GAUSSTM

Language Reference

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Introduction

This GAUSS[™] Language Reference describes each of the commands, procedures, and functions available in the GAUSS programming language. These functions can be divided into four categories:

- Mathematical, statistical, and scientific functions.
- Data handling routines, including data matrix manipulation and description routines, and file I/O.
- Programming statements, including branching, looping, display features, error checking, and shell commands.
- Graphics functions.

The first category contains those functions to be expected in a high-level mathematical language: trigonometric functions and other transcendental functions, distribution functions, random number generators, numerical differentiation and integration routines, Fourier transforms, Bessel functions, and polynomial evaluation routines. And, as a matrix programming language, GAUSS includes a variety of routines that perform standard matrix operations. Among these are routines to calculate determinants, matrix inverses, decompositions, eigenvalues and eigenvectors, and condition numbers.

Data handling routines include functions that return dimensions of matrices, and information about elements of data matrices, including functions to locate values lying in specific ranges or with certain values. Also under data handling routines fall all

those functions that create, save, open, and read from and write to GAUSS data sets. A variety of sorting routines that will operate on both numeric and character data are also available.

Programming statements are all of the commands that make it possible to write complex programs in GAUSS. These include conditional and unconditional branching, looping, file I/O, error handling, and system-related commands to execute OS shells and access directory and environment information.

The graphics functions of GAUSS Publication Quality Graphics (PQG) are a set of routines built on the graphics functions in GraphiC by Scientific Endeavors Corporation. GAUSS PQG consists of a set of main graphing procedures and several additional procedures and global variables for customizing the output.

Documentation Conventions

The following table describes how text formatting is used to identify GAUSS programming elements.

Text Style	Use	Example
regular text	narrative	"text formatting is used"
bold text	emphasis	"not supported under UNIX."
italic text	variables	" If <i>vnames</i> is a string or has fewer elements than <i>x</i> has columns, it will be"
monospace	code example	<pre>if scalerr(cm);</pre>
		cm = inv(x);
		endif;
monospace bold	Refers to a GAUSS programming element within a narrative paragraph.	"as explained under create"

Using This Manual

Users who are new to GAUSS should make sure they have familiarized themselves with "Language Fundamentals" in the *User's Guide* before proceeding here. That chapter contains the basics of GAUSS programming.

In all, there are over 400 routines described in this GAUSS Language Reference. We suggest that new GAUSS users skim through Chapter 2, and then browse through Chapter 3, the main part of this manual. Here, users can familiarize themselves with the kinds of tasks that GAUSS can handle easily.

Chapter 2 gives a categorical listing of all functions in this GAUSS Language Reference, and a short discussion of the functions in each category. Complete syntax, description of input and output arguments, and general remarks regarding each function are given in Chapter 3.

If a function is an "extrinsic" (that is, part of the Run-Time Library), its source code can be found on the .src subdirectory. The name of the file containing the source code is given in Chapter 3 under the discussion of that function.

Global Control Variables

Several GAUSS functions use global variables to control various aspects of their performance. The files gauss.ext, gauss.dec, and gauss.lcg contain the **external** statements, **declare** statements, and **library** references to these globals. All globals used by the GAUSS Run-Time library begin with an underscore '_'.

Default values for these common globals can be found in the file gauss.dec, located on the .src subdirectory. The default values can be changed by editing this file.

Changing the Default Values

To permanently change the default setting of a common global, two files need to be edited: gauss.dec and gauss.src.

To change the value of the common global __output from 2 to 0, for example, edit the file gauss.dec and change the statement

```
declare matrix __output = 2;
to read
  declare matrix __output = 0;
```

Also, edit the procedure **gausset**, located in the file gauss.src, and modify the statement

```
__output = 2; similarly.
```

The Procedure gausset

The global variables affect your program, even if you have not set them directly in a particular command file. If you have changed them in a previous run, they will retain their changed values until you exit GAUSS or execute the **new** command.

The procedure **gausset** will reset the Run-Time Library globals to their default values:

gausset;

If your program changes the values of these globals, you can use **gausset** to reset them whenever necessary. **gausset** resets the globals as a whole; you can write your own routine to reset specific ones.

Commands 2 by Category

Mathematical Functions

Scientific Functions

abs Returns absolute value of argument.

arccos Computes inverse cosine.

arcsin Computes inverse sine.

atan Computes inverse tangent.

atan2 Computes angle given a point x,y.besselj Computes Bessel function, first kind.bessely Computes Bessel function, second kind.

cos Computes cosine.

cosh Computes hyperbolic cosine.

digamma Computes the digamma function.

exp Computes the exponential function of x.

gamma Computes gamma function value.

In Computes the natural log of each element.Infact Computes natural log of factorial function.

log Computes the \log_{10} of each element.

pi Returns π .

sin Computes sine.

sinh Computes the hyperbolic sine.

spline1D Computes a smoothing spline for a curve.

spline2D Computes a smoothing spline for a surface.

sqrt Computes the square root of each element.

tan Computes tangent.

tanh Computes hyperbolic tangent trigamma Computes trigamma function.

All trigonometric functions take or return values in radian units.

Differentiation and Integration

gradMT Computes numerical gradient.

gradMTm Computes numerical gradient with mask.

Gradp Computes first derivative of a function.

hessMT Computes numerical Hessian.

hessMTg Computes numerical Hessian using gradient procedure.

hessMTgw Computes numerical Hessian using gradient procedure with

weights.

hessMTm Computes numerical Hessian with mask.

hessMTmw Computes numerical Hessian with mask and weights.

hessMTw Computes numerical Hessian with weights.

Computes second derivative of a function.

intgrat2 Integrates a 2-dimensional function over a user-defined region.intgrat3 Integrates a 3-dimensional function over a user-defined region.

intquad1 Integrates a 1-dimensional function.

intquad2 Integrates a 2-dimensional function over a user-defined

rectangular region.

intquad3 Integrates a 3-dimensional function over a user-defined

rectangular region.

intsimp Integrates by Simpson's method.

gradp and **hessp** use a finite difference approximation to compute the first and second derivatives. Use **gradp** to calculate a Jacobian.

intquad1, **intquad2**, and **intquad3** use Gaussian quadrature to calculate the integral of the user-defined function over a rectangular region.

To calculate an integral over a region defined by functions of x and y, use **intgrat2** and **intgrat3**.

To get a greater degree of accuracy than that provided by **intquad1**, use **intsimp** for 1-dimensional integration.

Linear Algebra

balance	Balances a matrix.
chol	Computes Cholesky decomposition, $X = Y'Y$.
choldn	Performs Cholesky downdate on an upper triangular matrix.
cholsol	Solves a system of equations given the Cholesky factorization of a matrix.
cholup	Performs Cholesky update on an upper triangular matrix.
cond	Computes condition number of a matrix.
crout	Computes Crout decomposition, $X = LU$ (real matrices only).
croutp	Computes Crout decomposition with row pivoting (real matrices only).
det	Computes determinant of square matrix.
detl	Computes determinant of decomposed matrix.
hess	Computes upper Hessenberg form of a matrix (real matrices only).
inv	Inverts a matrix.
invpd	Inverts a positive definite matrix.
invswp	Generalized sweep inverse.
lapeighb	Computes eigenvalues only of a real symmetric or complex Hermitian matrix selected by bounds.
lapeighi	Computes eigenvalues only of a real symmetric or complex Hermitian matrix selected by index.
lapeighvb	Computes eigenvalues and eigenvectors of a real symmetric or complex Hermitian matrix selected by bounds.
lapeighvi	Computes selected eigenvalues and eigenvectors of a real symmetric or complex Hermitian matrix.

lapgeig Computes generalized eigenvalues for a pair of real or complex general matrices. Computes generalized eigenvalues for a pair of real symmetric lapgeigh or Hermitian matrices. Computes generalized eigenvalues and eigenvectors for a pair lapgeighv of real symmetric or Hermitian matrices. lapgeigy Computes generalized eigenvalues, left eigenvectors, and right eigenvectors for a pair of real or complex general matrices. Compute the generalized singular value decomposition of a lapgsvds pair of real or complex general matrices. lapgsvdcst Compute the generalized singular value decomposition of a pair of real or complex general matrices. lapgsvdst Compute the generalized singular value decomposition of a pair of real or complex general matrices. lapschur Compute the generalized Schur form of a pair of real or complex general matrices. Computes the singular value decomposition a real or complex lapsvdcusv rectangular matrix, returns compact u and v. lapsvds Computes the singular values of a real or complex rectangular matrix lapsvdusv Computes the singular value decomposition a real or complex rectangular matrix. 1uComputes LU decomposition with row pivoting (real and complex matrices). null Computes orthonormal basis for right null space. null1 Computes orthonormal basis for right null space. orth Computes orthonormal basis for column space x. Generalized pseudo-inverse: Moore-Penrose. pinv QR decomposition: returns Q_1 and R. qqr QR decomposition: returns Q_1 , R, and a permutation vector, E. ggre QR decomposition with pivot control: returns Q_1 , R, and E. qqrep qr QR decomposition: returns R. QR decomposition: returns R and E. gre QR decomposition with pivot control: returns R and E. grep Solves a system of equations Rx=b given an upper triangular grsol matrix, typically the R matrix from a QR decomposition.

qrtsol	Solves a system of equations $R'x = b$ given an upper triangular matrix, typically the R matrix from a QR decomposition.
qtyr	QR decomposition: returns $Q'Y$ and R .
qtyre	QR decomposition: returns $Q'Y$, R , and E .
qtyrep	QR decomposition with pivot control: returns $Q'Y$, R , and E .
qyr	QR decomposition: returns QY and R .
qyre	QR decomposition: returns QY , R , and E .
qyrep	QR decomposition with pivot control: returns QY , R , and E .
rank	Computes rank of a matrix.
rcondl	Returns reciprocal of the condition number of last decomposed matrix.
rref	Computes reduced row echelon form of a matrix.
schur	Computes Schur decomposition of a matrix (real matrices only).
solpd	Solves a system of positive definite linear equations.
svd	Computes the singular values of a matrix.
svd1	Computes singular value decomposition, $X = USV'$.
svd2	Computes svd1 with compact U .

The decomposition routines are **chol** for Cholesky decomposition, **crout** and **croutp** for Crout decomposition, **qqr-qyrep** for QR decomposition, and **svd**, **svd1**, and **svd2** for singular value decomposition.

null, null1, and orth calculate orthonormal bases.

inv, invpd, solpd, cholsol, qrsol, and the "/" operator can all be used to
solve linear systems of equations.

rank and rref will find the rank and reduced row echelon form of a matrix.

det, **det1**, and **cond** will calculate the determinant and condition number of a matrix.

Eigenvalues

eig	Computes eigenvalues of general matrix.
eigh	Computes eigenvalues of complex Hermitian or real symmetric matrix.

eighv Computes eigenvalues and eigenvectors of complex Hermitian

or real symmetric matrix.

eigv Computes eigenvalues and eigenvectors of general matrix.

There are four eigenvalue-eigenvector routines. Two calculate eigenvalues only, and two calculate eigenvalues and eigenvectors. The types of matrices handled by these routines are:

General: eig, eigv
Symmetric or Hermitian: eigh, eighv

Polynomial Operations

polychar Computes characteristic polynomial of a square matrix.

polyeval Evaluates polynomial with given coefficients.

polyint Calculates N^{th} order polynomial interpolation given known

point pairs.

polymake Computes polynomial coefficients from roots.

polymat Returns sequence powers of a matrix.

polymult Multiplies two polynomials together.

polyroot Computes roots of polynomial from coefficients.

See also recserre, recsercp, and conv.

Fourier Transforms

rfftn

dfft	Computes discrete 1-D FFT.
dffti	Computes inverse discrete 1-D FFT.
fft	Computes 1- or 2-D FFT.
ffti	Computes inverse 1- or 2-D FFT.
fftm	Computes multi-dimensional FFT.
fftmi	Computes inverse multi-dimensional FFT.
fftn	Computes 1- or 2-D FFT using prime factor algorithm.
rfft	Computes real 1- or 2-D FFT.
rffti	Computes inverse real 1- or 2-D FFT.
rfftip	Computes inverse real 1- or 2-D FFT from packed format FFT.

Computes real 1- or 2-D FFT using prime factor algorithm.

rfftnp Computes real 1- or 2-D FFT using prime factor algorithm,

returns packed format FFT.

rfftp Computes real 1- or 2-D FFT, returns packed format FFT.

Random Numbers

rndbeta Computes random numbers with beta distribution.

rndcon Changes constant of the LC random number generator.
rndgam Computes random numbers with gamma distribution.

rndiReturns random integers, $0 \le y < 2^32$.rndKMbetaComputes beta pseudo-random numbers.rndKMgamComputes gamma pseudo-random numbers.rndKMiReturns random integers, $0 \le y < 2^32$.

rndKMnComputes standard normal pseudo-random numbers.rndKMnbComputes negative binomial pseudo-random numbers.

rndKMpComputes Poisson pseudo-random numbers.rndKMuComputes uniform pseudo-random numbers.rndKMvmComputes von Mises pseudo-random numbers.

rndLCbetaComputes beta pseudo-random numbers.rndLCgamComputes gamma pseudo-random numbers.rndLCiReturns random integers, 0 <= y < 2^32.</th>

rndLCnComputes standard normal pseudo-random numbers.rndLCnbComputes negative binomial pseudo-random numbers.

rndLCpComputes Poisson pseudo-random numbers.rndLCuComputes uniform pseudo-random numbers.rndLCvmComputes von Mises pseudo-random numbers.

rndmultChanges multiplier of the LC random number generator.rndnComputes random numbers with Normal distribution.rndnbComputes random numbers with negative binomial

distribution.

rndp Computes random numbers with Poisson distribution.rndseed Changes seed of the LC random number generator.rndu Computes random numbers with uniform distribution.

Fuzzy Conditional Functions

dotfeq	Fuzzy .==
dotfge	Fuzzy .>=
dotfgt	Fuzzy .>
dotfle	Fuzzy .<=
dotflt	Fuzzy .<
dotfne	Fuzzy ./=
feq	Fuzzy ==
fge	Fuzzy >=
fgt	Fuzzy >
fle	Fuzzy <=
flt	Fuzzy <
fne	Fuzzy /=

The global variable **_fcomptol** controls the tolerance used for comparison. By default, this is 1e-15. The default can be changed by editing the file fcompare.dec.

Statistical Functions

acf	Computes sample autocorrelations.
conv	Computes convolution of two vectors.
corrm	Computes correlation matrix of a moment matrix.
corrvc	Computes correlation matrix from a variance-covariance matrix.
corrx	Computes correlation matrix.
crossprd	Computes cross product.
design	Creates a design matrix of 0's and 1's.
dstat	Computes descriptive statistics of a data set or matrix.
loess	Computes coefficients of locally weighted regression.
meanc	Computes mean value of each column of a matrix.
median	Computes medians of the columns of a matrix.
moment	Computes moment matrix $(x'x)$ with special handling of missing values.
momentd	Computes moment matrix from a data set.
ols	Computes least squares regression of data set or matrix.

olsqr Computes OLS coefficients using QR decomposition.

olsqr2 Computes OLS coefficients, residuals, and predicted values

using QR decomposition.

pacf Computes sample partial autocorrelations.

princomp Computes principal components of a data matrix.

stdc Computes standard deviation of the columns of a matrix.

toeplitz Computes Toeplitz matrix from column vector.

varmall Computes the log-likelihood of a Vector ARMA model.

varmares Computes the residuals of a Vector ARMA model.

vcm Computes a variance-covariance matrix from a moment matrix.
vcx Computes a variance-covariance matrix from a data matrix.

Advanced statistics and optimization routines are available in the GAUSS Applications programs. (Contact Aptech Systems for more information.)

Optimization and Solution

eqsolve Solves a system of nonlinear equations.

QNewton Optimizes a function using the BFGS

descent algorithm.

QProg Solves the quadratic programming

problem.

sqpSolve Solves the nonlinear programming

problem using a sequential quadratic

programming method.

sqpSolveMT Solve the nonlinear programming

problem.

sqpSolveMTcontrolCreate Creates an instance of a structure of type

sqpSolveMTcontrol set to default values.

sqpSolveMTlagrangeCreate Creates an instance of a structure of type

sqpSolveMTlagrange set to default values.

sqpSolveMToutCreate Creates an instance of a structure of type

sqpSolveMTout set to default values.

Statistical Distributions

cdfbeta Computes integral of beta function.

cdfbvn Computes lower tail of bivariate Normal cdf.

cdfchic Computes complement of cdf of χ^2 .

cdfchii Computes χ^2 abscissae values given probability and degrees of

freedom.

cdfchinc Computes integral of noncentral χ^2 .

cdffc Computes complement of cdf of *F*. **cdffnc** Computes integral of noncentral *F*.

cdfgam Computes integral of incomplete Γ function.

cdfmvn Computes multivariate Normal cdf.

cdfn Computes integral of Normal distribution: lower tail, or cdf.

cdfn2 Computes interval of Normal cdf.

cdfnc Computes complement of cdf of Normal distribution (upper

tail).

cdftc Computes complement of cdf of *t*-distribution.

cdftnc Computes integral of noncentral *t*-distribution.

cdftvn Computes lower tail of trivariate Normal cdf.

erf Computes Gaussian error function.

computes complement of Gaussian error function.
 lncdfbvn
 lncdfmvn
 Computes natural log of bivariate Normal cdf.
 Computes natural log of multivariate Normal cdf.

lncdfn Computes natural log of Normal cdf.

lncdfn2Computes natural log of interval of Normal cdf.lncdfncComputes natural log of complement of Normal cdf.lnpdfmvnComputes multivariate Normal log-probabilities.

lnpdfn Computes Normal log-probabilities.

pdfn Computes standard Normal probability density function.

Series and Sequence Functions

recserar Computes autoregressive recursive series.

recsercp Computes recursive series involving products.
recserc Computes recursive series involving division.

seqa Creates an additive sequence.

seqm Creates a multiplicative sequence.

Precision Control

base10 Converts number to *x.xxx* and a power of 10.

ceil Rounds up towards $+\infty$.

floor Rounds down towards $-\infty$.

machEpsilon Returns the smallest number such that 1 + eps > 1. **prcsn** Sets computational precision for matrix operations.

round Rounds to the nearest integer.

trunc Truncates toward 0.

All calculations in GAUSS are done in double precision, with the exception of some of the intrinsic functions on OS/2 and DOS. These may use extended precision (18-19 digits of accuracy). Use **presn** to change the internal accuracy used in these cases.

round, trunc, ceil, and floor convert floating point numbers into integers. The internal representation for the converted integer is double precision (64 bits).

Each matrix element in memory requires 8 bytes of memory.

Finance Functions

AmericanBinomCall American binomial method Call.

AmericanBinomCall_Greeks American binomial method call Delta,

Gamma, Theta, Vega, and Rho.

AmericanBinomCall_ImpVol Implied volatilities for American binomial

method calls.

American BinomPut American binomial method Put.

AmericanBinomPut_Greeks American binomial method put Delta,

Gamma, Theta, Vega, and Rho.

AmericanBinomPut ImpVol Implied volatilities for American binomial

method puts.

American Black and Scholes Call.

AmericanBSCall_Greeks American Black and Scholes call Delta,

Gamma, Omega, Theta, and Vega.

AmericanBSCall ImpVol Implied volatilities for American Black

and Scholes calls.

American Black and Scholes Put.

AmericanBSPut_Greeks American Black and Scholes put Delta,

Gamma, Omega, Theta, and Vega.

AmericanBSPut_ImpVol Implied volatilities for American Black

and Scholes puts.

annualTradingDays Compute number of trading days in a

given year.

elapsedTradingDays Compute number of trading days between

two dates inclusively.

EuropeanBinomCall European binomial method call.

EuropeanBinomCall_Greeks European binomial method call Delta,

Gamma, Theta, Vega and Rho.

EuropeanBinomCall_ImpVol Implied volatilities for European binomial

method calls.

EuropeanBinomPut European binomial method Put.

EuropeanBinomPut_Greeks European binomial method put Delta,

Gamma, Theta, Vega, and Rho.

EuropeanBinomPut_ImpVol Implied volatilities for European binomial

method puts.

EuropeanBSCall European Black and Scholes Call.

European Black and Scholes call Delta,

Gamma, Omega, Theta, and Vega.

EuropeanBSCall_ImpVol Implied volatilities for European Black

and Scholes calls.

EuropeanBSPut European Black and Scholes Put.

EuropeanBSPut Greeks European Black and Scholes put Delta,

Gamma, Omega, Theta, and Vega.

EuropeanBSPut_ImpVol Implied volatilities for European Black

and Scholes puts.

getNextTradingDay Returns the next trading day.

getNextWeekDay Returns the next day that is not on a

weekend.

getPreviousTradingDay Returns the previous trading day.

getPreviousWeekDay Returns the previous day that is not on a

weekend.

Matrix Manipulation

Creating Vectors and Matrices

editm Simple matrix editor.

eye Creates identity matrix.

let Creates matrix from list of constants.

matalloc Allocates a matrix with unspecified contents.

matinit Allocates a matrix with unspecified contents.

medit Full-screen spreadsheet-like matrix editor.

ones Creates a matrix of ones.

zeros Creates a matrix of zeros.

Use **zeros** or **ones** to create a constant vector or matrix.

medit is a full-screen editor that can be used to create matrices to be stored in memory, or to edit matrices that already exist.

Matrices can also be loaded from an ASCII file, from a GAUSS matrix file, or from a GAUSS data set. (See "Procedures and Keywords" in the *User Guide* for more information.)

Loading and Storing Matrices

loadd Loads matrix from data set.

loadm Loads matrix from ASCII or matrix file.

save Saves matrix to matrix file.
saved Saves matrix to data set.

Size, Ranking, and Range

cols Returns number of columns in a matrix.

colsf Returns number of columns in an open data set.

counts Returns number of elements of a vector falling in specified

ranges.

countwts Returns weighted count of elements of a vector falling in

specified ranges.

cumprodc Computes cumulative products of each column of a matrix. **cumsumc** Computes cumulative sums of each column of a matrix.

indexcat Returns indices of elements falling within a specified range.

maxc Returns largest element in each column of a matrix.

maxindc Returns row number of largest element in each column of a

matrix.

minc Returns smallest element in each column of a matrix.

minindc Returns row number of smallest element in each column of a

matrix.

prodc Computes the product of each column of a matrix.

rankindx Returns rank index of Nx1 vector. (Rank order of elements in

vector.)

rows Returns number of rows in a matrix.

rowsf Returns number of rows in an open data set.

sumc Computes the sum of each column of a matrix.

These functions are used to find the minimum, maximum and frequency counts of elements in matrices.

Use **rows** and **cols** to find the number of rows or columns in a matrix. Use **rowsf** and **colsf** to find the numbers of rows or columns in an open GAUSS data set.

Sparse Matrix Functions

denseSubmat Returns dense submatrix of sparse matrix.

isSparse Tests whether a matrix is a sparse matrix.

sparseCols Returns number of columns in sparse matrix.

sparseEye Creates sparse identity matrix.

sparseFD Converts dense matrix to sparse matrix.

sparseFP Converts packed matrix to sparse matrix.

sparseHConcat Horizontally concatenates sparse matrices.

sparseNZE Returns the number of nonzero elements in sparse matrix.

sparseOnes Generates sparse matrix of ones and zeros.

sparseRows Returns number of rows in sparse matrix.

sparseSet Resets sparse library globals.

sparseSolve Solves Ax = B for x where A is a sparse matrix.

sparseSubmat Returns sparse submatrix of sparse matrix.

sparseTD Multiplies sparse matrix by dense matrix.

sparseTrTD Multiplies sparse matrix transposed by dense matrix.

sparseVConcat Vertically concatenates sparse matrices.

Miscellaneous Matrix Manipulation

complex Creates a complex matrix from two real matrices.

delif Deletes rows from a matrix using a logical expression.

diag Extracts the diagonal of a matrix.

diagrv Puts a column vector into the diagonal of a matrix.

exctsmpl Creates a random subsample of data set, with replacement.

imag Returns the imaginary part of a complex matrix.

indev Checks one character vector against another and returns the

indices of the elements of the first vector in the second vector.

indnv Checks one numeric vector against another and returns the

indices of the elements of the first vector in the second vector.

indsav Checks one string array against another and returns the indices

of the first string array in the second string array.

intrsect Returns the intersection of two vectors.

lowmat Returns the main diagonal and lower triangle.

lowmat1 Returns a main diagonal of 1's and the lower triangle.

real Returns the real part of a complex matrix.
reshape Reshapes a matrix to new dimensions.

rev Reverses the order of rows of a matrix.

rotater Rotates the rows of a matrix, wrapping elements as necessary.

selif Selects rows from a matrix using a logical expression.

setdif Returns elements of one vector that are not in another.

shifts Shifts rows of a matrix, filling in holes with a specified value.

submat Extracts a submatrix from a matrix.

trimr Trims rows from top or bottom of a matrix.

union Returns the union of two vectors.

upmat Returns the main diagonal and upper triangle.

upmat1 Returns a main diagonal of 1's and the upper triangle.vec Stacks columns of a matrix to form a single column.

vech Reshapes the lower triangular portion of a symmetric matrix

into a column vector.

vecr Stacks rows of a matrix to form a single column.

xpnd Expands a column vector into a symmetric matrix.

vech and **xpnd** are complementary functions. **vech** provides an efficient way to store a symmetric matrix; **xpnd** expands the stored vector back to its original symmetric matrix.

delif and **selif** are complementary functions. **delif** deletes rows of a matrix based on a logical comparison; **selif** selects rows based on a logical comparison.

lowmat, lowmat1, upmat, and upmat1 extract triangular portions of a matrix.

To delete rows that contain missing values from a matrix in memory, see packr.

Structures

dsCreate	Creates an instance of a structure of type DS set to default values.
pvCreate	Returns an initialized an instance of structure of type PV.
pvGetIndex	Gets row indices of a matrix in a parameter vector.
pvGetParNames	Generates names for parameter vector stored in structure of type PV.
pvGetParVector	Retrieves parameter vector from structure of type PV.
pvLength	Returns length of vector p.
pvList	Retrieves names of packed matrices in structure of type PV.
pvPack	Packs general matrix into a structure of type PV with matrix name.
pvPacki	Packs general matrix or array into a PV instance with name and index.
pvPackm	Packs general matrix into a structure of type PV with a mask and matrix name.
pvPackmi	Packs general matrix or array into a PV instance with a mask, name, and index.
pvPacks	Packs symmetric matrix into a structure of type PV.
pvPacksi	Packs symmetric matrix into a PV instance with matrix name and index.
pvPacksm	Packs symmetric matrix into a structure of type PV with a mask.
pvPacksmi	Packs symmetric matrix into a PV instance with a mask, matrix name, and index.

Inserts parameter vector into structure of type PV.

pvPutParVector

PVTest Tests an instance of structure of type PV to determine if it

is a proper structure of type PV.

pvUnpack Unpacks matrices stored in a structure of type PV.

N-Dimensional Array Handling

aconcat Concatenates conformable matrices and arrays in a user-

specified dimension.

amean Computes the mean across one dimension of an N-

dimensional array.

areshape Reshapes a scalar, matrix, or array into an array of user-

specified size.

arrayalloc Creates an N-dimensional array with unspecified contents.

arrayinit Creates an N-dimensional array with a specified fill value.

arraytomat Changes an array to type matrix.

asum Computes the sum across one dimension of an N-

dimensional array.

atranspose Transposes an N-dimensional array.

getarray Gets a contiguous subarray from an N-dimensional array.

getdims Gets the number of dimensions in an array.

getmatrix Gets a contiguous matrix from an N-dimensional array.

Gets a contiguous matrix from a 4-dimensional array.

Gets the vector of orders corresponding to an array.

getscalar3D Gets a scalar from a 3-dimensional array.

Gets a scalar form a 4-dimensional array.

loopnextindex Increments an index vector to the next logical index and

jumps to the specified label if the index did not wrap to the

beginning.

mattoarray Changes a matrix to a type array.

nextindex Returns the index of the next element or subarray in an

array.

previous index Returns the index of the previous element or subarray in an

array.

putarray Puts a contiguous subarray into an N-dimensional array and

returns the resulting array.

setarray Sets a contiguous subarray of an N-dimensional array.

walkindex Walks the index of an array forward or backward through a

specified dimension.

Data Handling (I/O)

Spreadsheets

SpreadsheetReadMReads and writes Excel files.SpreadsheetReadSAReads and writes Excel files.SpreadsheetWriteReads and writes Excel files.

xlsreadm Reads from an Excel spreadsheet, into a GAUSS

matrix.

xlsreadsa Reads from an Excel spreadsheet, into a GAUSS

string array or string.

xlswritem Writes a GAUSS matrix to an Excel spreadsheet. **xlswritesa** Writes a GAUSS string or string array to an Excel

spreadsheet.

Text Files

close Close a GAUSS file.

fcheckerr Gets the error status of a file.

fclearerr Gets the error status of a file, then clears it.

fflush Flushes a file's output buffer.

fgets Reads a line of text from a file.

fgetsa Reads lines of text from a file into a string array.

Reads lines of text from a file into a string array.

fgetst Reads a line of text from a file.

fopen Opens a file.

fputs Writes strings to a file.

fputst Writes strings to a file.

fseek Positions the file pointer in a file.

fstrerror Returns an error message explaining the cause of the most

recent file I/O error.

ftell Gets the position of the file pointer in a file.

Data Sets

close Closes an open data set (.dat file).

closeall Closes all open data sets.create Creates and opens a data set.

eof Tests for end of file.

iscplxf Returns whether a data set is real or complex.

Loadd Loads a small data set.open Opens an existing data set.readr Reads rows from open data set.

saved Creates small data sets.

seekr Moves pointer to specified location in open data set.

typef Returns the element size (2, 4, or 8 bytes) of data in open data

set.

writer Writes matrix to an open data set.

These functions all operate on GAUSS data sets (.dat files). (See "File I/O" in the *User's Guide* for more information.)

To create a GAUSS data set from a matrix in memory, use **saved**. To create a data set from an existing one, use **create**. To create a data set from a large ASCII file, use the utility **atog**. (See "Utilities" in the *User's Guide*.)

Data sets can be opened, read from, and written to using open, readr, seekr and writer. Test for the end of a file using eof, and close the data set using close or closeall.

The data in data sets may be specified as character or numeric. (See "File I/O" in the *User's Guide*.) See also **create** and **vartypef**.

typef returns the element size of the data in an open data set.

Data Set Variable Names

getname	Returns column vector of variable names in a data set.
getnamef	Returns string array of variable names in a data set.
indcv	Returns column numbers of variables within a data set.
indices	Retrieves column numbers and names from a data set.
indices2	Similar to indices , but matches columns with names for dependent and independent variables.
makevars	Decomposes matrix to create column vectors.
mergevar	Concatenates column vectors to create larger matrix.
setvars	Creates globals using the names in a data set.
vartype	Returns column vector of variable types (numeric/character) in a data set.
vartypef	Returns column vector of variable types (numeric/character) in a data set.

Use **getnamef** to retrieve the variable names associated with the columns of a GAUSS data set, and **vartypef** to retrieve the variable types. Use **makevars** and **setvars** to create global vectors from those names. Use **indices** and **indices2** to match names with column numbers in a data set.

getname and **vartype** are supported for backwards compatibility.

