alpha)% confidence interval for predictions.
	nlintool displays a "vector" of plots, one for each column of the matrix of inputs, x. The response variable, y, is a column vector that matches the number of rows in x.
	The default value for alpha is 0.05, which produces 95% confidence intervals.
	nlintool(x,y,FUN,beta0,alpha,'xname','yname') labels the plot using the string matrix, 'xname' for the x variables and the string 'yname' for the y variable.
	You can drag the dotted white reference line and watch the predicted values update simultaneously. Alternatively, you can get a specific prediction by typing the value for x into an editable text field. Use the pop-up menu labeled Export to move specified variables to the base workspace. You can change the type of confidence bounds using the Bounds menu.
Example	See "Nonlinear Regression Models" on page 5-1.
See Also	nlinfit, rstool

Purpose	Confidence intervals on estimates of parameters in nonlinear models
Syntax	ci = nlparci(beta,r,J)
Description	<pre>nlparci(beta,r,J) returns the 95% confidence interval ci on the nonlinear least squares parameter estimates beta, given the residuals r and the Jacobian matrix J at the solution. The confidence interval calculation is valid for systems where the number of rows of J exceeds the length of beta. nlparci uses the outputs of nlinfit for its inputs.</pre>
Example	<pre>Continuing the example from nlinfit: load reaction [beta,resids,J] = nlinfit(reactants,rate,'hougen',beta); ci = nlparci(beta,resids,J)</pre>
	ci =
	-1.07983.3445-0.05240.1689-0.04370.1145-0.08910.2941-1.17193.7321
See Also	nlinfit, nlintool, nlpredci

nlpredci

Purpose	Confidence intervals on predictions of nonlinear models
Syntax	ypred = nlpredci(FUN,inputs,beta,r,J) [ypred,delta] = nlpredci(FUN,inputs,beta,r,J) ypred = nlpredci(FUN,inputs,beta,r,J,alpha,' <i>simopt</i> ',' <i>predopt</i> ')
Description	<pre>ypred = nlpredci(FUN, inputs, beta, r, J) returns the predicted responses, ypred, given the fitted parameters beta, residuals r, and the Jacobian matrix J. inputs is a matrix of values of the independent variables in the nonlinear function.</pre>
	[ypred,delta] = nlpredci(FUN,inputs,beta,r,J) also returns the half-width, delta, of confidence intervals for the nonlinear least squares predictions. The confidence interval calculation is valid for systems where the length of r exceeds the length of beta and J is of full column rank. The interval [ypred-delta,ypred+delta] is a 95% non-simultaneous confidence interval for the true value of the function at the specified input values.
	<pre>ypred = nlpredci(FUN, inputs, beta, r, J, alpha, 'simopt', 'predopt') controls the type of confidence intervals. The confidence level is 100(1-alpha)%. 'simopt' can be 'on' for simultaneous intervals or 'off' (the default) for non-simultaneous intervals. 'predopt' can be 'curve' (the default) for confidence intervals for the function value at the inputs, or 'observation' for confidence intervals for a new response value.</pre>
	nlpredci uses the outputs of nlinfit for its inputs.
Example	Continuing the example from nlinfit, we can determine the predicted function value at [100 300 80] and the half-width of a confidence interval for it.
	load reaction [beta,resids,J] = nlinfit(reactants,rate,@hougen,beta); [ypred,delta] = nlpredci(@hougen,[100 300 80],beta,resids,J)
	ypred = 13
	delta = 1.4277

See Also nlinfit, nlintool, nlparci

normcdf

Purpose	Normal cumulative distribution function (cdf)
Syntax	<pre>P = normcdf(X,MU,SIGMA)</pre>
Description	normcdf(X,MU,SIGMA) computes the normal cdf at each of the values in X using the corresponding parameters in MU and SIGMA. Vector or matrix inputs for X, MU, and SIGMA must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in SIGMA must be positive.
	The normal cdf is
	$p = F(x \mu, \sigma) = \frac{1}{\sigma\sqrt{2\pi}} \int_{-\infty}^{x} e^{\frac{-(t-\mu)^2}{2\sigma^2}} dt$
	The result, p , is the probability that a single observation from a normal distribution with parameters μ and σ will fall in the interval (- ∞x].
	The <i>standard normal</i> distribution has $\mu = 0$ and $\sigma = 1$.
Examples	What is the probability that an observation from a standard normal distribution will fall on the interval [-1 1]?
	p = normcdf([-1 1]); p(2) - p(1)
	ans =
	0.6827
	More generally, about 68% of the observations from a normal distribution fall within one standard deviation, σ , of the mean, $\mu.$
See Also	cdf, normfit, norminv, normpdf, normplot, normrnd, normspec, normstat

Purpose	Parameter estimates and confidence intervals for normal data
Syntax	[muhat,sigmahat,muci,sigmaci] = normfit(X) [muhat,sigmahat,muci,sigmaci] = normfit(X,alpha)
Description	[muhat,sigmahat,muci,sigmaci] = normfit(X) returns estimates muhat and sigmahat of the normal distribution parameters μ and σ , given the matrix of data X. muci and sigmaci are 95% confidence intervals and have two rows and as many columns as matrix X. The top row is the lower bound of the confidence interval and the bottom row is the upper bound.
	[muhat,sigmahat,muci,sigmaci] = normfit(X,alpha) gives estimates and 100(1-alpha)% confidence intervals. For example, alpha = 0.01 gives 99% confidence intervals.
Example	In this example the data is a two-column random normal matrix. Both columns have $\mu = 10$ and $\sigma = 2$. Note that the confidence intervals below contain the "true values."
	r = normrnd(10,2,100,2); [mu,sigma,muci,sigmaci] = normfit(r)
	mu = 10.1455 10.0527
	sigma = 1.9072 2.1256
	muci = 9.7652 9.6288 10.5258 10.4766
	sigmaci = 1.6745 1.8663 2.2155 2.4693
See Also	normcdf, norminv, normlike, normpdf, normplot, normrnd, normspec, normstat, betafit, binofit, expfit, gamfit, poissfit, unifit, weibfit

norminv

Purpose Inverse of the normal cumulative distribution function (cdf)

Syntax X = norminv(P,MU,SIGMA)

DescriptionX = norminv(P,MU,SIGMA) computes the inverse of the normal cdf with
parameters MU and SIGMA at the corresponding probabilities in P. Vector or
matrix inputs for P, MU, and SIGMA must all have the same size. A scalar input
is expanded to a constant matrix with the same dimensions as the other inputs.
The parameters in SIGMA must be positive, and the values in P must lie on the
interval [0 1].

We define the normal inverse function in terms of the normal cdf as

$$x = F^{-1}(p|\mu, \sigma) = \{x: F(x|\mu, \sigma) = p\}$$

where

$$p = F(x|\mu, \sigma) = \frac{1}{\sigma\sqrt{2\pi}} \int_{-\infty}^{x} e^{\frac{-(t-\mu)^2}{2\sigma^2}} dt$$

The result, x, is the solution of the integral equation above where you supply the desired probability, p.

Examples Find an interval that contains 95% of the values from a standard normal distribution.

Note the interval x is not the only such interval, but it is the shortest.

```
xl = norminv([0.01 0.96],0,1)
xl =
    -2.3263 1.7507
```

The interval x1 also contains 95% of the probability, but it is longer than x.

See Also icdf, normfit, normfit, normpdf, normplot, normrnd, normspec, normstat

Purpose	Negative normal log-likelihood function
Syntax	logL = normlike(params,data) [logL,avar] = normlike(params,data)
Description	<pre>logL = normlike(params,data) returns the negative of the normal log-likelihood function for the parameters params(1) = mu and params(2) = sigma, given the vector data. The length of the vector logL is the length of data.</pre>
	<pre>[logL,avar] = normlike(params,data) also returns the inverse of Fisher's information matrix, avar. If the input parameter values in params are the maximum likelihood estimates, the diagonal elements of avar are their asymptotic variances. avar is based on the observed Fisher's information, not the expected information.</pre>
	normlike is a utility function for maximum likelihood estimation.
See Also	betalike, gamlike, mle, normfit, weiblike

normpdf

PurposeNormal probability density function (pdf)

Syntax Y = normpdf(X,MU,SIGMA)

Description normpdf(X,MU,SIGMA) computes the normal pdf at each of the values in X using the corresponding parameters in MU and SIGMA. Vector or matrix inputs for X, MU, and SIGMA must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in SIGMA must be positive.

The normal pdf is

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

The *likelihood function* is the pdf viewed as a function of the parameters. Maximum likelihood estimators (MLEs) are the values of the parameters that maximize the likelihood function for a fixed value of x.

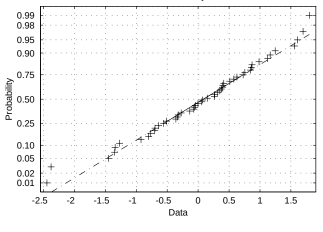
The *standard normal* distribution has $\mu = 0$ and $\sigma = 1$.

If *x* is standard normal, then $x\sigma + \mu$ is also normal with mean μ and standard deviation σ . Conversely, if *y* is normal with mean μ and standard deviation σ , then $x = (y-\mu) / \sigma$ is standard normal.

Examples	<pre>mu = [0:0.1:2]; [y i] = max(normpdf(1.5,mu,1)); MLE = mu(i)</pre>
	MLE =
	1.5000
See Also	<pre>mvnpdf, normfit, norminv, normplot, normrnd, normspec, normstat, pdf</pre>

normplot

Purpose	Normal probability plot for graphical normality testing
Syntax	<pre>normplot(X) h = normplot(X)</pre>
Description	normplot(X) displays a normal probability plot of the data in X. For matrix X, normplot displays a line for each column of X.
	The plot has the sample data displayed with the plot symbol '+'. Superimposed on the plot is a line joining the first and third quartiles of each column of X (a robust linear fit of the sample order statistics.) This line is extrapolated out to the ends of the sample to help evaluate the linearity of the data.
	If the data does come from a normal distribution, the plot will appear linear. Other probability density functions will introduce curvature in the plot.
	h = normplot(X) returns a handle to the plotted lines.
Examples	<pre>Generate a normal sample and a normal probability plot of the data. x = normrnd(0,1,50,1); h = normplot(x);</pre>
	Normal Probability Plot
	0.99 0.98 0.95



normplot

	The plot is linear, indicating that you can model the sample by a normal distribution.
See Also	cdfplot, hist, normfit, normfit, norminv, normpdf, normrnd, normspec, normstat

normrnd

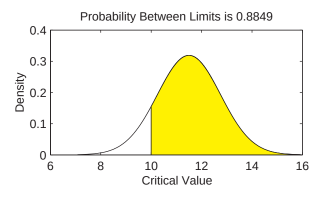
Purpose	Random numbers from the normal distribution
Syntax	R = normrnd(MU,SIGMA) R = normrnd(MU,SIGMA,m) R = normrnd(MU,SIGMA,m,n)
Description	R = normrnd(MU, SIGMA) generates normal random numbers with mean MU and standard deviation SIGMA. Vector or matrix inputs for MU and SIGMA must have the same size, which is also the size of R. A scalar input for MU or SIGMA is expanded to a constant matrix with the same dimensions as the other input.
	R = normrnd(MU, SIGMA, m) generates normal random numbers with parameters MU and SIGMA, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = normrnd(MU,SIGMA,m,n) generates normal random numbers with parameters MU and SIGMA, where scalars m and n are the row and column dimensions of R.
Examples	n1 = normrnd(1:6,1./(1:6))
	n1 = 2.1650 2.3134 3.0250 4.0879 4.8607 6.2827
	n2 = normrnd(0,1,[1 5])
	n2 = 0.0591 1.7971 0.2641 0.8717 -1.4462
	n3 = normrnd([1 2 3;4 5 6],0.1,2,3)
	n3 =
	0.9299 1.9361 2.9640 4.1246 5.0577 5.9864
See Also	normfit normfit norminy normodf normolot normspac normstat

See Also normfit, normfit, norminv, normpdf, normplot, normspec, normstat

normspec

Purpose	Plot normal density between specification limits
Syntax	p = normspec(specs,mu,sigma) [p,h] = normspec(specs,mu,sigma)
Description	<pre>p = normspec(specs,mu,sigma) plots the normal density between a lower and upper limit defined by the two elements of the vector specs, where mu and sigma are the parameters of the plotted normal distribution.</pre>
	<pre>[p,h] = normspec(specs,mu,sigma) returns the probability p of a sample falling between the lower and upper limits. h is a handle to the line objects.</pre>
	If specs(1) is -Inf, there is no lower limit, and similarly if specs(2) = Inf, there is no upper limit.
Example	Suppose a cereal manufacturer produces 10 ounce boxes of corn flakes. Variability in the process of filling each box with flakes causes a 1.25 ounce standard deviation in the true weight of the cereal in each box. The average box of cereal has 11.5 ounces of flakes. What percentage of boxes will have less than 10 ounces?

normspec([10 Inf],11.5,1.25)



See Also

capaplot, disttool, histfit, normfit, normfit, norminv, normpdf, normplot, normrnd, normstat

Purpose Mean and variance for the normal distribution

Syntax [M,V] = normstat(MU,SIGMA)

Description [M,V] = normstat(MU,SIGMA) returns the mean and variance for the normal distribution with parameters MU and SIGMA. Vector or matrix inputs for MU and SIGMA must have the same size, which is also the size of M and V. A scalar input for MU or SIGMA is expanded to a constant matrix with the same dimensions as the other input.

The mean of the normal distribution with parameters μ and σ is $\mu,$ and the variance is $\sigma^2.$

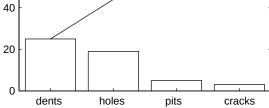
Examples	n = 1:5 [m,v] = [m,v] =	norms				
	m =					
	1	2	3	4	5	
	2	4	6	8	10	
	3	6	9	12	15	
	4	8	12	16	20	
	5	10	15	20	25	
	v =					
	1	4	9	16	25	
	4	16	36	64	100	
	9	36	81	144	225	
	16	64	144	256	400	
	25	100	225	400	625	

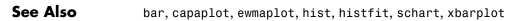
See Also

normfit, normfit, norminv, normpdf, normplot, normrnd, normspec

pareto

Purpose	Pareto charts for Statistical Process Control
Syntax	pareto(y) pareto(y,names) h = pareto()
Description	pareto(y,names) displays a Pareto chart where the values in the vector y are drawn as bars in descending order. Each bar is labeled with the associated value in the string matrix names. pareto(y) labels each bar with the index of the corresponding element in y.
	The line above the bars shows the cumulative percentage.
	pareto(y,names) labels each bar with the row of the string matrix names that corresponds to the plotted element of y.
	h = pareto() returns a combination of patch and line handles.
Example	Create a Pareto chart from data measuring the number of manufactured parts rejected for various types of defects.
	defects = ['pits';'cracks';'holes';'dents']; quantity = [5 3 19 25]; pareto(quantity,defects)
	60





Purpose	Principal Components Analysis (PCA) using the covariance matrix	
Syntax	pc = pcacov(X) [pc,latent,explained] = pcacov(X)	
Description	[pc,latent,explained] = pcacov(X) takes the covariance matrix X and returns the principal components in pc, the eigenvalues of the covariance matrix of X in latent, and the percentage of the total variance in the observations explained by each eigenvector in explained.	
Example	load hald covx = cov(ingredients); [pc,variances,explained] = pcacov(covx)	
	pc =	
	0.0678 -0.6460 0.5673 -0.5062	
	0.6785 -0.0200 -0.5440 -0.4933	
	-0.0290 0.7553 0.4036 -0.5156	
	-0.7309 -0.1085 -0.4684 -0.4844	
	variances =	
	517.7969	
	67.4964	
	12.4054	
	0.2372	
	explained =	
	86.5974	
	11.2882	
	2.0747	
	0.0397	
References	[1] Jackson, J. E., A User's Guide to Principal Components, John Wiley and Sons, Inc. 1991. pp. 1–25.	
See Also	barttest, pcares, princomp	

pcares

Purpose	Residuals from a Principal Components Analysis
Syntax	residuals = pcares(X,ndim)
Description	pcares(X,ndim) returns the residuals obtained by retaining ndim principal components of X. Note that ndim is a scalar and must be less than the number of columns in X. Use the data matrix, <i>not</i> the covariance matrix, with this function.
Example	This example shows the drop in the residuals from the first row of the Hald data as the number of component dimensions increase from one to three.
	<pre>load hald r1 = pcares(ingredients,1); r2 = pcares(ingredients,2); r3 = pcares(ingredients,3);</pre>
	r11 = r1(1,:) r11 =
	2.0350 2.8304 -6.8378 3.0879
	r21 = r2(1,:) r21 =
	-2.4037 2.6930 -1.6482 2.3425
	r31 = r3(1,:) r31 =
	0.2008 0.1957 0.2045 0.1921
Reference	[1] Jackson, J. E., <i>A User's Guide to Principal Components</i> , John Wiley and Sons, Inc. 1991. pp. 1–25.
See Also	barttest, pcacov, princomp

Purpose	Probability density function (pdf) for a specified distribution		
Syntax	Y = pdf('name',X,A1,A2,A3)		
Description	pdf(' <i>name</i> ',X,A1,A2,A3) returns a matrix of densities, where ' <i>name</i> ' is a string containing the name of the distribution. X is a matrix of values, and A1, A2, and A3 are matrices of distribution parameters. Depending on the distribution, some of the parameters may not be necessary.		
	Vector or matrix inputs for X, A1, A2, and A3 must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.		
	pdf is a utility routine allowing access to all the pdfs in the Statistics Toolbox using the name of the distribution as a parameter. See "Overview of the Distributions" on page 2-11 for the list of available distributions.		
Examples	p = pdf('Normal',-2:2,0,1)		
	p = 0.0540 0.2420 0.3989 0.2420 0.0540		
	p = pdf('Poisson',0:4,1:5)		
	p = 0.3679 0.2707 0.2240 0.1954 0.1755		
See Also	betapdf, binopdf, cdf, chi2pdf, exppdf, fpdf, gampdf, geopdf, hygepdf, lognpdf, nbinpdf, ncfpdf, nctpdf, ncx2pdf, normpdf, poisspdf, raylpdf, tpdf, unidpdf, unifpdf, weibpdf		

pdist

Purpose	Pairwise distance	between observations		
Syntax	Y = pdist(X) Y = pdist(X,'mer Y = pdist(X,dist Y = pdist(X,'min	tfun,p1,p2,)		
Description	<i>m</i> -by- <i>n</i> matrix X, v	putes the Euclidean distance between pairs of objects in which is treated as m vectors of size n . For a dataset made up are $(m-1) \cdot m/2$ pairs.		
	The output, Y, is a vector of length $(m-1) \cdot m/2$, containing the distance information. The distances are arranged in the order $(1,2)$, $(1,3)$,, $(1,m)$, $(2,3)$,, $(2,m)$,, $(m-1,m)$. Y is also commonly known as a similarity mat or dissimilarity matrix.			
	To save space and computation time, Y is formatted as a vector. However, you can convert this vector into a square matrix using the squareform function so that element i, j in the matrix, where $i < j$, corresponds to the distance between objects i and j in the original dataset.			
	matrix, X, using th	tric') computes the distance between objects in the data he method specified by 'metric', where 'metric' can be any haracter strings that identify ways to compute the distance.		
	'euclidean'	Euclidean distance (default)		
	'seuclidean'	Standardized Euclidean distance. Each coordinate in the sum of squares is inverse weighted by the sample variance of that coordinate.		
	'mahalanobis'	Mahalanobis distance		
	'cityblock'	City Block metric		
	'minkowski'	Minkowski metric		
	'cosine'	One minus the cosine of the included angle between points (treated as vectors)		
	'correlation'	One minus the sample correlation between points (treated as sequences of values).		

'hamming'	Hamming distance, the percentage of coordinates that differ
'jaccard'	One minus the Jaccard coefficient, the percentage of nonzero coordinates that differ

Y = pdist(X,distfun,p1,p2,...) accepts a function handle to a distance function of the form

d = distfun(XI,XJ,p1,p2,...)

taking as arguments two q-by-n matrices XI and XJ each of which contains rows of X, plus zero or more additional arguments, and returning a q-by-1 vector of distances d, whose kth element is the distance between the observations XI(k,:) and XJ(k,:). The arguments p1,p2,... are passed directly to the function distfun.

Y = pdist(X, 'minkowski', p) computes the distance between objects in the data matrix, X, using the Minkowski metric. p is the exponent used in the Minkowski computation which, by default, is 2.

Mathematical Definitions of Methods

Given an *m*-by-*n* data matrix X, which is treated as m (1-by-*n*) row vectors x_1 , x_2 , ..., x_m , the various distances between the vector x_r and x_s are defined as follows:

• Euclidean distance

$$d_{rs}^2 = (x_r - x_s)(x_r - x_s)'$$

• Standardized Euclidean distance

$$d_{rs}^2 = (x_r - x_s)D^{-1}(x_r - x_s)'$$

where *D* is the diagonal matrix with diagonal elements given by v_j^2 , which denotes the variance of the variable X_j over the *m* objects.

• Mahalanobis distance

$$d_{rs}^2 = (x_r - x_s) V^{-1} (x_r - x_s)'$$

where V is the sample covariance matrix.

• City Block metric

$$d_{rs} = \sum_{j=1}^{n} |x_{rj} - x_{sj}|$$

• Minkowski metric

$$d_{rs} = \left\{ \sum_{j=1}^{n} \left| x_{rj} - x_{sj} \right|^{p} \right\}^{\frac{1}{p}}$$

Notice that for the special case of p = 1, the Minkowski metric gives the City Block metric, and for the special case of p = 2, the Minkowski metric gives the Euclidean distance.

• Cosine distance

$$d_{rs} = \left(1 - x_r x'_s / (x'_r x_r)^{\frac{1}{2}} (x'_s x_s)^{\frac{1}{2}}\right)$$

• Correlation distance

$$d_{rs} = 1 - \frac{(x_r - \bar{x}_r)(x_s - \bar{x}_s)'}{[(x_r - \bar{x}_r)(x_r - \bar{x}_r)']^{\frac{1}{2}} [(x_s - \bar{x}_s)(x_s - \bar{x}_s)']^{\frac{1}{2}}}$$

where

$$\bar{x}_r = \frac{1}{n} \sum_j x_{rj}$$
 and $\bar{x}_s = \frac{1}{n} \sum_j x_{sj}$

• Hamming distance

$$d_{rs} = (\#(x_{rj} \neq x_{sj})/n)$$

• Jaccard distance

	$d_{rs} = \frac{\#[(x_{rj} \neq \frac{1}{4})]}{\#}$	$= x_{sj}) \land ((x_{rj}) \land ((x_{rj}) \land ((x_{rj} \neq 0) \land (x_{rj} \land 0) \land (x_{rj} \land 0)$	$\neq 0) \lor (x_{sj} \neq 0)$	0))]		
Examples	X = [1 2; 1 3	3;22;3	1]			
	X = 1 2 1 3 2 2 3 1					
	Y = pdist(X,' Y = 2.3452	'mahal') 2.0000	2.3452	1.2247	2.4495	1.2247
	Y = pdist(X) Y =					
	1.0000	1.0000	2.2361	1.4142	2.8284	1.4142
	squareform(Y))				
	ans = O	1.0000	1.0000	2.2361		
	1.0000		1.4142			
	1.0000 2.2361	1.4142 2.8284	0 1.4142	1.4142 0		

See Also cluster, clusterdata, cmdscale, cophenet, dendrogram, inconsistent, linkage, silhouette, squareform

perms

Purpose	All permutat	ions	
Syntax	P = perms(v)	
Description		possible	v is a row vector of length n, creates a matrix whose rows permutations of the n elements of v. The matrix P n columns.
	perms is only	practica	al when n is less than 8 or 9.
Example	perms([2	4 6])	
	ans =		
	6	4	2
	4	6	2
	6	2	4
	2	6	4
	4	2	6
	2	4	6

Purpose Poisson cumulative distribution function (cdf)

Syntax P = poisscdf(X,LAMBDA)

Description poisscdf (X, LAMBDA) computes the Poisson cdf at each of the values in X using the corresponding parameters in LAMBDA. Vector or matrix inputs for X and LAMBDA must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in LAMBDA must be positive.

The Poisson cdf is

$$p = F(x|\lambda) = e^{-\lambda} \sum_{i=0}^{floor(x)} \frac{\lambda^i}{i!}$$

Examples For example, consider a Quality Assurance department that performs random tests of individual hard disks. Their policy is to shut down the manufacturing process if an inspector finds more than four bad sectors on a disk. What is the probability of shutting down the process if the mean number of bad sectors (λ) is two?

```
probability = 1 - poisscdf(4,2)
probability =
0.0527
```

About 5% of the time, a normally functioning manufacturing process will produce more than four flaws on a hard disk.

Suppose the average number of flaws (λ) increases to four. What is the probability of finding fewer than five flaws on a hard drive?

```
probability = poisscdf(4,4)
probability =
    0.6288
```

This means that this faulty manufacturing process continues to operate after this first inspection almost 63% of the time.

poisscdf

See Also cdf, poissfit, poissinv, poisspdf, poissrnd, poisstat

Purpose	Parameter estimates and confidence intervals for Poisson data
Syntax	lambdahat = poissfit(X) [lambdahat,lambdaci] = poissfit(X) [lambdahat,lambdaci] = poissfit(X,alpha)
Description	poissfit(X) returns the maximum likelihood estimate (MLE) of the parameter of the Poisson distribution, λ , given the data X.
	[lambdahat,lambdaci] = poissfit(X) also gives 95% confidence intervals in lamdaci.
	<pre>[lambdahat,lambdaci] = poissfit(X,alpha) gives 100(1-alpha)% confidence intervals. For example alpha = 0.001 yields 99.9% confidence intervals.</pre>
	The sample average is the MLE of λ .
	$\hat{\lambda} = \frac{1}{n} \sum_{i=1}^{n} x_i$
Example	<pre>r = poissrnd(5,10,2); [1,lci] = poissfit(r)</pre>
	1 = 7.4000 6.3000
	lci = 5.8000 4.8000 9.1000 7.9000
See Also	betafit, binofit, expfit, gamfit, poisscdf, poissinv, poisspdf, poissrnd, poisstat, unifit, weibfit

poissinv

Purpose	Inverse of the Poisson cumulative distribution function (cdf)	
Syntax	X = poissinv(P,LAMBDA)	
Description	poissinv(P,LAMBDA) returns the smallest value X such that the Poisson cdf evaluated at X equals or exceeds P.	
Examples	If the average number of defects (λ) is two, what is the 95th percentile of the number of defects?	
	poissinv(0.95,2)	
	ans =	
	5	
	What is the median number of defects?	
	<pre>median_defects = poissinv(0.50,2)</pre>	
	median_defects =	
	2	
See Also	icdf, poisscdf, poissfit, poisspdf, poissrnd, poisstat	

Purpose	Poisson probabilit	y density function (pdf)
---------	--------------------	--------------------------

Syntax Y = poisspdf(X,LAMBDA)

Description poisspdf (X, LAMBDA) computes the Poisson pdf at each of the values in X using the corresponding parameters in LAMBDA. Vector or matrix inputs for X and LAMBDA must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in LAMBDA must all be positive.

The Poisson pdf is

$$y = f(x|\lambda) = \frac{\lambda^{x}}{x!}e^{-\lambda}I_{(0, 1, \ldots)}(x)$$

where x can be any nonnegative integer. The density function is zero unless x is an integer.

Examples A computer hard disk manufacturer has observed that flaws occur randomly in the manufacturing process at the average rate of two flaws in a 4 Gb hard disk and has found this rate to be acceptable. What is the probability that a disk will be manufactured with no defects?

In this problem, $\lambda = 2$ and x = 0. p = poisspdf(0,2)p =

- 0.1353
- See Also pdf, poisscdf, poissfit, poissinv, poissrnd, poisstat

poissrnd

Purpose	Random numbers from the Poisson distribution
Syntax	<pre>R = poissrnd(LAMBDA) R = poissrnd(LAMBDA,m) R = poissrnd(LAMBDA,m,n)</pre>
Description	R = poissrnd(LAMBDA) generates Poisson random numbers with mean LAMBDA. The size of R is the size of LAMBDA.
	R = poissrnd(LAMBDA,m) generates Poisson random numbers with mean LAMBDA, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = poissrnd(LAMBDA,m,n) generates Poisson random numbers with mean LAMBDA, where scalars m and n are the row and column dimensions of R.
Examples	Generate a random sample of 10 pseudo-observations from a Poisson distribution with $\lambda = 2$.
	lambda = 2;
	random_sample1 = poissrnd(lambda,1,10) random_sample1 =
	1 0 1 2 1 3 4 2 0 0
	random_sample2 = poissrnd(lambda,[1 10]) random_sample2 =
	1 1 1 5 0 3 2 2 3 4
	random_sample3 = poissrnd(lambda(ones(1,10))) random_sample3 =
	3 2 1 1 0 0 4 0 2 0
See Also	poisscdf, poissfit, poissinv, poisspdf, poisstat

poisstat

Purpose	Mean and variance for the Poisson distribution	
Syntax	M = poisstat(LAMBDA) [M,V] = poisstat(LAMBDA)	
Description	M = poisstat(LAMBDA) returns the mean of the Poisson distribution with parameter LAMBDA. The size of M is the size of LAMBDA.	
	[M,V] = poisstat(LAMBDA) also returns the variance V of the Poisson distribution.	
	For the Poisson distribution with parameter $\lambda,$ both the mean and variance are equal to $\lambda.$	
Examples	Find the mean and variance for the Poisson distribution with $\lambda = 2$.	
	[m,v] = poisstat([1 2; 3 4])	
	m =	
	1 2	
	3 4	
	v =	
	1 2	
	3 4	
See Also	poisscdf, poissfit, poissinv, poisspdf, poissrnd	

polyconf

Purpose	Polynomial evaluation and confidence interval estimation
Syntax	[Y,DELTA] = polyconf(p,X,S) [Y,DELTA] = polyconf(p,X,S,alpha)
Description	[Y, DELTA] = polyconf(p, X, S) uses the optional output S generated by polyfit to give 95% confidence intervals Y ± DELTA. This assumes the errors in the data input to polyfit are independent normal with constant variance.
	[Y,DELTA] = polyconf(p,X,S,alpha) gives 100(1-alpha)% confidence intervals. For example, alpha = 0.1 yields 90% intervals.
	If p is a vector whose elements are the coefficients of a polynomial in descending powers, such as those output from polyfit, then polyconf(p,X) is the value of the polynomial evaluated at X. If X is a matrix or vector, the polynomial is evaluated at each of the elements.
Examples	This example gives predictions and 90% confidence intervals for computing time for LU factorizations of square matrices with 100 to 200 columns.
	<pre>n = [100 100:20:200]; for i = n A = rand(i,i); tic B = lu(A); t(ceil((i-80)/20)) = toc; end</pre>
	[p,S] = polyfit(n(2:7),t,3); [time,delta_t] = polyconf(p,n(2:7),S,0.1)
	time =
	0.0829 0.1476 0.2277 0.3375 0.4912 0.7032
	delta_t =
	0.0064 0.0057 0.0055 0.0055 0.0057 0.0064

<pre>[p,S] = polyfit(x,y,n)</pre>
p = polyfit(x,y,n) finds the coefficients of a polynomial $p(x)$ of degree n that fits the data, $p(x(i))$ to $y(i)$, in a least-squares sense. The result p is a row vector of length n+1 containing the polynomial coefficients in descending powers.
$p(x) = p_1 x^n + p_2 x^{n-1} + \dots + p_n x + p_{n+1}$
[p,S] = polyfit(x,y,n) returns polynomial coefficients p and matrix S for use with polyval to produce error estimates on predictions. If the errors in the data, y, are independent normal with constant variance, polyval will produce error bounds which contain at least 50% of the predictions.
You may omit S if you are not going to pass it to polyval or polyconf for calculating error estimates.
The polyfit function is part of the standard MATLAB language.
<pre>[p,S] = polyfit(1:10,[1:10] + normrnd(0,1,1,10),1) p =</pre>

See Also polyval, polytool, polyconf

polytool

Purpose	Interactive plot for prediction of fitted polynomials
Syntax	polytool(x,y) polytool(x,y,n) polytool(x,y,n,alpha)
Description	polytool(x,y) fits a line to the column vectors x and y and displays an interactive plot of the result. This plot is graphic user interface for exploring the effects of changing the polynomial degree of the fit. The plot shows the fitted curve and 95% global confidence intervals on a new predicted value for the curve. Text with current predicted value of y and its uncertainty appears to the left of the y -axis.
	polytool(x,y,n) initially fits a polynomial of order n.
	polytool(x,y,n,alpha) plots 100(1-alpha)% confidence intervals on the predicted values.
	polytool fits by least-squares using the regression model
	$y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \ldots + \beta_n x_i^n + \varepsilon_i$
	$\varepsilon_i \sim N(0, \sigma^2) \forall i$
	$Cov(\varepsilon_i, \varepsilon_j) = 0 \forall i, j$
	Evaluate the function by typing a value in the <i>x</i> -axis edit box or by dragging the vertical reference line on the plot. The shape of the pointer changes from an arrow to a cross hair when you are over the vertical line to indicate that the line can be dragged. The predicted value of y will update as you drag the reference line.

The argument n controls the degree of the polynomial fit. To change the degree of the polynomial, choose from the pop-up menu at the top of the figure. To change the type of confidence intervals, use the **Bounds** menu. To change from least squares to a robust fitting method, use the **Method** menu.

When you are done, press the **Close** button.

polyval

Purpose	Polynomial evaluation	
Syntax	Y = polyval(p,X) [Y,DELTA] = polyval(p,X,S)	
Description	Y = polyval(p,X) returns the predicted value of a poly coefficients, p, at the values in X.	momial given its
	[Y,DELTA] = polyval(p,X,S) uses the optional output polyfit to generate error estimates, Y ± DELTA. If the en- to polyfit are independent normal with constant varian at least 50% of the predictions.	rrors in the data input
	If p is a vector whose elements are the coefficients of a p descending powers, then polyval(p,X) is the value of t evaluated at X. If X is a matrix or vector, the polynomial the elements.	he polynomial
	The polyval function is part of the standard MATLAB	language.
Examples	Simulate the function $y = x$, adding normal random error deviation of 0.1. Then use polyfit to estimate the polyno that predicted Y values are within DELTA of the integer x	omial coefficients. Note
	[p,S] = polyfit(1:10,(1:10) + normrnd(0,0.1,1 X = magic(3); [Y,D] = polyval(p,X,S)	,10),1);
	Y =	
	8.0696 1.0486 6.0636 3.0546 5.0606 7.0666	
	4.0576 9.0726 2.0516	
	D =	
	0.0889 0.0951 0.0861	
	0.0889 0.0861 0.0870 0.0870 0.0916 0.0916	
	0.0070 0.0010 0.0010	
See Also	polyfit, polytool, polyconf	

prctile

Purpose	Percentiles	s of a sa	mple					
Syntax	Y = prcti	le(X,p)	1					
Description	-					at is greater he interval	than p percent of the [0 100].	
	For vectors instance, i				_		e elements in X. For	
							ctor containing the pth v of Y is p(i) of X.	
Examples	x = (1	:5)'*(1	:5)					
	x =							
	1	2	3	4	5			
	2	4	6	8	10			
	3	6	9	12	15			
	4	8	12	16	20			
	5	10	15	20	25			
	y = pro	ctile(x	,[25 50	75])				
	y =							
	1.7	7500	3.5000	5.	2500	7.0000	8.7500	
	3.0	0000	6.0000	9.	0000	12.0000	15.0000	
	4.2	2500	8.5000	12.	7500	17.0000	21.2500	

Purpose	Principal Components Analysis (PCA)
Syntax	<pre>PC = princomp(X) [PC,SCORE,latent,tsquare] = princomp(X)</pre>
Description	[PC,SCORE,latent,tsquare] = princomp(X) takes a data matrix X and returns the principal components in PC, the so-called Z-scores in SCORE, the eigenvalues of the covariance matrix of X in latent, and Hotelling's T^2 statistic for each data point in tsquare.
	The Z-scores are the data formed by transforming the original data into the space of the principal components. The values of the vector, latent, are the variance of the columns of SCORE. Hotelling's T^2 is a measure of the multivariate distance of each observation from the center of the data set.
Example	Compute principal components for the ingredients data in the Hald dataset, and the variance accounted for by each component.
	load hald; [pc,score,latent,tsquare] = princomp(ingredients); pc,latent
	pc =
	0.0678 -0.6460 0.5673 -0.5062
	0.6785 -0.0200 -0.5440 -0.4933
	-0.0290 0.7553 0.4036 -0.5156
	-0.7309 -0.1085 -0.4684 -0.4844
	latent =
	517.7969
	67.4964
	12.4054
	0.2372
Reference	[1] Jackson, J. E., A User's Guide to Principal Components, John Wiley and Sons, Inc. 1991. pp. 1–25.
See Also	barttest, canoncorr, factoran, pcacov, pcares

procrustes

Purpose	Procrustes Analysis	
Syntax	d = procrustes(X,Y) [d,Z] = procrustes(X,Y) [d,Z,transform] = procrustes(X,Y)	
Description	d = procrustes(X,Y) determines a linear transformation (translation, reflection, orthogonal rotation, and scaling) of the points in matrix Y to best conform them to the points in matrix X. The "goodness-of-fit" criterion is the sum of squared errors. procrustes returns the minimized value of this dissimilarity measure in d. d is standardized by a measure of the scale of X, given by	
	<pre>sum(sum((X-repmat(mean(X,1),size(X,1),1)).^2,1))</pre>	
	i.e., the sum of squared elements of a centered version of X. However, if X comprises repetitions of the same point, the sum of squared errors is not standardized.	
	X and Y must have the same number of points (rows), and procrustes matches the ith point in Y to the ith point in X. Points in Y can have smaller dimension (number of columns) than those in X. In this case, procrustes adds columns of zeros to Y as necessary.	
	[d,Z] = procrustes(X,Y) also returns the transformed Y values.	
	[d,Z,transform] = procrustes(X,Y) also returns the transformation that maps Y to Z. transform is a structure with fields:	
	c Translation component	
	T Orthogonal rotation and reflection component	
	b Scale component	
	That is, Z = transform.b * Y * transform.T + transform.c.	
Examples	This example creates some random points in two dimensions, then rotates, scales, translates, and adds some noise to those points. It then uses procrustes to conform Y to X, and plots the original X and Y, and the transformed Y.	

```
X = normrnd(0,1,[10 2]);
S = [0.5 -sqrt(3)/2; sqrt(3)/2 0.5];
Y = normrnd(0.5*X*S + 2,0.05,size(X));
[d,Z,tr] = procrustes(X,Y);
plot(X(:,1),X(:,2),'rx',...
Y(:,1),Y(:,2),'b.',...
Z(:,1),Z(:,2),'bx');
```