Data Coding

code	Codes the data in a vector by applying a logical set of rules to assign each data value to a category.
dummy	Creates a dummy matrix, expanding values in vector to rows with ones in columns corresponding to true categories and zeros elsewhere.

dummybr Similar to dummy.
dummydn Similar to dummy.

ismiss Returns 1 if matrix has any missing values, 0 otherwise.
isinfnanmiss Returns true if the argument contains an infinity, NaN, or

missing value.

miss Changes specified values to missing value code.

missex Changes elements to missing value using logical expression.

missrv Changes missing value codes to specified values.

msym Sets symbol to be interpreted as missing value.

packr Deletes rows with missing values.

recode Similar to code, but leaves the original data in place if no

condition is met.

scalinfnanmis Returns true if the argument is a scalar infinity, NaN, or

S

missing value.

scalmiss Tests whether a scalar is the missing value code.

subscat Simpler version of recode, but uses ascending bins instead

of logical conditions.

substute Similar to **recode**, but operates on matrices.

code, **recode**, and **subscat** allow the user to code data variables and operate on vectors in memory. **substute** operates on matrices, and **dummy**, **dummybr**, and **dummydn** create matrices.

missex, missrv, and miss should be used to recode missing values.

Sorting and Merging

intrleav Produces one large sorted data file from two smaller sorted files

having the same keys.

mergeby Produces one large sorted data file from two smaller sorted files

having a single key column in common.

sortc Quick-sorts rows of matrix based on numeric key.sortcc Quick-sorts rows of matrix based on character key.

sortd Sorts data set on a key column.

sorthc Heap-sorts rows of matrix based on numeric key.

Heap-sortsrows of matrix based on character key.

sortind Returns a sorted index of a numeric vector.

sortindc Returns a sorted index of a character vector.

sortmc Sorts rows of matrix on the basis of multiple columns.

unique Returns a sorted unique index of a vector.

Removes duplicate elements of a vector.

sortc, **sorthc**, and **sortind** operate on numeric data only. **sortcc**, **sorthcc**, and **sortindc** operate on character data only.

Sortd, sortmc, unique, and uniqindx operate on both numeric and character data.

Use **sortd** to sort the rows of a data set on the basis of a key column.

Both intrleav and mergeby operate on data sets.

Compiler Control

#define	Defines a case-insensitive text-replacement or flag variable.	
#definecs	Defines a case-sensitive text-replacement or flag variable.	
#else	Alternates clause for #if-#else-#endif code block.	
#endif	End of #if-#else-#endif code block.	
#ifdef	Compiles code block if a variable has been #define'd.	
#ifdos	Compiles code block if running DOS.	
#iflight	Compiles code block if running GAUSS Light.	
#ifndef	Compiles code block if a variable has not been #define'd.	
#ifos2win	Compiles code block if running OS/2 or Windows.	
#ifunix	Compiles code block if running UNIX.	
#include	Includes code from another file in program.	
#linesoff	Compiles program without line number and file name records	
#lineson	Compiles program with line number and file name records.	
#srcfile	Inserts source file name record at this point (currently used when doing data loop translation).	
#srcline	Inserts source file line number record at this point (currently used when doing data loop translation).	
#undef	Undefines a text-replacement or flag variable.	

These commands are compiler directives. That is, they do not generate GAUSS program instructions; rather, they are instructions that tell GAUSS how to process a program during compilation. They determine what the final compiled form of a

program will be. They are not executable statements and have no effect at run-time. (See "Language Fundamentals" in the *User's Guide* for more information.)

Program Control

Execution Control

Terminates a program and close all files.
Pauses for the specified time.
Runs a program in a text file.
Sleeps for the specified time.
Stops a program and leave files open.
Quits and returns to the OS.

Both stop and end will terminate the execution of a program; end will close all open files, and stop will leave those files open. Neither stop nor end is required in a GAUSS program.

Branching

```
pop errmsg;
print errmsg;
end;
```

Looping

```
break
                             Jump out the bottom of a do or for loop.
            continue
                             Jump to the top of a do or for loop.
            do while .. endo
                        Loop if TRUE.
                              Loop if FALSE.
            do until..endo
                              Loop with integer counter.
            for.. endfor
            iter = 0;
            do while dif > tol;
               \{x,x0\} = eval(x,x0);
               dif = abs(x-x0);
               iter = iter + 1;
               if iter > maxits;
                  break;
               endif;
               if not prtiter;
                  continue;
               endif;
               format /rdn 1,0;
               print "Iteration: " iter;
               format /re 16,8;
print ", Error: Files needed to be included at the top of
               programs that use the function. "maxc(dif);
            endo;
```

```
for i (1, cols(x), 1);
  for j (1, rows(x), 1);
    x[i,j] = x[i,j] + 1;
  endfor;
endfor;
```

Subroutines

gosub Branch to subroutine.

pop Retrieve gosub arguments.

return Return from subroutine.

Arguments can be passed to subroutines in the branch to the subroutine label and then popped, in first-in-last-out order, immediately following the subroutine label definition. See Chapter 3, "Command Reference", for details.

Arguments can then be returned in an analogous fashion through the **return** statement.

Procedures

endp Terminates a procedure definition.
 local Declares variables local to a procedure.
 proc Begins definition of multi-line procedure.
 retp Returns from a procedure.

Here is an example of a GAUSS procedure:

```
proc (3) = crosprod(x,y);
  local r1, r2, r3;
  r1 = x[2,.].*y[3,.]-x[3,.].*y[2,.];
  r2 = x[3,.].*y[1,.]-x[1,.].*y[3,.];
  r3 = x[1,.].*y[2,.]-x[2,.].*y[1,.];
  retp( r1,r2,r3 );
endp;
```

The "(3) = " indicates that the procedure returns three arguments. All local variables, except those listed in the argument list, must appear in the local statement. Procedures may reference global variables. There may be more than one retp per procedure definition; none is required if the procedure is defined to return 0 arguments. The endp is always necessary and must appear at the end of the procedure definition. Procedure definitions cannot be nested. The syntax for using this example function is

```
{a1,a2,a3} = crosprod(u,v);
```

See "Procedures and Keywords" and "Libraries" in the *User's Guide* for details.

Libraries

call Calls function and discard return values.
 declare Initializes variables at compile time.
 external symbol definitions.
 lib Builds or updates a GAUSS library.
 library Sets up list of active libraries.

call allows functions to be called when return values are not needed. This is especially useful if a function produces printed output (**dstat**, **ols** for example) as well as return values.

Compiling

compile	Compiles and saves a program to a.	gcg file.
---------	------------------------------------	-----------

loadp Loads compiled procedure.

save Saves the compiled image of a procedure to disk.saveall Saves the contents of the current workspace to a file.

use Loads previously compiled code.

GAUSS procedures and programs may be compiled to disk files. By then using this compiled code, the time necessary to compile programs from scratch is eliminated. Use **compile** to compile a command file. All procedures, matrices and strings referenced by that program will be compiled as well.

Stand-alone applications may be created by running compiled code under the Run-Time Module. (Contact Aptech Systems for more information on this product.)

To save the compiled images of procedures that do not make any global references, use **save**. This will create an .fcg file. To load the compiled procedure into memory, use **loadp**. (This is not recommended because of the restriction on global

references and the need to explicitly load the procedure in each program that references it. It is included here to maintain backward compatibility with previous versions.)

OS Functions

cdir Returns current directory.

ChangeDir Changes directory in program.

chdir Changes directory interactively.

DeleteFile Deletes files.

dfree Returns free space on disk.
envget Gets an environment string.

exec Executes an executable program file.

fileinfo Takes a file specification, returns names and information of

files that match.

Takes a file specification, returns names of files that match.

Takes a file specification, returns names of files that match.

shell Shells to OS.

Workspace Management

clear Sets matrices equal to 0.
clearg Sets global symbols to 0.

delete Deletes specified global symbols.

hasimagExamines matrix for nonzero imaginary part.iscplxReturns whether a matrix is real or complex.

maxvec Returns maximum allowed vector size.

new Clears current workspace.show Displays global symbol table.

type Returns types of argument (matrix or string).

typecv Returns types of symbol (argument contains the names of the

symbols to be checked).

When working with limited workspace, it is a good idea to **clear** large matrices that are no longer needed by your program.

coreleft is most commonly used to determine how many rows of a data set may be read into memory at one time.

Error Handling and Debugging

#linesoff Omits line number and file name records from program.
#lineson Includes line number and file name records in program.
debug Executes a program under the source level debugger.
disable Disabls invalid operation interrupt of coprocessor.
enable Enables invalid operation interrupt of coprocessor.

error Creates user-defined error code.

errorlogSendserror message to screen and log file.ndpchkExamines status word of coprocessor.ndpclexClears coprocessor exception flags.ndpcntrlSets and gets coprocessor control word.

scalerr Tests for a scalar error code.

trace Traces program execution for debugging.
trap Controls trapping of program errors.

trapchk Examines the trap flag.

To trace the execution of a program, use **trace**.

User-defined error codes may be generated using **error**.

String Handling

chrs Converts ASCII values to a string.

ftocv Converts an NxK matrix to a character matrix.

ftos Converts a floating point scalar to string.

ftostrC Converts a matrix to a string array using a C language format

specification.

getf Loads ASCII or binary file into string.

loads Loads a string file (.fst file).lower Converts a string to lowercase.putf Writes a string to disk file.

stof Converts a string to floating point numbers.

strindx Finds starting location of one string in another string.

strlen Returns length of a string.

strrindx Finds starting location of one string in another string, searching

from the end to the start of the string.

strsect Extracts a substring of a string.

strsplit Splits an Nx1 string vector into an NxK string array of the

individual tokens.

strsplitPad Splits a string vector into a string array of the individual tokens.

Pads on the right with null strings.

strtof Converts a string array to a numeric matrix.

strtofcplx Converts a string array to a complex numeric matrix.

upper Changes a string to uppercase.vals Converts a string to ASCII values.

varget Accesses the global variable named by a string.

varget1 Accesses the local variable named by a string.

varput Assigns a global variable named by a string.

varput1 Assigns a local variable named by a string.

strlen, strindx, strrindx, and strsect can be used together to parse strings.

Use ftos to print to a string.

To create a list of generic variable names (X1, X2, X3, X4... for example), use **ftocv**.

Time and Date Functions

date Returns current system date.

datestr Formats date as "mm/dd/yy".

datestring Formats date as "mm/dd/yyyy".

datestrymd Formats date as "yyyymmdd".

Returns day number of a date.

dayofweek Returns day of week.

dtdate Creates a matrix in DT scalar format.

dtday Creates a matrix in DT scalar format containing only the year,

month and day. Time of day information is zeroed out.

dttime	Creates a matrix in DT scalar format containing only the hour,	
--------	--	--

minute and second. The date information is zeroed out.

dttodtv Converts DT scalar format to DTV vector format.

dttostr Converts a matrix containing dates in DT scalar format to a

string array.

dttoutc Converts DT scalar format to UTC scalar format.

dtvnormal Normalizes a date and time (DTV) vector.

dtvtodt Converts DT vector format to DT scalar format.

etdays Difference between two times in days.

ethsec Difference between two times in 100ths of a second.

etstr Converts elapsed time to string.

hsec Returns elapsed time since midnight in 100ths of a second.

strtodt Converts a string array of dates to a matrix in DT scalar format.

time Returns current system time.

timedt Returns system date and time in DT scalar format.

timestr Formats time as "hh:mm:ss".

todaydt Returns system date in DT scalar format. The time returned is

always midnight (00:00:00), the beginning of the returned day.

utctodt Converts UTC scalar format to DT scalar format.

utctodtv Converts UTC scalar format to DTV vector format.

Use hsec to time segments of code. For example,

et = hsec; x = y*y; et = hsec - et;

will time the GAUSS multiplication operator.

Console I/O

con Requests console input, create matrix.cons Requests console input, create string.

key Gets the next key from the keyboard buffer. If buffer is empty,

returns a 0.

keyav Check if keystroke is available.

keyw Gets the next key from the keyboard buffer. If buffer is empty,

waits for a key.

wait Waits for a keystroke.

waitc Flushes buffer, then waits for a keystroke.

key can be used to trap most keystrokes. For example, the following loop will trap the ALT-H key combination:

```
kk = 0;
do until kk == 1035;
    kk = key;
endo;
```

Other key combinations, function keys, and cursor key movement can also be trapped. See **key**.

cons and con can be used to request information from the console. keyw, wait, and waitc will wait for a keystroke.

Output Functions

cls

Text Output

color	Sets pixel, text, background colors.
comlog	Controls interactive command logging.
csrcol	Gets column position of cursor on window.
csrlin	Gets row position of cursor on window.
ed	Accesses an alternate editor.

Clears the window.

Edits a file with the GAUSS editor.

Format Defines format of matrix printing.

Positions the cursor on the window.

1pos Returns print head position in printer buffer.

lprint Prints expression to the printer.

lprint [[Switches auto printer mode on and off.
on|off]]

lpwidth Specifies printer width.

1show Prints global symbol table on the printer.

output Redirects print statements to auxiliary output.

outwidth Sets line width of auxiliary output.

Plots elements of two matrices in text mode.

plotsym Controls data symbol used by plot.

print Prints to window.

print [[Turns auto window print on and off.

on off]]

printdos Prints a string for special handling by the OS.

printfm Prints matrices using a different format for each

column.

screen [[Directs/suppresses **print** statements to window.

on|off]]

screen out Dumps snapshot of window to auxiliary output.

scroll Scrolls a section of the window.

Positions the cursor on the current line.

The results of all printing can be sent to an output file using **output**. This file can then be printed or ported as an ASCII file to other software.

printdos can be used to print in reverse video, or using different colors. It requires that ansi.sys be installed.

To produce boxes, etc. using characters from the extended ASCII set, use chrs.

Window Graphics

color Sets color.graph Sets pixels.line Draws lines.

setvmode Sets video mode.

graph allows the user to plot individual pixels.

Graphics

This section summarizes all procedures and global variables available within the Publication Quality Graphics (PQG) System. A general usage description will be found in "Publications Quality Graphics" in the *User's Guide*.

Graph Types

bar Generates bar graph.

box Graphs data using the box graph percentile method.

contour Graphs contour data.

draw Supplies additional graphic elements to graphs.

hist Computes and graphs frequency histogram.

histf Graphs a histogram given a vector of frequency counts.

histp Graphs a percent frequency histogram of a vector.

loglog Graphs X,Y using logarithmic X and Y axes.

logx Graphs X,Y using logarithmic X axis.

Graphs X,Y using logarithmic Y axis.

surface Graphs a 3-D surface.

xy Graphs X,Y using Cartesian coordinate system.

xyz Graphs X, Y, Z using 3-D Cartesian coordinate system.

Axes Control and Scaling

_pxpmax Controls precision of numbers on X axis.

paxes Turns axes on or off.

_pcross Controls where axes intersect.

_pgrid Controls major and minor grid lines.
_pticout Controls direction of tick marks on axes.
_pxsci Controls use of scientific notation on X axis.
_pypmax Controls precision of numbers on Y axis.
_pysci Controls use of scientific notation on Y axis.
_pzpmax Controls precision of numbers on Z axis.
_pzsci Controls use of scientific notation on Z axis.

scale Scales X,Y axes for 2-D plots.

scale3d Scales X,Y, and Z axes for 3-D plots.

xtics Scales X axis and control tick marks.

ytics Scales Y axis and control tick marks.

ztics Scales Z axis and control tick marks.

Text, Labels, Titles, and Fonts

pnumht Controls size of axes numeric labels.

_ptitlht Controls main title size.

paxht Controls size of axes labels.

_pdate Dates string contents and control.
_plegctl Sets location and size of plot legend.

_plegstr Specifies legend text entries.
_pmsgctl Controls message position.
_pmsgstr Specifies message text.

_pnum Axes numeric label control and orientation.

asclabel Defines character labels for tick marks.

fonts Loads fonts for labels, titles, messages and legend.

title Specifies main title for graph.

xlabelX axis label.ylabelY axis label.zlabelZ axis label.

Main Curve Lines and Symbols

_pboxctl Controls box plotter.

_pboxlim Outputs percentile matrix from box plotter.

_pcolor Controls line color for main curves.

_plctrl Controls main curve and frequency of data symbols.

_pltype Controls line style for main curves.

_plwidth Controls line thickness for main curves.
_pstype Controls symbol type for main curves.
psymsiz Controls symbol size for main curves.

_pzclr Z level color control for contour and surface.

Extra Lines and Symbols

_parrow Creates arrows.

_parrow3 Creates arrows for 3-D graphs.

_perrbar Plots error bars.

_pline Plots extra lines and circles.

_pline3d Plots extra lines for 3-D graphs.

_psym Plots extra symbols.

_psym3d Plots extra symbols for 3-D graphs.

Graphic Panel, Page, and Plot Control

_pageshf Shifts the graph for printer output.

_pagesiz Controls size of graph for printer output.

_plotshf Controls plot area position.
_plotsiz Controls plot area size.

_protate Rotates the graph 90 degrees.

axmargin Controls axes margins and plot size. **begwind** Graphic panel initialization procedure.

endwind End graphic panel manipulation, display graphs.

getwind Gets current graphic panel number.

loadwind Loads a graphic panel configuration from a file.

makewind Creates graphic panel with specified size and position.

margin Controls graph margins.

nextwindSets to next available graphic panel number.savewindSaves graphic panel configuration to a file.setwindSets to specified graphic panel number.windowCreates tiled graphic panels of equal size.

axmargin is preferred to the older **_plotsiz** and **_plotshf** globals for establishing an absolute plot size and position.

Output Options

_pscreen Controls graphics output to window.

_psilent Controls final beep.

_ptek Controls creation and name of graphics.tkf file.

_pzoom Specifies zoom parameters.
graphprt Generates print, conversion file.
pqgwin Sets the graphics viewer mode.
setvwrmode Sets the graphics viewer mode.

tkf2eps Converts .tkf file to Encapsulated PostScript file.

tkf2ps Converts .tkf file to PostScript file.

Miscellaneous

_pbox Draws a border around graphic panel/window.

_pcrop Controls cropping of graphics data outside axes area.

_pframe Draws a frame around 2-D, 3-D plots.

_pmcolor Controls colors to be used for axes, title, x and y labels,

date, box, and background.

graphset Resets all PQG globals to default values.
rerun Displays most recently created graph.

viewSets 3-D observer position in workbox units.viewxyzSets 3-D observer position in plot coordinates.

volume Sets length, width, and height ratios of 3-D workbox.

Command Reference

Command Components

The following list describes each of the components used in the GAUSS Language Command Reference.

Purpose Describes what the command or function does.

Library Lists the library that needs to be activated to access

the function.

Format Illustrates the syntax of the command or function.

Input Describes the input parameters of the function.

Global Input Describes the global variables that are referenced by

the function.

Output Describes the return values of the function.

Global Describes the global variables that are updated by

Output the function.

Command Reference

Portability Describes differences under various operating

systems.

Remarks Explanatory material pertinent to the command.

Example Sample code using the command or function.

Source The source file in which the function is defined, if

applicable.

Globals Global variables that are accessed by the command.

See also Other related commands.

Technical Technical discussion and reference source citations. **Notes**

References Reference material citations.

3-2

a

0

d

е

i

j

K

a

1.

S

L

X V Z

abs

abs

Purpose Returns the absolute value or complex modulus of x.

Format y = abs(x);

Input x NxK matrix.

Output y NxK matrix containing absolute values of x.

Example x = rndn(2,2);

y = abs(x);

 $x = \begin{array}{c} 0.675243 & 1.053485 \\ -0.190746 & -1.229539 \end{array}$

 $y = \begin{array}{c} 0.675243 & 1.053485 \\ 0.190746 & 1.229539 \end{array}$

In this example, a 2x2 matrix of Normal random numbers is generated and the absolute value of the matrix is computed.

a

b

c d

A

f g

h

i

k

m

n

0

p q

r

S

t

u

V

W

хух

acf

a

b

С

d

е

g

h

k

m

n

0

p

q

S

t

u

V

W

acf

```
Purpose
             Computes sample autocorrelations.
 Format
             rk = acf(y,k,d);
    Input
                    Nx1 vector, data.
             k
                    scalar, maximum number of autocorrelations to compute.
                    scalar, order of differencing.
             d
  Output
                    Kx1 vector, sample autocorrelations.
             rk
Example
             x = \{20.80,
                    18.58,
                    23.39,
                    20.47,
                    21.78,
                    19.56,
                    19.58,
                    18.91,
                    20.08,
                    21.88 };
             rk = acf(x,4,2);
             print rk;
                    -0.74911771
                     0.48360914
                    -0.34229330
                     0.17461180
  Source
             tsutil.src
```

хух

aconcat

a

b

C

d

f

g

h

m

n

0

q

V

хух

aconcat

Purpose	Concatenates conformable matrices and arrays in a user-specified dimension.	
Format	y = aconcat(a,b,dim);	
Input	 a matrix or N-dimensional array. b matrix or K-dimensional array, conformable with a. dim scalar, dimension in which to concatenate. 	
Output	y M-dimensional array, the result of the concatenation.	
Remarks	a and b are conformable only if all of their dimensions except dim have the same sizes. If a or b is a matrix, then the size of dimension 1 is the number of columns in the matrix, and the size of dimension 2 is the number of rows in the matrix.	
Example	a = arrayinit(2 3 4,0);	
	b = 3*ones(3,4);	
	<pre>y = aconcat(a,b,3);</pre>	
	y will be a $3x3x4$ array, where [1,1,1] through [2,3,4] are zeros and [3,1,1] through [3,2,4] are threes.	
	<pre>a = reshape(seqa(1,1,20),4,5);</pre>	
	b = zeros(4,5);	
	y = aconcat(a,b,3);	
	y will be a 2x4x5 array, where [1,1,1] through [1,4,5] are sequential integers beginning with 1, and [2,1,1] through [2,4,5] are zeros.	
	a = arrayinit(2 3 4,0);	
	b = seqa(1,1,24);	

b = areshape(b, 2 | 3 | 4);

y = aconcat(a,b,5);

aconcat

a

b

d

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q

y will be a 2x1x2x3x4 array, where [1,1,1,1,1] through [1,1,2,3,4] are zeros, and [2,1,1,1,1] through [2,1,2,3,4] are sequential integers beginning with 1.

$$a = arrayinit(2|3|4,0);$$

$$b = seqa(1,1,6);$$

$$b = areshape(b, 2|3|1);$$

$$y = aconcat(a,b,1);$$

y will be a 2x3x5 array, such that:

$$[1,1,1]$$
 through $[1,3,5] =$

$$[2,1,1]$$
 through $[2,3,5] =$

$$0\ 0\ 0\ 0\ 6$$

See also areshape

w x y z

u

V

amean

amean

Purpose Computes the mean across one dimension of an N-dimensional array.

Format y = amean(x, dim);

Input x N-dimensional array.

dim scalar, number of dimension to compute the mean across.

Output y [N-1]-dimensional array.

Remarks The output y, will be have the same sizes of dimensions as x, except that the dimension indicated by dim will be collapsed to 1.

Example x = seqa(1,1,24);

x = areshape(x, 2 | 3 | 4);

y = amean(x,3);

x is a 2x3x4 array, such that:

[1,1,1] through [1,3,4] =

1 2 3 4

5 6 7 8

9 10 11 12

[2,1,1] through [2,3,4] =

13 14 15 16

17 18 19 20

21 22 23 24

y will be a 1x3x4 array, such that:

[1,1,1] through [1,3,4] =

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$$y = amean(x,1);$$

Using the same array x as the above example, this example computes the mean across the first dimension. y will be a 2x3x1 array, such that:

2.5

6.5

10.5

14.5

18.5

22.5

See also

asum

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W X y Z

AmericanBinomCall

AmericanBinomCall

```
American binomial method Call.
Purpose
  Format
             c = AmericanBinomCall(SO, K, r, div, tau, sigma, N);
    Input
                    scalar, current price
              SO.
              K
                    Mx1 vector, strike prices
              r
                    scalar, risk free rate
                    continuous dividend yield
              div
                    scalar, elapsed time to exercise in annualized days of
              tau
                    trading
             sigma scalar, volatility
              N
                    number of time segments
  Output
                    Mx1 vector, call premiums
Example
              S0 = 718.46;
             K = \{ 720, 725, 730 \};
              r = .0498;
              sigma = .2493;
              t0 = dtday(2001, 1, 30);
              t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              c = AmericanBinomCall(S0,K,r,0,tau,sigma,60);
              print c;
              17.190224
              14.905054
              12.673322
```

AmericanBinomCall

Source

finprocs.src

Technical Notes

The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is

the basis of this procedure.

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AmericanBinomCall Greeks

AmericanBinomCall_Greeks

American binomial method call Delta, Gamma, Theta, Vega, and Rho. **Purpose Format** $\{d,g,t,v,rh\} =$ AmericanBinomCall Greeks(SO, K, r, div, tau, sigma, N); Input scalar, current price *S0* K Mx1 vector, strike price scalar, risk free rate r continuous dividend yield div scalar, elapsed time to exercise in annualized days of tau trading sigma scalar, volatility N number of time segments **Output** d Mx1 vector, delta Mx1 vector, gamma g Mx1 vector, theta Mx1 vector, vega rhMx1 vector, rho **Example** S0 = 305;K = 300;r = .08;sigma = .25;tau = .33;div = 0;print AmericanBinomcall_Greeks (S0,K,r,0,tau,sigma,30); 0.706312 0.000764

AmericanBinomCall Greeks

-17.400851

68.703849

76.691829

Source finprocs.src

Globals __fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal.

Default = 0.

_fin_epsilon scalar, finite difference stepsize. Default = 1e-8.

Technical Notes The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is the basis of this procedure.

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V

x y z

AmericanBinomCall_ImpVol

```
Implied volatilities for American binomial method calls.
Purpose
 Format
             sigma =
             AmericanBinomCall ImpVol(c,S0,K,r,div,tau,N);
    Input
                    Mx1 vector, call premiums
             c
             SO
                    scalar, current price
                    Mx1 vector, strike prices
             K
                    scalar, risk free rate
              r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
              tau
                    trading
             N
                    number of time segments
  Output
             sigma Mx1 vector, volatility
Example
             c = \{ 13.70, 11.90, 9.10 \};
              S0 = 718.46;
             K = \{ 720, 725, 730 \};
             r = .0498;
             div = 0;
              t0 = dtday(2001, 1, 30);
              t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              sigma = AmericanBinomCall_ImpVol
                    (c,S0,K,r,0,tau,30);
             print sigma;
              0.1981
```

AmericanBinomCall_ImpVol

0.1715

0.1301

Source

finprocs.src

Technical Notes

The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is the basis of this procedure.

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AmericanBinomPut

AmericanBinomPut

```
American binomial method Put.
 Purpose
  Format
              c = AmericanBinomPut(SO, K, r, div, tau, sigma, N);
    Input
                     scalar, current price
              SO.
              K
                     Mx1 vector, strike prices
              r
                     scalar, risk free rate
                     continuous dividend yield
              div
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma scalar, volatility
              Ν
                     number of time segments
  Output
                     Mx1 vector, put premiums
Example:
              S0 = 718.46;
              K = \{ 720, 725, 730 \};
              r = .0498;
              sigma = .2493;
              t0 = dtday(2001, 1, 30);
              t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                     annualTradingDays(2001);
              c = AmericanBinomPut(S0,K,r,0,tau,sigma,60);
              print c;
              16.862683
              19.606573
              22.433590
```

AmericanBinomPut

Source

finprocs.src

Technical Notes

The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is the basis of this procedure.

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AmericanBinomPut Greeks

AmericanBinomPut_Greeks

American binomial method put Delta, Gamma, Theta, Vega, and Rho. **Purpose Format** $\{d,g,t,v,rh\} =$ AmericanBinomPut Greeks(SO, K, r, div, tau, sigma, N); Input scalar, current price *S0* K Mx1 vector, strike price scalar, risk free rate r continuous dividend yield div scalar, elapsed time to exercise in annualized days of tau trading sigma scalar, volatility N number of time segments **Output** d Mx1 vector, delta Mx1 vector, gamma g Mx1 vector, theta t Mx1 vector, vega rhMx1 vector, rho **Example** S0 = 305;K = 300;r = .08;div = 0;sigma = .25;tau = .33;print AmericanBinomPut_Greeks (S), K, r, 0, tau, sigma, 60); -0.38324908 0.00076381912

AmericanBinomPut Greeks

8.1336630

68.337294

-27.585043

Source finprocs.src

Globals __fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal.

Default = 0.

_fin_epsilon scalar, finite difference stepsize. Default = 1e-8.

Technical Notes The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is

the basis of this procedure.

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AmericanBinomPut ImpVol

AmericanBinomPut_ImpVol

```
Implied volatilities for American binomial method puts.
Purpose
  Format
             sigma = AmericanBinomPut ImpVol(c,S0,K,r,div,tau,N);
    Input
                    Mx1 vector, put premiums
             c
              S0
                    scalar, current price
              K
                    Mx1 vector, strike prices
                    scalar, risk free rate
              r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
              tau
                    trading
             N
                    number of time segments
  Output
             sigma Mx1 vector, volatility
             p = \{ 14.60, 17.10, 20.10 \};
Example
             S0 = 718.46;
             K = \{ 720, 725, 730 \};
             r = .0498;
             div = 0;
              t0 = dtday(2001, 1, 30);
              t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              sigma = AmericanBinomPut ImpVol
                    (p,S0,K,r,0,tau,30);
             print sigma;
              0.1254
              0.1668
```

AmericanBinomPut_ImpVol

0.2134

Source

finprocs.src

Technical Notes The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is

the basis of this procedure.

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AmericanBSCall

AmericanBSCall

```
American Black and Scholes Call.
Purpose
  Format
             c = AmericanBSCall(SO, K, r, div, tau, sigma);
    Input
                    scalar, current price
             SO.
              K
                    Mx1 vector, strike prices
              r
                    scalar, risk free rate
                    continuous dividend yield
             div
                    scalar, elapsed time to exercise in annualized days of
             tau
                    trading
             sigma scalar, volatility
 Output:
                    Mx1 vector, call premiums
Example
             S0 = 718.46;
             K = \{ 720, 725, 730 \};
              r = .0498;
              sigma = .2493;
              t0 = dtday(2001, 1, 30);
              t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              c = AmericanBSCall(S0,K,r,0,tau,sigma);
             print c;
              16.093640
              13.846830
              11.829059
  Source
              finprocs.src
```

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AmericanBSCall Greeks

AmericanBSCall_Greeks

```
Purpose
              American Black and Scholes call Delta, Gamma, Omega, Theta, and
              Vega.
  Format
              \{d,g,t,v,rh\} =
              AmericanBSCall_Greeks(SO,K,r,div,tau,sigma);
    Input
                     scalar, current price
              S0
              K
                     Mx1 vector, strike price
              r
                     scalar, risk free rate
              div
                     continuous dividend yield
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma scalar, volatility
  Output
              d
                     Mx1 vector, delta
                     Mx1 vector, gamma
              g
              t
                     Mx1 vector, theta
                     Mx1 vector, vega
              ν
              rh
                     Mx1 vector, rho
Example
              S0 = 305;
              K = 300;
              r = .08;
              sigma = .25;
              tau = .33;
              print AmericanBSCall Greeks (S0,K,r,0,tau,sigma);
              0.40034039
              0.016804021
              -55.731079
              115.36906
              46.374528
```

AmericanBSCall_Greeks

Source finprocs.src

Globals __fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal.

Default=0.

_fin_epsilon scalar, finite difference stepsize. Default = 1e-8.

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AmericanBSCall ImpVol

AmericanBSCall_ImpVol

```
Implied volatilities for American Black and Scholes calls.
Purpose
  Format
             sigma = AmericanBSCall ImpVol(c,S0,K,r,div,tau);
    Input
                    Mx1 vector, call premiums
              SO.
                    scalar, current price
              K
                    Mx1 vector, strike prices
                    scalar, risk free rate
              r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
              tau
                    trading
  Output
             sigma Mx1 vector, volatility
Example
             c = \{ 13.70, 11.90, 9.10 \};
              S0 = 718.46i
             K = \{ 720, 725, 730 \};
              r = .0498;
              t0 = dtday(2001, 1, 30);
              t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              sigma = AmericanBSCall ImpVol
                    (c,S0,K,r,0,tau);
             print sigma;
              0.10350708
              0.089202881
              0.066876221
```

AmericanBSCall_ImpVol

Source finprocs.src

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AmericanBSPut

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AmericanBSPut

```
Purpose
              American Black and Scholes Put.
  Format
             c = AmericanBSPut(SO, K, r, div, tau, sigma);
    Input
                    scalar, current price
             SO.
              K
                    Mx1 vector, strike prices
                    scalar, risk free rate
              r
                    continuous dividend yield
             div
                    scalar, elapsed time to exercise in annualized days of
             tau
                    trading
             sigma scalar, volatility
  Output
                    Mx1 vector, put premiums
Example
             S0 = 718.46;
             K = \{ 720, 725, 730 \};
              r = .0498;
              sigma = .2493;
              t0 = dtday(2001, 1, 30);
             t1 = dtday(2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              c = AmericanBSPut(S0,K,r,0,tau,sigma);
             print c;
              16.748987
              19.41627
              22.318856
  Source
              finprocs.src
```

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AmericanBSPut Greeks

AmericanBSPut_Greeks

```
American Black and Scholes put Delta, Gamma, Omega, Theta, and
Purpose
              Vega.
  Format
              \{d,g,t,v,rh\} =
              AmericanBSPut_Greeks(SO, K, r, div, tau, sigma);
    Input
                     scalar, current price
              S0
              K
                     Mx1 vector, strike price
              r
                     scalar, risk free rate
              div
                     continuous dividend yield
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma
                     scalar, volatility
  Output
              d
                     Mx1 vector, delta
              g
                     Mx1 vector, gamma
              t
                     Mx1 vector, theta
                     Mx1 vector, vega
              ν
              rh
                     Mx1 vector, rho
Example
              S0 = 305;
              K = 300;
              r = .08;
              sigma = .25;
              tau = .33;
              print AmericanBSPut Greeks (S0,K,r,0,tau,sigma);
              -0.33296721
              0.0091658294
              -17.556118
              77.614238
              -40.575963
```

AmericanBSPut_Greeks

Source finprocs.src

Globals __fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal.

Default = 0.

_fin_epsilon scalar, finite difference stepsize. Default = 1e-8.

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AmericanBSPut ImpVol

AmericanBSPut_ImpVol

```
Implied volatilities for American Black and Scholes puts.
Purpose
  Format
             sigma = AmericanBSPut ImpVol(c,S0,K,r,div,tau);
    Input
                    Mx1 vector, put premiums
             c
             S0
                    scalar, current price
             K
                    Mx1 vector, strike prices
                    scalar, risk free rate
             r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
             tau
                    trading
  Output
             sigma Mx1 vector, volatility
Example
             p = \{ 14.60, 17.10, 20.10 \};
             S0 = 718.46i
             K = \{ 720, 725, 730 \};
             r = .0498;
             t0 = dtday(2001, 1, 30);
             t1 = dtday(2001, 2, 16);
             tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
             sigma = AmericanBSPut ImpVol(p,S0,K,r,0,tau);
             print sigma;
             0.12829346
             0.16885986
             0.21544312
  Source
             finprocs.src
```

annualTradingDays

annualTradingDays

Purpose Compute number of trading days in a given year.

Format n = annualTradingDays(a);

Input *a* scalar, year.

Output n number of trading days in year

Remarks A trading day is a weekday that is not a holiday as defined by the New

York Stock Exchange from 1888 through 2004. Holidays are defined in holidays. asc. You may edit that file to modify or add holidays.

Source finutils.src

V W

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arccos

arccos

Purpose Computes the inverse cosine.

Format $y = \arccos(x)$;

Input x NxK matrix.

Output y NxK matrix containing the angle in radians whose cosine is x.

Remarks If *x* is complex or has any elements whose absolute value is greater than 1, complex results are returned.

Example $x = \{ -1, -0.5, 0, 0.5, 1 \};$

y = arccos(x);

-1.000000

-0.500000

x = 0.000000

0.500000

1.000000

3.141593

2.094395

y = 1.570796

1.047198

0.000000

Source trig.src

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arcsin

arcsin

a b d е h k m n 0 p q t

u

V

W

```
Computes the inverse sine.
Purpose
 Format
              y = \arcsin(x);
    Input
                    NxK matrix.
  Output
                    NxK matrix, the angle in radians whose sine is x.
Remarks
              If x is complex or has any elements whose absolute value is greater than
              1, complex results are returned.
Example
              x = \{ -1, -0.5, 0, 0.5, 1 \};
              y = arcsin(x);
                   -1.000000
                   -0.500000
              x =
                     0.000000
                     0.500000
                     1.000000
                   -1.570796
                   -0.523599
                     0.000000
              y =
                     0.523599
                     1.570796
  Source
              trig.src
```

areshape

areshape

Purpose Reshapes a scalar, matrix, or array into an array of user-specified size.

Format y = areshape(x, o);

Input x scalar, matrix, or N-dimensional array.

o Mx1 vector of orders, the sizes of the dimensions of the new array.

Output y M-dimensional array, created from data in x.

Remarks If there are more elements in x than in y, the remaining elements are discarded. If there are not enough elements in x to fill y, then when **areshape** runs out of elements, it goes back to the first element of x and

starts getting additional elements from there.

```
Example x = 3i
```

```
orders = { 2,3,4 };
y = areshape(x,orders);
y will be a 2x3x4 array of threes.
x = reshape(seqa(1,1,90),30,3);
orders = { 2,3,4,5 };
y = areshape(x,orders);
```

y will be a 2x3x4x5 array. Since y contains 120 elements and x contains only 90, the first 90 elements of y will be set to the sequence of integers from 1 to 90 that are contained in x, and the last 30 elements of y will be set to the sequence of integers from 1 to 30 contained in the first 30 elements of x.

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x = reshape(seqa(1,1,60),20,3);
orders = { 3,2,4 };
y = areshape(x,orders);

y will be a 3x2x4 array. Since y contains 24 elements, and x contains 60, the elements of y will be set to the sequence of integers from 1 to 24 contained in the first 24 elements of x.

See also aconcat

arrayalloc

arrayalloc

Purpose Creates an N-dimensional array with unspecified contents.

Format y = arrayalloc(o,cf);

Input o Nx1 vector of orders, the sizes of the dimensions of the array.

cf scalar, 0 to allocate real array, or 1 to allocate complex array.

Output y N-dimensional array.

Remarks The contents are unspecified. This function is used to allocate an array

that will be written to in sections using **setarray**.

Example orders = { 2,3,4 };

y = arrayalloc(orders, 1);

y will be a complex 2x3x4 array with unspecified contents.

See also arrayinit

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arrayindex

arrayindex

a b

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m

11

o p

q

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```
Purpose Converts a scalar vector index to a vector of indices for an N-dimensional array.
```

Format i = arrayindex(si, o);

Input *si* scalar, index into vector or 1-dimensional array.

o Nx1 vector of orders of an N-dimensional array.

Output *i* Nx1 vector of indices, index of corresponding element in N-dimensional array.

Remarks This function and its opposite, **singleindex**, allow you to easily convert between an N-dimensional index and its corresponding location in a 1-dimensional object of the same size.

Example orders = { 2,3,4,5 };
v = rndu(prodc(orders),1);
a = areshape(v,orders);
vi = 50;
ai = arrayindex(vi,orders);
print vi;
print ai;
print v[vi];

print getarray(a,ai);

vi = 50

 $ai = \frac{3}{2}$

arrayindex

$$v[vi] = 0.13220899$$

$$getarray(a, ai) = 0.13220899$$

This example allocates a vector of random numbers and creates a 4-dimensional array using the same data. The 50th element of the vector v corresponds to the element of array a that is indexed with ai.

See also singleindex

a

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arrayinit

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k

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0

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q

arrayinit

a **Purpose** Creates an N-dimensional array with a specified fill value. **Format** y = arrayinit(o,v);Input Nx1 vector of orders, the sizes of the dimensions of the array. scalar, value to initialize. If v is complex the result will be complex. **Output** N-dimensional array with each element equal to the value of v. **Example** orders = { 2,3,4 }; y = arrayinit(orders, 0); y will be a 2x3x4 array of zeros. See also arrayalloc m

W хуг

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V

arraytomat

arraytomat

Purpose Changes an array to type matrix.

Format y = arraytomat(a);

Input *a* N-dimensional array.

Output y KxL or 1xL matrix or scalar, where L is the size of the fastest moving dimension of the array and K is the size of the second fastest moving dimension.

Remarks arraytomat will take an array of 1 or 2 dimensions or an N-dimensional array, in which the N-2 slowest moving dimensions each have a size of 1.

Example a = arrayinit(3|4,2); y = arraytomat(a); 2 2 2 2 y = 2 2 2 2

y = 2 2 2 2 2 2 2 2 2 2 2 2

See also mattoarray

a

b

c d

e f

g

h

j

K

m

n

0

p q

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S

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V

asclabel

a

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хух

asclabel

Purpose Sets up character labels for the X and Y axes. Library pgraph **Format** asclabel(xl,yl); Input xlstring or Nx1 character vector, labels for the tick marks on the X axis. Set to 0 if no character labels for this axis are desired. yl string or Mx1 character vector, labels for the tick marks on the Y axis. Set to 0 if no character labels for this axis are desired. **Example** This illustrates how to label the X axis with the months of the year: let lab = JAN FEB MAR APR MAY JUN JUL AUG SEP OCT NOV DEC; asclabel(lab,0); This will also work: lab = "JAN FEB MAR APR MAY JUN JUL AUG SEP OCT NOV DEC"; asclabel(lab,0); If the string format is used, then escape characters may be embedded in the labels. For example, the following produces character labels that are multiples of λ . The font Simgrma must be previously loaded in a **fonts** command. (See Chapter 13 in *Using GAUSS for Windows95*.) fonts("simplex simgrma"); $lab = "\2010.25\2021 \2010.5\2021$ \2010.75\2021 1"; asclabel(lab,0); Here, the **2021** produces the " λ " symbol from Simgrma. Source pgraph.src

asclabel

See also xtics, ytics, scale, scale3d, fonts

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asum

asum

a b d е g h k m 0 p q u V W

```
Purpose
              Computes the sum across one dimension of an N-dimensional array.
  Format
               y = asum(x, dim);
    Input
                      N-dimensional array.
              x
                      scalar, number of dimension to sum across.
               dim
  Output
                      N-dimensional array.
Remarks
               The output y, will have the same sizes of dimensions as x, except that the
               dimension indicated by dim will be collapsed to 1.
Example
              x = seqa(1,1,24);
               x = areshape(x, 2|3|4);
               y = asum(x,3);
              x is a 2x3x4 array, such that:
              [1,1,1] through [1,3,4] =
                         3
                    5 6
                         7 8
                   9 10 11 12
              [2,1,1] through [2,3,4] =
                    13 14 15 16
                    17 18 19 20
                   21 22 23 24
```

asum

y will be a 1x3x4 array, such that:

$$[1,1,1]$$
 through $[1,3,4] =$

14 16 18 20

22 24 26 28

30 32 34 36

$$y = asum(x,1);$$

Using the same array x as the above example, this example computes the sum across the first dimension. y will be a 2x3x1 array, such that:

$$[1,1,1]$$
 through $[1,3,1] =$

10

26

42

58

74

90

See also amean

a

b

С

d

f

g

h .

J

m

n o

n

q

40

S

t

u

V

atan

atan

a

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k

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p q

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хух

Purpose Returns the arctangent of its argument.

Format y = atan(x);

Input x NxK matrix.

Output y NxK matrix containing the arctangent of x in radians.

Remarks y will be a matrix the same size as x, containing the arctangents of the corresponding elements of x.

For real x, the arctangent of x is the angle whose tangent is x. The result is a value in radians in the range $\frac{-\pi}{2}$ to $\frac{+\pi}{2}$. To convert radians to degrees,

multiply by $\frac{180}{\pi}$.

For complex x, the arctangent is defined everywhere except i and -i. If x is complex, y will be complex.

Example $x = \{ 2, 4, 6, 8 \};$

z = x/2;

y = atan(z);

0.785398

 $y = \begin{array}{c} 1.107149 \\ 1.249046 \end{array}$

1.325818

See also atan2, sin, cos, pi, tan

atan2

atan2

Purpose Computes an angle from an *x*,*y* coordinate.

Format z = atan2(y,x);

Input y NxK matrix, the *Y* coordinate.

x LxM matrix, ExE conformable with y, the X coordinate.

Output $z = \max(N,L)$ by $\max(K,M)$ matrix.

Remarks Given a point *x*, *y* in a Cartesian coordinate system, **atan2** will give the correct angle with respect to the positive X axis. The answer will be in radians from -pi to +pi.

To convert radians to degrees, multiply by $\frac{180}{\pi}$.

atan2 operates only on the real component of *x*, even if *x* is complex.

Example x = 2i

$$y = \{ 2, 4, 6, 8 \};$$

$$z = atan2(y,x);$$

0.785398

= 1.107149

1.249046

1.325818

See also atan, sin, cos, pi, tan, arcsin, arccos

a

b

c d

е

f g

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p

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V

atranspose

atranspose

a b

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u

V

W

x y z

Purpose Transposes an N-dimensional array.

Format y = atranspose(x, nd);

Input x N-dimensional array.

nd Nx1 vector of dimension indices, the new order of dimensions.

Output y N-dimensional array, transposed according to *nd*.

Remarks The vector of dimension indices must be a unique vector of integers, 1-N, where 1 corresponds to the first element of the vector of orders.

Example x = seqa(1,1,24); x = areshape(x,2|3|4); $nd = \{ 2,1,3 \};$ y = atranspose(x,nd);

This example transposes the dimensions of x that correspond to the first and second elements of the vector of orders. x is a 2x3x4 array, such that:

[1,1,1] through [1,3,4] =

1 2 3 4

5 6 7 8

9 10 11 12

[2,1,1] through [2,3,4] =

13 14 15 16

17 18 19 20

21 22 23 24

atranspose

y will be a 3x2x4 array such that:

$$[1,1,1]$$
 through $[1,2,4] =$

13 14 15 16

$$[2,1,1]$$
 through $[2,2,4] =$

17 18 19 20

$$[3,1,1]$$
 through $[3,2,4] =$

21 22 23 24

$$nd = \{ 2,3,1 \};$$

$$y = atranspose(x,nd);$$

Using the same array x as the example above, this example transposes all three dimensions of x, returning a 3x4x2 array y, such that:

$$[1,1,1]$$
 through $[1,4,2] =$

1 13

2 14

3 15

4 16

$$[2,1,1]$$
 through $[2,4,2] =$

5 17

6 18

7 19

8 20

a

b

c d

e

f g

h

i

k

m

n

0

р

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S

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atranspose

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[3,1,1] through [3,4,2] =

9 21

10 22

11 23

12 24

See also areshape

axmargin

axmargin

Purpose Set absolute margins for the plot axes which control placement and size of plot.

•

Library pgraph

Format axmargin(l,r,t,b);

Input *l* scalar, the left margin in inches.

r scalar, the right margin in inches.

t scalar, the top margin in inches.

b scalar, the bottom margin in inches.

Remarks

axmargin sets an absolute distance from the axes to the edge of the graphic panel. Note that the user is responsible for allowing enough space in the margin if axes labels, numbers, and title are used on the graph, since **axmargin** does not size the plot automatically as in the case of **margin**.

All input inch values for this procedure are based on a full size window of 9 x 6.855 inches. If this procedure is used within a graphic panel, the values will be scaled to window inches automatically.

If both margin and axmargin are used for a graph, axmargin will override any sizes specified by margin.

Example The statement

```
axmargin(1,1,.5,.855);
```

will create a plot area of 7 inches horizontally by 5.5 inches vertically, and positioned 1 inch right and .855 up from the lower left corner of the graphic panel/page.

Source pgraph.src

a

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q

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5

t

u

V

balance

balance

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x y z

Purpose Balances a square matrix.

Format $\{b,z\}$ = balance(x)

Input x KxK matrix.

Output *b* KxK matrix, balanced matrix.

z KxK matrix, diagonal scale matrix.

Remarks

balance returns a balanced matrix b and another matrix z with scale factors in powers of two on its diagonal. b is balanced in the sense that the absolute sums of the magnitudes of elements in corresponding rows and columns are nearly equal.

balance is most often used to scale matrices to improve the numerical stability of the calculation of their eigenvalues. It is also useful in the solution of matrix equations.

In particular,

$$b = z^{-1}xz$$

balance uses the BALANC function from EISPACK.

Example

band

band

Purpose Extracts bands from a symmetric banded matrix.

Format a = band(y,n);

Input y KxK symmetric banded matrix.

n scalar, number of subdiagonals.

Output a Kx(N+1) matrix, 1 subdiagonal per column.

Remarks y can actually be a rectangular PxQ matrix. K is then defined as min(P,Q). It will be assumed that a is symmetric about the principal diagonal for

y[1:K,1:K].

The subdiagonals of y are stored right to left in a, with the principal diagonal in the rightmost (N+1'th) column of a. The upper left corner of a is unused; it is set to 0.

This compact form of a banded matrix is what **bandchol** expects.

Example

```
x = \{ 1 2 0 0, \\ 2 8 1 0, \\ 0 1 5 2, \\ 0 0 2 3 \};
```

bx = band(x,1);

0.0000000 1.0000000

 $bx = \begin{cases} 2.0000000 & 8.0000000 \\ 1.0000000 & 5.0000000 \end{cases}$

2.0000000 3.0000000

a

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bandchol

bandchol

a

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```
Purpose Computes the Cholesky decomposition of a positive definite banded matrix.
```

Format l = bandchol(a);

Input a KxN compact form matrix.

Output l KxN compact form matrix, lower triangle of the Cholesky decomposition of a.

Remarks Given a positive definite banded matrix A, there exists a matrix L, the lower triangle of the Cholesky decomposition of A, such that $A = L \times L'$. a is the compact form of A. See **band** for a description of the format of a.

l is the compact form of L. This is the form of matrix that **bandcholsol** expects.

Example

```
x = \{ 1 2 0 0,
      2 8 1 0,
      0 1 5 2,
      0 0 2 3 };
bx = band(x,1);
      0.0000000 1.0000000
      2.0000000 8.0000000
bx =
      1.0000000 5.0000000
      2.0000000 3.0000000
cx = bandchol(bx);
       0.0000000 1.0000000
       2.0000000 2.0000000
cx =
      0.50000000 2.1794495
      0.91766294 1.4689774
```

bandcholsol

bandcholsol

Purpose

Solves the system of equations Ax = b for x, given the lower triangle of the Cholesky decomposition of a positive definite banded matrix A.

Format

$$x = bandcholsol(b,l);$$

Input

b KxM matrix.

l KxN compact form matrix.

Output

x KxM matrix.

Remarks

Given a positive definite banded matrix A, there exists a matrix L, the lower triangle of the Cholesky decomposition of A, such that A = L*L'. l is the compact form of L; see **band** for a description of the format of l.

b can have more than one column. If so, Ax = b is solved for each column. That is,

$$A*x[.,i] = b[.,i]$$

Example

$$x = \{ 1 2 0 0, \\
2 8 1 0, \\
0 1 5 2, \\
0 0 2 3 \};$$

$$bx = band(x,1);$$

$$bx = \begin{cases} 0.0000000 & 1.0000000 \\ 2.0000000 & 8.0000000 \\ 1.0000000 & 5.0000000 \end{cases}$$

2.0000000 3.0000000

a

b

С

d

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k

1 m

n

0

q

r

S

t

u

V

w x y z

bandcholsol

b

d

h

k

m

n

0

p

q

cx = bandchol(bx);0.0000000 1.0000000 2.0000000 2.0000000 cx =0.50000000 2.1794495 0.91766294 1.4689774 xi = bandcholsol(eye(4),cx); 2.0731707 -0.05365854 0.14634146 0.09756098 $xi = -0.53658537 \quad 0.26829268 -0.07317073$ 0.04878049 $0.14634146 - 0.07317073 \ 0.29268293 - 0.19512195$ -0.09756098 0.04878049 -0.195121950.46341463

w x y z

u

V

bandltsol

bandltsol

Purpose Solves the system of equations Ax = b for x, where A is a lower triangular banded matrix.

Format x = bandltsol(b,A);

Input b KxM matrix.

A KxN compact form matrix.

Output x KxM matrix.

Remarks A is a lower triangular banded matrix in compact form. See **band** for a description of the format of A.

b can have more than one column. If so, Ax = b is solved for each column. That is,

$$A*x[.,i] = b[.,i]$$

Example 0.0000000 1.0000000

 $bx = \begin{cases} 2.0000000 & 8.0000000 \\ 1.0000000 & 5.0000000 \\ 2.0000000 & 3.0000000 \end{cases}$

cx = bandchol(bx);

 $cx = \begin{cases} 0.0000000 & 1.0000000 \\ 2.0000000 & 2.0000000 \\ 0.50000000 & 2.1794495 \\ 0.91766294 & 1.4689774 \end{cases}$

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bandltsol

xci = bandltsol(eye(4),cx);

 $xci = egin{array}{ccccccc} 1.0000000 & 0.0000000 & 0.0000000 & 0.0000000 \\ -1.0000000 & 0.50000000 & 0.0000000 & 0.0000000 \\ \end{array}$

 $0.22941573 - 0.11470787 \ 0.45883147 \ 0.0000000$

-0.14331487 0.07165744 -0.28662975 0.68074565

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bandry

bandry

Purpose Creates a symmetric banded matrix, given its compact form.

Format y = bandrv(a);

Input *a* KxN compact form matrix.

Output y KxK symmetrix banded matrix.

Remarks

a is the compact form of a symmetric banded matrix, as generated by **band**. a stores subdiagonals right to left, with the principal diagonal in the rightmost (Nth) column. The upper left corner of a is unused. **bandchol** expects a matrix of this form.

y is the fully expanded form of a, a KxK matrix with N-1 subdiagonals.

Example

 $bx = \begin{cases} 0.0000000 & 1.0000000 \\ 2.0000000 & 8.0000000 \\ 1.0000000 & 5.0000000 \\ 2.0000000 & 3.0000000 \end{cases}$

x = bandrv(bx);

1.0000000 2.0000000 0.0000000 0.0000000 2.0000000 8.0000000 1.0000000 0.0000000 \boldsymbol{x} 0.0000000 1.0000000 5.0000000 2.0000000 0.0000000 0.0000000 2.0000000 3.0000000 a

b

c d

e

f g

h

j

k

m

n o

р

q

1

t

u

V

W

bandsolpd

bandsolpd

Purpose

Solves the system of equations Ax = b for x, where A is a positive definite banded matrix.

Format

x = bandsolpd(b,A);

Input

b KxM matrix.

A KxN compact form matrix.

Output

x KxM matrix.

Remarks

A is a positive definite banded matrix in compact form. See **band** for a description of the format of A.

b can have more than one column. If so, Ax = b is solved for each column. That is,

$$A*x[.,i] = b[.,i]$$

a

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bar

Purpose Bar graph. Library pgraph **Format** bar(val,ht); Input val Nx1 numeric vector, bar labels. If scalar 0, a sequence from 1 to **rows** (ht) will be created. ht NxK numeric vector, bar heights. K overlapping or side-by-side sets of N bars will be graphed. For overlapping bars, the first column should contain the set of bars with the greatest height and the last column should contain the set of bars with the least height. Otherwise, the bars that are drawn first may be obscured by the bars drawn last. This is not a problem if the bars are plotted side-by-side. **Global Input** global scalar, width and positioning of bars in bar graphs pbarwid and histograms. The valid range is 0-1. If this is 0, the bars will be a single pixel wide. If this is 1, the bars will touch each other. If this value is positive, the bars will overlap. If negative, the bars will be plotted side-by-side. The default is 0.5. pbartyp Kx2 matrix. The first column controls the bar shading: no shading. 1 dots. vertical cross-hatch. diagonal lines with positive slope. 4 diagonal lines with negative slope. 5 diagonal cross-hatch. 6 solid. The second column controls the bar color. See "Colors Appendix" on page B-1.

bar

b

d

h

k

m

n

0

p

q

Remarks Use scale or ytics to fix the scaling for the bar heights.

Example In this example, three overlapping sets of bars will be created. The three heights for the *i*th bar are stored in *x*[*i*,.].

library pgraph;

graphset;

t = seqa(0,1,10);

x = (t^2/2).*(1~0.7~0.3);

Source pbar.src

 $_{pnum} = 2;$

See also asclabel, xy, logx, logy, loglog, scale, hist

bar(t,x); /* Use t vector to label X axis */

_

1

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t

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base10

base10

Purpose Break number into a number of the form #.####... and a power of 10.

Format $\{M,P\}$ = base10(x);

Input *x* scalar, number to break down.

Output M scalar, in the range -10 < M < 10.

P scalar, integer power such that:

 $M * 10^P = x$

Example $\{ b, e \} = base10(4500);$

b = 4.5000000

e = 3.0000000

Source base10.src

a

b

С

d

g

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1

1-

1

m

n

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begwind

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e

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u

V

W

begwind

Purpose Initialize global graphic panel variables.

Library pgraph

Format begwind;

Remarks This procedure must be called before any other graphic panel functions

are called.

Source pwindow.src

See also endwind, window, makewind, setwind, nextwind,

getwind

хуг

Z

besselj

besselj

```
Purpose
              Computes a Bessel function of the first kind, J_n(x).
  Format
              y = besselj(n,x);
    Input
              n
                     NxK matrix, the order of the Bessel function. Nonintegers will
                     be truncated to an integer.
                     LxM matrix, ExE conformable with n.
              \boldsymbol{x}
  Output
                     max(N,L) by max(K,M) matrix.
Example
              n = \{ 0, 1 \};
              x = \{ 0.1 1.2, 2.3 3.4 \};
              y = besselj(n,x);
                    0.99750156
                                 0.67113274
                    0.53987253
                                0.17922585
See also
              bessely, mbesseli
```

a

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k

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О

p

q

r

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bessely

bessely

a
b
c
d
e
f
g
h
i
j
k
l
m
n

```
Purpose To compute a Bessel function of the second kind (Weber's function), Y_n(x).

Format y = bessely(n,x);
```

Input n NxK matrix, the order of the Bessel function. Nonintegers will be truncated to an integer.

x LxM matrix, ExE conformable with n.

Output $y = \max(N,L)$ by $\max(K,M)$ matrix.

Example $n = \{ 0, 1 \};$ $x = \{ 0.1 1.2, 2.3 3.4 \};$ y = bessely(n,x); $y = \begin{cases} -1.5342387 & 0.22808351 \\ 0.05227732 & 0.40101529 \end{cases}$

See also besselj, mbesseli

хух

u

V

W

0

p

q

box

box

Purpose Graph data using the box graph percentile method.

Library pgraph

Format box(grp,y);

Input grp

1xM vector. This contains the group numbers corresponding to each column of y data. If scalar 0, a sequence from 1 to cols(y) will be generated automatically for the X axis.

y NxM matrix. Each column represents the set of y values for an individual percentiles box symbol.

Global Input _pboxctl 5x1 vector, controls box style, width, and color.

- [1] box width between 0 and 1. If zero, the box plot is drawn as two vertical lines representing the quartile ranges with a filled circle representing the 50th percentile.
- [2] box color. If this is set to 0, the colors may be individually controlled using the global variable _pcolor.
- [3] Min/max style for the box symbol. One of the following:
 - 1 Minimum and maximum taken from the actual limits of the data. Elements 4 and 5 are ignored.
 - 2 Statistical standard with the minimum and maximum calculated according to interquartile range as follows:

$$intqrange = 75^{th} - 25^{th}$$

 $min = 25^{th} - 1.5intqrange$
 $max = 75^{th} + 1.5intqrange$

Elements 4 and 5 are ignored.

- Minimum and maximum percentiles taken from elements 4 and 5.
- 4] Minimum percentile value (0-100) if $_{\mathbf{pboxctl}[3]} = 3$.

a

b

c d

e

f g

h

.

k

1

m n

0

p

q

C

f

u

V

W

box

h d h m n 0 p q u V W хух [5] Maximum percentile value (0-100) if $_{\mathbf{pboxctl}[3]} = 3$.

_plctrl 1xM vector or scalar as follows:

- O Plot boxes only, no symbols.
- 1 Plot boxes and plot symbols that lie outside the min and max box values.
- 2 Plot boxes and all symbols.
- **-1** Plot symbols only, no boxes.

These capabilities are in addition to the usual line control capabilities of _plctrl.

_pcolor 1xM vector or scalar for symbol colors. If scalar, all symbols will be one color. See "Colors Appendix" on page B-1.

Remarks If missing values are encountered in the *y* data, they will be ignored during calculations and will not be plotted.

Source pbox.src

boxcox

boxcox

Purpose Computes the Box-Cox function.

Format y = boxcox(x, lambda);

Input x MxN matrix.

lambda KxL matrix, ExE conformable to x.

Output $y = \max(M,L) \times \max(N,K)$.

Remarks Allowable range for x is:

x > 0

The **boxcox** function computes

$$boxcox(x) = \frac{x^{\lambda} - 1}{\lambda}$$

Example $x = \{.2.3, 1.5.2.5\};$

lambda = $\{.4, 2\};$

y = boxcox(x, lambda)

 $y = \begin{array}{r} -1.1867361 -0.95549787 \\ 0.62500000 \ \ 2.62500000 \end{array}$

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break

Purpose Breaks out of a **do** or **for** loop. b **Format** break; Example x = rndn(4,4);d r = 0;do while r < rows(x); r = r + 1;c = 0;h do while c < cols(x); c = c + 1;if c == r;x[r,c] = 1;elseif c > r; break; /* terminate inner do loop */ m else; n x[r,c] = 0;0 endif; p endo; /* break jumps to the statement */ q /* after this endo */ endo; $1.000 \ 0.326 \ -2.682 \ -0.594$ $0.000 \ 1.000 \ -0.879 \ 0.056$ x =u $0.000 \ 0.000 \ 1.000 \ -0.688$ 0.000 0.000 0.000 1.000 V Remarks This command works just like in C. W

break

See also continue, do, for

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Purpose Calls a function or procedure when the returned value is not needed and can be ignored, or when the procedure is defined to return nothing.

Format call function_name(argument_list);

call function_name;

This is useful when you need to execute a function or procedure and do not need the value that it returns. It can also be used for calling procedures

that have been defined to return nothing.

function_name can be any intrinsic GAUSS function, a procedure (**proc**), or any valid expression.

Example call chol(x);
v = detl;

The above example is the fastest way to compute the determinant of a positive definite matrix. The result of **chol** is discarded and **detl** is used to retrieve the determinant that was computed during the call to **chol**.

See also proc

cdfbeta

cdfbeta

Purpose Computes the incomplete beta function (i.e., the cumulative distribution function of the beta distribution).

Format y = cdfbeta(x,a,b);

Input x NxK matrix.

a LxM matrix, ExE conformable with x.

b PxQ matrix, ExE conformable with x and a.

Output $y = \max(N,L,P)$ by $\max(K,M,Q)$ matrix.

Remarks y is the integral from 0 to x of the beta distribution with parameters a and b. Allowable ranges for the arguments are:

$$0 <= x <= 1$$

A -1 is returned for those elements with invalid inputs.