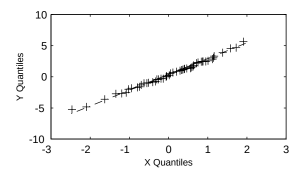
See Also cmdscale, factoran

References [1] Seber, G.A.F., *Multivariate Observations*, Wiley, 1984

[2] Bulfinch, T., *The Age of Fable; or, Stories of Gods and Heroes*, Sanborn, Carter, and Bazin, Boston, 1855.

qqplot

Purpose	Quantile-quantile plot of two samples
Syntax	<pre>qqplot(X) qqplot(X,Y) qqplot(X,Y,pvec) h = qqplot()</pre>
Description	qqplot(X) displays a quantile-quantile plot of the sample quantiles of X versus theoretical quantiles from a normal distribution. If the distribution of X is normal, the plot will be close to linear.
	qqplot(X,Y) displays a quantile-quantile plot of two samples. If the samples do come from the same distribution, the plot will be linear.
	For matrix X and Y, qqplot displays a separate line for each pair of columns. The plotted quantiles are the quantiles of the smaller dataset.
	The plot has the sample data displayed with the plot symbol '+'. Superimposed on the plot is a line joining the first and third quartiles of each distribution (this is a robust linear fit of the order statistics of the two samples). This line is extrapolated out to the ends of the sample to help evaluate the linearity of the data.
	Use $qqplot(X,Y,pvec)$ to specify the quantiles in the vector pvec.
	h = qqplot(X,Y,pvec) returns handles to the lines in h.
Examples	Generate two normal samples with different means and standard deviations. Then make a quantile-quantile plot of the two samples.
	<pre>x = normrnd(0,1,100,1); y = normrnd(0.5,2,50,1); qqplot(x,y);</pre>





normplot

random

Purpose	Random numbers from a specified distribution	
Syntax	y = random(' <i>name</i> ',A1,A2,A3,m,n)	
Description	y = random('name', A1, A2, A3, m, n) returns a matrix of random numbers, where 'name' is a string containing the name of the distribution, and A1, A2, and A3 are matrices of distribution parameters. Depending on the distribution some of the parameters may not be necessary.	
	Vector or matrix inputs must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.	
	The last two parameters, d and e, are the size of the matrix y. If the distribution parameters are matrices, then these parameters are optional, but they must match the size of the other matrix arguments (see second example).	
	random is a utility routine allowing you to access all the random number generators in the Statistics Toolbox using the name of the distribution as a parameter. See "Overview of the Distributions" on page 2-11 for the list of available distributions.	
Examples	<pre>rn = random('Normal',0,1,2,4)</pre>	
	<pre>rn = 1.1650 0.0751 -0.6965 0.0591 0.6268 0.3516 1.6961 1.7971 rp = random('Poisson',1:6,1,6) rp =</pre>	
	0 0 1 2 5 7	
See Also	betarnd, binornd, cdf, chi2rnd, exprnd, frnd, gamrnd, geornd, hygernd, icdf, lognrnd, nbinrnd, ncfrnd, nctrnd, ncx2rnd, normrnd, pdf, poissrnd, raylrnd,	

trnd, unidrnd, unifrnd, weibrnd

Purpose	Interactive random number generation using histograms for display
Syntax	randtool r = randtool('output')
Description	The randtool command sets up a graphic user interface for exploring the effects of changing parameters and sample size on the histogram of random samples from the supported probability distributions.
	The M-file calls itself recursively using the action and flag parameters. For general use call randtool without parameters.
	To output the current set of random numbers, press the Output button. The results are stored in the variable ans. Alternatively, use the following command.
	r = randtool('output') places the sample of random numbers in the vector r .
	To sample repetitively from the same distribution, press the Resample button.
	To change the distribution function, choose from the pop-up menu of functions at the top of the figure.
	To change the parameter settings, move the sliders or type a value in the edit box under the name of the parameter. To change the limits of a parameter, type a value in the edit box at the top or bottom of the parameter slider.
	To change the sample size, type a number in the Sample Size edit box.
	When you are done, press the Close button.
	For an extensive discussion, see "The randtool Demo" on page 11-18.
See Also	disttool

range

Purpose	Sample range	
Syntax	y = range(X)	
Description	<pre>range(X) returns the difference between the maximum and the minimum of a sample. For vectors, range(x) is the range of the elements. For matrices, range(X) is a row vector containing the range of each column of X.</pre>	
	The range is an easily calculated estimate of the spread of a sample. Outliers have an undue influence on this statistic, which makes it an unreliable estimator.	
Example	The range of a large sample of standard normal random numbers is approximately six. This is the motivation for the process capability indices C_{pk} in statistical quality control applications.	
	<pre>rv = normrnd(0,1,1000,5); near6 = range(rv)</pre>	
	near6 =	
	6.1451 6.4986 6.2909 5.8894 7.0002	
See Also	std, iqr, mad	

Purpose	Wilcoxon rank sum test that two populations are identical
Syntax	p = ranksum(x,y,alpha) [p,h] = ranksum(x,y,alpha) [p,h,stats] = ranksum(x,y,alpha)
Description	<pre>p = ranksum(x,y,alpha) returns the significance probability that the populations generating two independent samples, x and y, are identical. x and y are both vectors, but can have different lengths. alpha is the desired level of significance and must be a scalar between zero and one.</pre>
	[p,h] = ranksum(x,y,alpha) also returns the result of the hypothesis test, h. h is zero if the populations of x and y are not significantly different. h is one if the two populations are significantly different.
	p is the probability of observing a result equally or more extreme than the one using the data (x and y) if the null hypothesis is true. If p is near zero, this casts doubt on this hypothesis.
	<pre>[p,h,stats] = ranksum(x,y,alpha) also returns a structure containing the field stats.ranksum whose value is equal to the rank sum statistic. For large samples, it also contains stats.zval that is the value of the normal (Z) statistic used to compute p.</pre>
Example	This example tests the hypothesis of equality of means for two samples generated with poissrnd.
	<pre>x = poissrnd(5,10,1); y = poissrnd(2,20,1); [p,h] = ranksum(x,y,0.05)</pre>
	p = 0.0027
	h = 1
See Also	signrank, signtest, ttest2

raylcdf

 Purpose
 Rayleigh cumulative distribution function (cdf)

Syntax P = raylcdf(X,B)

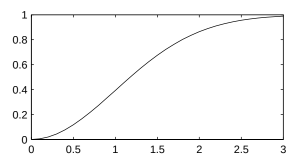
Description P = raylcdf(X,B) computes the Rayleigh cdf at each of the values in X using the corresponding parameters in B. Vector or matrix inputs for X and B must have the same size, which is also the size of P. A scalar input for X or B is expanded to a constant matrix with the same dimensions as the other input.

The Rayleigh cdf is

$$y = F(x|b) = \int_0^x \frac{t}{b^2} e^{\left(\frac{-t^2}{2b^2}\right)} dt$$

Example

x = 0:0.1:3; p = raylcdf(x,1); plot(x,p)



Reference [1] Evans, M., N. Hastings, and B. Peacock, *Statistical Distributions, Second Edition*, Wiley 1993. pp. 134–136.

See Also cdf, raylinv, raylpdf, raylrnd, raylstat

Purpose	Inverse of the Rayleigh cumulative distribution function
Syntax	X = raylinv(P,B)
Description	X = raylinv(P,B) returns the inverse of the Rayleigh cumulative distribution function with parameter B at the corresponding probabilities in P. Vector or matrix inputs for P and B must have the same size, which is also the size of X. A scalar input for P or B is expanded to a constant matrix with the same dimensions as the other input.
Example	<pre>x = raylinv(0.9,1) x = 2.1460</pre>
See Also	icdf, raylcdf, raylpdf, raylrnd, raylstat

raylpdf

Purpose Rayleigh probability density function

Syntax Y = raylpdf(X,B)

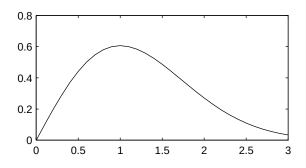
Description Y = raylpdf(X,B) computes the Rayleigh pdf at each of the values in X using the corresponding parameters in B. Vector or matrix inputs for X and B must have the same size, which is also the size of Y. A scalar input for X or B is expanded to a constant matrix with the same dimensions as the other input.

The Rayleigh pdf is

$$y = f(x|b) = \frac{x}{b^2} e^{\left(\frac{-x^2}{2b^2}\right)}$$

Example

x = 0:0.1:3; p = raylpdf(x,1); plot(x,p)



See Also

pdf, raylcdf, raylinv, raylrnd, raylstat

raylrnd

Purpose	Random matrices from the Rayleigh distribution
Syntax	R = raylrnd(B) R = raylrnd(B,m) R = raylrnd(B,m,n)
Description	 R = raylrnd(B) returns a matrix of random numbers chosen from the Rayleigh distribution with parameter B. The size of R is the size of B. R = raylrnd(B,m) returns a matrix of random numbers chosen from the Rayleigh distribution with parameter B, where m is a 1-by-2 vector that contains the row and column dimensions of R. R = raylrnd(B,m,n) returns a matrix of random numbers chosen from the Rayleigh distribution with parameter B, where scalars m and n are the row and column dimensions of R.
Example	r = raylrnd(1:5) r = 1.7986 0.8795 3.3473 8.9159 3.5182
See Also	random, raylcdf, raylinv, raylpdf, raylstat

raylstat

Purpose	Mean and variance for the Rayleigh distribution
Syntax	M = raylstat(B) [M,V] = raylstat(B)
Description	[M,V] = raylstat(B) returns the mean and variance of the Rayleigh distribution with parameter B.
	The mean of the Rayleigh distribution with parameter b is $b\sqrt{\pi/2}$ and the variance is
	$rac{4-\pi}{2}b^2$
Example	[mn,v] = raylstat(1)
	mn = 1.2533
	v = 0.4292
See Also	raylcdf, raylinv, raylpdf, raylrnd

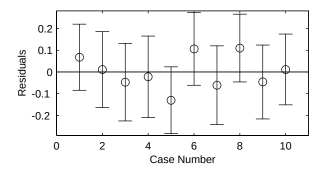
rcoplot

Purpose	Residual case order plot
---------	--------------------------

Syntax rcoplot(r,rint)

Description rcoplot(r,rint) displays an errorbar plot of the confidence intervals on the residuals from a regression. The residuals appear in the plot in case order. Inputs r and rint are outputs from the regress function.

Example X = [ones(10,1) (1:10)']; y = X * [10;1] + normrnd(0,0.1,10,1); [b,bint,r,rint] = regress(y,X,0.05); rcoplot(r,rint);

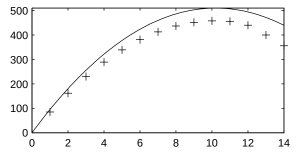


The figure shows a plot of the residuals with error bars showing 95% confidence intervals on the residuals. All the error bars pass through the zero line, indicating that there are no outliers in the data.

See Also regress

refcurve

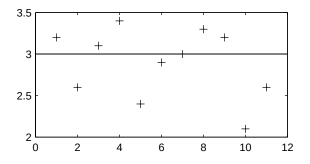
Purpose	Add a polynomial curve to the current plot		
Syntax	h = refcurve(p)		
Description	refcurve adds a graph of the polynomial p to the current axes. The function for a polynomial of degree n is: $y = p_1 x^n + p_2 x^{(n-1)} + \dots + p_n x + p_{n+1}$ Note that p_1 goes with the highest order term. h = refcurve(p) returns the handle to the curve.		
Example	<pre>Plot data for the height of a rocket against time, and add a reference curve showing the theoretical height (assuming no air friction). The initial velocity of the rocket is 100 m/sec. h = [85 162 230 289 339 381 413 437 452 458 456 440 400 356]; plot(h, '+') refcurve([-4.9 100 0])</pre>		



See Also

polyfit, polyval, refline

Purpose	Add a reference line to the current axes
Syntax	refline(slope,intercept) refline(slope) h = refline(slope,intercept) refline
Description	refline(slope,intercept) adds a reference line with the given slope and intercept to the current axes.
	refline(slope), where slope is a two-element vector, adds the line
	y = slope(2) + slope(1)*x
	to the figure.
	h = refline(slope, intercept) returns the handle to the line.
	refline with no input arguments superimposes the least squares line on each line object in the current figure (except LineStyles '-','',''). This behavior is equivalent to lsline.
Example	y = [3.2 2.6 3.1 3.4 2.4 2.9 3.0 3.3 3.2 2.1 2.6]'; plot(y,'+') refline(0,3)



See Also

lsline, polyfit, polyval, refcurve

regress

Iultiple linear regression		
= regress(y,X) b,bint,r,rint,stats] = regress(y,X) b,bint,r,rint,stats] = regress(y,X,alpha)		
<pre>b = regress(y,X) returns the least squares fit of y on X by solving the linear model</pre>		
$y = X\beta + \varepsilon$ $\varepsilon \sim N(0, \sigma^2 I)$		
or β , where:		
y is an <i>n</i> -by-1 vector of observations X is an <i>n</i> -by-p matrix of regressors β is a p-by-1 vector of parameters ϵ is an <i>n</i> -by-1 vector of random disturbances		
b,bint,r,rint,stats] = regress(y,X) returns an estimate of β in b, a 95% onfidence interval for β in the <i>p</i> -by-2 vector bint. The residuals are returned in r and a 95% confidence interval for each residual is returned in the <i>n</i> -by-2 ector rint. The vector stats contains the R^2 statistic along with the <i>F</i> and <i>p</i> alues for the regression.		
b,bint,r,rint,stats] = regress(y,X,alpha) gives 100(1-alpha)% onfidence intervals for bint and rint. For example, alpha = 0.2 gives 80% onfidence intervals.		
uppose the true model is		
$y = 10 + x + \varepsilon$ $\varepsilon \sim N(0, 0.01I)$ where I is the identity matrix. X = [ones(10,1) (1:10)'] $X = 1 \qquad 1$		

```
1
               2
        1
               3
        1
               4
        1
               5
               6
        1
               7
        1
        1
               8
        1
               9
        1
              10
  y = X * [10;1] + normrnd(0,0.1,10,1)
  y =
      11.1165
      12.0627
      13.0075
      14.0352
      14.9303
      16.1696
      17.0059
      18.1797
      19.0264
      20.0872
  [b,bint] = regress(y,X,0.05)
  b =
      10.0456
       1.0030
  bint =
                 10.1747
       9.9165
       0.9822
                  1.0238
Compare b to [10 \ 1]'. Note that bint includes the true model values.
```

Reference [1] Chatterjee, S. and A. S. Hadi. *Influential Observations, High Leverage Points, and Outliers in Linear Regression.* Statistical Science, 1986. pp. 379–416.

regstats

Purpose	Regression diagnostics for linear models			
Syntax		regstats(responses,DATA,' <i>model</i> ')' stats = regstats(responses,DATA,' <i>model</i> ',' <i>whichstats</i> ')		
Description	<pre>regstats(responses,data,'model') fits a multiple regression of the measurements in the vector, responses, on the values in the matrix, DATA. The function creates a UI that displays a group of checkboxes that save diagnostic statistics to the base workspace using specified variable names. 'model' controls the order of the regression model. By default, regstats uses a linear additive model with a constant term.</pre>			
	' <i>model</i> ' can be	one of the following strings		
	'linear'	Includes constant and linear terms (default).		
	'interaction' Includes constant, linear, and cross product ter			
	'quadratic'	Includes interactions and squared terms.		
	'purequadrati	ratic' Includes constant, linear, and squared terms.		
	The order of the coefficients is the order defined by the x2fx function. stats = regstats(responses,DATA,model,whichstats) creates an outp structure stats containing the statistics listed in ' <i>whichstats</i> '. ' <i>whichstat</i> can be a single name such as 'leverage' or a cell array of names such as {'leverage' 'standres' 'studres'}. Valid names are:			
	'Q'	Q from the QR Decomposition of X		
	'R'	R from the QR Decomposition of X		
	'beta'	Regression Coefficients		
	'covb'	Covariance of Regression Coefficients		
	'vbat'	Fitted Veluce of the Persona Date		

- 'yhat' Fitted Values of the Response Data
- 'r' Residuals
- 'mse' Mean Squared Error

'leverage' Leverage

'hatmat'	Hat (Projection) Matrix
's2_i'	Delete-1 Variance
'beta_i'	Delete-1 Coefficients
'standres'	Standardized Residuals
'studres'	Studentized Residuals
'dfbetas'	Scaled Change in Regression Coefficients
'dffit'	Change in Fitted Values
'dffits'	Scaled Change in Fitted Values
'covratio'	Change in Covariance
'cookd'	Cook's Distance
'all'	Create all of the above statistics

For more detail press the **Help** button in the regstats window. This provides formulae and interpretations for each of these regression diagnostics.

Algorithm The usual regression model is $y = X\beta + \varepsilon$, where:

- *y* is an *n*-by-1 vector of responses
- X is an *n*-by-*p* matrix of predictors
- β is an *p*-by-1 vector of parameters
- ε is an *n*-by-1 vector of random disturbances

Let X = Q R where Q and R come from a QR Decomposition of X. Q is orthogonal and R is triangular. Both of these matrices are useful for calculating many regression diagnostics (Goodall 1993).

The standard textbook equation for the least squares estimator of β is

 $\hat{\beta} = b = (XX)^{-1}Xy$

However, this definition has poor numeric properties. Particularly dubious is the computation of $(X'X)^{-1}$, which is both expensive and imprecise.

Numerically stable MATLAB code for $\boldsymbol{\beta}$ is

regstats

 $b = R \setminus (Q'*y);$

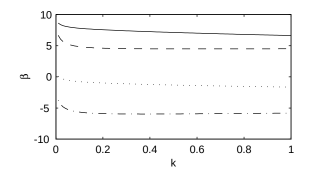
See Also leverage, stepwise, regress

Reference [1] Belsley, D.A., E. Kuh, and R.E. Welsch, *Regression Diagnostics*, New York: Wiley, 1980.

[2] Cook, R.D., and S. Weisberg, *Residuals and Influence in Regression*, New York: Wiley, 1982.

[3] Goodall, C. R., "Computation using the QR decomposition," *Statistical Computing* (C. R. Rao, ed.), Handbook in Statistics, Volume 9. Amsterdam, NL Elsevier/North-Holland, 1993.