Example $x = \{ .1, .2, .3, .4 \};$

$$a = 0.5;$$

$$b = 0.3;$$

y = cdfbeta(x,a,b);

0.260575

0.310875

See also cdfchic, cdffc, cdfn, cdfnc, cdftc, gamma

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Technical Notes

cdfbeta has the following approximate accuracy:

$max(a,b) \le 500$	the absolute error is approx. ±5e-13
$500 < \max(a,b) \le 10,000$	the absolute error is approx. ±5e-11
$10,000 < \max(a,b) \le 200,000$	the absolute error is approx. ±1e-9
$200,000 < \max(a,b)$	Normal approximations are used and
	the absolute error is approx. ±2e-9

References

Bol'shev, L.N. "Asymptotically Pearson's Transformations." Teor. Veroyat. Primen. (Theory of Probability and its Applications). Vol. 8 No. 2, 1963, 129-55.

Bosten, N. E., and E.L. Battiste. "Remark on Algorithm 179 Incomplete Beta Ratio." Comm. ACM. Vol. 17 No. 3, March 1974, 156-57.

Ludwig, O.G. "Algorithm 179 Incomplete Beta Ratio." Comm. ACM. Vol. 6 No. 6, June 1963, 314.

Mardia, K.V., and P.J. Zemroch. "Tables of the F- and related distributions with algorithms." Academic Press, NY, 1978. ISBN 0-12-471140-5

Peizer, D.B., and J.W. Pratt. "A Normal Approximation for Binomial, F, Beta, and Other Common, Related Tail Probabilities, I." Journal of American Statistical Association. Vol. 63, Dec. 1968, 1416-56.

Pike, M.C., and I.D. Hill. "Remark on Algorithm 179 Incomplete Beta Ratio." Comm. ACM. Vol. 10 No. 6, June 1967, 375-76.

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cdfbvn

Purpose

Computes the cumulative distribution function of the standardized bivariate Normal density (lower tail).

Format c = cdfbvn(h, k, r);

Input h NxK matrix, the upper limits of integration for variable 1.

> kLxM matrix, ExE conformable with h, the upper limits of integration for variable 2.

> PxQ matrix, ExE conformable with h and k, the correlation r coefficients between the two variables.

Output max(N,L,P) by max(K,M,Q) matrix, the result of the double integral from $-\infty$ to h and $-\infty$ to k of the standardized bivariate Normal density f(x,y,r).

Remarks The function integrated is:

$$f(x, y, r) = \frac{e^{-0.5w}}{2\pi\sqrt{1-r^2}}$$

with

$$w = \frac{x^2 - 2rxy + y^2}{1 - r^2}$$

Thus, x and y have 0 means, unit variances, and correlation = r.

Allowable ranges for the arguments are:

$$-\infty$$
 $< h$ $< +\infty$

$$-\infty$$
 < k < $+\infty$

A -1 is returned for those elements with invalid inputs.

To find the integral under a general bivariate density, with x and y having nonzero means and any positive standard deviations, use the transformation equations:

$$h = (ht - ux) ./ sx;$$

$$k = (kt - uy) ./ sy;$$

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where $\mathbf{u}\mathbf{x}$ and $\mathbf{u}\mathbf{y}$ are the (vectors of) means of x and y, $\mathbf{s}\mathbf{x}$ and $\mathbf{s}\mathbf{y}$ are the (vectors of) standard deviations of x and y, and $\mathbf{h}\mathbf{t}$ are the (vectors of) upper integration limits for the untransformed variables, respectively.

See also

cdfn, cdftvn

Technical Notes

The absolute error for **cdfbvn** is approximately $\pm 5.0e-9$ for the entire range of arguments.

References

Daley, D.J. "Computation of Bi- and Tri-variate Normal Integral." *Appl. Statist.* Vol. 23 No. 3, 1974, 435-38.

Owen, D.B. "A Table of Normal Integrals." *Commun. Statist.-Simula. Computa.*, B9(4). 1980, 389-419.

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cdfbvn2

Returns cdfbvn of a bounded rectangle.			
y = cdfbvn2(h,dh,k,dk,r);			
 h Nx1 vector, starting points of integration for variable 1. dh Nx1 vector, increments for variable 1. k Nx1 vector, starting points of integration for variable 2. dk Nx1 vector, increments for variable 2. r Nx1 vector, correlation coefficients between the two variables. 			
y Nx1 vector, the integral over the rectangle bounded by h , $h+dh$, k , and $k+dk$ of the standardized bivariate Normal distribution.			
Scalar input arguments are okay; they will be expanded to Nx1 vectors. ${\tt cdfbvn2}$ computes: ${\tt cdfbvn}(h+dh,k+dk,r)+{\tt cdfbvn}(h,k,r)-{\tt cdfbvn}(h,k+dk,r)-{\tt cdfbvn}(h+dh,k,r).$ ${\tt cdfbvn2}$ computes an error estimate for each set of inputs. The size of the error depends on the input arguments. If ${\tt trap}\ 2$ is set, a warning message is displayed when the error reaches $0.01*{\tt abs}(y)$. For an estimate of the actual error, see ${\tt cdfbvn2e}$.			
Example 1 cdfbvn2(1,-1,1,-1,0.5); produces: 1.4105101488974692e-001 Example 2 cdfbvn2(1,-1e-15,1,-1e-15,0.5); produces: 4.9303806576313238e-32			

```
Example 3
                       cdfbvn2(1,-1e-45,1,-1e-45,0.5);
                       produces:
b
                       0.0000000000000000e+000
                       Example 4
d
                       trap 2,2;
                       cdfbvn2(1,-1e-45,1,1e-45,0.5);
                       produces:
                       WARNING: Dubious accuracy from cdfbvn2:
h
                       0.000e+000 +/- 2.8e-060
                       0.0000000000000000e+000
             Source
                       lncdfn.src
k
           See also cdfbvn2e, lncdfbvn2
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cdfbvn2e

Purpose	Returns cdfbvn of a bounded rectangle.		
Format	$\{y, e\} = cdfbvn2e(h,dh,k,dk,r);$		
Input	 h Nx1 vector, starting points of integration for variable 1. dh Nx1 vector, increments for variable 1. k Nx1 vector, starting points of integration for variable 2. dk Nx1 vector, increments for variable 2. r Nx1 vector, correlation coefficients between the two variables. 		
Output	 Nx1 vector, the integral over the rectangle bounded by h, h+dh, k, and k+dk of the standardized bivariate Normal distribution. e Nx1 vector, an error estimate. 		
Remarks	Scalar input arguments are okay; they will be expanded to Nx1 vectors. $\texttt{cdfbvn2e}$ computes $\texttt{cdfbvn}(h+dh,k+dk,r) + \texttt{cdfbvn}(h,k,r) - \texttt{cdfbvn}(h,k+dk,r)$. The real answer is $y\pm e$. The size of the error depends on the input arguments.		
Example	Example 1 cdfbvn2e(1,-1,1,-1,0.5); produces: 1.4105101488974692e-001 1.9927918166193113e-014 Example 2 cdfbvn2e(1,-1e-15,1,-1e-15,0.5); produces:		

cdfbvn2e

7.3955709864469857e-032

2.8306169312687801e-030

Example 3

cdfbvn2e(1,-1e-45,1,-1e-45,0.5);

produces:

0.00000000000000000e+000

2.8306169312687770e-060

See also cdfbvn2, lncdfbvn2

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cdfchic

cdfchic

Purpose Computes the complement of the cdf of the chi-square distribution.

Format y = cdfchic(x,n)

Input x NxK matrix.

n LxM matrix, ExE conformable with x.

Output $y = \max(N,L)$ by $\max(K,M)$ matrix.

Remarks y is the integral from x to ∞ of the chi-square distribution with n degrees of freedom.

The elements of *n* must all be positive integers. The allowable ranges for the arguments are:

$$x >= 0$$
$$n > 0$$

A -1 is returned for those elements with invalid inputs.

This equals 1-F(x,n), where F is the chi-square cdf with n degrees of freedom. Thus, to get the chi-square cdf, subtract **cdfchic**(x, n) from 1. The complement of the cdf is computed because this is what is most commonly needed in statistical applications, and because it can be computed with fewer problems of roundoff error.

See also cdfbeta, cdffc, cdfn, cdfnc, cdftc, gamma

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cdfchic

Technical Notes

For n <= 1000, the incomplete gamma function is used and the absolute error is approx. $\pm 6e$ -13. For n > 1000, a Normal approximation is used and the absolute error is $\pm 2e$ -8.

For higher accuracy when n > 1000, use:

$$1-cdfgam(0.5*x, 0.5*n);$$

References

Bhattacharjee, G.P. "Algorithm AS 32, The Incomplete Gamma Integral." *Applied Statistics*. Vol. 19, 1970, 285-87.

Mardia, K.V., and P.J. Zemroch. "Tables of the F- and related distributions with algorithms." Academic Press, NY, 1978. ISBN 0-12-471140-5

Peizer, D.B., and J.W. Pratt. "A Normal Approximation for Binomial, F, Beta, and Other Common, Related Tail Probabilities, I." *Journal of American Statistical Association*. Vol. 63, Dec. 1968, 1416-56.

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cdfchii

cdfchii

Purpose Compute chi-square abscissae values given probability and degrees of freedom.

Format c = cdfchii(p,n);

Input *p* MxN matrix, probabilities.

n LxK matrix, ExE conformable with p, degrees of freedom.

Output c $\max(M,L)$ by $\max(N,K)$ matrix, abscissae values for chi-square distribution.

Example The following generates a 3x3 matrix of pseudo-random numbers with a chi-squared distribution with expected value of 4:

rndseed 464578; x = cdfchii(rndu(3,3),4+zeros(3,3)); 2.1096456 1.9354989 1.7549182 x = 4.4971008 9.2643386 4.3639694 4.5737473 1.3706243 2.5653688

Source cdfchii.src

See also gammaii

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Purpose

The integral under noncentral chi-square distribution, from 0 to x. It can return a vector of values, but the degrees of freedom and noncentrality parameter must be the same for all values of x.

Format

y = cdfchinc(x,v,d);

Input

Nx1 vector, values of upper limits of integrals, must be greater than 0.

v scalar, degrees of freedom, v > 0.

d scalar, noncentrality parameter, d > 0.

This is the square root of the noncentrality parameter that sometimes goes under the symbol lambda. (See Scheffe, *The Analysis of Variance*, App. IV. 1959.)

Output

Nx1 vector, integrals from 0 to x of noncentral chi-square.

Example

```
x = { .5, 1, 5, 25 };
p = cdfchinc(x,4,2);
     0.0042086234
     0.016608592
     0.30954232
     0.99441140
```

Source

cdfnonc.src

See also

cdffnc, cdftnc

Technical Notes Relation to cdfchic:

```
cdfchic(x,v) = 1 - cdfchinc(x,v,0);
```

The formula used is taken from Abramowitz and Stegun, *Handbook of Mathematical Functions*. Formula 26.4.25. 1970, 942.

cdffc

Purpose Computes the complement of the cdf of the F distribution.

Format y = cdffc(x,n1,n2);

Input x NxK matrix.

n1 LxM matrix, ExE conformable with x.

n2 PxQ matrix, ExE conformable with x and n1.

Output $y = \max(N,L,P)$ by $\max(K,M,Q)$ matrix.

Remarks y is the integral from x to ∞ of the F distribution with n1 and n2 degrees of freedom.

Allowable ranges for the arguments are:

$$x >= 0$$

$$n1 > 0$$

$$n2 > 0$$

A -1 is returned for those elements with invalid inputs.

This equals 1-G(x,n1,n2), where G is the F cdf with n1 and n2 degrees of freedom. Thus, to get the F cdf, subtract $\mathtt{cdffc}(x,n1,n2)$ from 1. The complement of the cdf is computed because this is what is most commonly needed in statistical applications, and because it can be computed with fewer problems of roundoff error.

 $x = \{ .1, .2, .3, .4 \};$

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cdffc

See also

cdfbeta, cdfchic, cdfn, cdfnc, cdftc, gamma

Technical Notes

For $max(n1,n2) \le 1000$, the absolute error is approximately $\pm 5e-13$. For max(n1,n2) > 1000, Normal approximations are used and the absolute error is approximately ±2e-6.

For higher accuracy when max(n1,n2) > 1000, use:

$$cdfbeta(n2/(n2+n1*x), n2/2, n1/2);$$

References

Bol'shev, L.N. "Asymptotically Pearson's Transformations." Teor. Veroyat. Primen. (Theory of Probability and its Applications). Vol. 8 No. 2, 1963, 129-55.

Bosten, N. E., and E.L. Battiste. "Remark on Algorithm 179 Incomplete Beta Ratio." Comm. ACM. Vol. 17 No. 3, March 1974, 156-57.

Kennedy, W.J., Jr., and J.E. Gentle. Statistical Computing. Marcel Dekker, Inc., NY, 1980.

Ludwig, O.G. "Algorithm 179 Incomplete Beta Ratio." Comm. ACM. Vol. 6 No. 6, June 1963, 314.

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Pike, M.C., and I.D. Hill. "Remark on Algorithm 179 Incomplete Beta Ratio." Comm. ACM. Vol. 10 No. 6, June 1967, 375-76.

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cdffnc

cdffnc

Purpose The integral under noncentral F distribution, from 0 to x.

Format y = cdffnc(x, v1, v2, d);

Input x Nx1 vector, values of upper limits of integrals, x > 0.

v1 scalar, degrees of freedom of numerator, v1 > 0.

v2 scalar, degrees of freedom of denominator, v2 > 0.

d scalar, noncentrality parameter, d > 0.

This is the square root of the noncentrality parameter that sometimes goes under the symbol lambda. (See Scheffe, *The Analysis of Variance*, App. IV. 1959.)

Output y Nx1 vector of integrals from 0 to x of noncentral F.

Source cdfnonc.src

See also cdftnc, cdfchinc

Technical Relation to cdffc:

cdffc(x,v1,v2) = 1 - cdffnc(x,v1,v2,0);

The formula used is taken from Abramowitz and Stegun, *Handbook of Mathematical Functions*. Formula 26.6.20. 1970, 947.

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Purpose Incomplete gamma function.

Format

g = cdfgam(x, intlim);

Input

x NxK matrix of data.

intlim LxM matrix, ExE compatible with *x*, containing the integration limit.

Output

 $g = \max(N,L)$ by $\max(K,M)$ matrix.

Remarks

The incomplete gamma function returns the integral

$$\int_0^{intlim} \frac{e^{-t}t^{(x-1)}}{gamma(x)} dt$$

The allowable ranges for the arguments are:

$$x > 0$$
 $intlim >= 0$

A -1 is returned for those elements with invalid inputs.

Example

intlim = seqa(0,.2,6);

 $x = \{ 0.5 1 3 10 \};$

g = cdfgam(x,intlim);

 $x = 0.500000 \ 1.00000 \ 3.00000 \ 10.00000$

0.000000

0.200000

intlim = 0.400000

0.600000

0.800000

1.000000

cdfgam

	0.000000	0.000000	0.000000	0.000000
	0.472911	0.181269	0.00114848	2.35307E - 014
g =	0.628907	0.329680	0.00792633	2.00981E - 011
O	0.726678	1.451188	0.0231153	9.66972E - 010
	0.794097	0.550671	0.0474226	1.43310E - 008
	0.842701	0.632120	0.0803014	1.11425E - 007

This computes the integrals over the range from 0 to 1, in increments of .2, at the parameter values 0.5, 1, 3, 10.

Technical Notes

cdfgam has the following approximate accuracy:

x < 500	the absolute error is approx. ±6e-13
$500 \le x \le 10,000$	the absolute error is approx. ±3e-11
10,000 < x	a Normal approximation is used and the
	absolute error is approx. ±3e-10

References

Bhattacharjee, G.P. "Algorithm AS 32, The Incomplete Gamma Integral." *Applied Statistics*. Vol. 19, 1970, 285-87.

Peizer, D.B., and J.W. Pratt. "A Normal Approximation for Binomial, F, Beta, and Other Common, Related Tail Probabilities, I." *Journal of American Statistical Association*. Vol. 63, Dec. 1968, 1416-56.

Pike, M.C., and I.D. Hill. "Remark on Algorithm 179 Incomplete Beta Ratio." *Comm. ACM.* Vol. 10 No. 6, June 1967, 375-76.

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cdfmvn

Purpose Computes multivariate Normal cumulative distribution function.

Format y = cdfmvn(x,r);

Input *x* KxL matrix, abscissae.

r KxK matrix, correlation matrix.

Output y Lx1 vector, $Pr(X < x \mid r)$.

Source lncdfn.src

See also cdfbvn, cdfn, cdftvn, lncdfmvn

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cdfn, cdfnc

cdfn, cdfnc

Purpose

cdfn computes the cumulative distribution function (cdf) of the Normal distribution. **cdfnc** computes 1 minus the cdf of the Normal distribution.

Format

```
n = \text{cdfn}(x);
nc = \text{cdfnc}(x);
```

Input

x NxK matrix.

Output

n NxK matrix.

nc NxK matrix.

Remarks

n is the integral from $-\infty$ to *x* of the Normal density function, and *nc* is the integral from x to $+\infty$.

Note that: $\mathbf{cdfn}(x) + \mathbf{cdfnc}(x) = 1$. However, many applications expect $\mathbf{cdfn}(x)$ to approach 1, but never actually reach it. Because of this, we have capped the return value of \mathbf{cdfn} at 1 - machine epsilon, or approximately 1 - 1.11e-16. As the relative error of \mathbf{cdfn} is about $\pm 5\text{e-}15$ for $\mathbf{cdfn}(x)$ around 1, this does not invalidate the result. What it does mean is that for $\mathbf{abs}(x) > (approx.)$ 8.2924, the identity does not hold true. If you have a need for the uncapped value of \mathbf{cdfn} , the following code will return it:

```
n = cdfn(x);
if n >= 1-eps;
    n = 1;
endif;
```

where the value of machine epsilon is obtained as follows:

```
x = 1;
do while 1-x /= 1;
    eps = x;
    x = x/2;
endo;
```

Note that this is an alternate definition of machine epsilon. Machine epsilon is usually defined as the smallest number such that 1 + machine

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cdfn, cdfnc

epsilon > 1, which is about 2.23e-16. This defines machine epsilon as the smallest number such that 1 - machine epsilon < 1, or about 1.11e-16.

The **erf** and **erfc** functions are also provided, and may sometimes be more useful than **cdfn** and **cdfnc**.

Example

$$x = \{ -2 -1 \ 0 \ 1 \ 2 \};$$

 $n = cdfn(x);$
 $nc = cdfnc(x);$
 $x = -2.00000 \ -1.00000 \ 0.00000 \ 1.00000 \ 2.00000$
 $n = 0.02275 \ 0.15866 \ 0.50000 \ 0.84134 \ 0.97725$
 $nc = 0.97725 \ 0.84134 \ 0.50000 \ 0.15866 \ 0.02275$

See also

erf, erfc, cdfbeta, cdfchic, cdftc, cdffc, gamma

Technical Notes

For the integral from $-\infty$ to x:

$x \le -37$,	cdfn underflows and 0.0 is returned
-36 < x < -10,	cdfn has a relative error of approx. ±5e-12
-10 < x < 0,	cdfn has a relative error of approx. ±1e-13
0 < x,	cdfn has a relative error of approx. ±5e-15

For **cdfnc**, i.e., the integral from x to $+\infty$, use the above accuracies but change x to -x.

References

Adams, A.G. "Remark on Algorithm 304 Normal Curve Integral." *Comm. ACM.* Vol. 12 No. 10, Oct. 1969, 565-66.

Hill, I.D., and S.A. Joyce. "Algorithm 304 Normal Curve Integral." *Comm. ACM.* Vol. 10 No. 6, June 1967, 374-75.

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cdfn2

Computes the integral over a Normal density function interval. **Purpose Format** y = cdfn2(x,dx);Input MxN matrix, abscissae. х dxKxL matrix, ExE conformable to x, intervals. **Output** max(M,K) by max(N,L) matrix, the integral from x to x+dx of the Normal distribution, i.e., $Pr(x \le X \le x + dx)$. Remarks The relative error is: |x| <= 1 and dx <= 1+1e-141 < |x| < 37 and |dx| < 1/|x| $\pm 1e-13$ min(x,x+dx) > -37 and y > 1e-300 $\pm 1e$ -11 or better A relative error of $\pm 1e$ -14 implies that the answer is accurate to better than ± 1 in the 14th digit. **Example** print cdfn2(1,0.5); 9.1848052662599017e-02 print cdfn2(20,0.5); 2.7535164718736454e-89 print cdfn2(20,1e-2); 5.0038115018684521e-90 print cdfn2(-5,2);1.3496113800582164e-03 print cdfn2(-5,0.15); 3.3065580013000255e-07 Source lncdfn.src See also lncdfn2

cdfni

cdfni

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Purpose Computes the inverse of the cdf of the Normal distribution.

0

Format x = cdfni(p);

p

Input

NxK real matrix, Normal probability levels, $0 \le p \le 1$.

Output

x NxK real matrix, Normal deviates, such that cdfn(x) = p

Remarks

cdfn(cdfni(p**)**) = p to within the errors given below:

 $p \le 4.6..e-308$

4.6..e-308

5e-24

0.5

p >= 1 - 2.22045e-16

-37.5 is returned

accurate to ± 5 in 12th digit accurate to ± 1 in 13th digit

accurate to ± 5 in 15th digit 8.12589... is returned

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cdftc

cdftc

Purpose Computes the complement of the cdf of the Student's *t* distribution.

Format y = cdftc(x,n);

Input x NxK matrix.

n LxM matrix, ExE conformable with *x*.

Output $y = \max(N,L)$ by $\max(K,M)$ matrix.

Remarks y is the integral from x to ∞ of the t distribution with n degrees of freedom.

Allowable ranges for the arguments are:

$$-\infty < x < +\infty$$
 $n > 0$

A -1 is returned for those elements with invalid inputs.

This equals 1-F(x,n), where F is the t cdf with n degrees of freedom. Thus, to get the t cdf, subtract cdftc(x,n) from 1. The complement of the cdf is computed because this is what is most commonly needed in statistical applications, and because it can be computed with fewer problems of roundoff error.

See also cdfbeta, cdfchic, cdffc, cdfn, cdfnc, gamma

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cdftc

Technical Notes

For results greater than 0.5e-30, the absolute error is approximately $\pm 1e$ -14 and the relative error is approximately $\pm 1e$ -12. If you multiply the relative error by the result, then take the minimum of that and the absolute error, you have the maximum actual error for any result. Thus, the actual error is approximately $\pm 1e$ -14 for results greater than 0.01. For results less than 0.01, the actual error will be less. For example, for a result of 0.5e-30, the actual error is only $\pm 0.5e$ -42.

References

Abramowitz, M., and I. A. Stegun, eds. *Handbook of Mathematical Functions*. 7th ed. Dover, NY, 1970. ISBN 0-486-61272-4

Hill, G.W. "Algorithm 395 Student's t-Distribution." *Comm. ACM*. Vol. 13 No. 10, Oct. 1970.

Hill, G.W. "Student's t-Distribution Quantiles to 20D." *Division of Mathematical Statistics Technical Paper No. 35*. Commonwealth Scientific and Industrial Research Organization, Australia, 1972.

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cdftci

cdftci

Purpose Computes the inverse of the complement of the Student's t cdf.

Format x = cdftci(p,n);

Input p NxK real matrix, complementary Student's t probability levels, $0 \le p \le 1$.

n LxM real matrix, degrees of freedom, $n \ge 1$, *n* need not be integral. ExE conformable with *p*.

Output $x = \max(N,L)$ by $\max(K,M)$ real matrix, Student's t deviates, such that $\mathbf{cdftc}(x,n) = p$.

Remarks cdftc(cdftci(p,n)) = p to within the errors given below:

 $0.5e-30 accurate to <math>\pm 1$ in 12th digit 0.01 < p accurate to $\pm 1e-14$

Extreme values of arguments can give rise to underflows, but no overflows are generated.

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cdftnc

cdftnc

Purpose

The integral under noncentral Student's t distribution, from $-\infty$ to x. It can return a vector of values, but the degrees of freedom and noncentrality parameter must be the same for all values of x.

Format

y = cdftnc(x, y, d);

Input

x Nx1 vector, values of upper limits of integrals.

v scalar, degrees of freedom, v > 0.

d scalar, noncentrality parameter.

This is the square root of the noncentrality parameter that sometimes goes under the symbol lambda. (See Scheffe, *The Analysis of Variance*, App. IV. 1959.)

Output

Nx1 vector, integrals from $-\infty$ to x of noncentral t.

Source

cdfnonc.src

See also

cdffnc, cdfchinc

Technical Notes Relation to cdftc:

```
cdftc(x,v) = 1 - cdftnc(x,v,0);
```

The formula used is based on the formula in *SUGI Supplemental Library User's Guide*. SAS Institute. 1983, 232 (which is attributed to Johnson and Kotz, 1970).

The formula used here is a modification of that formula. It has been tested against direct numerical integration, and against simulation experiments in which noncentral **t** random variates were generated and the cdf found directly.

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cdftvn

cdftvn

Purpose

Computes the cumulative distribution function of the standardized trivariate Normal density (lower tail).

Format

$$c = \text{cdftvn}(x1,x2,x3,rho12,rho23,rho31);$$

Input

- x1 Nx1 vector of upper limits of integration for variable 1.
- *x*2 Nx1 vector of upper limits of integration for variable 2.
- *x3* Nx1 vector of upper limits of integration for variable 3.
- *rho12* scalar or NX1 vector of correlation coefficients between the two variables x1 and x2.
- *rho23* scalar or Nx1 vector of correlation coefficients between the two variables *x*2 and *x*3.
- *rho31* scalar or Nx1 vector of correlation coefficients between the two variables x1 and x3.

Output

Nx1 vector containing the result of the triple integral from $-\infty$ to x1, $-\infty$ to x2, and $-\infty$ to x3 of the standardized trivariate Normal density:

Remarks

Allowable ranges for the arguments are:

$$-\infty < x1 < +\infty$$

$$-\infty < x2 < +\infty$$

$$-\infty < x3 < +\infty$$

$$-1 < rho 12 < 1$$

$$-1 < rho23 < 1$$

In addition, *rho12*, *rho23*, and *rho31* must come from a legitimate positive definite matrix. A -1 is returned for those rows with invalid inputs.

A separate integral is computed for each row of the inputs.

The first 3 arguments (x1,x2,x3) must be the same length, N. The second 3 arguments (rho12,rho23,rho31) must also be the same length, and this

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cdftvn

length must be N or 1. If it is 1, then these values will be expanded to apply to all values of x1,x2,x3. All inputs must be column vectors.

To find the integral under a general trivariate density, with x1, x2, and x3 having nonzero means and any positive standard deviations, transform by subtracting the mean and dividing by the standard deviation. For example:

```
x1 = (x1 - meanc(x1)) / stdc(x1);
```

See also

cdfn, cdfbvn

Technical Notes

The absolute error for **cdftvn** is approximately $\pm 2.5e-8$ for the entire range of arguments.

References

Daley, D.J. "Computation of Bi- and Tri-variate Normal Integral." *Appl. Statist.* Vol. 23 No. 3, 1974, 435-38.

Steck, G.P. "A Table for Computing Trivariate Normal Probabilities." *Ann. Math. Statist.* Vol. 29, 780-800.

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cdir

cdir

Purpose Returns the current directory.

Format y = cdir(s);

string, if the first character is 'A'-'Z' and the second character is a colon ':' then that drive will be used. If not, the current default drive will be used.

Output y string containing the drive and full path name of the current directory on the specified drive.

Remarks If the current directory is the root directory, the returned string will end with a backslash, otherwise it will not.

A null string or scalar zero can be passed in as an argument to obtain the current drive and path name.

Example x = cdir(0);
y = cdir("d:");
print x;
print y;

C:\GAUSS
D:\

See also files

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Purpose Round up toward $+\infty$.

Format y = ceil(x);

Input x NxK matrix.

Output y NxK matrix.

Remarks This rounds every element in the matrix x to an integer. The elements are rounded up toward $+\infty$.

Example x = 100*rndn(2,2);

y = ceil(x);

 $x = \begin{array}{rrr} 77.68 & -14.10 \\ 4.73 & -158.88 \end{array}$

 $y = \begin{array}{cc} 78.00 & -14.00 \\ 5.00 & -158.00 \end{array}$

See also floor, trunc

ChangeDir

ChangeDir

Purpose Changes the working directory.

Format d = ChangeDir(s);

Input *s* string, directory to change to.

Output d string, new working directory, or null string if change failed.

See also chdir

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Purpose Changes working directory.

Format chdir dirstr;

Input *dirstr* literal or ^string, directory to change to.

Remarks This is for interactive use. Use **ChangeDir** in a program.

If the directory change fails, **chdir** prints an error message.

chol

Purpose Computes the Cholesky decomposition of a symmetric, positive definite matrix.

Format y = chol(x);

Input x NxN matrix.

Output y NxN matrix containing the Cholesky decomposition of x.

Remarks y is the "square root" matrix of x. That is, it is an upper triangular matrix such that x = y'y.

chol does not check to see that the matrix is symmetric. **chol** will look only at the upper half of the matrix including the principal diagonal.

If the matrix *x* is symmetric but not positive definite, either an error message or an error code will be generated, depending on the lowest order bit of the trap flag:

trap 0 Print error message and terminate program.

trap 1 Return scalar error code 10.

See **scalerr** and **trap** for more details about error codes.

Example x = moment(rndn(100,4),0);

$$y = chol(x);$$

$$ypy = y'y;$$

$$x = \begin{cases} 90.746566 & -6.467195 & -1.927489 & -15.696056 \\ -6.467195 & 87.806557 & 6.319043 & -2.435953 \\ -1.927489 & 6.319043 & 101.973276 & 4.355520 \\ -15.696056 & -2.435953 & 4.355520 & 99.042850 \end{cases}$$

$$y = \begin{cases} 9.526099 - 0.678892 & -0.202338 - 1.647690 \\ 0.000000 & 9.345890 & 0.661433 - 0.380334 \\ 0.000000 & 0.000000 & 10.074465 & 0.424211 \\ 0.000000 & 0.000000 & 0.000000 & 9.798130 \end{cases}$$

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 $ypy = \begin{cases} 90.746566 & -6.467195 & -1.927489 & -15.696056 \\ -6.467195 & 87.806557 & 6.319043 & -2.435953 \\ -1.927489 & 6.319043 & 101.973276 & 4.355520 \\ -15.696056 & -2.435953 & 4.355520 & 99.042850 \end{cases}$

See also crout, solpd

choldn

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choldn

Purpose Performs a Cholesky downdate of one or more rows on an upper triangular matrix.

Format r = choldn(C,x);

Input *C* KxK upper triangular matrix.

x NxK matrix, the rows to downdate C with.

Output r KxK upper triangular matrix, the downdated matrix.

Remarks *C* should be a Cholesky factorization.

choldn(C,x) is equivalent to chol(C'C - x'x), but choldn is numerically much more stable.

Warning: it is possible to render a Cholesky factorization non-positive definite with **choldn**. You should keep an eye on the ratio of the largest diagonal element of r to the smallest — if it gets very large, r may no longer be positive definite. This ratio is a rough estimate of the condition number of the matrix.

Example

See also cholup

3-105

cholsol

cholsol

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Purpose Solves a system of linear equations given the Cholesky factorization of the system.

Format x = cholsol(b,C);

Input b NxK matrix.

C NxN matrix.

Output x NxK matrix.

Remarks C is the Cholesky factorization of a linear system of equations A. x is the solution for Ax = b. b can have more than one column. If so, the system is solved for each column, i.e., A*x[.,i] = b[.,i].

cholsol(eye(N),C) is equivalent to invpd(A). Thus, if you have the Cholesky factorization of A, cholsol is the most efficient way to obtain the inverse of A.

Example

x = cholsol(b,C);

x = -1.52686768

3.21579513

-1.94396905

3.00506436 2.65577048 3.08742844

 $A0 = 2.65577048 \ 3.55545737 \ 3.42362593$

3.08742844 3.42362593 4.02095978

cholup

cholup

Purpose Performs a Cholesky update of one or more rows on an upper triangular matrix.

Format r = cholup(C,x);

Input *C* KxK upper triangular matrix.

x NxK matrix, the rows to update C with.

Output r KxK upper triangular matrix, the updated matrix.

Remarks *C* should be a Cholesky factorization.

cholup(C,x) is equivalent to chol(C'C + x'x), but cholup is numerically much more stable.

Example let C[3,3] = 18.87055964 15.32294435 8.04947012

0 9.30682813 -2.12009339 0 0 7.62878355;

let x[2,3] = 1.76644971 7.49445820 9.79114666 6.87691156 4.41961438 4.32476921;

r = cholup(C,x);

20.16210005 16.50544413 9.86676135

r = 0.000000000 11.16601462 2.97761666

0.00000000 0.00000000 11.65496052

See also choldn

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chrs

chrs

Purpose b

Converts a matrix of ASCII values into a string containing the appropriate characters.

Format

```
y = chrs(x);
```

Input

NxK matrix.

Output

string of length N*K containing the characters whose ASCII values are equal to the values in the elements of x.

Remarks

This function is useful for embedding control codes in strings and for creating variable length strings when formatting printouts, reports, etc.

Example

```
n = 5i
```

```
print chrs(ones(n,1)*42);
```

Since the ASCII value of the asterisk character is 42, the program above will print a string of **n** asterisks.