Purpose	Parameter estimates for ridge regression		
Syntax	<pre>b = ridge(y,X,k)</pre>		
Description	b = ridge(y,X,k) returns the ridge regression coefficients b for the linear model $y = X\beta + \varepsilon$, where:		
	• X is an n -by- p matrix		
	• y is the <i>n</i> -by-1 vector of observations		
	• k is a scalar constant (the ridge parameter)		
	The ridge estimator of β is $b = (XX + kI)^{-1}Xy$.		
	When $k = 0$, b is the least squares estimator. For increasing k, the bias of b increases, but the variance of b falls. For poorly conditioned X, the drop in the variance more than compensates for the bias.		
Example	This example shows how the coefficients change as the value of k increases, using data from the hald dataset.		
	load hald; b = zeros(4,100); kvec = 0.01:0.01:1; count = 0;		
	<pre>for k = 0.01:0.01:1 count = count + 1; b(:,count) = ridge(heat,ingredients,k); end</pre>		
	plot(kvec',b'),xlabel('k'),ylabel('b','FontName','Symbol')		





regress, stepwise

robustdemo

Purpose	Demo of robust regression		
Syntax	robustdemo robustdemo(X,Y)		
Description	<pre>rsmdemo demonstrates robust regression and ordinary least squares regression on a sample dataset. The function creates a figure window containing a scatter plot of sample data vectors X and Y, along with two fitted lines calculated using least squares and the robust bisquare method. The bottom of the figure shows the equations of the lines and the estimated error standard deviations for each fit. If you use the left mouse button to select an point and move it to a new location, both fits will update. If you hold down the right mouse button over any point, the point will be labeled with the leverage of that point on the least squares fit, and the weight of that point in the robust fit. rsmdemo(X,Y) performs the same demonstration using the X and Y values that you specify.</pre>		
Example	See "The robustdemo Demo" on page 11-22.		
See Also	robustfit, leverage		

robustfit

Purpose	Robust regression		
Syntax	<pre>b = robustfit(X,Y) [b,stats] = robustfit(X,Y) [b,stats] = robustfit(X,Y,'wfun',tune,'const')</pre>		
Description	<pre>b = robustfit(X,Y) uses robust regression to fit Y as a function of the columns of X, and returns the vector b of coefficient estimates. The robustfit function uses an iteratively reweighted least squares algorithm, with the weights at each iteration calculated by applying the bisquare function to the residuals from the previous iteration. This algorithm gives lower weight to points that do not fit well. The results are less sensitive to outliers in the data as compared with ordinary least squares regression.</pre>		
	<pre>[b,stats] = robustfit(X,Y) also returns a stats structure with the following fields:</pre>		
	• stats.ols_s sigma estimate (rmse) from least squares fit		
	• stats.robust_s robust estimate of sigma		
	• stats.mad_s estimate of sigma computed using the median absolute deviation of the residuals from their median; used for scaling residuals during the iterative fitting		
	 stats.s final estimate of sigma, the larger of robust_s and a weighted average of ols_s and robust_s 		
	• stats.se standard error of coefficient estimates		
	• stats.t ratio of b to stats.se		
 stats.p p-values for stats.t 			
	• stats.coeffcorr estimated correlation of coefficient estimates		
	• stats.w vector of weights for robust fit		
	• stats.h vector of leverage values for least squares fit		
	• stats.dfe degrees of freedom for error		
	• stats.R R factor in QR decomposition of X matrix		
	The robustfit function estimates the variance-covariance matrix of the coefficient estimates as $V = inv(X'*X)*stats.s^2$. The standard errors and correlations are derived from V.		

[b,stats] = robustfit(X,Y, 'wfun',tune, 'const') specifies a weight function, a tuning constant, and the presence or absence of a constant term. The weight function 'wfun' can be any of the names listed in the following table.

Weight function	Meaning	Tuning constant
'andrews'	w = (abs(r) <pi) .="" .*="" r<="" sin(r)="" td=""><td>1.339</td></pi)>	1.339
'bisquare'	w = (abs(r)<1) .* (1 - r.^2).^2	4.685
'cauchy'	w = 1 ./ (1 + r.^2)	2.385
'fair'	w = 1 . / (1 + abs(r))	1.400
'huber'	w = 1 ./ max(1, abs(r))	1.345
'logistic'	w = tanh(r) . / r	1.205
'talwar'	w = 1 * (abs(r)<1)	2.795
'welsch'	$w = exp(-(r.^{2}))$	2.985

Table 12-2:

The value r in the weight function expression is equal to

```
resid/(tune*s*sqrt(1-h))
```

where resid is the vector of residuals from the previous iteration, tune is the tuning constant, h is the vector of leverage values from a least squares fit, and s is an estimate of the standard deviation of the error term.

```
s = MAD/0.6745
```

The quantity MAD is the median absolute deviation of the residuals from their median. The constant 0.6745 makes the estimate unbiased for the normal distribution. If there are p columns in the X matrix (including the constant term, if any), the smallest p-1 absolute deviations are excluded when computing their median.

In addition to the function names listed above, 'wfun' can be 'ols' to perform unweighted ordinary least squares.

robustfit

The argument tune overrides the default tuning constant from the table. A smaller tuning constant tends to downweight large residuals more severely, and a larger tuning constant downweights large residuals less severely. The default tuning constants, shown in the table, yield coefficient estimates that are approximately 95% as efficient as least squares estimates, when the response has a normal distribution with no outliers. The value of 'const' can be 'on' (the default) to add a constant term or 'off' to omit it. If you want a constant term, you should set 'const' to 'on' rather than adding a column of ones to your X matrix.

As an alternative to specifying one of the named weight functions shown above, you can write your own weight function that takes a vector of scaled residuals as input and produces a vector of weights as output. You can specify 'wfun' using @ (for example, @myfun) or as an inline function.

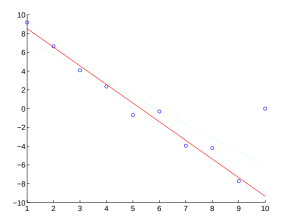
Example Let's see how a single erroneous point affects least squares and robust fits. First we generate a simple dataset following the equation y = 10-2*x plus some random noise. Then we change one y value to simulate an outlier that could be an erroneous measurement.

```
x = (1:10)';
y = 10 - 2*x + randn(10,1);
y(10) = 0;
```

We use both ordinary least squares and robust fitting to estimate the equations of a straight line fit.

A scatter plot with both fitted lines shows that the robust fit (solid line) fits most of the data points well but ignores the outlier. The least squares fit (dotted line) is pulled toward the outlier.

```
scatter(x,y)
hold on
plot(x,bls(1)+bls(2)*x,'g:')
plot(x,brob(1)+brob(2)*x,'r-')
```



See Also regress, robustdemo

References [1] DuMouchel, W.H., and F.L. O'Brien (1989), "Integrating a Robust Option into a Multiple Regression Computing Environment," *Computer Science and Statistics: Proceedings of the 21st Symposium on the Interface*, Alexandria, VA: American Statistical Association.

> [2] Holland, P.W., and R.E. Welsch (1977), "Robust Regression Using Iteratively Reweighted Least-Squares," *Communications in Statistics: Theory and Methods*, A6, 813-827.

[3] Huber, P.J. (1981), Robust Statistics, New York: Wiley.

[4] Street, J.O., R.J. Carroll, and D. Ruppert (1988), "A Note on Computing Robust Regression Estimates via Iteratively Reweighted Least Squares," *The American Statistician*, 42, 152-154

rowexch

Purpose	D-optimal design of experiments – row exchange algorithm		
Syntax	<pre>settings = rowexch(nfactors,nruns) [settings,X] = rowexch(nfactors,nruns) [settings,X] = rowexch(nfactors,nruns,'model') [settings,X] = rowexch(,'param1',value1,'param2',value2,)</pre>		
Description	settings = rowexch(nfactors,nruns) generates the factor settings matrix, settings, for a D-Optimal design using a linear additive model with a constant term. settings has nruns rows and nfactors columns.		
			wexch(nfactors,nruns) also generates the associated tings, often called the design matrix.
	<pre>[settings,X] = rowexch(nfactors,nruns,'model') produces a design for fitting a specified regression model. The input, 'model', can be one of these strings:</pre>		
	'linear'		Includes constant and linear terms (the default)
			Includes constant, linear, and cross product terms.
			Includes interactions plus squared terms.
	'purequadrat	urequadratic' Includes constant, linear and squared t	
	<pre>[settings,X] = rowexch(,'param1',value1,'param2',val provides more control over the design generation through a set of parameter/value pairs. Valid parameters are:</pre>		ol over the design generation through a set of
	'display' Either 'on' or 'off' to control display of iteration counter The default is 'on'.		
	'init'	Initial design as an nruns-by-nfactors matrix. The default is a randomly selected set of points. Maximum number of iterations. The default is 10.	
	'maxiter'		
Example	This example illustrates that the D-optimal design for three factors in eight runs, using an interactions model, is a two level full-factorial design.		

```
s = rowexch(3,8,'interaction')
s =
     - 1
            - 1
                     1
            - 1
      1
                    -1
      1
            - 1
                     1
     - 1
            - 1
                    -1
     - 1
             1
                     1
      1
              1
                     1
             1
     - 1
                    -1
      1
             1
                    - 1
```

Algorithm The rowexch function searches for a D-optimal design using a row-exchange algorithm. It first generates a candidate set of points that are eligible to be included in the design, and then iteratively exchanges design points for candidate points in an attempt to reduce the variance of the coefficients that would be estimated using this design. If you need to use a candidate set that differs from the default one, call the candgen and candexch functions in place of rowexch.

See Also bbdesign, candexch, candgen, ccdesign, cordexch, x2fx

rsmdemo

Purpose	Demo of design of experiments and surface fitting		
Syntax	rsmdemo		
Description	 rsmdemo creates a GUI that simulates a chemical reaction. To start, you have a budget of 13 test reactions. Try to find out how changes in each reactant affect the reaction rate. Determine the reactant settings that maximize the reaction rate. Estimate the run-to-run variability of the reaction. Now run a designed experiment using the model pop-up. Compare your previous results with the output from response surface modeling or nonlinear modeling of the reaction. The GUI has the following elements: A Run button to perform one reactor run at the current settings An Export button to export the <i>x</i> and <i>y</i> data to the base workspace Three sliders with associated data entry boxes to control the partial pressures of the chemical reactants: Hydrogen, n-Pentane, and Isopentane A text box to keep track of the number of test reactions you have left 		
Example	See "The rsmdemo Demo" on page 11-19.		
See Also	rstool, nlintool, cordexch		

Purpose	Interactive fitting and visualization of a response surface
Syntax	rstool(x,y) rstool(x,y,' <i>model</i> ') rstool(x,y,' <i>model</i> ',alpha,'xname','yname')
Description	rstool(x,y) displays an interactive prediction plot with 95% global confidence intervals. This plot results from a multiple regression of (x,y) data using a linear additive model.
	<pre>rstool(x,y,'model') allows control over the initial regression model, where 'model' can be one of the following strings:</pre>
	 'interaction' – includes constant, linear, and cross product terms 'quadratic' – includes interactions and squared terms 'purequadratic' – includes constant, linear and squared terms
	<pre>rstool(x,y,'model',alpha) plots 100(1-alpha)% global confidence interval for predictions as two red curves. For example, alpha = 0.01 gives 99% confidence intervals.</pre>
	rstool displays a "vector" of plots, one for each column of the matrix of inputs x. The response variable, y, is a column vector that matches the number of rows in x.
	<pre>rstool(x,y,'model',alpha,'xname','yname') labels the graph using the string matrix 'xname' for the labels to the x-axes and the string, 'yname', to label the y-axis common to all the plots.</pre>
	Drag the dotted white reference line and watch the predicted values update simultaneously. Alternatively, you can get a specific prediction by typing the value of <i>x</i> into an editable text field. Use the pop-up menu labeled Model to interactively change the model. Use the pop-up menu labeled Export to move specified variables to the base workspace.
Example	See "Quadratic Response Surface Models" on page 4-22.
See Also	nlintool

schart

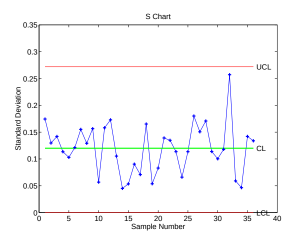
Purpose	Chart of standard deviation for Statistical Process Control
Syntax	<pre>schart(DATA,conf) schart(DATA,conf,specs) schart(DATA,conf,specs) [outliers,h] = schart(DATA,conf,specs)</pre>
Description	<pre>schart(data) displays an S chart of the grouped responses in DATA. The rows of DATA contain replicate observations taken at a given time. The rows must be in time order. The graph contains the sample standard deviation s for each group, a center line at the average s value, and upper and lower control limits. The limits are placed at a three-sigma distance on either side of the center line, where sigma is an estimate of the standard deviation of s. If the process is in control, fewer than 3 out of 1000 observations would be expected to fall outside the control limits by random chance. So, if you observe points outside the limits, you can take this as evidence that the process is not in control. schart(DATA, conf) allows control of the confidence level of the upper and lower plotted control limits. The default conf = 0.9973 produces three-sigma limits.</pre>
	k = 2; 1-2*(1-normcdf(k)) ans = 0.9545
	schart(DATA,conf,specs) plots the specification limits in the two element vector specs.
	[outliers,h] = schart(data,conf,specs) returns outliers, a vector of

[outliers,h] = schart(data,conf,specs) returns outliers, a vector of indices to the rows where the mean of DATA is out of control, and h, a vector of handles to the plotted lines.

Example

This example plots an S chart of measurements on newly machined parts, taken at one hour intervals for 36 hours. Each row of the runout matrix contains the measurements for 4 parts chosen at random. The values indicate, in thousandths of an inch, the amount the part radius differs from the target radius.

load parts
schart(runout)



All points are within the control limits, so the variability within subgroups is consistent with what would be expected by random chance. There is no evidence that the process is out of control.

Reference [1] Montgomery, D., *Introduction to Statistical Quality Control*, John Wiley and Sons 1991. p. 235.

See Also capaplot, ewmaplot, histfit, xbarplot

signrank

Purpose	Wilcoxon signed rank test of equality of medians
Syntax	p = signrank(x,y,alpha) [p,h] = signrank(x,y,alpha) [p,h,stats] = signrank(x,y,alpha)
Description	<pre>p = signrank(x,y,alpha) returns the significance probability that the medians of two matched samples, x and y, are equal. x and y must be vectors of equal length. y may also be a scalar; in this case, signrank computes the probability that the median of x is different from the constant y. alpha is the desired level of significance, and must be a scalar between zero and one.</pre>
	<pre>[p,h] = signrank(x,y,alpha) also returns the result of the hypothesis test, h. h is zero if the difference in medians of x and y is not significantly different from zero. h is one if the two medians are significantly different.</pre>
	p is the probability of observing a result equally or more extreme than the one using the data (x and y) if the null hypothesis is true. p is calculated using the rank values for the differences between corresponding elements in x and y. If p is near zero, this casts doubt on this hypothesis.
	<pre>[p,h,stats] = signrank(x,y,alpha) also returns a structure stats containing the field stats.signedrank whose value is the signed rank statistic. For large samples, it also contains stats.zval, the value of the normal (Z) statistic used to compute p.</pre>
Example	This example tests the hypothesis of equality of means for two samples generated with normrnd. The samples have the same theoretical mean but different standard deviations.
	<pre>x = normrnd(0,1,20,1); y = normrnd(0,2,20,1); [p,h] = signrank(x,y,0.05)</pre>
	p = 0.2959
	h = 0

See Also ranksum, signtest, ttest

signtest

Purpose	Sign test for paired samples
Syntax	p = signtest(x,y,alpha) [p,h] = signtest(x,y,alpha) [p,h,stats] = signtest(x,y,alpha)
Description	<pre>p = signtest(x,y,alpha) returns the significance probability that the medians of two matched samples, x and y, are equal. x and y must be vectors of equal length. y may also be a scalar; in this case, signtest computes the probability that the median of x is different from the constant y. alpha is the desired level of significance and must be a scalar between zero and one.</pre>
	<pre>[p,h] = signtest(x,y,alpha) also returns the result of the hypothesis test, h. h is 0 if the difference in medians of x and y is not significantly different from zero. h is 1 if the two medians are significantly different.</pre>
	p is the probability of observing a result equally or more extreme than the one using the data (x and y) if the null hypothesis is true. p is calculated using the signs (plus or minus) of the differences between corresponding elements in x and y. If p is near zero, this casts doubt on this hypothesis.
	<pre>[p,h,stats] = signtest(x,y,alpha) also returns a structure stats containing the field stats.sign whose value is the sign statistic. For large samples, it also contains stats.zval, the value of the normal (Z) statistic used to compute p.</pre>
Example	This example tests the hypothesis of equality of medians for two samples generated with normrnd. The samples have the same theoretical median but different standard deviations. (For the normal distribution, the mean and median are the same.)
	<pre>x = normrnd(0,1,20,1); y = normrnd(0,2,20,1); [p,h] = signtest(x,y,0.05)</pre>
	p = 0.2632
	h = 0

See Also ranksum, signrank, ttest

silhouette

Purpose	Silhouette plot for clustered data		
Syntax	<pre>silhouette(X,clust) s = silhouette(X,clust) [s,h] = silhouette(X,clust) [] = silhouette(X,clust,distance) [] = silhouette(X,clust,distfun,p1,p2,)</pre>		
Description	<pre>silhouette(X,clust) plots cluster silhouettes for the n-by-p data matrix X, with clusters defined by clust. Rows of X correspond to points, columns correspond to coordinates. clust can be a numeric vector containing a cluster index for each point, or a character matrix or cell array of strings containing a cluster name for each point. silhouette treats NaNs or empty strings in clust as missing values, and ignores the corresponding rows of X. By default, silhouette uses the squared Euclidean distance between points in X.</pre>		
	<pre>s = silhouette(X,clust) returns the silhouette values in the n-by-1 vector s, but does not plot the cluster silhouettes.</pre>		
	<pre>[s,h] = silhouette(X,clust) plots the silhouettes, and returns the silhouette values in the n-by-1 vector s, and the figure handle in h.</pre>		
	[] = silhouette(X,clust,distance) plots the silhouettes using the inter-point distance measure specified in distance. Choices for distance are:		
	'Euclidean' Euclidean distance		
	'sqEuclidean'Squared Euclidean distance (default)'cityblock'Sum of absolute differences, i.e., L1		
	'cosine'	One minus the cosine of the included angle between points (treated as vectors)	
	'correlation' One minus the sample correlation between points (treated as sequences of values)		
	'Hamming'	Percentage of coordinates that differ	

	'Jaccard'	Percentage of non-zero coordinates that differ	
	Vector	A numeric distance matrix in upper triangular vector form, such as is created by pdist. X is not used in this case, and can safely be set to [].	
	[] = silhoue function of the for	tte(X,clust,distfun,p1,p2,) accepts a distance m	
	d = distfun(X0,X,p1,p2,)		
	where X0 is a 1-by-p point, X is an n-by-p matrix of points, and p1,p2, are optional additional arguments. The function distfun returns an n-by-1 vector d of distances between X0 and each point (row) in X. The arguments p1, p2, are passed directly to the function distfun.		
Remarks		ue for each point is a measure of how similar that point is to luster compared to points in other clusters, and ranges from ned as	
	S(i) = (min(b))	(i,:),2) - a(i)) ./ max(a(i),min(b(i,:),2))	
		average distance from the ith point to the other points in its () is the average distance from the ith point to points in	
Examples	randn(10 cidx = kmeans	,2)+ones(10,2); ,2)-ones(10,2)]; (X,2,'distance','sqeuclid'); e(X,cidx,'sqeuclid');	
See Also	dendrogram, kmea	ns, linkage, pdist	
References		d P.J. Rousseeuw, <i>Finding Groups in Data: An</i> uster Analysis, Wiley, 1990	

skewness

Purpose	Sample skewness
Syntax	y = skewness(X) y = skewness(X,flag)
Description	<pre>y = skewness(X) returns the sample skewness of X. For vectors, skewness(x) is the skewness of the elements of x. For matrices, skewness(X) is a row vector containing the sample skewness of each column.</pre>
	Skewness is a measure of the asymmetry of the data around the sample mean. If skewness is negative, the data are spread out more to the left of the mean than to the right. If skewness is positive, the data are spread out more to the right. The skewness of the normal distribution (or any perfectly symmetric distribution) is zero.
	The skewness of a distribution is defined as

$$y = \frac{E(x-\mu)^3}{\sigma^3}$$

where μ is the mean of x, σ is the standard deviation of x, and E(t) represents the expected value of the quantity t.

y = skewness(X, flag) specifies whether to correct for bias (flag = 0) or not (flag = 1, the default). When X represents a sample from a population, the skewness of X is biased; that is, it will tend to differ from the population skewness by a systematic amount that depends on the size of the sample. You can set flag = 0 to correct for this systematic bias.