```
y = chrs(67~65~84);
print y;
```

CAT

See also

```
vals, ftos, stof
```

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clear

clear

Purpose Clears space in memory by setting matrices equal to scalar zero.

Format clear x, y;

Remarks clear x; is equivalent to x = 0;.

Matrix names are retained in the symbol table after they are cleared.

Matrices can be **clear**'ed even though they have not previously been

defined. clear can be used to initialize matrices to scalar 0.

Example clear x;

See also clearg, new, show, delete

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clearg

clearg

Purpose This command clears global symbols by setting them equal to scalar zero.

Format clearg a,b,c;

Output a,b,c scalar global matrices containing 0.

Remarks clearg x; is equivalent to x = 0;, where x is understood to be a

global symbol. **clearg** can be used to initialize symbols not previously referenced. This command can be used inside procedures to clear global

matrices. It will ignore any locals by the same name.

Example x = 45;

clearg x;

x = 0.0000000

See also clear, delete, new, show, local

3-110

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close

Purpose Close a GAUSS file.

Format y = close(handle);

Input handle scalar, the file handle given to the file when it was opened with the open, create, or fopen command.

Output y scalar, 0 if successful, -1 if unsuccessful.

Remarks

handle is the scalar file handle created when the file was opened. It will contain an integer which can be used to refer to the file.

close will close the file specified by handle, and will return a 0 if successful and a -1 if not successful. The handle itself is not affected by **close** unless the return value of **close** is assigned to it.

If fI is a file handle and it contains the value 7, then after:

call
$$close(fl)$$
;

the file will be closed but fI will still have the value 7. The best procedure is to do the following:

$$fI = close(fI);$$

This will set fI to 0 upon a successful close.

It is important to set unused file handles to zero because both **open** and **create** check the value that is in a file handle before they proceed with the process of opening a file. During **open** or **create**, if the value that is in the file handle matches that of an already open file, the process will be aborted and a "File already open" message will be given. This gives you some protection against opening a second file with the same handle as a currently open file. If this happened, you would no longer be able to access the first file.

An advantage of the **close** function is that it returns a result which can be tested to see if there were problems in closing a file. The most common reason for having a problem in closing a file is that the disk on which the file is located is no longer in the disk drive — or the handle was invalid. In both of these cases, **close** will return a -1.

Files are not automatically closed when a program terminates. This allows users to run a program that opens files, and then access the files

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close

from interactive mode after the program has been run. Files are automatically closed when GAUSS exits to the operating system or when a program is terminated with the **end** statement. **stop** will terminate a program but not close files.

As a rule it is good practice to make **end** the last statement in a program, unless further access to the open files is desired from interactive mode. You should close files as soon as you are done writing to them to protect against data loss in the case of abnormal termination of the program due to a power or equipment failure.

The danger in not closing files is that anything written to the files may be lost. The disk directory will not reflect changes in the size of a file until the file is closed and system buffers may not be flushed.

```
Example
```

```
open f1 = dat1 for append;
y = writer(f1,x);
f1 = close(f1);
```

See also

closeall

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closeall

closeall

Purpose Close all currently open GAUSS files.

Format closeall;

closeall list of handles;

Remarks list

list_of_handles is a comma-delimited list of file handles.

closeal1 with no specified list of handles will close all files. The file handles will not be affected. The main advantage of using **closeal1** is ease of use; the file handles do not have to be specified, and one statement will close all files.

When a list of handles follows **closeall**, all files are closed and the file handles listed are set to scalar 0. This is safer than **closeall** without a list of handles because the handles are cleared.

It is important to set unused file handles to zero because both **open** and **create** check the value that is in a file handle before they proceed with the process of opening a file. During **open** or **create**, if the value that is in the file handle matches that of an already open file, the process will be aborted and a "File already open" message will be given. This gives you some protection against opening a second file with the same handle as a currently open file. If this happened, you would no longer be able to access the first file.

Files are not automatically closed when a program terminates. This allows users to run a program that opens files, and then access the files from interactive mode after the program has been run. Files are automatically closed when GAUSS exits to the operating system or when a program is terminated with the **end** statement. **stop** will terminate a program but not close files.

As a rule it is good practice to make **end** the last statement in a program, unless further access to the open files is desired from interactive mode. You should close files as soon as you are done writing to them to protect against data loss in the case of abnormal termination of the program due to a power or equipment failure.

The danger in not closing files is that anything written to the files may be lost. The disk directory will not reflect changes in the size of a file until the file is closed and system buffers may not be flushed.

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Example open f1 = dat1 for read; open f2 = dat1 for update; x = readr(f1, rowsf(f1));x = sqrt(x);call writer(f2,x); closeall f1,f2; See also close, open

cls

Purpose Clear the window.

Format cls;

Portability UNIX 3.2 only

cls clears the active graphic panel. For Text graphic panels, this means the graphic panel buffer is cleared to the background color. For TTY graphic panels, the current output line is panned to the top of the graphic panel, effectively clearing the display. The output log is still intact. To clear the output log of a TTY graphic panel, use WinClearTTYLog. For PQG graphic panels, the graphic panel is cleared to the background color, and the related graphics file is truncated to zero length.

UNIX 3.5+

cls will clear the screen on some terminals.

Windows

cls clears the Command window if you're in Cmnd I/O mode, the Output window if you're in Split I/O mode.

OS/2

cls clears the Main window.

Remarks

This command will cause the window to clear and will locate the cursor at the upper left hand corner of the window.

See also locate

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```
Purpose Allows a new variable to be created (coded) with different values depending upon which one of a set of logical expressions is true.
```

Format y = code(e, v);

Input

- NxK matrix of 1's and 0's. Each column of this matrix is created by a logical expression using "dot" conditional and boolean operators. Each of these expressions should return a column vector result. The columns are horizontally concatenated to produce *e*. If more than one of these vectors contains a 1 in any given row, the **code** function will terminate with an error message.
- v (K+1)x1 vector containing the values to be assigned to the new variable.

Output

y Nx1 vector containing the new values.

Remarks

If none of the K expressions is true, the new variable is assigned the default value, which is given by the last element of v.

Example

```
let x1 = 0 /* column vector of original values */
5
```

10

15

20;

```
let v = 1 /* column vector of new values */
2
```

3; /* the last element of v is the

:: "default"

* /

code

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For every row in e, if a 1 is in the first column, the first element of v is used. If a 1 is in the second column, the second element of v is used, and

code

so on. If there are only zeros in the row, the last element of v is used. This is the default value.

If there is more than one 1 in any row of e, the function will terminate with an error message.

Source datatran.src

See also recode, substute

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code (dataloop)

code (dataloop)

Purpose

Creates new variables with different values based on a set of logical expressions.

Format

code [#] [\$] var [default defval] with
 val_1 for expression_1,
 val_2 for expression_2,

.

val_n for expression_n;

Input

var literal, the new variable name.

defval scalar, the default value if none of the expressions are

TRUE.

val scalar, value to be used if corresponding expression is

TRUE.

expression logical scalar-returning expression that returns nonzero

TRUE or zero FALSE.

Remarks

If '\$' is specified, the new variable will be considered a character variable. If '#' or nothing is specified, the new variable will be considered numeric.

The logical expressions must be mutually exclusive; i.e., only one may return *TRUE* for a given row (observation).

Any variables referenced must already exist, either as elements of the source data set, as externs, or as the result of a previous **make**, **vector**, or **code** statement.

If no default value is specified, 999 is used.

a

b

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g

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u

V

X y Z

code (dataloop)

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е

g

h

k

m

n

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q

t

u

V

W

Example code agecat default 5 with 1 for age < 21,

"FEMALE" for gender == 0;

See also recode

color

color

Purpose Set pixel, text, background color, or VGA palette color registers.

Format y = color(cv)

Input *cv* scalar, 2x1 or 3x1 vector of color values or Nx4 matrix of palette color values. See Portability, below, for platform specifics.

If the input vector is smaller than 3x1 or the corresponding element in the input vector is -1, the corresponding color will be left unchanged.

If the input is an Nx4 matrix, it will initialize the VGA palette (DOS) or active graphic panel's colormap (UNIX) with user-defined RGB colors interpreted as follows:

[N,1] palette register index 0-255

[N,2] red value 0-63

[N,3] green value 0-63

[N,4] blue value 0-63

Output

vector, or Nx4 matrix the same size as the input which contains the original color values or palette values.

Portability DOS

v

[1] pixel color

[2] text color

[3] ignored

UNIX 3.2 only

color affects the active graphic panel. X supports foreground and background colors. The **color** command makes no distinction between text and pixel colors; both affect the foreground color of the active graphic panel. If both a pixel color and text color are specified, the pixel color will be ignored, and the text color will be used to set the foreground color. Thus:

[1] foreground

or

[1] ignored

a

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c d

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color

[2] foreground

or

- [1] ignored
- [2] foreground
- [3] background

OS/2, Windows, UNIX 3.5+

This function is not supported under OS/2 or Windows.

Remarks This changes the window colors for your program's output. The editor and interactive mode will not be affected.

See "Colors Appendix" on page B-1 for a color value table.

Under DOS, the VGA color palette registers may be set only if the display adapter has been already been initialized to VGA graphics mode 19 (320x200, 256 colors) with the **setvmode** command. The registers will retain the new values until the adapter is reset to text mode, which resets the palette to the default VGA colors.

This function is useful for obtaining 64 shades of a single color and/or mixing colors to user-specification.

See also graph, line, setvmode

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cols, colsf

Purpose cols returns the number of columns in a matrix.

colsf returns the number of columns in a GAUSS data (.dat) file or GAUSS matrix (.fmt) file.

Format y = cols(x);yf = colsf(fh);

Input x any valid expression that returns a matrix.

fh file handle of an open file.

Output y number of columns in x.

yf number of columns in the file that has the handle fh.

Remarks If x is an empty matrix, rows(x) and cols(x) return 0.

For **colsf**, the file must be open.

Example x = rndn(100,3);

y = cols(x);

y = 3.000000

create fp = myfile with x,10,4;

b = colsf(fp);

b = 10.000000

See also rows, rowsf, show, lshow

a

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f

g h

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comlog

comlog

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u v

W

хуг

Purpose Controls logging of interactive mode commands to a disk file.

Format

comlog [file=filename] [on|off|reset];

Input

filename literal or ^string.

The **file**=*filename* subcommand selects the file to log interactive mode statements to. This can be any legal file name.

If the name of the file is to be taken from a string variable, the name of the string must be preceded by the ^ (caret) operator.

There is no default file name.

on, off, reset

literal, mode command:

on turns on command logging to the current log file.

If the file already exists, subsequent commands

will be appended.

off closes the log file and turns off command logging.

reset similar to the on subcommand, except that it resets

the log file by deleting any previous commands.

Remarks

Interactive mode statements are always logged into the file specified in the log_file configuration variable, regardless of the state of comlog.

The command **comlog file**=*filename* selects the file but does not turn on logging.

The command **comlog off** will turn off logging. The filename will remain the same. A subsequent **comlog on** will cause logging to resume. A subsequent **comlog reset** will cause the existing contents of the log file to be destroyed and a new file created.

The command **comlog** by itself will cause the name and status of the current log file to be printed in the window.

In interactive mode under DOS, **F10** will load the current log file into the editor if logging is **on**. If logging is **off**, the default log file listed in the **log_file** configuration variable will be loaded into the editor.

compile

Purpose

Compiles a source file to a compiled code file. See also "Compiler" in the *User's Guide*.

Format

compile source fname;

Input

source literal or ^string, the name of the file to be compiled.

fname literal or *string, optional, the name of the file to be created. If

not given, the file will have the same filename and path as

source. It will have a .gcg extension.

Remarks

The *source* file will be searched for in the **src_path** if the full path is not specified and it is not present in the current directory.

The source file is a regular DOS text file containing a GAUSS program. There can be references to global symbols, Run-Time Library references, etc.

If there are **library** statements in *source*, they will be used during the compilation to locate various procedures and symbols used in the program. Since all of these library references are resolved at compile time, the **library** statements are not transferred to the compiled file. The compiled file can be run without activating any libraries.

If you do not want extraneous matter saved in the compiled image, put a **new** at the top of the *source file* or execute a **new** from interactive level before compiling.

The program saved in the compiled file can be run with the **run** command. If no extension is given, the **run** command will look for a file with the correct extension for the version of GAUSS. The **src_path** will be used to locate the file if the full path name is not given and it is not located on the current directory.

When the compiled file is **run**, all previous symbols and procedures are deleted before the program is loaded. It is therefore unnecessary to execute a **new** before **run**'ning a compiled file.

If you want line number records in the compiled file you can put a **#lineson** statement in the *source* file or turn line tracking on from the Options menu.

Do not try to include compiled files with **#include**.

a

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b h

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v

W

compile

Example

compile qxy.e;

In this example, the **src_path** would be searched for qxy.e, which would be compiled to a file called qxy.gcg on the same subdirectory qxy.e was found.

compile qxy.e xy;

In this example, the **src_path** would be searched for qxy.e which would be compiled to a file called xy.gcg on the current subdirectory.

See also r

run, use, saveall

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V

W

x y z

complex

complex

See also

imag, real

Converts a pair of real matrices to a complex matrix. Purpose Format z = complex(xr,xi);Input NxK real matrix, the real elements of z. xr NxK real matrix or scalar, the imaginary elements of z. xi Output NxK complex matrix. Example $x = \{ 46,$ 98 }; $y = \{ 35,$ 1 7 }; t = complex(x,y);4.0000000 + 3.0000000i 6.0000000 + 5.0000000i $9.0000000 + 1.0000000i \ 8.0000000 + 7.0000000i$

a

b

c

е

f g

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V

con

con

a b d h m n 0 p q u V

Purpose Requests input from the keyboard, and returns it in a matrix. **Format** x = con(r,c);Input scalar, row dimension of matrix. scalar, column dimension of matrix. c**Output** RxC matrix. **Remarks** Enter ? to get a help screen at the **con** function prompt. The following commands are available: Up one row IJ First row u d Down one row D Last row 1 Left one column L First column Last column Right one column R r t First element Last element b Goto element Goto element of vector g Move horizontally, default h Move vertically V Move diagonally Show size of matrix s Display element as numeric, default n Display element as character С exp(1)е р pi missing value

help

?

W

x y z

con

x exit

Use a leading single quote for character input.

See also cons, let, load

a

b

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d

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f g

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XX7

cond

cond

a b

c

d

e

g

h

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k

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m

0

р

q

1

f

u

V

W

хух

```
Purpose This procedure will compute the condition number of a matrix using the singular value decomposition.
```

Format c = cond(x);

Input x NxK matrix.

Output c scalar, an estimate of the condition number of x. This equals the ratio of the largest singular value to the smallest. If the smallest singular value is zero or not all of the singular values can be computed, the return value is 10^{300} .

Example $x = \{ 4 \ 2 \ 6, \\ 8 \ 5 \ 7, \\ 3 \ 8 \ 9 \ \};$ y = cond(x);y = 9.8436943

Source svd.src

conj

conj

```
Purpose
              Returns the complex conjugate of a matrix.
 Format
             y = conj(x);
    Input
                    NxK matrix.
  Output
                    NxK matrix, the complex conjugate of x.
              v
Remarks
             Compare conj with the transpose (') operator.
Example
                    { 1+9i 2,
              x =
                       4+4i 5i,
                       7i 8-2i };
              y = conj(x);
                   1.0000000 + 9.0000000i
                                                      2.0000000
                   4.0000000 + 4.0000000i \ 0.0000000 + 5.0000000i
                   0.0000000 + 7.0000000i \ 8.0000000 - 2.0000000i
                   1.0000000 - 9.0000000i
                                                      2.0000000
                   4.0000000 - 4.0000000i \ 0.0000000 - 5.0000000i
              y =
                   0.0000000 - 7.0000000i \ 8.0000000 + 2.0000000i
```

a

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e.

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cons

cons

Purpose Retrieves a character string from the keyboard.

Format x = cons;

Output The characters entered from the keyboard. The output will be of type string.

Remarks If you are working in terminal mode GAUSS will not "see" any input until you press ENTER. *x* is assigned the value of a character string typed in at the keyboard. The program will pause to accept keyboard input. The maximum length of the string that can be entered is 254 characters. The program will resume execution when the ENTER key is pressed. The standard DOS editing keys will be in effect.

Example x = cons;

At the cursor enter:

probability

x = "probability"

See also con

d

g

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k

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p

q

1

t

u

V

W

continue

a

b

c

d

e

f

g

h

j

k

m

n

0

p

q

t

u

V

хух

continue

```
Purpose
             Jumps to the top of a do or for loop.
 Format
             continue;
Remarks
             This command works just like in C.
Example
             x = rndn(4,4);
             r = 0;
             do while r < rows(x);
                r = r + 1;
                c = 0;
                do while c < cols(x); /* continue jumps here */</pre>
                    c = c + 1;
                    if c == ri
                       continue;
                    endif;
                    x[r,c] = 0;
                 endo;
             endo;
                   -1.032195
                              0000000 0.000000
                                                 0.000000
                    0.000000 - 1.033763 \ 0.000000
                                                 0.000000
             x =
                    0.000000 0.000000 0.061205
                                                 0.000000
                    0.000000 \quad 0.000000 \quad 0.000000 \quad -0.225936
```

contour

contour

a **Purpose** To graph a matrix of contour data. h Library pgraph Format contour (x,y,z); d Input 1xK vector, the X axis data. K must be odd. Nx1 vector, the Y axis data. N must be odd. y Z. NxK matrix, the matrix of height data to be plotted. **Global Input** plev KX1 vector, user-defined contour levels for **contour**. h Default 0. Nx1 or Nx2 vector. This controls the Z level colors. See pzclr **surface** for a complete description of how to set this global. k Remarks A vector of evenly spaced contour levels will be generated automatically from the z matrix data. Each contour level will be labeled. For unlabeled contours, use ztics. m To specify a vector of your own unequal contour levels, set the vector plev before calling contour. To specify your own evenly spaced contour levels, see **ztics**. 0

Source pcontour.src

See also surface

p

q

t

u

V

W

conv

Purpose Computes the convolution of two vectors.

Format c = conv(b, x, f, l);

Input b Nx1 vector.

x Lx1 vector.

f scalar, the first convolution to compute.

l scalar, the last convolution to compute.

Output c Qx1 result, where Q = (l - f + 1).

If f is 0, the first to the l'th convolutions are computed. If l is 0, the f'th to the last convolutions are computed. If f and l are both zero, all the convolutions are computed.

Remarks If x and b are vectors of polynomial coefficients, this is the same as multiplying the two polynomials.

Example $x = \{ 1, 2, 3, 4 \};$

$$y = \{ 5,6,7,8 \};$$

z1 = conv(x,y,0,0);

z2 = conv(x,y,2,5);

5

16

34

z1 = 60

61

52

12

32

a

b

c

d e

f

g h

i

k

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n

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conv

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61

See also polymult

coreleft

coreleft

Purpose Returns the amount, in bytes, of free workspace memory.

Format y = coreleft;

Output y scalar, number of bytes free.

Portability DOS only

All others will return the **coreleft** value specified in the GAUSS configuration (.cfg) file.

Remarks

The amount of free memory is dynamic and can change rapidly as expressions and procedures are being executed. **coreleft** returns the amount of workspace memory free at the time it is called. Workspace memory is used for storing matrices, strings, procedures, and for manipulating matrices and strings.

This function can be used to write programs that automatically adjust their use of memory so they do not crash with the "Insufficient memory" error if they are used on machines with less free memory than the one used for development, or if the size of the data used becomes larger. A common use is to adjust the number of rows that are read per iteration of a read loop in programs that access data from a disk.

Example

```
open fp = myfile;
k = colsf(fp); /* columns in file */
fac = 4;
/* check amount of memory available */
nr = coreleft/(fac*k*8);
```

In this example, **nr**, the number of rows to read, is computed by taking the number of bytes free (**coreleft**) divided by **fac*k*8**. **fac** is a guesstimate of the number of copies of the data read each iteration that the algorithm we are using will require, plus a little. **k*8** is the number of columns times the number of bytes per element.

See also dfree, new

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V

W

corrm, corrvc, corrx

corrm, corrvc, corrx

Purpose Computes a correlation matrix.

Format cx = corrm(m);

cx = corrvc(vc);

cx = corrx(x);

Input m KxK moment (x'x) matrix. A constant term MUST have been the

first variable when the moment matrix was computed.

vc KxK variance-covariance matrix (of data or parameters).

x NxK matrix of data.

Output cx PxP correlation matrix. For corrm, P = K-1. For corrvc

and corrx, P = K.

Source corr.src

See also momentd

COS

Purpose Returns the cosine of its argument.

Format y = cos(x);

Input x NxK matrix.

Output y NxK matrix.

Remarks For real matrices, *x* should contain angles measured in radians.

To convert degrees to radians, multiply the degrees by $\frac{\pi}{180}$.

Example $x = \{ 0, .5, 1, 1.5 \};$

y = cos(x);

1.00000000

 $y = \begin{array}{c} 0.87758256 \\ 0.54030231 \end{array}$

0.07073720

See also atan, atan2, pi

a

b

c

A

f g

h

1

k

m

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q

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V

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cosh

a

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хух

cosh

Purpose Computes the hyperbolic cosine. **Format** $y = \cosh(x);$ Input NxK matrix. **Output** NxK matrix containing the hyperbolic cosines of the elements of $x = \{ -0.5, -0.25, 0, 0.25, 0.5, 1 \};$ Example x = x * pi;y = cosh(x);-1.570796-0.7853980.000000 x =0.785398 1.570796 3.141593 2.509178 1.324609 1.000000 v =1.324609 2.509178 11.591953 Source trig.src

3-140

counts

counts

Purpose Count the numbers of elements of a vector that fall into specified ranges.

Format c = counts(x, v);

Input x Nx1 vector containing the numbers to be counted.

v Px1 vector containing breakpoints specifying the ranges within which counts are to be made. The vector *v* MUST be sorted in ascending order.

Output c Px1 vector, the counts of the elements of x that fall into the regions:

$$x \le v[1],$$
 $v[1] < x \le v[2],$
 \vdots
 $v[p-1] < x \le v[p].$

Remarks If the maximum value of x is greater than the last element (the maximum value) of v, the sum of the elements of the result, c, will be less than N, the total number of elements in x.

If

b

c

d e

f

g h

i

K

m

n

0

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r

1

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V

W

counts

then

 $c = \frac{4}{1}$

The first category can be a missing value if you need to count missings directly. Also $+\infty$ or $-\infty$ are allowed as breakpoints. The missing value must be the first breakpoint if it is included as a breakpoint and infinities must be in the proper location depending on their sign. $-\infty$ must be in the [2,1] element of the breakpoint vector if there is a missing value as a category as well, otherwise it has to be in the [1,1] element. If $+\infty$ is included, it must be the last element of the breakpoint vector.

Example

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u

V

W

countwts

countwts

Purpose

Returns a weighted count of the numbers of elements of a vector that fall into specified ranges.

Format

c = countwts(x, v, w);

Input

x Nx1 vector, the numbers to be counted.

- v Px1 vector, the breakpoints specifying the ranges within which counts are to be made. This MUST be sorted in ascending order (lowest to highest).
- w Nx1 vector, containing weights.

Output

Px1 vector, the counts of the elements of x that fall into the regions:

$$x \le v[1],$$
 $v[1] < x \le v[2],$
 \vdots
 $v[p-1] < x \le v[p].$

That is, when x[i] falls into region j, the weight w[i] is added to the j^{th} counter.

Remarks

If any elements of x are greater than the last element of v, they will not be counted.

Missing values are not counted unless there is a missing in v. A missing value in v MUST be the first element in v.

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b

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е

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g h

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k

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n

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p q

42

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u

V

countwts

Example

 $x = \{ 1, 3, 2, 4, 1, 3 \};$ $w = \{ .25, 1, .333, .1, .25, 1 \};$

 $v = \{ 0, 1, 2, 3, 4 \};$

c = countwts(x,v,w);

0.000000

0.500000

c = 0.333000

2.00000

0.100000

a

b

c

d

е

f g

h

J

k

m

n

0

p

q

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V

W

create

Creates and opens a GAUSS data set for subsequent writing. **Purpose Format** create $\lceil vflag \rceil \rceil$ $\lceil complex \rceil$ fh = filename with vnames, col, dtyp, vtyp; create [vflag] [complex] fh = filename using comfile; Input vflag version flag. -v89 supported for read -v92 for read/write -v96 for read/write For details on the various versions, see "File I/O" in the *User's* Guide. The default format can be specified in gauss.cfg by setting the dat_fmt_version configuration variable. If dat fmt version is not set, the default is v96.

filename literal or ^string.

filename is the name to be given the file on the disk. The name can include a path if the directory to be used is not the current directory. This file will automatically be given the extension .dat. If an extension is specified, the .dat will be overridden. If the name of the file is to be taken from a string variable, the name of the string must be preceded by the ^ (caret) operator.

create... with...

vnames literal or *string or *character matrix.

vnames controls the names to be given to the columns of the data file. If the names are to be taken from a string or character matrix, the ^ (caret) operator must be placed before the name of the string or character matrix. The number of columns parameter, col, also has an effect on the way the names will be created. See below and see the examples for details on the ways names are assigned to a data file.

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W

хух

col scalar expression.

col is a scalar expression containing the number of columns in the data file. If col is 0, the number of columns will be controlled by the contents of vnames. If col is positive, the file will contain col columns and the names to be given each column will be created as necessary depending on the vnames parameter. See the examples.

dtyp scalar expression.

dtyp is the precision used to store the data. This is a scalar expression containing 2, 4, or 8, which is the number of bytes per element.

2 signed integer

4 single precision

8 double precision

Data Type	Digits	Range
integer	4	-32768 <= X <= 32767
single	6-7	$8.43x10^{-37} \le X \le 3.37x10^{+38}$
double	15-16	$4.19x10^{-307} \le X \le 1.67x10^{+308}$

If the integer type is specified, numbers will be rounded to the nearest integer as they are written to the data set. If the data to be written to the file contains character data, the precision must be 8 or the character information will be lost.

vtyp matrix, types of variables.

The types of the variables in the data set. If rows(vtyp)*cols(vtyp) < col only the first element is used. Otherwise nonzero elements indicate a numeric variable and zero elements indicate character variables. vtyp is ignored for v89 files.

create... using...

comfile literal or *string.

comfile is the name of a command file that contains the information needed to create the file. The default extension for the command file is .gcf, which can be overridden.

There are three possible commands in this file:

numvar *n str*:

outvar varlist;

outtyp dtyp;

numvar and **outvar** are alternate ways of specifying the number and names of the variables in the data set to be created.

When **numvar** is used, *n* is a constant which specifies the number of variables (columns) in the data file and *str* is a string literal specifying the prefix to be given to all the variables. Thus:

says that there are 10 variables and that they are to be named **xx01** through **xx10**. The numeric part of the names will be padded on the left with zeros as necessary so the names will sort correctly:

xx1,	 xx9	1-9 names
xx01,	 xx10	10-99 names
xx001,	 xx100	100-999 names
xx0001,	 xx1000	1000-8100 names

If *str* is omitted, the variable prefix will be "X".

When **outvar** is used, *varlist* is a list of variable names, separated by spaces or commas. For instance:

specifies that there are to be 3 variables per row of the data set, and that they are to be named **x1**, **x2**, **zed**, in that order.

outtyp specifies the precision. It can be a constant: 2, 4, or 8, or it can be a literal: I, F, or D. For an explanation of the available data types, see *dtyp* in **create...** with..., previously.

The **outtyp** statement does not have to be included. If it is not, then all data will be stored in 4 bytes as single precision floating point numbers.

a b

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k

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n

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p q

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S

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V

Output *fh* scalar.

fh is the file handle which will be used by most commands to refer to the file within GAUSS. This file handle is actually a scalar containing an integer value that uniquely identifies each file. This value is assigned by GAUSS when the **create** (or **open**) command is executed.

Remarks

If the **complex** flag is included, the new data set will be initialized to store complex number data. Complex data is stored a row at a time, with the real and imaginary halves interleaved, element by element.

Example

```
let vnames = age sex educat wage occ;
create f1 = simdat with ^vnames,0,8;
obs = 0;
nr = 1000;
do while obs < 10000;
  data = rndn(nr,colsf(f1));
  if writer(f1,data) /= nr;
     print "Disk Full";
  end;
  endif;
  obs = obs+nr;
endo;
closeall f1;</pre>
```

uses create... with... to create a double precision data file called simdat.dat on the default drive with 5 columns. The writer command is used to write 10000 rows of Normal random numbers into the file. The variables (columns) will be named: AGE, SEX, EDUCAT, WAGE, OCC.

Following are examples of the variable names that will result when using a character vector of names in the argument to the **create** function.

a

b

А

Е

ø

h

:

k

m

0

р

q

r

0

u

V

W

```
vnames = { AGE PAY SEX JOB };
typ = { 1, 1, 0, 0 };
create fp = mydata with ^vnames,0,2,typ;
```

The names in this example will be: AGE PAY SEX JOB

AGE and PAY are numeric variables, SEX and JOB are character variables.

```
create fp = mydata with ^vnames,3,2;
```

The names will be: AGE PAY SEX

```
create fp = mydata with ^vnames,8,2;
```

The names will now be: AGE PAY SEX JOB1 JOB2 JOB3 JOB4 JOB5

If a literal is used for the *vnames* parameter, the number of columns should be explicitly given in the *col* parameter and the names will be created as follows:

```
create fp = mydata with var,4,2;
```

giving the names: var1 var2 var3 var4

The next example assumes a command file called comd.gcf containing the following lines created using a text editor:

```
outvar age, pay, sex;
outtyp i;
```

Then the following could be used to write 100 rows of random integers into a file called smpl.dat in the subdirectory called /gauss/data:

```
filename = "/gauss/data/smpl";
create fh = ^filename using comd;
x = rndn(100,3)*10;
if writer(fh,x) /= rows(x);
  print Disk Full;
  end;
endif;
closeall fh;
```

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For platforms using the backslash as a path separator, remember that two backslashes (\\) are required to enter one backslash inside double quotes. This is because a backslash is the escape character used to embed special characters in strings.

See also open, readr, writer, eof, close, output, iscplxf

a

b

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u

V

w x y z

crossprd

crossprd

Purpose Computes the cross-products (vector products) of sets of 3x1 vectors.

Format z = crossprd(x,y);

Input x 3xK matrix, each column is treated as a 3x1 vector.

y 3xK matrix, each column is treated as a 3x1 vector.

Output z 3xK matrix, each column is the cross-product (sometimes called vector product) of the corresponding columns of x

and y.

Remarks The cross-product vector z is orthogonal to both x and y. sumc $(x \cdot *z)$ and sumc $(y \cdot *z)$ will be Kx1 vectors all of whose elements are 0

(except for rounding error).

Example

 $x = \{ 10 \ 4, \\ 11 \ 13, \\$

14 13 };

 $y = \{ 311, 512,$

z = crossprd(x,y);

7 9 };

7.0000000 -39.000000

z = -28.000000 107.00000

17.000000 -95.000000

Source

crossprd.src

a

b

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е

f

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О

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t

u

V

W

crout

crout

a

b

c

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0

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V

W

хуг

Purpose Computes the Crout decomposition of a square matrix without row pivoting, such that: X = LU.

Format y = crout(x);

Input x NxN square nonsingular matrix.

Output y NxN matrix containing the lower (L) and upper (U) matrices of the Crout decomposition of x. The main diagonal of y is the main diagonal of the lower matrix L. The upper matrix has an implicit main diagonal of ones. Use **lowmat** and **upmat1** to extract the L and U matrices from y.

Remarks Since it does not do row pivoting, it is intended primarily for teaching purposes. (See **croutp** for a decomposition with pivoting.)

Example $X = \{ 1 \ 2 \ -1, \\ 2 \ 3 \ -2, \\ 1 \ -2 \ 1 \};$

y = crout(x);
L = lowmat(y);
U = upmat1(y);

 $y = \begin{array}{ccc} 1 & 2 & -1 \\ 2 & -1 & 0 \\ 1 & -4 & 2 \end{array}$

 $L = \begin{array}{ccc} 1 & 0 & 0 \\ 2 & -1 & 0 \\ 1 & -4 & 2 \end{array}$

crout

$$U = \begin{array}{cccc} 1 & 2 & -1 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{array}$$

See also croutp, chol, lowmat, lowmat1, lu, upmat, upmat1

a

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croutp

croutp

а

Purpose

Computes the Crout decomposition of a square matrix with partial (row) pivoting.

Format

y = croutp(x);

Input

x NxN square nonsingular matrix.

Output

(N+1)xN matrix containing the lower (L) and upper (U) matrices of the Crout decomposition of a permuted x. The N+1 row of the matrix y gives the row order of the y matrix. The matrix must be reordered prior to extracting the L and U matrices. Use **lowmat** and **upmat1** to extract the L and U matrices from the reordered y matrix.

Example

This example illustrates a procedure for extracting L and U of the permuted x matrix. It continues by sorting the result of LU to compare with the original matrix x.

8

b

d

e

1

h

. .

j

k

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m

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р

q

r

S

u

V

W

croutp

$$X = \begin{array}{ccc} 1 & 2 & -1 \\ 2 & 3 & -2 \\ 1 & -2 & 1 \end{array}$$

$$y = \begin{array}{cccc} 1 & 0.5 & 0.2857 \\ 2 & 1.5 & -1 \\ 1 & -3.5 & -0.5714 \\ 2 & 3 & 1 \end{array}$$

$$r = 4$$

$$indx = \begin{array}{c} 2\\ 3\\ 1 \end{array}$$

$$z = 1.5 -1$$

$$z = 1 -3.5 -0.5714$$

$$1 0.5 0.2857$$

$$L = \begin{array}{ccc} 2 & 0 & 0 \\ 1 & -3.5 & 0 \\ 1 & 0.5 & 0.2857 \end{array}$$

$$U = \begin{array}{ccc} 1 & 1.5 & -1 \\ 0 & 1 & -0.5714 \\ 0 & 0 & 1 \end{array}$$

$$q = \begin{array}{cccc} 1 & 1 & 2 & -1 \\ 2 & 2 & 3 & -2 \\ 3 & 1 & -2 & 1 \end{array}$$

a

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V

croutp

 $x2 = \begin{array}{rrr} 1 & 2 - 1 \\ 2 & 3 - 2 \\ 1 - 2 & 1 \end{array}$

See also crout, chol, lowmat, lowmat1, lu, upmat, upmat1

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csrcol, csrlin

csrcol, csrlin

Purpose Returns the position of the cursor.

Format y = csrcol;

y = csrlin;

Portability UNIX 3.2 only

csrcol returns the cursor column for the active graphic panel. For Text graphic panels, this value is the cursor column with respect to the text buffer. For TTY graphic panels, this value is the cursor column with respect to the current output line, i.e., it will be the same whether the text is wrapped or not. For PQG graphic panels, this value is meaningless.

csrlin returns the cursor line for the active graphic panel. For Text graphic panels, this value is the cursor row with respect to the text buffer. For TTY graphic panels, this value is the current output line number (i.e., the number of lines logged + 1). For PQG graphic panels, this value is meaningless.

UNIX 3.5+

csrcol and csrlin always return 1.

OS/2, Windows

csrcol returns the cursor column with respect to the current output line, i.e., it will return the same value whether the text is wrapped or not. **csrlin** returns the cursor line with respect to the top line in the graphic panel.

DOS

Under DOS, columns are usually numbered 1-80, rows are usually numbered 1-25. **setymode** will return the current window dimensions.

Remarks

y will contain the current column or row position of the cursor in the graphic panel. The upper left corner is (1,1).

csrcol returns the column position of the cursor. **csrlin** returns the row position.

The **locate** statement allows the cursor to be positioned at a specific row and column.

Example r = csrlin;

a

b

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0

p q

r

3

u

V

W

csrcol, csrlin

c = csrcol;
cls;
locate r,c;

In this example the window is cleared without affecting the cursor position.

See also cls, locate, lpos

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csrtype

csrtype

Purpose To set the cursor shape.

Format old = csrtype(mode);

Portability UNIX

This function is not supported in terminal mode.

OS/2, Windows

This function is not supported under OS/2 or Windows.

Input *mode* scalar, cursor type to set.

DOS

0 cursor off

1 normal cursor

2 large cursor

UNIX 3.2

0 cursor off

1 normal cursor

2 large cursor

3 triangular cursor

Output *old* scalar, original cursor type.

Remarks Under DOS, this function will set the same cursor shape that GAUSS is

already using for its three modes.

Example x = csrtype(2);

See also csrcol, csrlin

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cumprodc

cumprodc

a b

Purpose

Computes the cumulative products of the columns of a matrix.

Format

y = cumprodc(x);

Input

x NxK matrix.

e

d

h

m

n

0

p

q

Output

NxK matrix containing the cumulative products of the columns of *x*.

Remarks

This is based on the recursive series **recsercp**. **recsercp** could be called directly as follows:

recserp(x,zeros(1,cols(x)))

to accomplish the same thing.

Example

 $x = \{ 1 -3, 2 2,$

3 -1 };

y = cumprodc(x);

1.00 - 3.00

y = 2.00 - 6.00

6.00 6.00

Source

cumprodc.src

See also

cumsumc, recsercp, recserar

u

V

W

cumsumc

cumsumc

Purpose Computes the cumulative sums of the columns of a matrix.

Format y = cumsumc(x);

Input x NxK matrix.

Output y NxK matrix containing the cumulative sums of the columns of x.

Remarks This is based on the recursive series function recserar. recserar could be called directly as follows:

recserar(x,x[1,.], ones(1,cols(x)))

to accomplish the same thing.

Example $x = \{ 1 - 3, \}$

2 2,
3 -1 };

y = cumsumc(x);

1 -3

y = 3 - 1

6 - 2

Source cumsumc.src

See also cumprode, recsercp, recserar

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curve

curve

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a

u V W

x y z

Purpose Computes a one-dimensional smoothing curve.

Format $\{u,v\} = curve(x,y,d,s,sigma,G);$

Input Kx1 vector, x-abscissae (x-axis values).

> Kx1 vector, y-ordinates (y-axis values). v

d Kx1 vector or scalar, observation weights.

scalar, smoothing parameter. If s = 0, curve performs an 2. interpolation. If d contains standard deviation estimates, a reasonable value for s is K.

sigma scalar, tension factor. Gscalar, grid size factor.

Output K*Gx1 vector, x-abscissae, regularly spaced. и

> ν K*Gx1 vector, y-ordinates, regularly spaced.

Remarks

sigma contains the tension factor. This value indicates the curviness desired. If sigma is nearly zero (e.g., .001), the resulting curve is approximately the tensor product of cubic curves. If sigma is large, (e.g., 50.0) the resulting curve is approximately bi-linear. If sigma equals zero, tensor products of cubic curves result. A standard value for sigma is approximately 1.

G is the grid size factor. It determines the fineness of the output grid. For G=1, the input and output vectors will be the same size. For G=2, the output grid is twice as fine as the input grid, i.e., u and v will have twice as many rows as x and y.

Source spline.src

cvtos

Purpose Converts a character vector to a string.

Format s = cvtos(v);

Input v Nx1 character vector, to be converted to a string.

Output s string, contains the contents of v.

Remarks cvtos in effect appends the elements of *v* together into a single string.

cvtos was written to operate in conjunction with stocv. If you pass it a character vector that does not conform to the output of stocv, you may get unexpected results. For example, cvtos DOES NOT look for 0 terminating bytes in the elements of v; it assumes every element except the last is 8 characters long. If this is not true, there will be 0's in the middle of s.

If the last element of v does not have a terminating 0 byte, **cvtos** supplies one for s.

s = cvtos(v);

s = "Now is the time for all good men"

See also stocv, vget, vlist, vput, vread

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datalist

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datalist

Purpose List selected variables from a data set.

Format datalist dataset [var1 [var2...]];

Input *dataset* literal, name of the dataset.

var# literal, the names of the variables to list.

Global Input range global scalar, the range of rows to list. The default is all

rows

miss global scalar, controls handling of missing values.

0 display rows with missing values.

1 do not display rows with missing values.

The default is 0.

_prec global scalar, the number of digits to the right of the

decimal point to display. The default is 3.

Remarks The variables are listed in an interactive mode. As many rows and

columns as will fit in the window are displayed. You can use the cursor

keys to pan and scroll around in the listing.

Example datalist freq age sex pay;

This command will display the variables age, sex, and pay from the

data set freq.dat.

Source datalist.src

dataloop (dataloop)

dataloop (dataloop)

Purpose Specifies the beginning of a data loop.

src = "source";

Format dataloop infile outfile;

Input *infile* string variable or literal, the name of the source data set.

Output outfile string variable or literal, the name of the output data set.

Remarks

The statements between the **dataloop... endata** commands are assumed to be metacode to be translated at compile time. The data from *infile* is manipulated by the specified statements, and stored to the data set *outfile*. Case is not significant within the **dataloop... endata** section, except for within quoted strings. Comments can be used as in any GAUSS code.

Example

```
dataloop ^src dest;
make newvar = x1 + x2 + log(x3);
x6 = sqrt(x4);
keep x6, x5, newvar;
endata;
```

Here, **src** is a string variable requiring the caret operator (^), while **dest** is a string literal.

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с **d**

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date

date

a Purpose

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Returns the current date in a 4-element column vector, in the order: year, month, day, and hundredths of a second since midnight.

Format y = date;

Remarks The hundredths of a second since midnight can be accessed using **hsec**.

Example print date;

1998.0000

6.0000000

15.000000

4011252.7

See also time, timestr, ethsec, hsec, etstr

w x y z

u

V

datestr

datestr

```
Purpose
             Returns a date in a string.
 Format
             str = datestr(d);
    Input
             d
                    4x1 vector, like the date function returns. If this is 0, the
                    date function will be called for the current system date.
  Output
             str
                    8 character string containing current date in the form: mo/
                    dy/yr
Example
             d = \{ 1998, 6, 15, 0 \};
             y = datestr(d);
             print y;
                 6/15/98
 Source
             time.src
See also
             date, datestring, datestrymd, time, timestr,
              ethsec
```

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datestring

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V

W

datestring

Purpose Returns a date in a year-2000-compliant string. Format str = datestring(d); Input d 4x1 vector, like the **date** function returns. If this is 0, the date function will be called for the current system date. **Output** str 10 character string containing current date in the form: mm/dd/yyyy Example y = datestring(0); print y; 6/15/1998 Source time.src See also date, datestr, datestrymd, time, timestr, ethsec

datestrymd

datestrymd

```
Purpose
             Returns a date in a string.
 Format
             str = datestrymd(d);
    Input
                    4x1 vector, like the date function returns. If this is 0, the
             d
                    date function will be called for the current system date.
  Output
             str
                    8 character string containing current date in the form:
                    yyyymmdd
             d = \{ 1998, 6, 15, 0 \};
Example
             y = datestrymd(d);
             print y;
                 19980615
 Source
             time.src
See also
             date, datestr, datestring, time, timestr, ethsec
```

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dayinyr

dayinyr

a Returns day number in the year of a given date. Purpose b **Format** daynum = dayinyr(dt);С Input 3x1 or 4x1 vector, date to check. The date should be in the form dtd returned by date. Output daynum scalar, the day number of that date in that year, 1-366. Example $x = \{ 1998, 6, 15, 0 \};$ y = dayinyr(x);print y; 166.00000 Source time.src Globals _isleap

е g h k m n 0 p q t u V W

dayofweek

dayofweek

Purpose Returns day of week.

Format d = dayofweek(a);

Input *a* Nx1 vector, dates in DT format.

Output d Nx1 vector, integers indicating day of week of each date:

[1] Sunday

[2] Monday

[3] Tuesday

[4] Wednesday

[5] Thursday

[6] Friday

[7] Saturday

Remarks The DT scalar format is a double precision representation of the date and

time. In the DT scalar format, the number

20010421183207

represents 18:32:07 or 6:32:07 PM on April 21, 2001.

Source time.src

a

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XX7

debug

debug

Purpose Runs a program under the source level debugger.

Format debug filename;

Input filename Literal or name of file to debug.

Remarks See "Debugging" in the User's Guide.

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3-172

declare

Purpose To initialize matrices and strings at compile time. **Format** declare [type] symbol [aop clist]; Input type optional literal, specifying the type of the symbol. matrix string if *type* is not specified, **matrix** is assumed. the name of the symbol being declared. symbol the type of assignment to be made. aop if not initialized, initialize. if already initialized, reinitialize. if not initialized, initialize. ! = if already initialized, reinitialize. if not initialized, initialize. :=

if already initialized, leave as is.

if not initialized, initialize.

If *aop* is specified, *clist* must be also.

clist a list of constants to assign to symbol.

?=

If *aop clist* is not specified, *symbol* is initialized as a scalar 0 or a null string.

if already initialized, redefinition error.

Remarks

The **declare** syntax is similar to the **let** statement.

declare generates no executable code. This is strictly for compile time initialization. The data on the right-hand side of the equal sign must be constants. No expressions or variables are allowed.

declare statements are intended for initialization of global matrices and strings that are used by procedures in a library system.

It is best to place **declare** statements in a separate file from procedure definitions. This will prevent redefinition errors when rerunning the same program without clearing your workspace.

Complex numbers can be entered by joining the real and imaginary parts with a sign (+ or -); there should be no spaces between the numbers and

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W X y Z

the sign. Numbers with no real part can be entered by appending an 'i' to the number.

There should be only one declaration for any symbol in a program. Multiple declarations of the same symbol should be considered a programming error. When GAUSS is looking through the library to reconcile a reference to a matrix or a string, it will quit looking as soon as a symbol with the correct name is found. If another symbol with the same name existed in another file, it would never be found. Only the first one in the search path would be available to programs.

Here are some of the possible uses of the three forms of declaration:

- !=, = Interactive programming or any situation where a global by the same name will probably be listed in the symbol table when the file containing the **declare** statement is compiled. The symbol will be reset.
 - This allows mixing **declare** statements with the procedure definitions that reference the global matrices and strings, or placing them in your main file.
- Redefinition is treated as an error. This will not allow you to assign one symbol with another value already in your program. Rename one of them.
 - Place **declare** statements in a separate file from the rest of your program and procedure definitions.
- ?= Interactive programming where some global defaults were set when you started and you do not want them reset for each successive run even if the file containing the declare's gets recompiled. Be careful when using.

CTRL+W controls the **declare** statement warning level. If **declare** warnings are on, you will be warned whenever a **declare** statement encounters a symbol that is already initialized. This happens when you declare a symbol that is already initialized when **declare** warnings are turned on:

```
declare != Reinitialize and warn.
declare := Crash with fatal error.
declare ?= Leave as is and warn.
```

If **declare** warnings are off, no warnings are given for the != and ?= cases.

Example

declare matrix x,y,z;

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е

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.

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V

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С

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q

V

```
x = 0
  y = 0
  z = 0
declare string x = "This string.";
  x = "This string."
declare matrix x;
x = 0
declare matrix x != { 1 2 3, 4 5 6, 7 8 9 };
    1 2 3
x = 456
    7 8 9
declare matrix x[3,3] = 1 2 3 4 5 6 7 8 9;
    1 2 3
x = 456
    789
declare matrix x[3,3] = 1;
    1 1 1
x = 1 1 1
    1 1 1
declare matrix x[3,3];
    0 0 0
x = 0.00
    0.00
declare matrix x = 1 2 3 4 5 6 7 8 9;
```

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$$\begin{array}{r}
1\\2\\3\\4\\x=5\\6\\7
\end{array}$$

8

9

declare matrix x = dog cat;

$$x = \begin{array}{c} DOG \\ CAT \end{array}$$

declare matrix x = "dog" "cat";

$$x = \frac{\log}{\cot}$$

See also let, external

delete

a

b

С

d

g

h

m

n o

q

хух

delete

Purpose	Deletes global symbols from the symbol table.	
r ui pose	Deletes global symbols from the symbol table.	
Format	<pre>delete [-flags] [symbol] [symbol2] [symbol3];</pre>	
Input	flags	specify the type(s) of symbols to be deleted
		p procedures
		k keywords
		f fn functions
		m matrices
		s strings
		g only symbols with global references
		only symbols with all local references
	, ,	n no pause for confirmation
	symbol	literal, name of symbol to be deleted. If symbol ends in an asterisk, all symbols matching the
		leading characters will be deleted.
Remarks	This completely and irrevocably deletes symbols from GAUSS's memory and workspace. Flags must be preceded by a slash (e.g., -pfk). If the n (no pause) flag is used, you will not be asked for confirmation for each symbol. This command is supported only from interactive level. Since the interpreter executes a compiled pseudo-code, this command would invalidate a previously compiled code image and therefore would destroy any program it was a part of. If any symbols are deleted, all procedures, keywords, and functions with global references will be deleted as well.	
Example	print x;	
	96.000000	
	6.000000	
	14.000000	
	3502965.9	
	delete -m	ı x;

delete

At the Delete? [Yes No Previous Quit] prompt, enter y.

show x;

x no longer exists.

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ι

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V

w x y z

delete (dataloop)

delete (dataloop)

Purpose Removes specific rows in a data loop based on a logical expression.

Format delete logical expression;

Remarks Deletes only those rows for which *logical expression* is *TRUE*. Any variables referenced must already exist, either as elements of the source data set, as externs, or as the result of a previous **make**, **vector**, or

code statement.

GAUSS expects *logical expression* to return a row vector of 1's and 0's. The relational and other operators (e.g., <) are already interpreted in terms of their dot equivalents (•<), but it is up to the user to make sure that

function calls within logical expression result in a vector.

Example delete age < 40 or sex == 'FEMALE';

See also select

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DeleteFile

DeleteFile

Purpose Deletes files.

Format ret = DeleteFile(name);

Input *name* string or NxK string array, name of file or files to delete.

Output *ret* scalar or NxK matrix, 0 if successful.

Remarks The return value, *ret*, is scalar if name is a string. If name is an NxK string array, *ret* will be an NxK matrix reflecting the success or failure of each separate file deletion.

DeleteFile calls the C library **unlink** function for each file. If **unlink** fails it sets the C library errno value. **DeleteFile** returns the value of errno if **unlink** fails, otherwise it returns zero. If you want detailed information about the reason for failure, consult the C library **unlink** documentation for your platform for details.

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delif

Purpose

Deletes rows from a matrix. The rows deleted are those for which there is a 1 in the corresponding row of e.

Format

$$y = delif(x,e);$$

Input

x NxK data matrix.

e Nx1 logical vector (vector of 0's and 1's).

Output

y MxK data matrix consisting of the rows of y for which there is a 0 in the corresponding row of e. If no rows remain, **delif** will return a scalar missing.

Remarks

The input *e* will usually be generated by a logical expression using dot operators. For instance:

```
y = delif(x, x[.,2] .> 100);
```

will delete all rows of *x* whose second element is greater than 100. The remaining rows of *x* will be assigned to *y*.

Example

Here is the resulting matrix *y*:

0 10 20

All rows for which the elements in column 1 are greater than 0 and the elements in column 3 are less than 100 are deleted.

Source

datatran.src

See also selif

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denseSubmat

denseSubmat

Purpose Returns dense submatrix of sparse matrix.

Format e = denseSubmat(x,r,c);

Input x MxN sparse matrix.

r Kx1 vector, row indices.

c Lx1 vector, column indices.

Output e KxL dense matrix.

Remarks If *r* or *c* are scalar zeros, all rows or columns will be returned.

Source sparse.src

See also sparseFd, sparseFp

3-182

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design

design

Purpose Creates a design matrix of 0's and 1's from a column vector of numbers specifying the columns in which the 1's should be placed.

Format y = design(x);

Input x Nx1 vector.

Output y NxK matrix, where K = maxc(x); each row of y will contain a single 1, and the rest 0's. The one in the i^{th} row will be in the round(x[i,1]) column.

Remarks Note that *x* does not have to contain integers: it will be rounded to nearest if necessary.

0001

Source design.src

See also cumprodc, cumsumc, recserrc

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Input ,

y = det(x);x NXN square matrix.

Output

Purpose

Format

y determinant of x.

Remarks

x may be any valid expression that returns a square matrix (number of rows equals number of columns).

det computes a LU decomposition.

detl can be much faster in many applications.

Returns the determinant of a square matrix.

Example

$$x = \begin{cases} 3 & 2 & 1, \\ 0 & 1 & -2, \\ 1 & 3 & 4 \end{cases};$$

$$y = det(x);$$

$$x = \begin{array}{rrr} 3 & 2 & 1 \\ 0 & 1 & -2 \\ 1 & 3 & 4 \end{array}$$

$$y = 25$$

See also detl

detl

Purpose

Returns the determinant of the last matrix that was passed to one of the intrinsic matrix decomposition routines.

Format

y = detl;

Remarks

Whenever one of the following functions is executed, the determinant of the matrix is also computed and stored in a system variable. This function will return the value of that determinant and, because the value has been computed in a previous instruction, this will require no computation.

The following functions will set the system variable used by **det1**:

Example

If both the inverse and the determinant of the matrix are needed, the following two commands will return both with the minimum amount of computation:

```
xi = inv(x);

xd = detl;
```

The function det(x) returns the determinant of a matrix using the Crout decomposition. If you only want the determinant of a positive definite matrix, the following code will be the fastest for matrices larger than 10x10:

```
call chol(x);
xd = detl;
```

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detl

The Cholesky decomposition is computed and the result from that is discarded. The determinant saved during that instruction is retrieved using $\mathtt{det1}$. This can execute up to 2.5 times faster than $\mathtt{det}(x)$ for large positive definite matrices.

See also det

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dfft

dfft

Purpose Computes a discrete Fourier transform.

Format y = dfft(x);

Input x Nx1 vector.

Output y Nx1 vector.

Remarks The transform is divided by N.

This uses a second-order Goertzel algorithm. It is considerably slower than **fft**, but it may have some advantages in some circumstances. For one thing, N does not have to be an even power of 2.

Source dfft.src

See also dffti, fft, ffti

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dffti

dffti

Purpose Computes inverse discrete Fourier transform.

Format y = dffti(x);

Input x Nx1 vector.

Output y Nx1 vector.

Remarks The transform is divided by N.

This uses a second-order Goertzel algorithm. It is considerably slower than **ffti**, but it may have some advantages in some circumstances. For one thing, *N* does not have to be an even power of 2.

Source dffti.src

See also fft, dffti, ffti

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3-188

dfree (DOS only)

dfree (DOS only)

Purpose Returns the amount of room left on a diskette or hard disk.

Format y = dfree(drive);

Input *drive* scalar, valid disk drive number.

Output y number of bytes free.

Portability All others return -1

Remarks Valid disk drive numbers are 0 = default, 1 = A, 2 = B, etc. If an error is

encountered, **dfree** will return -1.

See also coreleft

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Purpose Creates a column vector from the diagonal of a matrix.

Format y = diag(x);

Input x NxK matrix.

Output $y = \min(N,K)x1$ vector.

Remarks The matrix need not be square.

diagrv reverses the procedure and puts a vector into the diagonal of a matrix.

Example x = rndu(3,3);

y = diag(x);

0.660818 0.367424 0.302208

 $x = 0.204800 \ 0.077357 \ 0.145755$ $0.712284 \ 0.353760 \ 0.642567$

0.660818

y = 0.077357

0.642567

See also

diagrv

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x y z

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diagrv

Purpose Inserts a vector into the diagonal of a matrix. **Format** y = diagrv(x, v);Input NxK matrix. x min(N,K) vector. ν **Output** NxK matrix equal to x with its principal diagonal elements equal to those of v. Remarks **diag** reverses the procedure and pulls the diagonal out of a matrix. **Example** x = rndu(3,3);v = ones(3,1);y = diagrv(x,v);0.660818 0.367424 0.302208 0.204800 0.077357 0.145755 0.712284 0.353760 0.642567 1.000000 1.000000 v =1.000000 1.000000 0.367424 0.302208 y =0.204800 1.000000 0.145755 0.712284 0.353760 1.000000 See also diag

3-191

digamma

digamma

Purpose Computes the digamma function.

Format y = digamma(x);

Input x MxN matrix.

Output y MxN matrix, digamma.

Remarks The digamma function is the first derivative of the log of the gamma

function with respect to argument.

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dlibrary

dlibrary

Purpose Dynamically links and unlinks shared libraries.

Format dlibrary *lib1* [*lib2*...];

dlibrary -a lib1 [lib2...];

dlibrary -d dlibrary

Input

lib1 lib2... literal, the base name of the library or the pathed name of the library.

dlibrary takes two types of arguments, "base" names and file names. Arguments without any "\" path separators are assumed to be library base names, and are expanded by adding the suffix .dll. They are searched for in the default dynamic library directory. Arguments that include "\" path separators are assumed to be file names, and are not expanded. Relatively pathed file names are assumed to be specified relative to the current working directory, not relative to the dynamic library directory.

- a append flag, the DLL's listed are added to the current set of DLL's rather than replacing them. For search purposes, the new DLL's follow the already active ones. Without the -a flag, any previously linked libraries are dumped.
- d dump flag, ALL DLL's are unlinked and the functions they contain are no longer available to your programs. If you use
 dllcall to call one of your functions after executing a
 dlibrary -d, your program will terminate with an error.

Remarks

If no flags are used, the DLL's listed are linked into GAUSS and any previously linked libraries are dumped. When you call **dllcall**, the DLL's will be searched in the order listed for the specified function. The first instance of the function found will be called.

dlibrary with no arguments prints out a list of the currently linked DLL's. The order in which they are listed is the order in which they are searched for functions.

dlibrary recognizes a default directory in which to look for dynamic libraries. You can specify this by setting the variable dlib_path in gauss.cfg. Set it to point to a single directory, not a sequence of

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dlibrary

directories. A new case (case 24) has also been added to **sysstate** for getting and setting this default.

GAUSS maintains its own DLL, gauss.dll. gauss.dll is listed when you execute **dlibrary** with no arguments, and searched when you call **dllcall**. By default, gauss.dll is searched last, after all other DLL's but you can force it to be searched earlier by listing it explicitly in a **dlibrary** statement. gauss.dll is always active. It is not unlinked when you execute **dlibrary** -d. gauss.dll is located in the gauss.exe directory.

See also dllcall, sysstate-case 24

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dllcall

Purpose

Calls functions located in dynamic libraries.

Format

dllcall [-r] [-v] func[(arg1[,arg2...])];

dllcall works in conjunction with dlibrary. dlibrary is used to link dynamic-link libraries (DLL's) into GAUSS; dllcall is used to access the functions contained in those DLL's. dllcall searches the DLL's (see dlibrary for an explanation of the search order) for a function named *func*, and calls the first instance it finds. The default DLL, gauss.dll, is searched last.

Input

func the name of a function contained in a DLL (linked into GAUSS with dlibrary). If func is not specified or cannot be located in a DLL. dllcall will fail.

arg# arguments to be passed to *func*; optional. These must be elementary variable; they cannot be expressions.

- -r optional flag. If -r is specified, **dllcall** examines the value returned by *func*, and fails if it is nonzero.
- optional flag. Normally, dllcall passes parameters to func in a list. If -v is specified, dllcall passes them in a vector. See below for more details.

Remarks

func should be written to:

- 1. Take 0 or more pointers to doubles as arguments.
- 2. Take arugments either in a list of a vector.
- 3. Return an integer.

In C syntax, func should take one of the following forms:

- int func (void);
- int func (double *arg1[,double *arg2...]);
- 3. int func (double *argv[]);

dllcall can pass a list of up to 100 arguments to *func*; if it requires more arguments than that, you MUST write it to take a vector of arguments, and you MUST specify the -v flag when calling it. dllcall can pass up to 1000 arguments in vector format. In addition, in vector format dllcall appends a null pointer to the vector, so you can write

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dllcall

func to take a variable number of arguments and just test for the null pointer.

Arguments are passed to *func* by reference. This means you can send back more than just the return value, which is usually jsut a success/failure code. (It also means that you need to be careful not to overwrite the contents of matrices or strings you want to preserve.) To return data from *func*, simply set up one or more of its arguments as return matrices (basically, by making them the size of what you intend to return), and inside *func* assign the results to them before returning.

See also dlibrary, sysstate-case 24

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do while, do until

do while, do until

Purpose

Executes a series of statements in a loop as long as a given expression is true (or false).

Format

do while expression;

or

do until expression;

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statements in loop

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.

endo;

Remarks

expression is any expression that returns a scalar. It is TRUE if it is nonzero and FALSE if it is zero.

In a **do while** loop, execution of the loop will continue as long as the expression is TRUE.

In a **do until** loop, execution of the loop will continue as long as the expression is FALSE.

The condition is checked at the top of the loop. If execution can continue, the statements of the loop are executed until the **endo** is encountered. Then GAUSS returns to the top of the loop and checks the condition again.

The **do** loop does not automatically increment a counter. See the first example, following.

do loops may be nested.

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do while, do until

It is often possible to avoid using loops in GAUSS by using the appropriate matrix operator or function. It is almost always preferable to avoid loops when possible, since the corresponding matrix operations can be much faster.

Example

```
format /rdn 1,0;
space = " ";
comma = ",";
i = 1;
do while i <= 4;
   j = 1;
  do while j <= 3;
     print space i comma j;;
      j = j+1;
   endo;
   i = i+1;
  print;
endo;
1, 1 1, 2 1, 3
2, 1 2, 2 2, 3
3, 1 3, 2 3, 3
4, 1 4, 2 4, 3
```

In the example above, two nested loops are executed and the loop counter values are printed out. Note that the inner loop counter must be reset inside the outer loop before entering the inner loop. An empty print statement is used to print a carriage return/line feed sequence after the inner loop finishes.