Example X =

X = randn([5 4])

X = 1.1650 1.6961 -1.4462 -0.3600 0.6268 0.0591 -0.7012 -0.1356 0.0751 1.7971 1.2460 -1.34930.3516 0.2641 -0.6390 -1.2704 -0.6965 0.8717 0.5774 0.9846 y = skewness(X)y =

-0.2933 0.0482 0.2735 0.4641

See Also kurtosis, mean, moment, std, var

squareform

Purpose	Reformat the output of pdist into a square matrix.		
Syntax	S = squareform(Y)		
Description	S = squareform(Y) reformats the distance information returned by pdist from a vector into a square matrix. In this format, $S(i,j)$ denotes the distance between the <i>i</i> and <i>j</i> observations in the original data.		
See Also	pdist		

Purpose	Standard deviation of a sample		
Syntax	y = std(X)		
Description	$y = std(X)$ computes the sample standard deviation of the data in X. For vectors, $std(x)$ is the standard deviation of the elements in x. For matrices, $std(X)$ is a row vector containing the standard deviation of each column of X. std normalizes by <i>n</i> -1 where <i>n</i> is the sequence length. For normally distributed data, the square of the standard deviation is the minimum variance unbiased estimator of σ^2 (the second parameter).		
	The standard deviation is		
	$s = \left(\frac{1}{n-1}\sum_{i=1}^{n} (x_i - \bar{x})^2\right)^{\frac{1}{2}}$ where the sample average is $\bar{x} = \frac{1}{n}\sum x_i$. The std function is part of the standard MATLAB language.		
Examples	<pre>In each column, the expected value of y is one. x = normrnd(0,1,100,6); y = std(x)</pre>		
	y = stu(x) y =		
	0.9536 1.0628 1.0860 0.9927 0.9605 1.0254		
	y = std(-1:2:1)		
	y = 1.4142		
See Alco			

See Also

cov, var

stepwise

Purpose	Interactive environment for stepwise regression		
Syntax	stepwise(X,y) stepwise(X,y,inmodel) stepwise(X,y,inmodel,alpha)		
Description	stepwise(X,y) fits a regression model of y on the columns of X. It displays three figure windows for interactively controlling the stepwise addition and removal of model terms.		
	<pre>stepwise(X,y,inmodel) allows control of the terms in the original regression model. The values of vector, inmodel, are the indices of the columns of the matrix X to include in the initial model.</pre>		
	stepwise(X,y,inmodel,alpha) allows control of the length confidence intervals on the fitted coefficients. alpha is the significance for testing each term in the model. By default, alpha = $1 - (1 - 0.025)^{(1/p)}$ where p is the number of columns in X. This translates to plotted 95% simultaneous confidence intervals (Bonferroni) for all the coefficients.		
	The least squares coefficient is plotted with a green filled circle. A coefficient is not significantly different from zero if its confidence interval crosses the white zero line. Significant model terms are plotted using solid lines. Terms not significantly different from zero are plotted with dotted lines.		
	Click on the confidence interval lines to toggle the state of the model coefficients. If the confidence interval line is green, the term is in the model. If the confidence interval line is red, the term is not in the model.		
	Use the Export menu to move variables to the base workspace.		
Example	See "Stepwise Regression" on page 4-24.		
Reference	[1] Draper, N. and H. Smith, <i>Applied Regression Analysis, Second Edition</i> , John Wiley and Sons, Inc. 1981 pp. 307–312.		
See Also	regstats, regress, rstool		

surfht

Purpose	Interactive contour plot		
Syntax	<pre>surfht(Z) surfht(x,y,Z)</pre>		
Description	surfht(Z) is an interactive contour plot of the matrix Z treating the values in Z as height above the plane. The x-values are the column indices of Z while the y-values are the row indices of Z. surfht(x,y,Z) where x and y are vectors specify the x and y-axes on the contour plot. The length of x must match the number of columns in Z, and the length of y must match the number of rows in Z.		
	There are vertical and horizontal reference lines on the plot whose intersection defines the current <i>x</i> -value and <i>y</i> -value. You can drag these dotted white reference lines and watch the interpolated <i>z</i> -value (at the top of the plot) update simultaneously. Alternatively, you can get a specific interpolated <i>z</i> -value by typing the <i>x</i> -value and <i>y</i> -value into editable text fields on the <i>x</i> -axis and <i>y</i> -axis respectively.		

tabulate

Purpose	Frequency table		
Syntax	<pre>table = tabulate(x) tabulate(x)</pre>		
Description	table = $tabulate(x)$ takes a vector of positive integers, x, and returns a matrix, table.		
	The first column of table contains the values of x. The second contains the number of instances of this value. The last column contains the percentage each value.		
	tabulate with no output arguments displays a formatted table in the command window.		
Example	tabulate([1 2 4 4 3 4])		
	Value Count Percent		
	1 1 16.67%		
	2 1 16.67%		
	3 1 16.67%		
	4 3 50.00%		
See Also	pareto		

 Purpose
 Read tabular data from the file system

 Syntax
 [data,varnames,casenames] = tblread

 [data,varnames,casenames] = tblread('filename')
 [data,varnames,casenames] = tblread('filename', 'delimiter')

Description [data,varnames,casenames] = tblread displays the **File Open** dialog box for interactive selection of the tabular data file. The file format has variable names in the first row, case names in the first column and data starting in the (2,2) position.

[data,varnames,casenames] = tblread(filename) allows command line specification of the name of a file in the current directory, or the complete pathname of any file.

[data,varnames,casenames] = tblread(filename,'delimiter') allows specification of the field 'delimiter' in the file. Accepted values are 'tab', 'space', or 'comma'.

tblread returns the data read in three values.

Return Value	Description
data	Numeric matrix with a value for each variable-case pair.
varnames	String matrix containing the variable names in the first row.
casenames	String matrix containing the names of each case in the first column.

Example

[data,varnames,casenames] = tblread('sat.dat')

data =

470	530
520	480
varnames	=
Ma] a	
Male	

tblread

Female casenames = Verbal Quantitative

See Also caseread, tblwrite, tdfread

Purpose	Writes tabular data to the file system	
Syntax	tblwrite(data,'varnames','casenames') tblwrite(data,'varnames','casenames','filename')	
Description	tblwrite(data, 'varnames', 'casenames') displays the File Open dialog box for interactive specification of the tabular data output file. The file format ha variable names in the first row, case names in the first column and data starting in the (2,2) position.	
	'varnames' is a string matrix containing the variable names. 'casenames' is a string matrix containing the names of each case in the first column. data is a numeric matrix with a value for each variable-case pair.	
	tblwrite(data,'varnames','casenames','filename') allows command lin specification of a file in the current directory, or the complete pathname of an file in the string 'filename'.	
Example	Continuing the example from tblread:	
	tblwrite(data,varnames,casenames,'sattest.dat') type sattest.dat	
	Male Female Verbal 470 530 Quantitative 520 480	
See Also	casewrite, tblread	

Purpose	Student's t cumulative distribution function (cdf)
---------	--

Syntax P = tcdf(X,V)

Description P = tcdf(X,V) computes Student's t cdf at each of the values in X using the corresponding degrees of freedom in V. Vector or matrix inputs for X and V must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in V must be positive integers.

The t cdf is

$$p = F(x|v) = \int_{-\infty}^{x} \frac{\Gamma\left(\frac{v+1}{2}\right)}{\Gamma\left(\frac{v}{2}\right)} \frac{1}{\sqrt{v\pi}} \frac{1}{\left(1 + \frac{t^2}{v}\right)^{\frac{v+1}{2}}} dt$$

The result, p, is the probability that a single observation from the t distribution with v degrees of freedom will fall in the interval (- ∞x].

Examples Suppose 10 samples of Guinness beer have a mean alcohol content of 5.5% by volume and the standard deviation of these samples is 0.5%. What is the probability that the true alcohol content of Guinness beer is less than 5%?

t = (5.0 - 5.5) / 0.5; probability = tcdf(t,10 - 1) probability = 0.1717

See Also cdf, tinv, tpdf, trnd, tstat

Purpose	Read file containing t	ab-delimited	numeric	and text values
Syntax	tdfread tdfread('filename' tdfread('filename'	-)	
Description	file. The file should co column names in the assigned to a variable numeric, the variable string matrix. After a	onsist of colum first line of the e with the spece is converted to all values are i	nns of va e file. Ea cified na to numb mported	for interactive selection of the data alues, separated by tabs, and with ach column is read from the file and ame. If all values for a column are ers; otherwise the variable is a l, tdfread displays information t of the whos command.
	tdfread('filename' in the current directo	-		e specification of the name of a file thname of any file.
	tdfread('filename' ' <i>delimiter</i> ' separate	-	-	tes that the character specified by Accepted values are:
	 ' ' or 'space' '\t' or 'tab' ',' or 'comma' ';' or 'semi' ' ' or 'bar' 			
	The default delimiter	'is 'tab'.		
Example	type sat2.dat			
	Test,Gender,Scor Verbal,Male,470 Verbal,Female,53 Quantitative,Mal Quantitative,Fem tdfread('sat2.da	30 Le,520 nale,480 at',',')		
	Name S	Size	Bytes	Class

tdfread

Gender	4x6	48 char array	
Score	4x1	32 double array	
Test	4x12	96 char array	

Grand total is 76 elements using 176 bytes

See Also tblread

Purpose Inverse of the Student's t cumulative distribution function (cdf)

Syntax X = tinv(P,V)

Description X = tinv(P,V) computes the inverse of Student's t cdf with parameter V for the corresponding probabilities in P. Vector or matrix inputs for P and V must be the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The degrees of freedom in V must be positive integers, and the values in P must lie on the interval [0 1].

The t inverse function in terms of the t cdf is

$$x = F^{-1}(p|v) = \{x:F(x|v)=p\}$$

where

$$p = F(x|v) = \int_{-\infty}^{x} \frac{\Gamma\left(\frac{v+1}{2}\right)}{\Gamma\left(\frac{v}{2}\right)} \frac{1}{\sqrt{v\pi}} \frac{1}{\left(1+\frac{t^{2}}{v}\right)^{\frac{v+1}{2}}} dt$$

The result, *x*, is the solution of the cdf integral with parameter v, where you supply the desired probability *p*.

Examples What is the 99th percentile of the t distribution for one to six degrees of freedom?

percentile = tinv(0.99,1:6)

percentile =

31.8205 6.9646 4.5407 3.7469 3.3649 3.1427

See Also icdf, tcdf, tpdf, trnd, tstat

Purpose	Student's t probability density function (pdf)
---------	--

Syntax Y = tpdf(X,V)

Description Y = tpdf(X,V) computes Student's t pdf at each of the values in X using the corresponding parameters in V. Vector or matrix inputs for X and V must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The degrees of freedom in V must be positive integers.

Student's t pdf is

$$y = f(x|v) = \frac{\Gamma\left(\frac{v+1}{2}\right)}{\Gamma\left(\frac{v}{2}\right)} \frac{1}{\sqrt{v\pi}} \frac{1}{\left(1 + \frac{x^2}{v}\right)^{\frac{v+1}{2}}}$$

Examples The mode of the t distribution is at x = 0. This example shows that the value of the function at the mode is an increasing function of the degrees of freedom.

tpdf(0,1:6)

ans =

0.3183 0.3536 0.3676 0.3750 0.3796 0.3827

The t distribution converges to the standard normal distribution as the degrees of freedom approaches infinity. How good is the approximation for v = 30?

See Also

pdf, tcdf, tinv, trnd, tstat

Purpose	Show classification or regression tree graphically	
Syntax	treedisp(T) treedisp(T,'param1',val1,'param2',val2,)	
Description	treedisp(T) takes as input a decision tree T as computed by the treefit function, and displays it in a figure window. Each branch in the tree is labeled with its decision rule, and each terminal node is labeled with the predicted value for that node.	
		ode, the left child node corresponds to the points that satisfy the right child node corresponds to the points that do not on.
	—	lay pop-up menu at the top of the figure enables you to rmation about each node:
	Identity	The node number, whether the node is a branch or a leaf, and the rule that governs the node
	Variable ranges	The range of each of the predictor variables for that node
	Node statistics	The number of elements of the data set that fit the node and (the mean and standard deviation of what?)
	After you select the type of information you want, click on any node to di the information for that node.	
	The Pruning level spin button displays the number of levels that have been cut from the tree, and the number of levels in the unpruned tree. For example, 1 of 6 indicates that the unpruned tree has six levels, and that one level has been cut from the tree. Use the spin button to change the pruning level.	
		am1',val1,'param2',val2,) specifies optional value pairs. Valid parameters are:
	0	A cell array of names for the predictor variables, in the order in which they appear in the X matrix from which the ree was created (see treefit)
	'nrunelevel' I	nitial pruning level to display

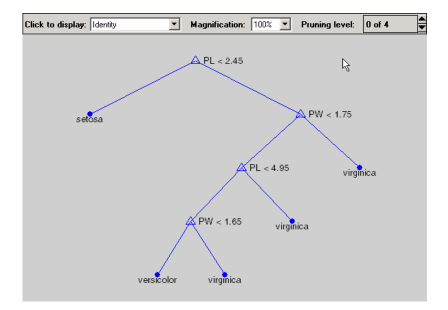
'prunelevel' Initial pruning level to display

treedisp

Examples

Create and graph classification tree for Fisher's iris data. The names in this example are abbreviations for the column contents (sepal length, sepal width, petal length, and petal width).

```
load fisheriris;
t = treefit(meas,species);
treedisp(t,'names',{'SL' 'SW' 'PL' 'PW'});
```



See Also

treefit, treeprune, treetest

Purpose	Fit a tree-bas	ed model for classification or regression
Syntax	T = treefit(T = treefit((X,y) (X,y,'param1',val1,'param2',val2,)
Description	function of provector of n res strings contai	(X,y) creates a decision tree T for predicting response y as a edictors X. X is an n-by-m matrix of predictor values. y is either a ponse values (for regression), or a character array or cell array of ining n class names (for classification). Either way, T is a binary ch non-terminal node is split based on the values of a column of X.
	T = treefit(X,y,'param1',val1,'param2',val2,) specifies optional parameter name-value pairs. Valid parameters are	
	For all trees:	
	'catidx'	Vector of indices of the columns of X. treefit treats these columns as unordered categorical values.
	'method'	Either 'classification' (default if y is text) or 'regression' (default if y is numeric)
	'splitmin'	A number n such that impure nodes must have n or more observations to be split (default 10)
	'prune'	'on' (default) to compute the full tree and a sequence of pruned subtrees, or 'off' for the full tree without pruning
	For classificat	tion trees only:

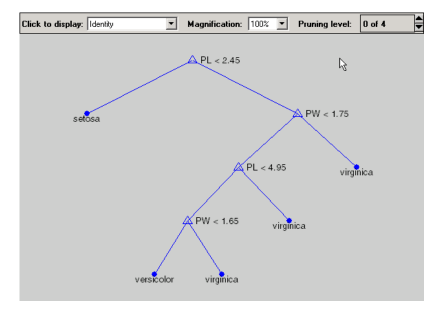
'cost' p-by-p matrix C, where C(i,j) is the cost of classifying a
point into class i if its true class is j (default has
C(i,j)=1 if i~=j, and C(i,j)=0 if i=j). C can also be a
structure S with two fields : S.group containing the
group names, and S.cost containing a matrix of cost
values.

treefit

'splitcriterion'	Criterion for choosing a split: either 'gdi' (default) for Gini's diversity index, 'twoing' for the twoing rule, or 'deviance' for maximum deviance reduction
'priorprob'	Prior probabilities for each class, specified as a vector (one value for each distinct group name) or as a structure S with two fields: S.group containing the group names, and S.prob containing a vector of corresponding probabilities

Examples Create a classification tree for Fisher's iris data.

```
load fisheriris;
t = treefit(meas,species);
treedisp(t,'names',{'SL' 'SW' 'PL' 'PW'});
```



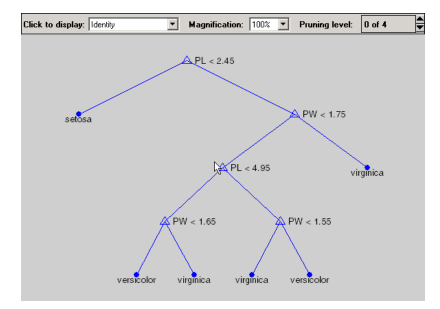
See Also treedisp, treetest

References [1] Breiman, et al., *Classification and Regression Trees*, Chapman and Hall, Boca Raton, 1993.

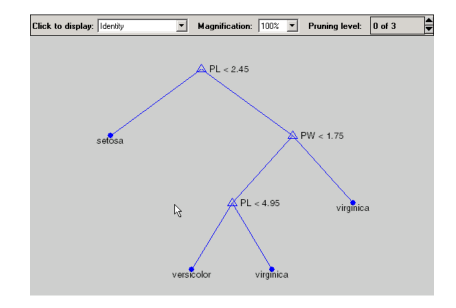
treeprune

Purpose	Produce a sequence of subtrees by pruning
Syntax	<pre>T2 = treeprune(T1, 'level',level) T2 = treeprune(T1, 'nodes',nodes) T2 = treeprune(T1)</pre>
Description	T2 = treeprune(T1, 'level', level) takes a decision tree T1 as created by the treefit function, and a pruning level level, and returns the decision tree T2 pruned to that level. The value level = 0 means no pruning. Trees are pruned based on an optimal pruning scheme that first prunes branches giving less improvement in error cost.
	<pre>T2 = treeprune(T1, 'nodes', nodes) prunes the nodes listed in the nodes vector from the tree. Any T1 branch nodes listed in nodes become leaf nodes in T2, unless their parent nodes are also pruned. The treedisp function can display the node numbers for any node you select.</pre>
	T2 = treeprune(T1) returns the decision tree T2 that is the same as T1, but with the optimal pruning information added. This is useful only if you created T1 by pruning another tree, or by using the treefit function with pruning set 'off'. If you plan to prune a tree multiple times, it is more efficient to create the optimal pruning sequence first.
	Pruning is the process of reducing a tree by turning some branch nodes into leaf nodes, and removing the leaf nodes under the original branch.
Examples	Display the full tree for Fisher's iris data, as well as the next largest tree from the optimal pruning sequence.
	load fisheriris; t = treefit(meas,species,'splitmin',5); treedisp(t,'names',{'SL' 'SW' 'PL' 'PW'});

treeprune



t1 = treeprune(t, 'level',1); treedisp(t1, 'names', {'SL' 'SW' 'PL' 'PW'});



treeprune

See Also treefit, treetest, treedisp

Purpose	Compute error rate for tree
Syntax	<pre>cost = treetext(T,'resubstitution') cost = treetest(T,'test',X,y) cost = treetest(T,'crossvalidate',X,y) [cost,secost,ntnodes,bestsize] = treetest() [] = treetest(,'param1',val1,'param2',val2,)</pre>
Description	<pre>cost = treetext(T, 'resubstitution') computes the cost of the tree T using a resubstitution method. T is a decision tree as created by the treefit function. The cost of the tree is the sum over all terminal nodes of the estimated probability of that node times the node's cost. If T is a classification tree, the cost of a node is the sum of of the misclassification costs of the observations in that node. If T is a regression tree, the cost of a node is the average squared error over the observations in that node. cost is a vector of cost values for each subtree in the optimal pruning sequence for T. The resubstitution cost is based on the same sample that was used to create the original tree, so it underestimates the likely cost of applying the tree to new data. cost = treetest(T, 'test',X,y) uses the predictor matrix X and response y as a test sample, applies the decision tree T to that sample, and returns a vector cost of cost values computed for the test sample. X and y should not be the same as the learning sample, which is the sample that was used to fit the tree T. cost = treetest(T, 'crossvalidate',X,y) uses 10-fold cross-validation to compute the cost vector. X and y should be the learning sample, which is the</pre>
	sample that was used to fit the tree T. The function partitions the sample into 10 subsamples, chosen randomly but with roughly equal size. For classification trees, the subsamples also have roughly the same class proportions. For each subsample, treetest fits a tree to the remaining data and uses it to predict the subsample. It pools the information from all subsamples to compute the cost for the whole sample.
	<pre>[cost,secost,ntnodes,bestlevel] = treetest() also returns the vector secost containing the standard error of each cost value, the vector ntnodes containing number of terminal nodes for each subtree, and the scalar</pre>

treetest

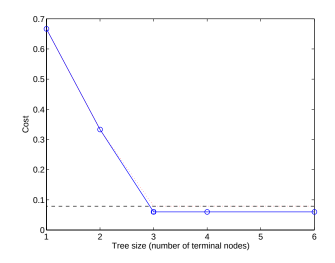
bestlevel containing the estimated best level of pruning. bestlevel = 0 means no pruning, i.e., the full unpruned tree. The best level is the one that produces the smallest tree that is within one standard error of the minimum-cost subtree.