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do while, do until

The following are examples of simple loops that execute a predetermined number of times. These loops will both have the result shown.

```
First loop
```

```
format /rd 1,0;
   i = 1;
  do while i <= 10;
  print i;;
   i = i+1;
   endo;
produces
12345678910
Second loop
  format /rd 1,0;
   i = 1;
  do until i > 10;
  print i;;
   i = i+1;
   endo;
produces
12345678910
```

See also continue, break

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dos

ao

Purpose Provides access to the operating system from within GAUSS.

Format dos [s];

Input *s* literal or ^string, the OS command to be executed.

Portability UNIX

Control and output go to the controlling terminal, if there is one.

This function may be used in terminal mode.

OS/2, Windows

The **dos** function opens a new terminal.

Running programs in the background is allowed in all three of the aforementioned platforms.

Remarks This allows all operating system commands to be used from within GAUSS. It allows other programs to be run even though GAUSS is still resident in memory.

If no operating system command (for instance, dir or copy) or program name is specified, a shell of the operating system will be entered which can be used just like the base level OS. The exit command must be given from the shell to get back into GAUSS. If a command or program name is included, the return to GAUSS is automatic after the DOS command has been executed.

All matrices are retained in memory when the OS is accessed in this way. This command allows the use of word processing, communications, and other programs from within GAUSS.

Do not execute programs that terminate and remain resident because they will be left resident inside GAUSS's workspace. Some examples are programs that create RAM disks or print spoolers.

If the command is to be taken from a string variable, the ^ (caret) must precede the string.

The shorthand ">" can be used in place of the word "DOS".

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dos

Example

comstr = "basic myprog";

dos ^comstr;

This will cause the BASIC program **myprog** to be run. When that program is finished, control will automatically return to GAUSS.

>dir *.prg;

This will use the DOS **dir** command to print a directory listing of all files with a .prg extension. When the listing is finished, control will be returned to GAUSS.

dos;

This will cause a second level OS shell to be entered. The OS prompt will appear and OS commands or other programs can be executed. To return to GAUSS, type exit.

See also exec

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doswin

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doswin

a

Purpose Opens the DOS compatibility window with default settings.

Format doswin;

Portability Windows only

Remarks Calling doswin is equivalent to:

call DOSWinOpen("",error(0));

Source gauss.src

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DOSWinCloseall

DOSWinCloseall

Purpose Closes the DOS compatibility window. **Format** DOSWinCloseall; **Portability** Windows only Remarks Calling DOSWinCloseall closes the DOS window immediately, without asking for confirmation. If a program is running, its I/O reverts to the Command window. **Example** let attr = 50 50 7 0 7; if not DOSWinOpen("Legacy Window", attr); errorlog "Failed to open DOS window, aborting"; stop; endif; DOSWinCloseall;

DOSWinOpen

DOSWinOpen

Purpose

Opens the DOS compatibility window and gives it the specified title and attributes.

Format

ret = DOSWinOpen(title,attr);

Portability

Windows 3.2 only

Input

title string, window title.

attr 5x1 vector or scalar missing, window attributes.

[1] window x position

[2] window y position

[3] text foreground color

[4] text background color

[5] close action bit flags

bit 0 (1's bit)] issue dialog bit 1 (2's bit)] close window bit 2 (4's bit)] stop program

Output

ret

scalar, success flag, 1 if successful, 0 if not.

Remarks

If *title* is a null string (""), the window will be titled "GAUSS-DOS".

Defaults are defined for the elements of *attr*. To use the default, set an element to a missing value. Set *attr* to a scalar missing to use all defaults

[1] varies use x position of previous DOS window
[2] varies use y position of previous DOS window
[3] 7 white foreground
[4] 0 black background
[5] 6 4+2: stop program and close window without confirming

If the DOS window is already open, the new title and attr will be applied to it. Elements of attr that are missing are not reset to the default values, but are left as is.

To set the close action flags value (*attr*[5]), just sum the desired bit values. For example:

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DOSWinOpen

```
stop program (4) + close window (2) + confirm close (1) = 7
```

The close action flags are only relevant when a user attempts to interactively close the DOS window while a program is running. If GAUSS is idle, the window will be closed immediately. Likewise, if a program calls <code>DOSWinCloseall</code>, the window is closed, but the program does not get terminated.

Example

```
let attr = 50 50 7 0 7;
if not DOSWinOpen("Legacy Window", attr);
  errorlog "Failed to open DOS window, aborting";
  stop;
endif;
```

This example opens the DOS window at screen location (50,50), with white text on a black background. The close action flags are 4 + 2 + 1 (stop program + close window + issue confirm dialog) = 7. Thus, if the user attempts to close the window while a program is running, he/she will be asked for confirmation. Upon confirmation, the window will be closed and the program terminated.

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dotfeg, dotfge, dotfgt, dotfle, dotflt, dotfne

dotfeq, dotfge, dotfgt, dotfle, dotfle,

Purpose Fuzzy comparison functions. These functions use **_fcmptol** to fuzz the comparison operations to allow for roundoff error.

Format y = dotfeq(a,b);
 y = dotfge(a,b);
 y = dotfgt(a,b);
 y = dotfle(a,b);
 y = dotflt(a,b);
 y = dotfne(a,b);

Input *a* NxK matrix, first matrix.

b LxM matrix, second matrix, ExE compatible with a.

Global Input _fcmptol global scalar, comparison tolerance. The default value is 1.0e-15.

Output $y = \max(N,L)$ by $\max(K,M)$ matrix of 1's and 0's.

Remarks The return value is 1 if true and 0 if false.

The statement:

y = dotfeq(a,b);
is equivalent to:
y = a .eq b;

The calling program can reset **_fcmptol** before calling these procedures.

 $_{fcmptol} = 1e-12;$

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dotfeq, dotfge, dotfgt, dotfle, dotflt, dotfne

Example x = rndu(2,2);

y = rndu(2,2);

t = dotfge(x,y);

 $x = 0.85115559 \ 0.98914218$

0.12703276 0.43365175

 $y = 0.41907226 \ 0.49648058$

0.58039125 0.98200340

 $t = 1.0000000 \ 1.0000000$

0.0000000 0.0000000

Source fcompare.src

Globals _fcmptol

See also feq-fne

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draw

draw

a b d е g h k m n 0 p q u V

```
Purpose
             Graphs lines, symbols, and text using the PQG global variables. This
             procedure does not require actual X, Y, or Z data since its main purpose is
              to manually build graphs using _pline, _pmsgctl, _psym, _paxes,
             _parrow and other globals.
 Library
             pgraph
  Format
             draw;
Remarks
             draw is especially useful when used in conjunction with transparent
             graphic panels.
Example
             library pgraph;
             graphset;
             begwind;
             /* make full size window for plot */
             makewind(9,6.855,0,0,0);
             /* make small overlapping window for text*/
             makewind(3,1,3,3,0);
                           setwind(1);
                 x = sega(.1, .1, 100);
                 y = \sin(x);
                 /* plot data in first window*/
                 xy(x,y);
```

W

x y z

draw

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drop (dataloop)

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W

x y z

drop (dataloop)

Purpose Specifies columns to be dropped from the output data set in a data loop.

Format drop variable_list;

Remarks Commas are optional in variable_list.

Deletes the specified variables from the output data set. Any variables referenced must already exist, either as elements of the source data set, or as the result of a previous **make**, **vector**, or **code** statement.

If neither **keep** nor **drop** is used, the output data set will contain all variables from the source data set, as well as any defined variables. The effects of multiple **keep** and **drop** statements are cumulative.

Example drop age, pay, sex;

See also keep

dsCreate

dsCreate

Purpose Creates an instance of a structure of type DS set to default values.

Format s = dsCreate;

Output *s* instance of structure of type DS.

Source ds.src

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dstat

dstat

a **Purpose** Compute descriptive statistics. h **Format** Input string, name of data set. dataset interpreted as: the variables. vars containing the data. h KX1 numeric vector data set will be used. NXK matrix m 0 procedure. p **altnam** global matrix, default 0. miss names for the output. global scalar, default 0. 0 1 in it. pairwise deletion. u row V the read loop. W internally. хуг

{ vnam, mean, var, std, min, max, valid, mis } = dstat(dataset, vars); If dataset contains the name of a GAUSS data set, vars will be If dataset is null or 0, *vars* will be assumed to be a matrix **KX1** character vector names of variables. indices of columns. These can be any size subset of the variables in the data set and can be in any order. If a scalar 0 is passed, all columns of the If dataset is null or 0, vars will be interpreted as: the data on which to compute the descriptive statistics. Defaults are provided for the following global input variables, so they can be ignored unless you need control over the other options provided by this This can be a Kx1 character vector of alternate variable there are no missing values (fastest). listwise deletion, drop a row if any missings occur global scalar, the number of rows to read per iteration of if 0, (default) the number of rows will be calculated

dstat

__output global scalar, controls output, default 1.

- **1** print output table.
- 0 do not print output.

Output

vnam Kx1 character vector, the names of the variables used in the

statistics.

mean Kx1 vector, means.

var Kx1 vector, variance.

std Kx1 vector, standard deviation.

min Kx1 vector, minima.max Kx1 vector, maxima.

valid Kx1 vector, the number of valid cases.

mis Kx1 vector, the number of missing cases.

Remarks

If pairwise deletion is used, the minima and maxima will be the true values for the valid data. The means and standard deviations will be computed using the correct number of valid observations for each variable.

Source

dstat.src

Globals

output, dstatd, dstatx

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Purpose Creates a matrix in DT scalar format.

Format dt = dtdate(year, month, day, hour, minute, second);

Input *year* NxK matrix of years.

month NxK matrix of months, 1-12.
day NxK matrix of days, 1-31.
hour NxK matrix of hours, 0-23.
minute NxK matrix of minutes, 0-59.
second NxK matrix of seconds, 0-59.

Output dt NxK matrix of DT scalar format dates.

Remarks The arguments must be ExE conformable.

Source time.src

See also dtday, dttime, utctodt, dttostr

dtday

dtday

Purpose Creates a matrix in DT scalar format containing only the year, month and

day. Time of day information is zeroed out.

Format dt = dtday(year, month, day);

Input *year* NxK matrix of years.

month NxK matrix of months, 1-12. day NxK matrix of days, 1-31.

Output dt NxK matrix of DT scalar format dates.

Remarks This amounts to 00:00:00 or midnight on the given day. The arguments

must be ExE conformable.

Source time.src

See also dttime, dtdate, utctodt, dttostr

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dttime

dttime

Purpose Creates a matrix in DT scalar format containing only the hour, minute and second. The date information is zeroed out.

Format dt = dttime(hour, minute, second);

Input *hour* NxK matrix of hours, 0-23.

minute NxK matrix of minutes, 0-59. *second* NxK matrix of seconds, 0-59.

Output *dt* NxK matrix of DT scalar format times.

Remarks The arguments must be ExE conformable.

Source time.src

See also dtday, dtdate, utctodt, dttostr

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dttodtv

Converts DT scalar format to DTV vector format. **Purpose** Format dtv = dttodtv(dt);Input dtNx1 vector, DT scalar format. Output dtv Nx8 matrix. DTV vector format. In DT scalar format, 11:06:47 on March 15, 2001 is 20010315110647. Remarks Each row of dtv, in DTV vector format, contains: [N,1]Year [N,2]Month in Year, 1-12 [N,3]Day of month, 1-31 [N,4]Hours since midnight, 0-23 [N,5]Minutes, 0-59 [N,6]Seconds, 0-59 [N,7]Day of week, 0-6, 0 = Sunday[N,8] Days since Jan 1 of current year, 0-365 Example dt = 20010326110722;print "dt = " dt; dtv = dttodtv(dt);print "dtv = " dtv; produces: dt = 20010326110722 $dtv = 2001 \ 3 \ 26 \ 11 \ 7 \ 22 \ 1 \ 84$ Source time.src See also dtvnormal, timeutc, utctodtv, dtvtodt, dttoutc, dtvtodt, strtodt, dttostr

dttostr

dttostr

Purpose Converts a matrix containing dates in DT scalar format to a string array. h **Format** sa = dttostr(x, fmt);Input NxK matrix containing dates in DT scalar format. string containing date/time format characters. fmt **Output** NxK string array. sa Remarks The DT scalar format is a double precision representation of the date and time. In the DT scalar format, the number h 20010421183207 represents 18:32:07 or 6:32:07 PM on April 21, 2001. **dttostr** converts a date in DT scalar format to a character string using the format string in fmt. The following formats are supported: YYYY 4 digit year YR Last two digits of year m MO Number of month, 01-12 n DD Day of month, 01-31 HH Hour of day, 00-23 0 ΜI Minute of hour, 00-59 p SS Second of minute, 00-59 q **Example** s0 = dttostr(utctodt(timeutc), "YYYY-MO-DD HH:MI:SS"); Print ("Date and Time are: " \$+ s0); produces: u Date and time are: 2001-03-25 14:59:40 V

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print dttostr(utctodt(timeutc),
                 "Today is DD-MO-YR")
           produces:
           Today is 25-03-01
           s = dttostr(x, "YYYY-MO-DD");
           if x = 20000317060424
                   20010427031213
                   20010517020437
                   20011117161422
                   20010717120448
                   20010817043451
                   20010919052320
                   20011017032203
                   20011107071418
           then s = 2000-03-17
                     2001-04-27
                     2001-05-17
                     2001-11-17
                     2001-07-17
                     2001-08-17
                     2001-09-19
                     2001-10-17
                     2001-11-07
See also strtodt, dttoutc, utctodt
```

dttoutc

dttoutc

Purpose b **Format** utc = dttoutc(dt);Input dtd **Output** utc Remarks Greenwich Mean Time. h **Example** produces: tc = 985633642;m Source time.src See also 0 p q

```
Converts DT scalar format to UTC scalar format.
      Nx1 vector, DT scalar format.
      Nx1 vector, UTC scalar format.
In DT scalar format, 11:06:47 on March 15, 2001 is 20010315110647. A
UTC scalar gives the number of seconds since or before January 1, 1970
dt = 20010326085118;
tc = dttoutc(dt);
print "utc = " utc;
dtvnormal, timeutc, utctodtv, dttodtv, dtvtodt,
dtvtoutc, dtvtodt, strtodt, dttostr
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dtvnormal

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dtvnormal

Normalizes a date and time (DTV) vector. **Purpose** Format d = dtvnormal(t);Input 1x8 date and time vector that has one or more elements outside t the normal range. **Output** Normalized 1x8 date and time vector. Remarks The date and time vector is a 1x8 vector whose elements consist of: Year: Year, four digit integer. Month: 1-12, Month in year. Day: 1-31, Day of month. Hour: 0-23, Hours since midnight. Min: 0-59, Minutes. Sec: 0-59, Seconds. 0-6, Day of week, 0 = Sunday. DoW: DiY: 0-365, Days since Jan 1 of year. The last two elements are ignored on input. **Example** format /rd 10,2; $x = \{ 1996 14 21 6 21 37 0 0 \};$ d = dtvnormal(x);d:97.00 2.00 21.00 6.00 21.00 37.00 2.00 51.00 See also date, ethsec, etstr, time, timestr, timeutc, utctodtv

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dtvtodt

dtvtodt

Purpose Converts DT vector format to DT scalar format. h Format dt = dtvtodt(dtv);Input Nx8 matrix. DTV vector format. dtv е Output dtNx1 vector, DT scalar format. Remarks In DT scalar format, 11:06:47 on March 15, 2001 is 20010315110647. Each row of dtv, in DTV vector format, contains: h [N,1]Year [N,2]Month in Year, 1-12 [N,3]Day of month, 1-31 [N,4]Hours since midnight, 0-23 [N,5]Minutes, 0-59 [**N,6**] Seconds, 0-59 [N,7]Day of week, 0-6, 0 = Sunday[N,8] Days since Jan 1 of current year, 0-365 m Example let dtv = { 2001 3 26 11 7 22 1 84 }; 0 print "dtv = " dtv; p dt = dtvtodt(dtv);print "dt = " dt; q produces: dtv = 2001 3 26 11 7 22 1 84; dt = 20010326110722u Source time.src V See also dtvnormal, timeutc, utctodtv, dttodtv, dtvtodt, dttoutc, dtvtodt, strtodt, dttostr W

dtvtoutc

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dtvtoutc

Purpose Converts DTV vector format to UTC scalar format. **Format** utc = dtvtoutc(dtv); Input dtv Nx8 matrix. DTV vector format. Output utc Nx1 vector, UTC scalar format. Remarks A UTC scalar gives the number of seconds since or before January 1, 1970 Greenwich Mean Time. Each row of *dtv*, in DTV vector format, contains: [N,1]Year [N,2]Month in Year, 1-12 [N,3]Day of month, 1-31 [N,4]Hours since midnight, 0-23 [N,5]Minutes, 0-59 [N,6]Seconds, 0-59 [N,7]Day of week, 0-6, 0 = Sunday[N,8] Days since Jan 1 of current year, 0-365 Example dtv = utctodtv(timeutc); print "dtv = " dtv; utc = dtvtoutc(dtv); print "utc = " utc; produces: $dtv = 2001 \ 3 \ 26 \ 11 \ 7 \ 22 \ 1$ utc = 84985633642See also dtvnormal, timeutc, utctodt, dttodtv, dttoutc, dtvtodt, dtvtoutc, strtodt, dttostr

dummy

dummy

Purpose

Creates a set of dummy (0/1) variables by breaking up a variable into specified categories. The highest (rightmost) category is unbounded on the right

the right.

Format y = dummy(x, v);

Input x Nx1 vector of data that is to be broken up into dummy variables.

(K-1)x1 vector specifying the K-1 breakpoints (these must be in ascending order) that determine the K categories to be used.
 These categories should not overlap.

Output y NxK matrix containing the K dummy variables.

Remarks Missings are deleted before the dummy variables are created.

All categories are open on the left (i.e., do not contain their left boundaries) and all but the highest are closed on the right (i.e., do contain their right boundaries). The highest (rightmost) category is unbounded on the right. Thus, only K-1 breakpoints are required to specify K dummy variables.

The function **dummybr** is similar to **dummy**, but in that function the highest category is bounded on the right. The function **dummydn** is also similar to **dummy**, but in that function a specified column of dummies is dropped.

Example $x = \{ 0, 2, 4, 6 \};$ $v = \{ 1, 5, 7 \};$ y = dummy(x, v);

The result y looks like this:

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dummy

The vector v will produce 4 dummies satisfying the following conditions:

$$x \leq 1$$

$$1 < x \le 5$$

$$5 < x \le 7$$

Source datatran.src

See also dummybr, dummydn

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Purpose Creates a set of dummy (0/1) variables. The highest (rightmost) category is bounded on the right.

Format y = dummybr(x, v);

Input x Nx1 vector of data that is to be broken up into dummy variables.

KX1 vector specifying the K breakpoints (these must be in ascending order) that determine the K categories to be used.
 These categories should not overlap.

Output y NxK matrix containing the K dummy variables. Each row will have a maximum of one 1.

Remarks Missings are deleted before the dummy variables are created.

All categories are open on the left (i.e., do not contain their left boundaries) and are closed on the right (i.e., do contain their right boundaries). Thus, K breakpoints are required to specify K dummy variables.

The function **dummy** is similar to **dummybr**, but in that function the highest category is unbounded on the right.

Example $x = \{0, 2, 2, \dots \}$

4,

6};

 $v = \{ 1, 5,$

7};

y = dummybr(x,v);

The resulting matrix y looks like this:

dummybr

The vector v = 1.5.7 will produce 3 dummies satisfying the following conditions:

$$x \leq 1$$

$$1 < x \le 5$$

$$5 < x \le 7$$

Source datatran.src

See also dummydn, dummy

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dummydn

dummydn

Input

Purpose

Creates a set of dummy (0/1) variables by breaking up a variable into specified categories. The highest (rightmost) category is unbounded on the right, and a specified column of dummies is dropped.

Format $y = \operatorname{dummydn}(x, v, p)$;

x Nx1 vector of data to be broken up into dummy variables.

Kx1 vector specifying the K-1 breakpoints (these must be in ascending order) that determine the K categories to be used.
 These categories should not overlap.

p positive integer in the range [1,K], specifying which column should be dropped in the matrix of dummy variables.

Output y Nx(K-1) matrix containing the K-1 dummy variables.

Remarks This is just like the function **dummy**, except that the p^{th} column of the matrix of dummies is dropped. This ensures that the columns of the matrix of dummies do not sum to 1, and so these variables will not be collinear with a vector of ones.

Missings are deleted before the dummy variables are created.

All categories are open on the left (i.e., do not contain their left boundaries) and all but the highest are closed on the right (i.e., do contain their right boundaries). The highest (rightmost) category is unbounded on the right. Thus, only K-1 breakpoints are required to specify K dummy variables.

Example $x = \{ 0, 2, 4, 6 \};$ $v = \{ 1, 5, 7 \};$ p = 2;

y = dummydn(x,v,p);

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dummydn

The resulting matrix y looks like this:

- 1 0 0
- 0 0 0
- $0 \ 0 \ 0$
- 0 1 0

The vector v = 1.5.7 will produce 4 dummies satisfying the following conditions:

$$x \leq 1$$

$$1 < x \le 5$$

$$5 < x \le 7$$

Source datatran.src

See also dummy, dummybr

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Purpose Accesses an alternate editor.

Format ed filename;

Input *filename* The name of the file to be edited.

Remarks The default name of the editor is set in gauss.cfg. To change the name of the editor used type:

ed = editor_name flags;

or

ed = "editor_name flags";

The flags are any command line flags you may want between the name of the editor and the filename when your editor is invoked. The quoted version will prevent the flags, if any, from being forced to uppercase.

This command can be placed in the startup file so it will be set for you automatically when you start GAUSS.

edit

edit

Purpose Edits a disk file.

Format edit filename;

Remarks The edit command does not follow the src_path to locate files. You must

specify the location in the filename. The default location is the current

directory.

Example edit test1.e;

See also run

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Purpose Computes the eigenvalues of a general matrix.

Format va = eig(x);

Input x NxN matrix.

Output va Nx1 vector, the eigenvalues of x.

Remarks If the eigenvalues cannot all be determined, va[1] is set to an error code. Passing va[1] to the **scalerr** function will return the index of the

eigenvalue that failed. The eigenvalues for indices scalerr(va[1])+1 to N should be correct.

Error handling is controlled with the low bit of the trap flag.

trap 0 set va[1] and terminate with message

trap 1 set va[1] and continue execution

The eigenvalues are unordered except that complex conjugate pairs of eigenvalues will appear consecutively with the eigenvalue having the positive imaginary part first.

Example $x = \{481,$

942,

5 5 7 };

va = eig(x);

-4.4979246

va = 14.475702

5.0222223

See also eigh, eighv, eigv

eigcg

eigcg

Purpose Computes the eigenvalues of a complex, general matrix. (Included for backwards compatibility — use eig instead.)

Format { var, vai } = eigcg(xr, xi);

Input *xr* NxN matrix, real part.

xi NxN matrix, imaginary part.

Output *var* Nx1 vector, real part of eigenvalues.

vai Nx1 vector, imaginary part of eigenvalues.

_eigerr global scalar, if all the eigenvalues can be determined _eigerr = 0, otherwise _eigerr is set to the index of the eigenvalue that failed. The eigenvalues for indices

_eigerr+1 to N should be correct.

Remarks Error handling is controlled with the low bit of the trap flag.

trap 0 set _eigerr and terminate with message
trap 1 set _eigerr and continue execution

The eigenvalues are unordered except that complex conjugate pairs of eigenvalues will appear consecutively with the eigenvalue having the positive imaginary part first.

Source eigcg.src

Globals _eigerr

See also eigcg2, eigch, eigrg, eigrs

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eigcg2

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Purpose Computes eigenvalues and eigenvectors of a complex, general matrix. (Included for backwards compatibility — use **eigv** instead.) **Format** $\{ var, vai, ver, vei \} = eigcg2(xr, xi);$ Input xr NxN matrix, real part. xiNxN matrix, imaginary part. **Output** Nx1 vector, real part of eigenvalues. var vai Nx1 vector, imaginary part of eigenvalues. ver NxN matrix, real part of eigenvectors. vei NxN matrix, imaginary part of eigenvectors. **Global Input** eigerr global scalar, if all the eigenvalues can be determined

eigerr = 0, otherwise eigerr is set to the index of the eigenvalue that failed. The eigenvalues for indices eigerr+1 to N should be correct. The eigenvectors are not computed.

Remarks Error handling is controlled with the low bit of the trap flag.

> trap 0 set _eigerr and terminate with message trap 1 set eigerr and continue execution

The eigenvalues are unordered except that complex conjugate pairs of eigenvalues will appear consecutively with the eigenvalue having the positive imaginary part first. The columns of *ver* and *vei* contain the real and imaginary eigenvectors of x in the same order as the eigenvalues. The eigenvectors are not normalized.

Source eigcg.src

Globals eigerr

See also eigcg, eigch, eigrg, eigrs

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eigch

eigch

Purpose Computes the eigenvalues of a complex, hermitian matrix. (Included for backwards compatibility — use eigh instead.)

Format va = eigch(xr,xi);

Input *xr* NxN matrix, real part.

xi NxN matrix, imaginary part.

Output *va* Nx1 vector, real part of eigenvalues.

_eigerr global scalar, if all the eigenvalues can be determined

_eigerr = 0, otherwise **_eigerr** is set to the index of the eigenvalue that failed. The eigenvalues for indices 1 to

_eigerr-1 should be correct.

Remarks Error handling is controlled with the low bit of the trap flag.

trap 0 set _eigerr and terminate with message
trap 1 set eigerr and continue execution

The eigenvalues are in ascending order. The eigenvalues for a complex hermitian matrix are always real so this procedure returns only one vector.

Source eigch.src

Globals _eigerr

See also eigch2, eigcg, eigrg, eigrs

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eigch2

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Purpose
                Computes eigenvalues and eigenvectors of a complex, hermitian matrix.
                (Included for backwards compatibility — use eighv instead.)
  Format
                \{ var, vai, ver, vei \} = eigch2(xr, xi);
     Input
                xr
                        NxN matrix, real part.
                xi
                        NxN matrix, imaginary part.
  Output
                           Nx1 vector, real part of eigenvalues.
                var
                vai
                           Nx1 vector, imaginary part of eigenvalues.
                ver
                           NxN matrix, real part of eigenvectors.
                vei
                           NxN matrix, imaginary part of eigenvectors.
                           global scalar, if all the eigenvalues can be determined
                           _eigerr = 0, otherwise _eigerr is set to the index of the
                           eigenvalue that failed. The eigenvalues for indices 1 to
                           eigerr-1 should be correct. The eigenvectors are not
                           computed.
Remarks
                Error handling is controlled with the low bit of the trap flag.
                    trap 0 set eigerr and terminate with message
                    trap 1 set eigerr and continue execution
```

The eigenvalues are in ascending order. The eigenvalues of a complex hermitian matrix are always real. This procedure returns a vector of zeros for the imaginary part of the eigenvalues so the syntax is consistent with other **eig***xx* procedure calls. The columns of *ver* and *vei* contain the real and imaginary eigenvectors of *x* in the same order as the eigenvalues. The eigenvectors are orthonormal.

Source eigch.src

Globals _eigerr

See also eigch, eigcg, eigrg, eigrs

W

eigh

Purpose Computes the eigenvalues of a complex hermitian or real symmetric matrix.

Format va = eigh(x);

Input x NxN matrix.

Output va Nx1 vector, the eigenvalues of x.

Remarks If the eigenvalues cannot all be determined, va[1] is set to an error code. Passing va[1] to the **scalerr** function will return the index of the

eigenvalue that failed. The eigenvalues for indices 1 to $\mathbf{scalerr}(va[1])$ -1 should be correct.

Error handling is controlled with the low bit of the trap flag.

trap 0 set va[1] and terminate with message

trap 1 set va[1] and continue execution

The eigenvalues are in ascending order.

The eigenvalues of a complex hermitian or real symmetric matrix are

always real.

See also eig, eighv, eigv

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eighv

eighv

Purpose

Computes eigenvalues and eigenvectors of a complex hermitian or real symmetric matrix.

Format $\{va, ve\} = eighv(x);$

Input x NxN matrix.

Output va Nx1 vector, the eigenvalues of x.

ve NxN matrix, the eigenvectors of x.

Remarks If the eigenvalues cannot all be determined, va[1] is set to an error code.

Passing va[1] to the **scalerr** function will return the index of the eigenvalue that failed. The eigenvalues for indices 1 to **scalerr**(va[1])-1 should be correct. The eigenvectors are not computed.

Error handling is controlled with the low bit of the trap flag.

trap 0 set va[1] and terminate with message

trap 1 set va[1] and continue execution

The eigenvalues are in ascending order. The columns of ve contain the eigenvectors of x in the same order as the eigenvalues. The eigenvectors are orthonormal.

The eigenvalues of a complex hermitian or real symmetric matrix are always real.

See also eig, eigh, eigv

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eigrg

Purpose Computes the eigenvalues of a real, general matrix. (Included for backwards compatibility — use eig instead.)

Format $\{ var, vai \} = eigrg(x);$

Input x NxN matrix.

Output *var* Nx1 vector, real part of eigenvalues.

vai Nx1 vector, imaginary part of eigenvalues.

_eigerr global scalar, if all the eigenvalues can be determined _eigerr = 0, otherwise _eigerr is set to the index of the eigenvalue that failed. The eigenvalues for indices

eigenvalue that failed. The eigenvalues for indic

_eigerr+1 to N should be correct.

Remarks Error handling is controlled with the low bit of the trap flag.

trap 0 set _eigerr and terminate with message
trap 1 set _eigerr and continue execution

The eigenvalues are unordered except that complex conjugate pairs of eigenvalues will appear consecutively with the eigenvalue having the positive imaginary part first.

Example

10.301956

$$n = -1.4598755$$

$$6.2306252$$

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Source eigrg.src

Globals _eigerr

See also eigrg2, eigcg, eigch, eigrs

eigrg2

Purpose Computes eigenvalues and eigenvectors of a real, general matrix. (Included for backwards compatibility — use eigv instead.)

Format { var, vai, ver, vei } = eigrg2(x);

Input x NxN matrix.

Output *var* Nx1 vector, real part of eigenvalues.

vai Nx1 vector, imaginary part of eigenvalues.

ver NxN matrix, real part of eigenvectors.

vei NxN matrix, imaginary part of eigenvectors.

_eigerr global scalar, if all the eigenvalues can be determined

_eigerr = 0, otherwise _eigerr is set to the index of the

eigenvalue that failed. The eigenvalues for indices

_eigerr+1 to N should be correct. The eigenvectors are

not computed.

Remarks Error handling is controlled with the low bit of the trap flag.

trap 0 set _eigerr and terminate with message
trap 1 set eigerr and continue execution

The eigenvalues are unordered except that complex conjugate pairs of eigenvalues will appear consecutively with the eigenvalue having the positive imaginary part first. The columns of *ver* and *vei* contain the real and imaginary eigenvectors of *x* in the same order as the eigenvalues. The eigenvectors are not normalized.

Source eigrg.src

Globals _eigerr

See also eigrg, eigcg, eigch, eigrs

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eigrs

eigrs

Remarks

a b d e h k m n 0 p q t u

V W x y z