[...] = treetest(..., 'param1', val1, 'param2', val2,...) specifies optional parameter name-value pairs chosen from the following:

nsamples'	The number of cross-validations samples (default 10).
treesize'	Either 'se' (default) to choose the smallest tree whose cost is within one standard error of the minimum cost, or 'min' to choose the minimal cost tree.

Examples Find the best tree for Fisher's iris data using cross-validation. The solid line shows the estimated cost for each tree size, the dashed line marks 1 standard error above the minimum, and the square marks the smallest tree under the dashed line.

```
% Start with a large tree.
load fisheriris;
t = treefit(meas,species','splitmin',5);
% Find the minimum-cost tree.
[c,s,n,best] = treetest(t,'cross',meas,species);
tmin = treeprune(t,'level',best);
% Plot smallest tree within 1 std. error of minimum cost tree.
[mincost,minloc] = min(c);
plot(n,c,'b-o', n,c+s,'r:', n(best+1),c(best+1),'bs',...
n,(mincost+s(minloc))*ones(size(n)),'k--');
xlabel('Tree size (number of terminal nodes)')
ylabel('Cost')
```





treefit, treedisp

treeval

Purpose	Compute fitted value for decision tree applied to data
Syntax	YFIT = treeval(T,X) YFIT = treeval(T,X,subtrees) [YFIT,NODE] = treeval() [YFIT,NODE,CNAME] = treeval()
Description	YFIT = treeval(T,X) takes a classification or regression tree T as produced by the treefit function, and a matrix X of predictor values, and produces a vector YFIT of predicted response values. For a regression tree, YFIT(j) is the fitted response value for a point having the predictor values $X(j,:)$. For a classification tree, YFIT(j) is the class number into which the tree would assign the point with data $X(j,:)$. To convert the number into a class name, use the third output argument, cname (below).
	YFIT = treeval(T,X,subtrees) takes an additional vector subtrees of pruning levels, with 0 representing the full, unpruned tree. T must include a pruning sequence as created by the treefit or prunetree function. If subtree has k elements and X has n rows, then the output YFIT is an n-by-k matrix, with the jth column containing the fitted values produced by the subtrees(j) subtree. subtrees must be sorted in ascending order.
	[YFIT,NODE] = treeval() also returns an array NODE of the same size as YFIT containing the node number assigned to each row of X. The treedisp function can display the node numbers for any node you select.
	[YFIT,NODE,CNAME] = treeval() is valid only for classification trees. It returns a cell array CNAME containing the predicted class names.
Examples	Find the predicted classifications for Fisher's iris data.
	<pre>load fisheriris; t = treefit(meas,species); % Create decision tree sfit = treeval(t,meas); % Find assigned class numbers sfit = t.classname(sfit); % Get class names mean(strcmp(sfit,species)) % Compute proportion correctly % classified ans =</pre>

See Also treefit, treeprune, treetest

trimmean

Purpose	Mean of a sample of data excluding extreme values
Syntax	<pre>m = trimmean(X,percent)</pre>
Description	m = trimmean(X, percent) calculates the mean of a sample X excluding the highest and lowest percent/2 of the observations. The trimmed mean is a robust estimate of the location of a sample. If there are outliers in the data, the trimmed mean is a more representative estimate of the center of the body of the data. If the data is all from the same probability distribution, then the trimmed mean is less efficient than the sample average as an estimator of the location of the data.
Examples	<pre>This example shows a Monte Carlo simulation of the efficiency of the 10% trimmed mean relative to the sample average for normal data. x = normrnd(0,1,100,100); m = mean(x); trim = trimmean(x,10); sm = std(m); strim = std(m); efficiency = (sm/strim).^2 efficiency = 0.9702</pre>
See Also	mean, median, geomean, harmmean

Purpose	Random numbers from Student's t distribution
Syntax	R = trnd(V) R = trnd(V,m) R = trnd(V,m,n)
Description	R = trnd(V) generates random numbers from Student's t distribution with V degrees of freedom. The size of R is the size of V.
	R = trnd(V,m) generates random numbers from Student's t distribution with V degrees of freedom, where m is a 1-by-2 vector that contains the row and column dimensions of R.
	R = trnd(V,m,n) generates random numbers from Student's t distribution with V degrees of freedom, where scalars m and n are the row and column dimensions of R.
Examples	<pre>noisy = trnd(ones(1,6))</pre>
	noisy =
	19.7250 0.3488 0.2843 0.4034 0.4816 -2.4190
	numbers = trnd(1:6,[1 6])
	numbers =
	-1.9500 -0.9611 -0.9038 0.0754 0.9820 1.0115
	numbers = trnd(3,2,6)
	numbers =
	-0.3177 -0.0812 -0.6627 0.1905 -1.5585 -0.0433 0.2536 0.5502 0.8646 0.8060 -0.5216 0.0891
See Also	tcdf, tinv, tpdf, tstat

tstat

Purpose	Mean and var	ianc	e for the S	Studer	nt's t di	stribution	
Syntax	[M,V] = tsta	t(N	U)				
Description	[M,V] = tsta with paramet	•					or Student's t distribution ne size as NU.
		. If	v is one, th			-	ter ν is zero for values of ν ne variance for values of ν
Examples	Find the mean $[m,v] = t$					legrees of fro	eedom.
	m =						
	NaN	0	0	0	0		
	0	0	0	0	0		
	0	0	0	0	0		
	0	0	0	0	0		
	0	0	0	0	0		
	0	0	0	0	0		
	v =						
	Na	N	1.4000	1.	1818	1.1176	1.0870
	Na		1.3333		1667	1.1111	1.0833
	3.000		1.2857		1538	1.1053	1.0800
	2.000		1.2500		1429	1.1000	1.0769
	1.666 1.500		1.2222		1333 1250	1.0952 1.0909	1.0741 1.0714
	1.500	0	1.2000	1.	1200	1.0909	1.0/14

Note that the variance does not exist for one and two degrees of freedom.

See Also tcdf, tinv, tpdf, trnd

Purpose	Hypothesis testing for a single sample mean
Syntax	h = ttest(x,m) h = ttest(x,m,alpha) [h,sig,ci] = ttest(x,m,alpha,tail)
Description	h = ttest(x,m) performs a t-test at significance level 0.05 to determine whether a sample from a normal distribution (in x) could have mean m when the standard deviation is unknown.
	h = ttest(x,m,alpha) gives control of the significance level, alpha. For example if alpha = 0.01, and the result, h, is 1 you can reject the null hypothesis at the significance level 0.01. If h is 0, you cannot reject the null hypothesis at the alpha level of significance.
	<pre>[h,sig,ci] = ttest(x,m,alpha,tail) allows specification of one- or two-tailed tests. tail is a flag that specifies one of three alternative hypotheses:</pre>
	• tail = 0 specifies the alternative $\bar{x} \neq m$ (default)
	• tail = 1 specifies the alternative $\bar{x} > m$
	• tail = -1 specifies the alternative $\bar{x} < m$
	Output sig is the p-value associated with the T-statistic
	$T = rac{ar{x} - m}{s \diagup \sqrt{n}}$
	where s is the sample standard deviation and n is the number of observations in the sample. sig is the probability that the observed value of T could be as large or larger <i>by chance</i> under the null hypothesis that the mean of x is equal to m .
	ci is a 1-alpha confidence interval for the true mean.
Example	This example generates 100 normal random numbers with theoretical mean zero and standard deviation one. The observed mean and standard deviation are different from their theoretical values, of course. We test the hypothesis that there is no true difference.

Normal random number generator test.

The result h = 0 means that we cannot reject the null hypothesis. The significance level is 0.4474, which means that by chance we would have observed values of T more extreme than the one in this example in 45 of 100 similar experiments. A 95% confidence interval on the mean is [-0.1165 0.2620], which includes the theoretical (and hypothesized) mean of zero.

PurposeHypothesis testing for the difference in means of two samplesSyntaxh = ttest2(x,y)
[h,significance,ci] = ttest2(x,y,alpha)
[h,significance,ci,stats] = ttest2(x,y,alpha)
[...] = ttest2(x,y,alpha,tail)Descriptionh = ttest2(x,y) performs a t-test to determine whether two samples from a
normal distribution (in x and y) could have the same mean when the standard
deviations are unknown but assumed equal.
The result, h, is 1 if you can reject the null hypothesis at the 0.05 significance
level alpha and 0 otherwise.
The significance is the p-value associated with the T-statistic

$$T = \frac{x - y}{s \sqrt{\frac{1}{n} + \frac{1}{m}}}$$

where *s* is the pooled sample standard deviation and *n* and *m* are the numbers of observations in the x and y samples. significance is the probability that the observed value of T could be as large or larger *by chance* under the null hypothesis that the mean of *x* is equal to the mean of *y*.

ci is a 95% confidence interval for the true difference in means.

[h, significance, ci] = ttest2(x, y, alpha) gives control of the significance level alpha. For example if alpha = 0.01, and the result, h, is 1, you can reject the null hypothesis at the significance level 0.01. ci in this case is a 100(1-alpha)% confidence interval for the true difference in means.

[h, significance, ci, stats] = ttest2(x, y, alpha) returns a structure stats with two elments, tstat and df. The tstat element is the value of the t statistic, and df is its degree of freedom.

 $[\ldots]$ = ttest2(x,y,alpha,tail) allows specification of one- or two-tailed tests, where tail is a flag that specifies one of three alternative hypotheses:

• tail = 0 specifies the alternative $\mu_x \neq \mu_y$ (default)

- tail = 1 specifies the alternative $\mu_x > \mu_v$
- tail = -1 specifies the alternative $\mu_x < \mu_v$

Examples

This example generates 100 normal random numbers with theoretical mean 0 and standard deviation 1. We then generate 100 more normal random numbers with theoretical mean 1/2 and standard deviation 1. The observed means and standard deviations are different from their theoretical values, of course. We test the hypothesis that there is no true difference between the two means. Notice that the true difference is only one half of the standard deviation of the individual observations, so we are trying to detect a signal that is only one half the size of the inherent noise in the process.

The result h = 1 means that we can reject the null hypothesis. The significance is 0.0017, which means that by chance we would have observed values of t more extreme than the one in this example in only 17 of 10,000 similar experiments! A 95% confidence interval on the mean is [-0.7352 -0.1720], which includes the theoretical (and hypothesized) difference of -0.5.

Purpose	Discrete uniform cumulative distribution (cdf) function
Syntax	P = unidcdf(X,N)
Description	P = unidcdf(X,N) computes the discrete uniform cdf at each of the values in X using the corresponding parameters in N. Vector or matrix inputs for X and N must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The maximum observable values in N must be positive integers.
	The discrete uniform cdf is
	$p = F(x N) = \frac{floor(x)}{N}I_{(1,,N)}(x)$
	The result, p , is the probability that a single observation from the discrete uniform distribution with maximum N will be a positive integer less than or equal to x . The values x do not need to be integers.
Examples	What is the probability of drawing a number 20 or less from a hat with the numbers from 1 to 50 inside?
	<pre>probability = unidcdf(20,50)</pre>
	probability =
	0.4000
See Also	cdf, unidinv, unidpdf, unidrnd, unidstat

unidinv

Purpose	Inverse of the discrete uniform cumulative distribution function
Syntax	<pre>X = unidinv(P,N)</pre>
Description	X = unidinv(P,N) returns the smallest positive integer X such that the discrete uniform cdf evaluated at X is equal to or exceeds P. You can think of P as the probability of drawing a number as large as X out of a hat with the numbers 1 through N inside.
	Vector or matrix inputs for N and P must have the same size, which is also the size of X. A scalar input for N or P is expanded to a constant matrix with the same dimensions as the other input. The values in P must lie on the interval $[0\ 1]$ and the values in N must be positive integers.
Examples	<pre>x = unidinv(0.7,20) x = 14 y = unidinv(0.7 + eps,20) y = 15 A small change in the first parameter produces a large jump in output. The cdf and its inverse are both step functions. The example shows what happens at a step.</pre>
See Also	icdf, unidcdf, unidpdf, unidrnd, unidstat

Purpose	Discrete uniform probability density function (pdf)		
Syntax	Y = unidpdf(X,N)		
Description	unidpdf(X,N) computes the discrete uniform pdf at each of the values in X using the corresponding parameters in N. Vector or matrix inputs for X and N must have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in N must be positive integers.		
	The discrete uniform pdf is		
	$y = f(x N) = \frac{1}{N}I_{(1,,N)}(x)$		
	You can think of y as the probability of observing any one number between 1 and n .		
Examples	For fixed n, the uniform discrete pdf is a constant.		
	y = unidpdf(1:6,10)		
	y = 0.1000 0.1000 0.1000 0.1000 0.1000 0.1000		
	Now fix x, and vary n.		
	likelihood = unidpdf(5,4:9)		
	likelihood =		
	0 0.2000 0.1667 0.1429 0.1250 0.1111		
See Also	pdf, unidcdf, unidinv, unidrnd, unidstat		

unidrnd

Purpose	Random numbers from the discrete uniform distribution
Syntax	R = unidrnd(N) R = unidrnd(N,mm) R = unidrnd(N,mm,nn)
Description	The discrete uniform distribution arises from experiments equivalent to drawing a number from one to N out of a hat.
	R = unidrnd(N) generates discrete uniform random numbers with maximum N. The parameters in N must be positive integers. The size of R is the size of N.
	R = unidrnd(N,mm) generates discrete uniform random numbers with maximum N, where mm is a 1-by-2 vector that contains the row and column dimensions of R.
	R = unidrnd(N,mm,nn) generates discrete uniform random numbers with maximum N, where scalars mm and nn are the row and column dimensions of R.
Examples	<pre>In the Massachusetts lottery, a player chooses a four digit number. Generate random numbers for Monday through Saturday. numbers = unidrnd(10000,1,6) - 1 numbers =</pre>
	2189 470 6788 6792 9346
See Also	unidcdf, unidinv, unidpdf, unidstat

unidstat

Purpose	Mean and variance for the discrete uniform distribution			
Syntax	[M,V] = unidstat(N)			
Description	[M,V] = unidstat(N) returns the mean and variance for the discrete uniform distribution with parameter N.			
	The mean of the discrete uniform distribution with parameter N is $(N+1)/2$. The variance is $(N^2-1)/12$.			
Examples	[m,v] = unidstat(1:6)			
	m = 1.0000 1.5000 2.0000 2.5000 3.0000 3.5000 v =			
	0 0.2500 0.6667 1.2500 2.0000 2.9167			
See Also	unidcdf, unidinv, unidpdf, unidrnd			

unifcdf

Purpose	Continuous uniform cumulative distribution function (cdf)
Syntax	P = unifcdf(X,A,B)
Description	P = unifcdf(X,A,B) computes the uniform cdf at each of the values in X using the corresponding parameters in A and B (the minimum and maximum values, respectively). Vector or matrix inputs for X, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.
	The uniform cdf is
	$p = F(x a,b) = \frac{x-a}{b-a}I_{[a,b]}(x)$
	The standard uniform distribution has $A = 0$ and $B = 1$.
Examples	What is the probability that an observation from a standard uniform distribution will be less than 0.75?
	<pre>probability = unifcdf(0.75)</pre>
	probability =
	0.7500
	What is the probability that an observation from a uniform distribution with $a = -1$ and $b = 1$ will be less than 0.75?
	<pre>probability = unifcdf(0.75,-1,1)</pre>
	probability =
	0.8750
See Also	cdf, unifinv, unifit, unifpdf, unifrnd, unifstat

Purpose	Inverse continuous uniform cumulative distribution function (cdf)
Syntax	X = unifinv(P,A,B)
Description	X = unifinv(P,A,B) computes the inverse of the uniform cdf with parameters A and B (the minimum and maximum values, respectively) at the corresponding probabilities in P. Vector or matrix inputs for P, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs.
	The inverse of the uniform cdf is
	$x = F^{-1}(p a, b) = a + p(a - b)I_{[0, 1]}(p)$
	The standard uniform distribution has $A = 0$ and $B = 1$.
Examples	What is the median of the standard uniform distribution? median_value = unifinv(0.5) median_value =
	0.5000
	What is the 99th percentile of the uniform distribution between -1 and 1?
	<pre>percentile = unifinv(0.99,-1,1)</pre>
	percentile =
	0.9800
See Also	icdf, unifcdf, unifit, unifpdf, unifrnd, unifstat

unifit

Purpose	Parameter estimates for uniformly distributed data
Syntax	[ahat,bhat] = unifit(X) [ahat,bhat,ACI,BCI] = unifit(X) [ahat,bhat,ACI,BCI] = unifit(X,alpha)
Description	[ahat,bhat] = unifit(X) returns the maximum likelihood estimates (MLEs) of the parameters of the uniform distribution given the data in X.
	[ahat,bhat,ACI,BCI] = unifit(X) also returns 95% confidence intervals, ACI and BCI, which are matrices with two rows. The first row contains the lower bound of the interval for each column of the matrix X. The second row contains the upper bound of the interval.
	[ahat,bhat,ACI,BCI] = unifit(X,alpha) allows control of the confidence level alpha. For example, if alpha = 0.01 then ACI and BCI are 99% confidence intervals.
Example	r = unifrnd(10,12,100,2); [ahat,bhat,aci,bci] = unifit(r)
	ahat =
	10.0154 10.0060
	bhat =
	11.9989 11.9743
	aci =
	9.9551 9.9461
	10.0154 10.0060
	bci =
	11.9989 11.9743
	12.0592 12.0341
See Also	betafit, binofit, expfit, gamfit, normfit, poissfit, unifcdf, unifinv, unifpdf, unifrnd, unifstat, weibfit

Purpose	Continuous uniform probability density function (pdf)
Syntax	Y = unifpdf(X,A,B)
Description	Y = unifpdf(X,A,B) computes the continuous uniform pdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in B must be greater than those in A.
	The continuous uniform distribution pdf is
	$y = f(x a, b) = \frac{1}{b-a}I_{[a, b]}(x)$
	The standard uniform distribution has $A = 0$ and $B = 1$.
Examples	For fixed a and b, the uniform pdf is constant.
	<pre>x = 0.1:0.1:0.6; y = unifpdf(x)</pre>
	y = 1 1 1 1 1 1
	What if x is not between a and b?
	y = unifpdf(-1,0,1)
	y = 0
See Also	pdf, unifcdf, unifinv, unifrnd, unifstat

unifrnd

Purpose	Random numbers from the continuous uniform distribution		
Syntax	R = unifrnd(A,B) R = unifrnd(A,B,m) R = unifrnd(A,B,m,n)		
Description	 R = unifrnd(A,B) generates uniform random numbers with parameters A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of R. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input. R = unifrnd(A,B,m) generates uniform random numbers with parameters A and B, where m is a 1-by-2 vector that contains the row and column dimensions of R. 		
	R = unifrnd(A,B,m,n) generates uniform random numbers with parameters A and B, where scalars m and n are the row and column dimensions of R.		
Examples	<pre>random = unifrnd(0,1:6) random =</pre>		
See Also	unifcdf, unifinv, unifpdf, unifstat		

Purpose	Mean and variance for the continuous uniform distribution		
Syntax	[M,V] = unifstat(A,B)		
Description	[M,V] = unifstat(A,B) returns the mean and variance for the continuous uniform distribution with parameters specified by A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of M and V. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input.		
	The mean of the continuous uniform distribution with parameters a and b is $(a+b)/2$, and the variance is $(b-a)^2/12$.		
Examples	a = 1:6; b = 2.*a; [m,v] = unifstat(a,b)		
	m = 1.5000 3.0000 4.5000 6.0000 7.5000 9.0000		
	v = 0.0833 0.3333 0.7500 1.3333 2.0833 3.0000		
See Also	unifcdf, unifinv, unifpdf, unifrnd		

Purpose	Variance of a sample
Syntax	y = var(X) y = var(X,1) y = var(X,w)
Description	y = var(X) computes the variance of the data in X. For vectors, $var(x)$ is the variance of the elements in x. For matrices, $var(X)$ is a row vector containing the variance of each column of X.
	y = var(x) normalizes by <i>n</i> -1 where <i>n</i> is the sequence length. For normally distributed data, this makes var(x) the minimum variance unbiased estimator MVUE of σ^2 (the second parameter).
	y = var(x,1) normalizes by <i>n</i> and yields the second moment of the sample data about its mean (moment of inertia).
	y = var(X,w) computes the variance using the vector of positive weights w. The number of elements in w must equal the number of rows in the matrix X. For vector x, w and x must match in length.
	var supports both common definitions of variance. Let <i>SS</i> be the sum of the squared deviations of the elements of a vector x from their mean. Then, $var(x) = SS/(n-1)$ is the MVUE, and $var(x,1) = SS/n$ is the maximum likelihood estimator (MLE) of σ^2 .
Examples	$x = [-1 \ 1];$ $w = [1 \ 3];$ v1 = var(x) v1 = 2 v2 = var(x,1) v2 = 1 v3 = var(x,w) v3 = var(x,w)

0.7500

See Also cov, std

weibcdf

Purpose	Weibull cumulative distribution function (cdf)				
Syntax	P = weibcdf(X,A,B)				
Description	P = weibcdf(X,A,B) computes the Weibull cdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must be positive.				
	The Weibull cdf is				
	$p = F(x a,b) = \int_0^x abt^{b-1} e^{-at^b} dt = 1 - e^{-ax^b} I_{(0,\infty)}(x)$				
Examples	What is the probability that a value from a Weibull distribution with parameters $a = 0.15$ and $b = 0.24$ is less than 500?				
	<pre>probability = weibcdf(500,0.15,0.24)</pre>				
	probability =				
	0.4865				
	How sensitive is this result to small changes in the parameters?				
	<pre>[A,B] = meshgrid(0.1:0.05:0.2,0.2:0.05:0.3); probability = weibcdf(500,A,B)</pre>				
	probability =				
	0.2929 0.4054 0.5000				
	0.3768 0.5080 0.6116				
	0.4754 0.6201 0.7248				
See Also	cdf, weibfit, weibinv, weiblike, weibpdf, weibplot, weibrnd, weibstat				

Purpose	Parameter estimates and confidence intervals for Weibull data		
Syntax	phat = weibfit(x) [phat,pci] = weibfit(x) [phat,pci] = weibfit(x,alpha)		
Description	phat = weibfit(x) returns the maximum likelihood estimates, phat, of the parameters of the Weibull distribution given the values in vector x, which must be positive. phat is a two-element row vector: phat(1) estimates the Weibull parameter a, and phat(2) estimates the Weibull parameter b in the pdf $y = f(x a, b) = abx^{b-1}e^{-ax^{b}}I_{(0,\infty)}(x)$		
	<pre>[phat,pci] = weibfit(x) also returns 95% confidence intervals in the two-row matrix pci. The first row contains the lower bound of the confidence interval, and the second row contains the upper bound. The columns of pci correspond to the columns of phat. [phat,pci] = weibfit(x,alpha) allows control over the confidence interval returned, 100(1-alpha)%.</pre>		
Example	<pre>r = weibrnd(0.5,0.8,100,1); [phat,pci] = weibfit(r) phat =</pre>		
See Also	betafit, binofit, expfit, gamfit, normfit, poissfit, unifit, weibcdf, weibinv, weiblike, weibpdf, weibplot, weibrnd, weibstat		

weibinv

Purpose

Syntax	X = weibinv(P,A,B)			
Description	X = weibinv(P,A,B) computes the inverse of the Weibull cdf with parameters A and B for the corresponding probabilities in P. Vector or matrix inputs for P, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other inputs. The parameters in A and B must be positive.			
	The inverse of the Weibull cdf is			
	$x = F^{-1}(p a, b) = \left[\frac{1}{a}\ln\left(\frac{1}{1-p}\right)\right]^{\frac{1}{b}}I_{[0, 1]}(p)$			
Examples	A batch of light bulbs have lifetimes (in hours) distributed Weibull with parameters a = 0.15 and b = 0.24. What is the median lifetime of the bulbs? life = weibinv(0.5,0.15,0.24)			
	life =			
	588.4721			
	What is the 90th percentile?			
	life = weibinv(0.9,0.15,0.24)			
	life =			
	8.7536e+04			
See Also	icdf, weibcdf, weibfit, weiblike, weibpdf, weibplot, weibrnd, weibstat			

Inverse of the Weibull cumulative distribution function

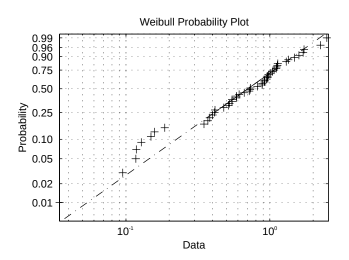
Purpose	Weibull negative log-likelihood function			
Syntax	logL = weiblike(params,data) [logL,avar] = weiblike(params,data)			
Description	logL = weiblike(params,data) returns the Weibull log-likelihood with parameters params(1) = a and params(2) = b given the data x_i .			
	[logL,avar] = weiblike(params,data) also returns avar, which is the asymptotic variance-covariance matrix of the parameter estimates if the values in params are the maximum likelihood estimates. avar is the inverse of Fisher's information matrix. The diagonal elements of avar are the asymptotic variances of their respective parameters.			
	The Weibull negative log-likelihood is			
	$(-\log L) = -\log \prod_{i=1}^{n} f(a, b x_i) = -\sum_{i=1}^{n} \log f(a, b x_i)$ weiblike is a utility function for maximum likelihood estimation.			
Example	This example continues the example from weibfit.			
	r = weibrnd(0.5,0.8,100,1); [logL,info] = weiblike([0.4746 0.7832],r)			
	logL =			
	203.8216			
	info =			
	0.0021 0.0022 0.0022 0.0056			
Reference	[1] Patel, J. K., C. H. Kapadia, and D. B. Owen, <i>Handbook of Statistical Distributions</i> , Marcel-Dekker, 1976.			
See Also	betalike, gamlike, mle, normlike, weibcdf, weibfit, weibinv, weibpdf, weibplot, weibrnd, weibstat			

weibpdf

Purpose	Weibull probability density function (pdf)					
Syntax	Y = weibpdf(X,A,B)					
Description	Y = weibpdf(X,A,B) computes the Weibull pdf at each of the values in X using the corresponding parameters in A and B. Vector or matrix inputs for X, A, and B must all have the same size. A scalar input is expanded to a constant matrix with the same dimensions as the other input. The parameters in A and B must all be positive.					
	The Weibull pdf is					
	$y = f(x a, b) = abx^{b-1}e^{-ax^{b}}I_{(0,\infty)}(x)$					
	Some references refer to the Weibull distribution with a single parameter. This corresponds to weibpdf with $A = 1$.					
Examples	The exponential distribution is a special case of the Weibull distribution.					
	lambda = 1:6; y = weibpdf(0.1:0.1:0.6,lambda,1)					
	y =					
	0.9048 1.3406 1.2197 0.8076 0.4104 0.1639					
	y1 = exppdf(0.1:0.1:0.6,1./lambda)					
	y1 =					
	0.9048 1.3406 1.2197 0.8076 0.4104 0.1639					
Reference	[1] Devroye, L., Non-Uniform Random Variate Generation. Springer-Verlag. New York, 1986.					
See Also	pdf, weibcdf, weibfit, weibinv, weiblike, weibplot, weibrnd, weibstat					

weibplot

Purpose	Weibull probability plot
Syntax	<pre>weibplot(X) h = weibplot(X)</pre>
Description	<pre>weibplot(X) displays a Weibull probability plot of the data in X. If X is a matrix, weibplot displays a plot for each column. h = weibplot(X) returns handles to the plotted lines. The purpose of a Weibull probability plot is to graphically assess whether the data in X could come from a Weibull distribution. If the data are Weibull the plot will be linear. Other distribution types may introduce curvature in the plot.</pre>
Example	r = weibrnd(1.2,1.5,50,1); weibplot(r)





normplot, weibcdf, weibfit, weibinv, weiblike, weibpdf, weibrnd, weibstat

weibrnd

Purpose	Random numbers from the Weibull distribution			
Syntax	R = weibrnd(A,B) R = weibrnd(A,B,m) R = weibrnd(A,B,m,n)			
Description	R = weibrnd(A,B) generates Weibull random numbers with parameters A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of R. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input.			
	R = weibrnd(A,B,m) generates Weibull random numbers with parameters A and B, where m is a 1-by-2 vector that contains the row and column dimensions of R.			
	 R = weibrnd(A,B,m,n) generates Weibull random numbers with parameter A and B, where scalars m and n are the row and column dimensions of R. Devroye refers to the Weibull distribution with a single parameter; this is weibrnd with A = 1. 			
Examples	n1 = weibrnd(0.5:0.5:2,0.5:0.5:2)			
	n1 = 0.0093 1.5189 0.8308 0.7541			
	n2 = weibrnd(1/2,1/2,[1 6])			
	n2 = 29.7822 0.9359 2.1477 12.6402 0.0050 0.0121			
Reference	[1] Devroye, L., <i>Non-Uniform Random Variate Generation</i> . Springer-Verlag. New York, 1986.			
See Also	weibcdf, weibfit, weibinv, weiblike, weibpdf, weibplot, weibstat			

Purpose Mean and variance for the Weibull distribution

Syntax [M,V] = weibstat(A,B)

Description [M,V] = weibstat(A,B) returns the mean and variance for the Weibull distribution with parameters specified by A and B. Vector or matrix inputs for A and B must have the same size, which is also the size of M and V. A scalar input for A or B is expanded to a constant matrix with the same dimensions as the other input.

The mean of the Weibull distribution with parameters a and b is

$$a^{-\frac{1}{b}}\Gamma(1+b^{-1})$$

and the variance is

	$a^{-\frac{2}{b}} \left[\Gamma(1+2b^{-1}) \right]$	$(1) - \Gamma^2(1 + b)$	-1)]	
Examples	[m,v] = weib	stat(1:4,1	:4)	
	m = 1.0000	0.6267	0.6192	0.6409
	v = 1.0000	0.1073	0.0506	0.0323
	weibstat(0.5	,0.7)		
	ans =			
	3.4073			
See Also	weibcdf, weibfit	,weibinv,v	weiblike,we	eibpdf,weibplot,weibrnd

wishrnd

Purpose	Generate Wishart random matrix
Syntax	W = wishrnd(SIGMA,df) W = wishrnd(SIGMA,df,D) [W,D] = wishrnd(SIGMA,df)
Description	W = wishrnd(SIGMA,df) generates a random matrix W having the Wishart distribution with covariance matrix SIGMA and with df degrees of freedom.
	W = wishrnd(SIGMA,df,D) expects D to be the Cholesky factor of SIGMA. If you call wishrnd multiple times using the same value of SIGMA, it's more efficient to supply D instead of computing it each time.
	[W,D] = wishrnd(SIGMA,df) returns D so you can provide it as input in future calls to wishrnd.
See Also	iwishrnd

Purpose	Transform a factor	settings matrix to a design matrix	
Syntax	D = x2fx(X) D = x2fx(X,'mode.	1')	
Description	D = x2fx(X) transforms a matrix of system inputs, X, to a design matrix for a linear additive model with a constant term.		
	<pre>D = x2fx(X, 'model') allows control of the order of the regression model. 'model' can be one of these strings:</pre>		
	'interaction'	Includes constant, linear, and cross product terms	
	'quadratic'	Includes interactions and squared terms	
	'purequadratic'	Includes constant, linear, and squared terms	
	 Alternatively model can be a matrix of terms. In this case, each row of model represents one term. The value in a column is the exponent to which the same column in X for that term is raised, D(i,j) = prod(x(i,:).^model(j,:)). This allows for models with polynomial terms of arbitrary order. The order of columns for a quadratic model is: 		
		ins for a quadratic model is.	
	 Constant term Linear terms (the input Y columns 1, 2,, k) 		
	 Linear terms (the input X columns 1,2,,k) Interaction terms formed by taking pairwise products of X columns (1,2), (1,3),, (1,k), (2,3),, (k-1,k) Squared terms in the order 1,2,,k 		
	Other models use a subset of these terms but keep them in this order.		
	x2fx is a utility function for rstool, regstats, and cordexch.		
Examples	<pre>Example 1. x = [1 2 3]'; model = [0 1 2]'; D = x2fx(x,model) D =</pre>		

1	1	1
1	2	4
1	3	9

The first column is *x* to the 0th power. The second column is *x* to the 1st power. And the last column is *x* squared.

Example 2.

```
x = [1 2 3;4 5 6]';
model = 'quadratic';
D = x2fx(x,model)
D =
1 1 4 4 1 16
1 2 5 10 4 25
1 3 6 18 9 36
```

Let x_1 be the first column of x and x_2 be the second. Then the first column of D is the constant term, the second column is x_1 , the third column is x_2 , the fourth column is x_1x_2 , the fifth column is x_1^2 , and the last columns is x_2^2 .

See Also rstool, candexch, candgen, cordexch, rowexch, regstats

Purpose	X-bar chart for Statistical Process Control	
Syntax	xbarplot(DATA) xbarplot(DATA,conf) xbarplot(DATA,conf,specs,' <i>sigmaest</i> ') [outlier,h] = xbarplot()	
Description	xbarplot (DATA) displays an x-bar chart of the grouped responses in DATA. The rows of DATA contain replicate observations taken at a given time, and must be in time order. The graph contains the sample mean \bar{x} for each group, a center line at the average \bar{x} value, and upper and lower control limits. The limits are placed at a three-sigma distance on either side of the center line, where sigma is an estimate of the standard deviation of \bar{x} . If the process is in control, fewer than 3 out of 1000 observations would be expected to fall outside the control limits by random chance. So if you observe points outside the limits, you can take this as evidence that the process is not in control. xbarplot (DATA, conf) allows control of the confidence level of the upper and lower plotted confidence limits. The default conf = 0.9973 produces three-sigma limits.	
	<pre>norminv(1 - (19973)/2) ans = 3 To get k-sigma limits, use the expression 1-2*(1-normcdf(k)). For example, the correct conf value for 2-sigma limits is 0.9545, as shown below. k = 2; 1-2*(1-normcdf(k)) ans =</pre>	
	0.9545	

xbarplot(DATA,conf,specs) plots the specification limits in the two element vector specs.

xbarplot(DATA,conf,specs,'sigmaest') specifies how xbarplot should estimate the standard deviation. Acceptable values are:

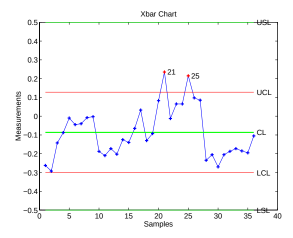
xbarplot

- 's' use the average of the group standard deviations (default)
- 'v' use the square root of a pooled variance estimate
- 'r' use the average range with each group; requires 25 or fewer observations per group

[outlier,h] = xbarplot(DATA,conf,specs) returns outlier, a vector of indices to the rows where the mean of DATA is out of control, and h, a vector of handles to the plotted lines.

Example Plot an x-bar chart of measurements on newly machined parts, taken at one hour intervals for 36 hours. Each row of the runout matrix contains the measurements for four parts chosen at random. The values indicate, in thousandths of an inch, the amount the part radius differs from the target radius.

load parts
xbarplot(runout,0.999,[-0.5 0.5])



The points in groups 21 and 25 are out of control, so the mean in those groups was higher than would be expected by random chance alone. There is evidence that the process was not in control when those measurements were collected.

See Also capaplot, histfit, ewmaplot, schart

Purpose	Standardized Z score
Syntax	Z = zscore(D)
Description	Z = zscore(D) returns the deviation of each column of D from its mean, normalized by its standard deviation. This is known as the Z score of D.
	For column vector V, the Z score is $Z = (V-mean(V))./std(V)$.

ztest

Purpose	Hypothesis testing for the mean of one sample with known variance
Syntax	h = ztest(x,m,sigma) h = ztest(x,m,sigma,alpha) [h,sig,ci,zval] = ztest(x,m,sigma,alpha,tail)
Description	<pre>h = ztest(x,m,sigma) performs a Z test at significance level 0.05 to determine whether a sample x from a normal distribution with standard deviation sigma could have mean m.</pre>
	h = ztest(x,m,sigma,alpha) gives control of the significance level alpha. For example, if alpha = 0.01 and the result is $h = 1$, you can reject the null hypothesis at the significance level 0.01. If $h = 0$, you cannot reject the null hypothesis at the alpha level of significance.
	<pre>[h,sig,ci] = ztest(x,m,sigma,alpha,tail) allows specification of one- or two-tailed tests, where tail is a flag that specifies one of three alternative hypotheses:</pre>
	• tail = 0 specifies the alternative $\bar{x} \neq m$ (default)
	• tail = 1 specifies the alternative $\bar{x} > m$
	• tail = -1 specifies the alternative $\bar{x} < m$
	zval is the value of the Z statistic
	$z = \frac{\bar{x} - m}{\sigma / \sqrt{n}}$
	where n is the number of observations in the sample.
	sig is the probability that the observed value of Z could be as large or larger by <i>chance</i> under the null hypothesis that the mean of x is equal to m .
	ci is a 1-alpha confidence interval for the true mean.
Example	This example generates 100 normal random numbers with theoretical mean zero and standard deviation one. The observed mean and standard deviation are different from their theoretical values, of course. We test the hypothesis that there is no true difference. x = normrnd(0,1,100,1);

The result, h = 0, means that we cannot reject the null hypothesis. The significance level is 0.4669, which means that by chance we would have observed values of Z more extreme than the one in this example in 47 of 100 similar experiments. A 95% confidence interval on the mean is [-0.1232 0.2687], which includes the theoretical (and hypothesized) mean of zero.

A

Selected Bibliography

[1] Atkinson, A.C., and A.N. Donev, *Optimum Experimental Designs*, Oxford Science Publications 1992.

[2] Bates, D. and D. Watts. *Nonlinear Regression Analysis and Its Applications*, John Wiley and Sons. 1988. pp. 271–272.

[3] Bernoulli, J., Ars Conjectandi, Basiliea: Thurnisius [11.19], 1713

[4] Box, G.E.P. and N.R. Draper, *Empirical Model-Building and Response Surfaces*, Wiley, New York. 1987.

[5] Box, G.E.P., W.G. Hunter, and J.S. Hunter. *Statistics for Experimenters*. Wiley, New York. 1978.

[6] Chatterjee, S. and A.S. Hadi. Influential Observations, High Leverage Points, and Outliers in Linear Regression. Statistical Science, 1986. pp. 379– 416.

[7] Dobson, A. J., An Introduction to Generalized Linear Models, 1990, CRC Press.

[8] Efron, B., and R.J. Tibshirani. An Introduction to the Bootstrap, Chapman and Hall, New York. 1993.

[9] Evans, M., N. Hastings, and B. Peacock. *Statistical Distributions, Second Edition*. John Wiley and Sons, 1993.

[10] Hald, A., *Statistical Theory with Engineering Applications*, John Wiley and Sons, 1960. p. 647.

[11] Hogg, R.V., and J. Ledolter. *Engineering Statistics*. MacMillan Publishing Company, 1987.

[12] Johnson, N., and S. Kotz. *Distributions in Statistics: Continuous Univariate Distributions*. John Wiley and Sons, 1970.

[13] McCullagh, P., and J. A. Nelder, *Generalized Linear Models*, 2nd edition, 1990, Chapman and Hall.

[14] Montgomery, D.C., *Design and Analysis of Experiments*, Wiley, New York. 1984.

[15] Moore, J., *Total Biochemical Oxygen Demand of Dairy Manures*. Ph.D. thesis. University of Minnesota, Department of Agricultural Engineering, 1975.

[16] Poisson, S.D., Recherches sur la Probabilité des Jugements en Matiere Criminelle et en Metière Civile, Précédées des Regles Générales du Calcul des Probabilitiés. Paris: Bachelier, Imprimeur-Libraire pour les Mathematiques, 1837.

[17] "Student," On the Probable Error of the Mean. Biometrika, 6:1908. pp. 1-25.

[18] Weibull, W., A Statistical Theory of the Strength of Materials. Ingeniors Vetenskaps Akademiens Handlingar, Royal Swedish Institute for Engineering Research. Stockholm, Sweden, No. 153. 1939.



Index

A

absolute deviation 3-5 additive effects 4-9 alternative hypothesis 6-3 analysis of variance 2-23 multivariate 7-20 N-way 4-12 one-way 4-4 two-way 4-9 ANOVA 4-2 anoval 12-19 anova2 12-25 anovan 12-29 aoctool 12-35 aoctool demo 11-10 average linkage 12-219

B

bacteria counts 4-4 barttest 12-38 baseball odds 12-48, 12-50 bbdesign 12-39 Bera-Jarque. See Jarque-Bera Bernoulli random variables 12-52 beta distribution 2-11 betacdf 12-40 betafit **12-41** betainv 12-43 betalike **12-44** betapdf 12-45 betarnd **12-46** betastat 12-47 binocdf 12-48 binofit 12-49 binoinv **12-50** binomial distribution 2-13

negative. See also negative binomial distribuiton binopdf **12-51** binornd **12-52** binostat **12-53** bootstrap **12-54** bootstrap sampling 3-19 box plots 8-3 Box-Behnken designs 10-9 generating 12-39 boxplot **12-56**

С

candexch 12-59 candgen 12-61 canoncorr 12-62 capability studies 9-6 capable **12-64** capaplot 12-66 casenames reading from file 12-68 writing to file 12-69 caseread 12-68 casewrite 12-69 ccdesign 12-70 cdf definition 2-5 cdf 12-72 cdfplot 12-73 central composite designs 10-8 generating 12-70 Central Limit Theorem 2-34 centroid linkage 12-219 Chatterjee and Hadi example 4-20 chi2cdf 12-75

chi2inv 12-76 chi2pdf 12-77 chi2rnd 12-78 chi2stat 12-79 chi-square distributions 2-16 circuit boards 12-51 City Block metric in cluster analysis 12-304 classical multidimensional scaling 7-47 cmdscale function 12-86 overview 7-47 reconstructing a map 7-49 simple example 7-48 classification trees 5-8 See also decision trees classify 12-80 cluster 12-82 cluster analysis 7-26 functions 12-13 hierarchical clustering 7-26 K-means clustering 7-40 cluster tree creation 12-218 from data 12-84 from linkage output 12-82 cluster trees inconsistency coefficient 12-188 plotting 12-106 clusterdata 12-84 cmdscale 12-86 coin 12-155 combnk 12-88 comparisons, multiple 4-6 complete linkage 12-219 confidence intervals hypothesis tests 6-3 nonlinear regression 5-6 control charts 9-3

EWMA charts 9-5 S charts 9-4 Xbar charts 9-3 cophenet 12-89 cophenetic correlation coefficient 12-89 defined 7-32 cordexch 12-91 corr 12-93 corrcoef 12-95 correlation coefficients 12-93, 12-95 cov 12-98 Cp index 9-6, 12-64 Cpk index 9-6, 12-64 crosstab 12-99 cumulative distribution functions 12-5 cumulative distribution function (cdf) 2-5 empirical 3-16 graphing an estimate 8-8

D

data partitioning K-means clustering 7-40 data sets statistical examples 12-16 daugment **12-101** dcovary **12-103** decision trees 5-8 computing error rate 12-383 computing response values 12-386 creating 12-377 creating subtrees 12-380 displaying 12-375 example 5-8 fitting 12-377 pruning 12-380 demos 11-1, 12-16 design of experiments 11-19 polynomial curve fitting 11-5 probability distributions 11-3 random number generation 11-18 dendrogram 12-106, 12-234 density estimation ksdensity function 12-202 descriptive statistics 3-1 functions 12-9 **Design of Experiments** functions 12-13 design of experiments 10-1 Box-Behnken designs 10-9 central composite designs 10-8 D-optimal designs 10-11 fractional factorial designs 10-6 full factorial designs 10-4 response surface designs 10-8 dimension reduction common factor analysis 12-122 PCA from covariance matrix 12-299 PCA from raw data matrix 12-319 PCA residuals 12-300 discrete uniform distribution 2-19 dissimilarity matrix creating 7-27 distance matrix creating 7-27 distribution testing functions 12-15 distributions 2-1 supported 2-11 disttool 12-109 disttool demo 11-3 DOE. See Design of Experiments D-optimal designs 10-11

creating from candidate set 12-59 generating candidate set 12-61 dummyvar **12-110**

E

ecdf 12-111 empirical cumulative distribution function 3-16 ecdf function 12-111 erf 2-34 error function 2-34 errorbar 12-113 estimate 11-6 Euclidean distance in cluster analysis 12-303 EWMA charts 9-5 ewmaplot **12-114** expcdf 12-116 expfit **12-117** expinv 12-118 exponential distribution 2-20 exppdf 12-119 exprnd 12-120 expstat **12-121** extrapolated 12-322

F

F distributions 2-22 F statistic 4-21 factor analysis maximum likelihood 12-122 factoran **12-122** factorial designs fractional 10-6 full 10-4 generating fractional 12-135 factorial designs (continued) generating full 12-147 fcdf 12-131 ff2n 12-132 file I/O functions 12-16 finv 12-133 floppy disks 12-182 fpdf 12-134 fracfact 12-135 fractional factorial designs 10-6 generating 12-135 friedman 12-139 Friedman's test 4-34 frnd 12-143 fstat 12-144 fsurfht 12-145 full factorial designs 10-4 generating 12-147 fullfact 12-147 furthest neighbor linkage 12-219

G

gamcdf 12-148 gamfit 12-149 gaminv 12-150 gamlike 12-151 gamma distribution 2-25 gampdf 12-152 gamrnd 12-153 gamstat 12-154 Gaussian 12-179 geocdf 12-155 geoinv 12-156 geomean 12-157 geometric distribution 2-27 geopdf 12-158 geornd 12-159 geostat 12-160 gline 12-161 glmdemo 12-162 glmdemo demo 11-21 glmfit 12-163 glmval 12-168 gname 12-170 gplotmatrix 12-172 group mean clusters, plot 7-25 grouped plot matrix 7-20 grpstats 12-175 gscatter 12-176 Guinness beer 2-39, 12-370

Η

harmmean 12-178 hat matrix 4-19 hierarchical clustering 7-26 basic procedure 7-27 computing inconsistency coefficient 12-188 creating cluster tree 12-218 creating clusters 7-37 creating clusters from data 12-84 creating clusters from linkage output 12-82 depth of comparison 7-33 determining proximity 12-302 evaluating cluster formation 12-89 finding dissimilarities betwen objects 7-33 finding similarities beween objects 7-27 formatting distance information 12-362 grouping objects 7-30 inconsistency coefficient 12-188 plotting cluster trees 12-106 hierarchiical clustering cophenetic correlation coefficient 12-89

hist 12-179 histfit 12-180 histogram 11-18 Hotelling's T squared 7-12 hougen 12-181 Hougen-Watson model 5-3 hygecdf 12-182 hygeinv 12-183 hygepdf 12-184 hygernd 12-185 hygestat 12-186 hypergeometric distribution 2-28 hypothesis tests 6-1 functions 12-15

I

icdf 12-187 incomplete beta function 2-11 incomplete gamma function 2-25 inconsistency coefficient 12-188 inconsistent 12-188 inspector 12-307 interaction 4-9 interpolated 12-365 interquartile range (iqr) 3-5 inverse cdf 2-5 inverse cdf 2-5 inverse cumulative distribution functions 12-6 iqr 12-190 iwishrnd 12-191

J

Jarque-Bera test 12-192 jbtest **12-192**

Κ

Kaplan-Meier cumulative distribution function 12-111 kernel bandwidth 3-13 kernel smoothing function 3-15 K-means clustering 7-40 cluster separation 7-41 creating clusters 12-194 example 7-41 local minima 7-45 number of clusters 7-43 overview 7-40 silhouette plot 12-358 kmeansdata partitioning K-means clustering 12-194 kruskalwallis 12-198 Kruskal-Wallis test 4-34 ksdensity 12-202 kstest 12-204 kstest2 12-208 kurtosis **12-211**

L

latin hypercube sample 12-214 normal distribution 12-215 least squares 12-315 leverage 12-213 lhsdesign 12-214 lhsnorm 12-215 light bulbs, life of 12-118 likelihood function 12-45 Lilliefors test 6-5 lillietest 12-216 linear models 4-1 functions 12-11 generalized 4-28 linear transformation Procrustes 12-320 linkage 12-218 logncdf 12-221 logninv 12-222 lognormal distribution 2-29 lognpdf 12-224 lognrnd 12-225 lognstat 12-226 lottery 12-398 lsline 12-227 LU factorizations 12-314

Μ

mad 12-228 mahal 12-229 Mahalanobis distance 12-229 in cluster analysis 12-303 MANOVA 7-20 manova1 12-230 manovacluster 12-234 maximum likelihood factor analysis 12-122 MDS See also multidimensional scaling mean 2-9 of probability distribution 2-9 mean 12-236 Mean Squares (MS) 12-19 measures of central tendency 3-3 dispersion 3-5 median 12-237 metric multidimensional scaling See also classical multidimensional scaling Minkowski metric

in cluster analysis 12-304 mle 12-238 models linear 4-1 nonlinear 5-1 moment 12-239 moments of distribution functions 12-8 Monte Carlo simulation 12-190 multcompare **12-240** multidimensional arrays classical (metric) scaling 12-86 multidimensional scaling (MDS) classical (metric) 7-47 multiple linear regression 4-18 multivariate analysis of variance 7-20 example 7-20 multivariate statistics 7-1 analysis of variance 7-20 cluster analysis 7-26 functions 12-13 hierarchical clustering 7-26 K-means clustering 7-40 Principal Components Analysis 7-3 mvnpdf 12-247 mvnrnd 12-248 mvtrnd 12-249

Ν

nanmax **12-250** nanmean **12-251** nanmedian **12-252** nanmin **12-253** NaNs 3-7 nanstd **12-254** nansum **12-255**

nbincdf 12-256 nbinfit 12-258 nbininy **12-259** nbinpdf 12-260 nbinrnd **12-262** nbinstat 12-263 ncfcdf 12-265 ncfinv 12-267 ncfpdf 12-268 ncfrnd 12-269 ncfstat 12-270 nctcdf 12-271 nctinv **12-272** nctpdf **12-273** nctrnd **12-274** nctstat 12-275 ncx2cdf 12-276 ncx2inv 12-278 ncx2pdf 12-279 ncx2rnd 12-280 ncx2stat **12-281** nearest neighbor linkage 12-219 negative binomial distribution 2-31 confidence intervals 12-258 cumulative distribution function (cdf) 12-256 definition 2-31 inverse cumulative distribution function (cdf) 12 - 259mean and variance 12-263 modeling number of auto accidents 2-32 nbincdf function 12-256 nbininv function 12-259 nbinpdf function 12-260 parameter estimates 12-258 probability density function (pdf) 12-260 random matrices 12-262 Newton's method 12-150

nlinfit **12-282** nlintool 12-284 nlintool demo 5-7 nlparci 12-285 nlpredci 12-286 noncentral F distribution 2-23 nonlinear regression functions 12-12 nonlinear regression models 5-1 nonparametric testing functions 12-15 normal distribution 2-34 normal probability plots 8-2, 8-4 normalizing a dataset 7-28 using zscore 12-421 normcdf 12-288 normdemo 12-296 normfit **12-289** norminy **12-290** normlike **12-291** normpdf **12-292** normplot **12-293** normrnd 12-295 normstat **12-297** notation, mathematical conventions 1-6 notches 12-56 null 6-3 null hypothesis 6-3

0

one-way analysis of variance (ANOVA) 4-2 outliers 3-3

P

parameter estimation functions 12-4 pareto 12-298 Pascal, Blaise 2-14 PCA. See Principal Components Analysis pcacov 12-299 pcares 12-300 pdf definition 2-4 pdf 12-301 pdist 12-302 percentiles 3-11 perms 12-306 plots 3-11 plotting statistical functions 12-10 poisscdf 12-307 poissfit 12-309 poissinv 12-310 Poisson distribution 2-36 poisspdf 12-311 poissrnd 12-312 poisstat 12-313 polyconf 12-314 polyfit **12-315** polynomial 11-5 polytool 12-316 polytool demo 11-5 polyval 12-317 popcorn 12-27, 12-141 prctile 12-318 Principal Components Analysis (PCA) 7-3 component scores 7-7 component variances 7-10 Hotelling's T squared 7-12 principal components 7-7

quality of life example 7-4 Scree plot 7-11 princomp 12-319 probability density functions 12-5 probability density estimation 3-13 comparing estimates 3-16 function 12-202 kernel bandwidth 3-13 kernel smoothing functioon 3-15 probability density function (pdf) definition 2-4 probability distribution mean and variance 2-9 probability distributions 2-1 functions 12-4 process control statistical 9-1 procrustes 12-320 Procrustes Analysis 12-320 p-value 4-11, 6-3

Q

qqplot **12-322** QR decomposition 4-18 quality assurance 12-51 quantile-quantile plots 8-2, 8-6

R

random **12-324** random number generation 2-7 direct 2-7 functions 12-7 inverted 2-7 rejection 2-8 random numbers 2-7 random samples inverse Wishart 12-191 latin hypercube 12-214 latin hypercube with normal distribution 12 - 215Wishart 12-416 randtool 12-325 randtool demo 11-18 range 12-326 ranksum 12-327 ravlcdf 12-328 raylinv 12-329 raylpdf 12-330 raylrnd **12-331** raylstat 12-332 rcoplot 12-333 reconstruction map from inter-city distances 7-49 refcurve **12-334** reference lines 11-3 references A-1 refline **12-335** regress 12-336 regression 2-23 nonlinear 5-1 robust 4-32 stepwise 4-24 regression trees 5-8 See also decision trees regstats 12-338 relative efficiency 12-190 residuals 4-21 response surface designs 10-8 Response Surface Methodology (RSM) 4-22 respoonse surface designs Box-Behnken 10-9

central composite 10-8 ridge 12-341 robust 3-3 robust linear fit 12-322 robustdemo 12-343 robustdemo demo 11-22 robustfit 12-344 rowexch 12-348 rsmdemo 12-350 rsmdemo demo 11-19 R-square 4-21 rstool 12-351 rstool demo 4-22

S

S charts 9-4 scaling arrays classical multidimensional 12-86 scatter plots 8-10 grouped 7-20 schart 12-352 Scree plot 7-11 segmentation analysis 7-26 significance level 6-3 signrank 12-354 signtest 12-356 silhouette 12-358 similarity matrix creating 7-27 simulation 12-190 single linkage 12-219 skewness 8-3 skewness 12-360 squareform 12-362 standard normal 12-292

Standardized Euclidean distance in cluster analysis 12-303 statistical plots 8-1 Statistical Process Control 9-1 capability studies 9-6 control charts 9-3 functions 12-11 statistical references A-1 statistically significant 12-19, 12-198, 12-230 stepwise 4-24, **12-364** stepwise regression 4-24 Sum of Squares (SS) 12-19 surfht **12-365** symmetric 12-148

T

t distributions 2-39 noncentral 2-40 tab-delimited data reading from file 12-371 tabular data reading from file 12-367 tabulate 12-366 taxonomy analysis 7-26 tblread 12-367 tblwrite **12-369** tcdf 12-370 tdfread **12-371** tinv **12-373** tpdf **12-374** treefit 12-377 treeprune **12-380** trees See also decision trees See also decision trees treeshow 12-375

treetest **12-383** treeval **12-386** trimmean **12-388** trnd **12-389** tstat **12-390** ttest **12-391** ttest2 **12-393** two-way ANOVA 4-9 typographical conventions (table) xiv

U

unbiased 12-363, 12-406 unidcdf **12-395** unidinv **12-396** unidpdf **12-397** unidrnd **12-398** unidstat **12-399** unifcdf **12-400** unifinv **12-401** unifit **12-402** uniform distribution 2-42 unifpdf **12-403** unifrnd **12-404** unifstat **12-405**

V

var **12-406** variance 2-9 of probability distribution 2-9

W

ward linkage 12-220 weibcdf **12-408** weibfit **12-409**

Index

weibinv 12-410
weiblike 12-411
weibpdf 12-412
weibplot 12-413
weibrnd 12-414
weibstat 12-415
Weibull distribution 2-43
Weibull probability plots 8-7
Weibull, Waloddi 2-43
whiskers 8-3, 12-56
Wishart random matrix 12-416
inverse 12-191
wishrnd 12-416

Х

x2fx **12-417** Xbar charts 9-3 xbarplot **12-419**

Ζ

zscore **12-421** ztest **12-422** Index