```
Purpose
               Computes the eigenvalues of a real, symmetric matrix. (Included for
               backwards compatibility — use eigh instead.)
 Format
               va = eigrs(x);
    Input
                      NxN matrix.
 Output
                          Nx1 vector, eigenvalues of x.
               va
               eigerr global scalar, if all the eigenvalues can be determined
                          _eigerr = 0, otherwise _eigerr is set to the index of the
                          eigenvalue that failed. The eigenvalues for indices 1 to
                          _eigerr-1 should be correct.
```

Error handling is controlled with the low bit of the trap flag. trap 0 set _eigerr and terminate with message

trap 1 set eigerr and continue execution

The eigenvalues are in ascending order. The eigenvalues for a real symmetric matrix are always real so this procedure returns only one vector.

Source eigrs.src

Globals eigerr

See also eigrs2, eigcg, eigch, eigrg

eigrs2

Purpose Computes eigenvalues and eigenvectors of a real, symmetric matrix. (Included for backwards compatibility — use **eighv** instead.)

Format $\{va, ve\} = eigrs2(x);$

Input x NxN matrix.

Output va Nx1 vector, eigenvalues of x.

ve NxN matrix, eigenvectors of x.

_eigerr global scalar, if all the eigenvalues can be determined

_eigerr = 0, otherwise **_eigerr** is set to the index of the eigenvalue that failed. The eigenvalues and eigenvectors for

indices 1 to **_eigerr**-1 should be correct.

Remarks Error handling is controlled with the low bit of the trap flag.

trap 0 set _eigerr and terminate with message
trap 1 set _eigerr and continue execution

The eigenvalues are in ascending order. The columns of ve contain the eigenvectors of x in the same order as the eigenvalues. The eigenvectors are orthonormal.

The eigenvalues and eigenvectors for a real symmetric matrix are always real so this procedure returns only the real parts.

Source eigrs.src

Globals _eigerr

See also eigrs, eigcg, eigch, eigrg

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Purpose Computes eigenvalues and eigenvectors of a general matrix.

Format $\{va, ve\} = eigv(x);$

Input x NxN matrix.

Output va Nx1 vector, the eigenvalues of x.

ve NxN matrix, the eigenvectors of x.

Remarks If the eigenvalues cannot all be determined, va[1] is set to an error code.

Passing va[1] to the **scalerr** function will return the index of the eigenvalue that failed. The eigenvalues for indices **scalerr**(va[1])+1 to N should be correct. The eigenvectors are not computed.

Error handling is controlled with the low bit of the trap flag.

trap 0 set va[1] and terminate with message **trap 1** set va[1] and continue execution

The eigenvalues are unordered except that complex conjugate pairs of eigenvalues will appear consecutively with the eigenvalue having the positive imaginary part first. The columns of ve contain the eigenvectors of x in the same order as the eigenvalues. The eigenvectors are not normalized.

Example

$$x = \{481, 942, 557\};$$

 $\{y,n\} = eigv(x);$
 -4.4979246
 $y = 14.475702$
 5.0222223
 $-0.66930459 -0.64076622 -0.40145623$
 $n = 0.71335708 -0.72488533 -0.26047487$
 $-0.01915672 -0.91339349 1.6734214$

eigv

See also eig, eigh, eighv

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elapsedTradingDays

elapsedTradingDays

Purpose Compute number of trading days between two dates inclusively.

Format n = elapsedTradingDays(a,b);

Input a scalar, date in DT scalar format.

b scalar, date in DT scalar format.

Output n number of trading days between dates inclusively, that is,

elapsed time includes the dates a and b.

Remarks A trading day is a weekday that is not a holiday as defined by the New

York Stock Exchange from 1888 through 2004. Holidays are defined in holidays. asc. You may edit that file to modify or add holidays.

Source finutils.src

3-246

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end

Format end;

Remarks end causes GAUSS to revert to interactive mode, and closes all open files. end also closes the auxiliary output file and turns the screen on. It is not necessary to put an end statement at the end of a program.

An end command can be placed above a label which begins a subroutine to make sure that a program does not enter a subroutine without a gosub.

stop also terminates a program but closes no files and leaves the screen setting as it is.

Example output on;

screen off;

end;
In this example, a matrix **x** is printed to the auxiliary output. The output to the screen is turned off to speed up the printing. The **end** statement is used to terminate the program so the output file will be closed and the screen will be turned back on.

See also new, stop, system

print x;

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endp

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x y z

endp

Purpose Closes a procedure or keyword definition. **Format** endp; Remarks **endp** marks the end of a procedure definition that began with a **proc** or **keyword** statement. (For details on writing and using procedures, see "Procedures and Keywords" in the *User's Guide*.) **Example** proc regress(y,x); retp(inv(x'x)*x'y);endp; $x = \{ 1 3 2, 7 4 9, 1 1 6, 3 3 2 \};$ $y = \{ 3, 5, 2, 7 \};$ b = regress(y,x);1.00000000 3.00000000 2.00000000 7.00000000 4.00000000 9.00000000 x =1.00000000 1.00000000 6.00000000 3.00000000 3.00000000 2.00000000 3.00000000 5.00000000 v =2.00000000 7.00000000 0.15456890 b =1.50276345 -0.12840825See also proc, keyword, retp

endwind

endwind

Purpose Ends graphic panel manipulation; display graphs with rerun.

Library pgraph

Format endwind;

Remarks This function uses **rerun** to display the most recently created .tkf file.

Source pwindow.src

See also begwind, window, makewind, setwind, nextwind,

getwind

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envget

envget

Searches the environment table for a defined name. **Purpose** h **Format** y = envget(s);Input string, the name to be searched for. d e **Output** v string, the string that corresponds to that name in the environment or a null string if it is not found. **Example** proc dopen(file); local fname, fp; h fname = envget("DPATH"); if fname \$== ""; fname = file; else; if strsect(fname,strlen(fname),1) \$== "\\"; m fname = fname \$+ file; else; 0 fname = fname \$+ "\\" \$+ file; endif; p endif; open fp = ^fname; retp(fp); endp; This is an example of a procedure which will open a data file using a path u stored in an environment string called DPATH. The procedure returns the file handle and is called as follows: V fp = dopen("myfile"); W

050

See also

cdir

3-250

x y z

eof

Purpose Tests if the end of a file has been reached.

Format y = eof(fh);

Input *fh* scalar, file handle.

Output y scalar, 1 if end of file has been reached, else 0.

Remarks This function is used with the **readr** and **fgets**xxx commands to test for the end of a file.

The **seekr** function can be used to set the pointer to a specific row position in a data set; the **fseek** function can be used to set the pointer to a specific byte offset in a file opened with **fopen**.

In this example, the data file dat1.dat is opened and given the handle **£1**. Then the data are read from this data set and are used to create the moment (**x'x**) matrix of the data. On each iteration of the loop, 100 additional rows of data are read in and the moment matrix for this set of rows is computed, and added to the matrix **xx**. When all the data have been read, **xx** will contain the entire moment matrix for the data set.

GAUSS will keep reading until **eof(f1)** returns the value 1, which it will when the end of the data set has been reached. On the last iteration of the loop, all remaining observations are read in if there are 100 or fewer left.

See also open, readr, seekr

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eqSolve

eqSolve

Purpose Solves a system of nonlinear equations h $\{x, retcode\} = eqSolve(&F, start);$ Format Input d start Kx1 vector, starting values. scalar, a pointer to a procedure which computes the value at x of &F the equations to be solved. **Global Input** The following are set by eqsolveset. egs JacobianProc pointer to a procedure which computes the analytical Jacobian. By default, egsolve will compute the Jacobian numerically. scalar, the maximum number of iterations. egs MaxIters Default = 100. scalar, the step tolerance. _eqs_StepTol Default = $macheps^{2}(2/3)$. Kx1 vector of the typical F(x) values at a point egs TypicalF not near a root, used for scaling. This becomes important when the magnitudes of the components of F(x) are expected to be very m different. By default, function values are not scaled. Kx1 vector of the typical magnitude of x, used egs TypicalX for scaling. This becomes important when the magnitudes of the components of x are p expected to be very different. By default, variable values are not scaled. eas IterInfo scalar, if nonzero, iteration information is printed. Default = 0. The following are set by gausset. scalar, the tolerance of the scalar function Tol $f = 0.5*||F(x)||^2$ required to terminate the u algorithm. Default = 1e-5. V

W

хуZ

egSolve

__altnam Kx1 character vector of alternate names to be used by the printed output. By default, the names X1, X2, X3... or X01, X02, X03 (depending on how __vpad is set) will be used.

__output scalar. If non-zero, final results are printed.

title string, a custom title to be printed at the top of

string, a custom title to be printed at the top of the iterations report. By default, only a generic title will be printed.

Output

Kx1 vector, solution.

retcode scalar, the return code:

- 1 Norm of the scaled function value is less than ___Tol. x given is an approximate root of F(x) (unless ___Tol is too large).
- **2** The scaled distance between the last two steps is less than the step-tolerance ($_{\tt eqs_StepTol}$). x may be an approximate root of F(x), but it is also possible that the algorithm is making very slow progress and is not near a root, or the step-tolerance is too large.
- 3 The last global step failed to decrease norm2(F(x)) sufficiently; either x is close to a root of F(x) and no more accuracy is possible, or an incorrectly coded analytic Jacobian is being used, or the secant approximation to the Jacobian is inaccurate, or the step-tolerance is too large.
- 4 Iteration limit exceeded.
- Five consecutive steps of maximum step length have been taken; either norm2(F(x)) asymptotes from above to a finite value in some direction or the maximum step length is too small.
- 6 x seems to be an approximate local minimizer of norm2(F(x)) that is not a root of F(x). To find a root of F(x), restart **eqSolve** from a different region.

Remarks

The equation procedure should return a column vector containing the result for each equation. For example:

Equation 1:
$$x1^2 + x2^2 - 2 = 0$$

Equation 2: $exp(x1-1) + x2^3 - 2 = 0$
proc f(var);

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eqSolve

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```
local x1,x2,eqns;
                  x1 = var[1];
                  x2 = var[2];
                  eqns[1] = x1^2 + x2^2 - 2; /* Equation 1 */
                  eqns[2] = \exp(x1-1) + x2^3 - 2; /*Equation*/
                                                       /* 2 */
                  retp( eqns );
               endp;
Example
           eqSolveset;
            proc f(x);
                  local f1, f2, f3;
                  f1 = 3*x[1]^3 + 2*x[2]^2 + 5*x[3] - 10;
                  f2 = -x[1]^3 - 3*x[2]^2 + x[3] + 5;
                  f3 = 3*x[1]^3 + 2*x[2]^2 - 4*x[3];
                  retp(f1|f2|f3);
            endp;
            proc fjc(x);
               local fjc1,fjc2, fjc3;
               fjc1 = 9*x[1]^2 \sim 4*x[2] \sim 5;
               fjc2 = -3*x[1]^2 \sim -6*x[2] \sim 1;
               fjc3 = 9*x[1]^2 \sim 4*x[2] \sim -4;
               retp(fjc1|fjc2|fjc3);
            endp;
            start = \{ -1, 12, -1 \};
            _eqs_JacobianProc = &fjc;
```

egSolve

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q

x y z

```
{ x,tcode } = eqSolve(&f,start);
Produces
______
EgSolve Version 3.2.22
                       2/24/97 9:54 am
______
||F(X)|| at final solution:
Termination Code = 1:
Norm of the scaled function value is less than
    __Tol;
VARIABLE START ROOTS F(ROOTS)
      -1.00000 0.54144351 4.4175402e-06
X1
X2 12.00000 1.4085912 -6.6263102e-06
      -1.00000 1.1111111 4.4175402e-06
X3
eqsolve.src
```

Source

eqSolveset

eqSolveset

Purpose Sets global input used by eqsolve to default values.

Format eqSolveset;

Global Output

```
\underline{\hspace{0.5cm}} \texttt{\_eqs\_TypicalX} = 0
```

 $__{\tt eqs_TypicalF} = 0$

 $__$ eqs_IterInfo = 0

 $__$ eqs $_$ MaxIters = 100

__eqs_StepTol __macheps^(2/3)

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erf, erfc

Computes the Gaussian error function (erf) and its complement (erfc). **Purpose**

Format y = erf(x);y = erfc(x);

Input NxK matrix.

Output NxK matrix.

Remarks The allowable range for x is:

$$x \ge 0$$

The **erf** and **erfc** functions are closely related to the Normal distribution:

$$cdfn(x) = \begin{cases} \frac{1}{2} \left(1 + erf\left(\frac{x}{\sqrt{2}}\right) \right) & x \ge 0\\ \frac{1}{2} erfc\left(\frac{-x}{\sqrt{2}}\right) & x < 0 \end{cases}$$

Example $x = \{ .5 .4 .3 ,$.6 .8 .3 }; y = erf(x);

> 0.52049988 0.42839236 0.32862676 0.60385609 0.74210096 0.32862676

 $x = \{ .5.4.3,$.6 .8 .3 }; y = erfc(x);

 $y = 0.47950012 \ 0.57160764 \ 0.67137324$ 0.39614391 0.25789904 0.67137324 a

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x y z

erf, erfc

See also cdfn, cdfnc

Technical erf and erfc are computed by summing the appropriate series and continued fractions. They are accurate to about 10 digits.

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error

Purpose

Allows the user to generate a user-defined error code which can be tested quickly with the **scalerr** function.

Format y = error(x);

Input x scalar, in the range 0-65535.

proc syminv(x);

Output y scalar error code which can be interpreted as an integer with the **scalerr** function.

Remarks

The user may assign any number in the range 0-65535 to denote particular error conditions. This number may be tested for as an error code by **scalerr**.

The **scalerr** function will return the value of the error code and so is the reverse of **error**. These user-generated error codes work in the same way as the intrinsic GAUSS error codes which are generated automatically when **trap 1** is on and certain GAUSS functions detect a numerical error such as a singular matrix.

error(0) is equal to the missing value code.

Example

```
local oldtrap,y;
if not x == x';
    retp(error(99));
endif;
oldtrap = trapchk(0xffff);
trap 1;
y = invpd(x);
if scalerr(y);
    y = inv(x);
endif;
```

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trap oldtrap,0xffff;
 retp(y);
endp;

The procedure **syminv** returns error code 99 if the matrix is not symmetric. If **invpd** fails, it returns error code 20. If **inv** fails, it returns error code 50. The original trap state is restored before the procedure returns.

See also scalerr, trap, trapchk

errorlog

errorlog

Purpose Prints an error message to the window and error log file.

Format errorlog str;

Input *str* string, the error message to print.

Remarks This function prints to the screen and the error log file.

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etdays

etdays

a **Purpose** Computes the difference between two times, as generated by the **date** b command, in days. **Format** days = etdays(tstart, tend); d Input 3x1 or 4x1 vector, starting date, in the order: yr, mo, day. (Only tstart e the first 3 elements are used.) tend 3x1 or 4x1 vector, ending date, in the order: yr, mo, day. (Only the first 3 elements are used.) MUST be later than *tstart*. **Output** scalar, elapsed time measured in days. days h Remarks This will work correctly across leap years and centuries. The assumptions are a Gregorian calendar with leap years on the years evenly divisible by 4 and not evenly divisible by 400. k Example let date1 = 1986 1 2; let date2 = 1987 10 25; d = etdays(date1,date2); m d = 661n 0 Source time.src p See also dayinyr q

x y z

u

V

W

ethsec

ethsec

Purpose Computes the difference between two times, as generated by the date command, in hundredths of a second.

Format hs = ethsec(tstart, tend);

Input *tstart* 4x1 vector, starting date, in the order: yr, mo, day, hundredths of

a second.

tend 4x1 vector, ending date, in the order: yr, mo, day, hundredths of a

second. MUST be later than tstart.

Output *hs* scalar, elapsed time measured in hundredths of a second.

Remarks This will work correctly across leap years and centuries. The assumptions

are a Gregorian calendar with leap years on the years evenly divisible by

4 and not evenly divisible by 400.

Example let date1 = 1986 1 2 0;

let date2 = 1987 10 25 0;

t = ethsec(date1,date2);

t = 5711040000

Source time.src

See also dayinyr

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3-263

etstr

etstr

a b d e h k m n 0 p q u

```
Purpose
              Formats an elapsed time, measured in hundredths of a second, to a string.
  Format
              str = etstr(tothsecs);
    Input
              tothsecs scalar, an elapsed time measured in hundredths of a second, as
                       given, for instance, by the ethsec function.
  Output
                     string containing the elapsed time in the form:
              str
                                  # hours # minutes #.## seconds
Example
              d1 = \{ 86, 1, 2, 0 \};
              d2 = \{ 86, 2, 5, 815642 \};
               t = ethsec(d1,d2);
               str = etstr(t);
               t = 2.9457564e + 08
              str = 34 \ days \ 2 \ hours \ 15 \ minutes \ 56.42 \ seconds
  Source
              time.src
```

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EuropeanBinomCall

EuropeanBinomCall

```
European binomial method call.
Purpose
  Format
              c = \text{EuropeanBinomCall}(SO, K, r, div, tau, sigma, N);
    Input
                    scalar, current price
              SO.
              K
                     Mx1 vector, strike prices
              r
                     scalar, risk free rate
                    continuous dividend yield
              div
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma scalar, volatility
              N
                     number of time segments
  Output
                     Mx1 vector, call premiums
Example
              S0 = 718.46;
             K = \{ 720, 725, 730 \};
              r = .0498;
              sigma = .2493;
              t0 = dtday (2001, 1, 30);
              t1 = dtday (2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                     annualTradingDays(2001);
              n = S0;
              c = EuropeanBinomCall(S0,K,r,div,tau,sigma,N);
              print c;
              17.1071
              15.0067
              12.9064
```

EuropeanBinomCall

Source

finprocs.src

Technical Notes

The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is the basis of this procedure.

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EuropeanBinomCall Greeks

EuropeanBinomCall_Greeks

```
European binomial method call Delta, Gamma, Theta, Vega and Rho.
Purpose
  Format
               \{d,g,t,v,rh\} =
              EuropeanBinomCall Greeks(SO, K, r, div, tau, sigma, N);
    Input
              S0
                      scalar, current price
              K
                      Mx1 vector, strike price
                      scalar, risk free rate
              r
                      continuous dividend yield
              div
                      scalar, elapsed time to exercise in annualized days of
               tau
                      trading
              sigma scalar, volatility
              N
                      number of time segments
  Output
              d
                      Mx1 vector, delta
                      Mx1 vector, gamma
              g
              t
                      Mx1 vector, theta
                      Mx1 vector, vega
               rh
                      Mx1 vector, rho
Example
              S0 = 305;
              K = 300;
              r = .08;
               sigma = .25;
              tau = .33;
              N = 30;
              print EuropeanBinomCall_Greeks
                      (S0,K,r,0,taou,sigma,N);
               0.6738
               0.0008
```

EuropeanBinomCall Greeks

-44.7874

69.0880

96.9225

Source finprocs.src

Globals _fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal.

Default = 0.

scalar, finite difference stepsize. Default = 1e-8. _fin_epsilon

Technical Notes The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is the basis of this procedure.

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EuropeanBinomCall_ImpVol

```
Implied volatilities for European binomial method calls.
Purpose
 Format
             sigma =
             EuropeanBinomCall ImpVol(c,S0,K,r,div,tau,N);
    Input
                    Mx1 vector, call premiums
             c
             SO
                    scalar, current price
                    Mx1 vector, strike prices
             K
                    scalar, risk free rate
              r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
              tau
                    trading
             N
                    number of time segments
  Output
             sigma Mx1 vector, volatility
Example
             c = \{ 13.70, 11.90, 9.10 \};
              S0 = 718.46;
             K = \{ 720, 725, 730 \};
             r = .0498;
              t0 = dtday (2001, 1, 30);
              t1 = dtday (2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) / annualTrading-
                    Days(2001);
             N = 30;
              sigma = EuropeanBinomCall_ImpVol
                    (c,S0,K,r,0,tau,N);
             print sigma;
              0.1982
```

EuropeanBinomCall_ImpVol

0.1716

0.1301

Source

finprocs.src

Technical Notes

The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is the basis of this procedure.

b

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u

V

хух

EuropeanBinomPut

EuropeanBinomPut

```
Purpose
              European binomial method Put.
  Format
              c = \text{EuropeanBinomPut}(SO, K, r, div, tau, sigma, N);
     Input
                     scalar, current price
              SO.
               K
                     Mx1 vector, strike prices
               r
                     scalar, risk free rate
                     continuous dividend yield
              div
              div
                     continuous dividend yield
                     scalar, elapsed time to exercise in annualized days of
               tau
                     trading
              sigma scalar, volatility
                     number of time segments
              N
  Output
                     Mx1 vector, put premiums
Example:
              S0 = 718.46;
              K = \{ 720, 725, 730 \};
              r = .0498;
               sigma = .2493;
               t0 = dtday (2001, 1, 30);
              t1 = dtday (2001, 2, 16);
               tau = elapsedTradingDays(t0,t1) /
                     annualTradingDays(2001);
              N = 30;
               c = EuropeanBinomPut
                     (S0,K,r,0,tau,sigma,N);
              print c;
               16.6927
```

EuropeanBinomPut

19.5266

22.3604

Source finprocs.src

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m

n

0

q

t

u

V

x y z

EuropeanBinomPut Greeks

EuropeanBinomPut_Greeks

European binomial method put Delta, Gamma, Theta, Vega, and Rho. **Purpose Format** $\{d,g,t,v,rh\} =$ EuropeanBinomPut Greeks (SO, K, r, div, tau, sigma, N); Input *S0* scalar, current price K Mx1 vector, strike price scalar, risk free rate r continuous dividend yield div scalar, elapsed time to exercise in annualized days of tau trading sigma scalar, volatility N number of time segments **Output** d Mx1 vector, delta Mx1 vector, gamma g t Mx1 vector, theta Mx1 vector, vega rhMx1 vector, rho **Example** S0 = 305;K = 300;r = .08;sigma = .25;tau = .33;N = 30;print EuropeanBinomPut_Greeks (S0,K,r,0,tau,sigma,N); -0.38040.0038

EuropeanBinomPut_Greeks

-17.9838

69.0880

-33.7666

Globals

_fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal.

Default = 0.

_fin_epsilon scalar, finite difference stepsize. Default = 1e-8.

Source

finprocs.src

Technical Notes The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is

the basis of this procedure.

a

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k

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n

0

p

q

u

V

x y z

EuropeanBinomPut ImpVol

EuropeanBinomPut_ImpVol

Implied volatilities for European binomial method puts. **Purpose Format** sigma = EuropeanBinomPut ImpVol(c, SO, K, r, div, tau, N);Input Mx1 vector, put premiums c*S0* scalar, current price K Mx1 vector, strike prices scalar, risk free rate r div continuous dividend yield scalar, elapsed time to exercise in annualized days of tau trading N number of time segments Output sigma Mx1 vector, volatility $p = \{ 14.60, 17.10, 20.10 \};$ Example: S0 = 718.46; $K = \{ 720, 725, 730 \};$ r = .0498;t0 = dtday (2001, 1, 30);t1 = dtday (2001, 2, 16);tau = elapsedTradingDays(t0,t1) / annualTradingDays(2001); N = 30;sigma = EuropeanBinomPut ImpVol (p,S0,K,r,0,tau,N); print sigma; 0.1307 0.1714

EuropeanBinomPut_ImpVol

0.2165

Source

finprocs.src

Technical Notes The binomial method of Cox, Ross, and Rubinstein ("Option pricing: a simplified approach", Journal of Financial Economics, 7:229:264) as described in Options, Futures, and other Derivatives by John C. Hull is

the basis of this procedure.

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EuropeanBSCall

EuropeanBSCall

```
European Black and Scholes Call.
Purpose
  Format
              c = \text{EuropeanBSCall}(SO, K, r, div, tau, sigma);
    Input
                    scalar, current price
              S0
              K
                     Mx1 vector, strike prices
                     scalar, risk free rate
              r
                    continuous dividend yield
              div
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma scalar, volatility
  Output
                     Mx1 vector, call premiums
Example
              S0 = 718.46;
              K = \{ 720, 725, 730 \};
              b = .0498;
              r = .0498;
              sigma = .2493;
              t0 = dtday (2001, 1, 30);
              t1 = dtday (2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              c = EuropeanBSCall(S0,K,r,0,tau,sigma);
              print c;
              17.0975
              14.7583
              12.6496
```

EuropeanBSCall

Source finprocs.src

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С

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m

n

0

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u

V

x y z

EuropeanBSCall Greeks

EuropeanBSCall_Greeks

```
European Black and Scholes call Delta, Gamma, Omega, Theta, and
Purpose
              Vega.
  Format
               \{d,g,t,v,rh\} =
              EuropeanBSCall_Greeks(SO, K, r, div, tau, sigma);
    Input
                     scalar, current price
              S0
              K
                      Mx1 vector, strike price
              r
                     scalar, risk free rate
              div
                     continuous dividend yield
                      scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma
                     scalar, volatility
  Output
              d
                      Mx1 vector, delta
              g
                      Mx1 vector, gamma
              t
                     Mx1 vector, theta
                     Mx1 vector, vega
              ν
              rh
                      Mx1 vector, rho
Example
              S0 = 305;
              K = 300;
              r = .08;
              sigma = .25;
              tau = .33;
              print EuropeanBSCall Greeks (S0,K,r,0,tau,sigma);
              0.6446
              0.0085
              -38.5054
              65.2563
              56.8720
```

EuropeanBSCall_Greeks

Source finprocs.src

Globals __fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal. Default = 0.

w x y z

u

V

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a

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m

n

0

p

q

t

u

V

x y z

EuropeanBSCall ImpVol

EuropeanBSCall_ImpVol

```
Implied volatilities for European Black and Scholes calls.
Purpose
  Format
             sigma = EuropeanBSCall ImpVol(c,S0,K,r,div,tau);
    Input
             c
                    Mx1 vector, call premiums
              S0
                    scalar, current price
              K
                    Mx1 vector, strike prices
                    scalar, risk free rate
              r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
              tau
                    trading
  Output
             sigma Mx1 vector, volatility
Example
             c = \{ 13.70, 11.90, 9.10 \};
             S0 = 718.46i
             K = \{ 720, 725, 730 \};
             r = .0498;
              t0 = dtday (2001, 1, 30);
              t1 = dtday (2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              sigma = EuropeanBSCall ImpVol
                    (c,S0,K,r,0,tau);
             print sigma;
              0.1991
              0.1725
              0.1310
```

EuropeanBSCall_ImpVol

Source finprocs.src

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m

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u

V

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EuropeanBSPut

EuropeanBSPut

```
European Black and Scholes Put.
Purpose
  Format
              c = \text{EuropeanBSPut}(SO, K, r, div, tau, sigma);
    Input
                    scalar, current price
              SO.
              K
                     Mx1 vector, strike prices
              r
                    scalar, risk free rate
                    continuous dividend yield
              div
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
              sigma scalar, volatility
  Output
                     Mx1 vector, put premiums
Example
              S0 = 718.46;
              K = \{ 720, 725, 730 \};
              r = .0498;
              sigma = .2493;
              div = 0;
              t0 = dtday (2001, 1, 30);
              t1 = dtday (2001, 2, 16);
              tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
              c = EuropeanBSPut(S0,K,r,0,tau,sigma);
              print c;
              16.6403
              19.2872
              22.1647
```

EuropeanBSPut

Source finprocs.src

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n

0

q

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u

V

x y z

EuropeanBSPut Greeks

EuropeanBSPut_Greeks

```
European Black and Scholes put Delta, Gamma, Omega, Theta, and Vega.
Purpose
  Format
              \{d,g,t,v,rh\} =
              EuropeanBSPut_Greeks(S0,K,r,div,tau,sigma);
    Input
                     scalar, current price
              S0
              K
                     Mx1 vector, strike price
                     scalar, risk free rate
              r
                     continuous dividend yield
              div
                     scalar, elapsed time to exercise in annualized days of
              tau
                     trading
                     scalar, volatility
              sigma
  Output
              d
                     Mx1 vector, delta
                     Mx1 vector, gamma
              g
              t
                     Mx1 vector, theta
                     Mx1 vector, vega
              rh
                     Mx1 vector, rho
Example
              S0 = 305;
              K = 300;
              r = .08;
              sigma = .25;
              tau = .33;
              print EuropeanBSPut_Greeks (S0,K,r,0,tau,sigma);
              -0.3554
              0.0085
              -15.1307
              65.2563
              -39.5486
```

EuropeanBSPut_Greeks

Source finprocs.src

Globals __fin_thetaType scalar, if 1, one day look ahead, else, infinitesmal. Default = 0.

хух

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V

x y z

EuropeanBSPut ImpVol

EuropeanBSPut_ImpVol

```
Implied volatilities for European Black and Scholes puts.
Purpose
  Format
             sigma = EuropeanBSPut ImpVol(c,S0,K,r,div,tau);
    Input
                    Mx1 vector, put premiums
             c
             S0
                    scalar, current price
             K
                    Mx1 vector, strike prices
                    scalar, risk free rate
             r
             div
                    continuous dividend yield
                    scalar, elapsed time to exercise in annualized days of
             tau
                    trading
  Output
             sigma Mx1 vector, volatility
Example
             p = \{ 14.60, 17.10, 20.10 \};
             S0 = 718.46i
             K = \{ 720, 725, 730 \};
             r = .0498;
             t0 = dtday (2001, 1, 30);
             t1 = dtday (2001, 2, 16);
             tau = elapsedTradingDays(t0,t1) /
                    annualTradingDays(2001);
             sigma = EuropeanBSPut ImpVol(p,S0,K,r,0,tau);
             print sigma;
             0.2123
             0.2493
             0.2937
  Source
             finprocs.src
```

exctsmpl

exctsmpl

h d e m n p u V

Purpose Computes a random subsample of a data set. **Format** n = exctsmpl(infile,outfile, percent); Input infile string, the name of the original data set. outfile string, the name of the data set to be created. scalar, the percentage random sample to take. This must be in percent the range 0-100. **Output** scalar, number of rows in output data set. Error returns are controlled by the low bit of the trap flag. trap 0 terminate with error message **trap 1** return scalar negative integer can't open input file -2 can't open output file -3 disk full Remarks Random sampling is done with replacement. Thus, an observation may be in the resulting sample more than once. If *percent* is 100, the resulting sample will not be identical to the original sample, though it will be the same size.

Example n = exctsmpl("freq.dat", "rout", 30); n = 30

freq.dat is an example data set provided with GAUSS. Switching to the GAUSS examples directory will make it possible to do the above example as shown. Otherwise substitute data set names will need to be used.

Source exctsmpl.src

W

хух

exec

Purpose Executes an executable program and returns the exit code to GAUSS. **Format** y = exec(program, comline);Input string, the name of the program, including the extension, to program be executed. comline string, the arguments to be placed on the command line of the program being executed. **Output** the exit code returned by program. v If **exec** cannot execute *program*, the error returns will be negative. -1 file not found -2 the file is not an executable file -3 not enough memory -4 command line too long **Example** y = exec("atog.exe", "comd1.cmd"); if y; errorlog "atog failed"; end; endif; In this Windows example, the ATOG ASCII conversion utility is executed under the exec function. The name of the command file to be used,

In this Windows example, the ATOG ASCII conversion utility is executed under the **exec** function. The name of the command file to be used, comd1.cmd, is passed to ATOG on its command line. The exit code **y** returned by **exec** is tested to see if ATOG was successful; if not, the program will be terminated after printing an error message. See "Utilities" in the *User Guide*.

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execbg

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```
Purpose Executes an executable program in the background and returns the process id to GAUSS.
```

Format

pid = execbg(program, comline);

Input

program string, the name of the program, including the extension, to

be executed.

comline

string, the arguments to be placed on the command line of

the program being executed.

Output

pid the process id of the executable returned by program.

If **exec** cannot execute *program*, the error returns will be negative.

- **-1** file not found
- **-2** the file is not an executable file
- **-3** not enough memory
- **-4** command line too long

Example

```
y = execbg("atog.exe","comd1.cmd");
if (y < 0);
  errorlog "atog failed";
  end;
endif;</pre>
```

In this Windows example, the ATOG ASCII conversion utility is executed under the **execbg** function. The name of the command file to be used, comd1.cmd, is passed to ATOG on its command line. The returned value, **y**, is tested to see whether ATOG was successful. If not successful the program terminates after printing an error message. See "Utilities" in the *User Guide*.

exp

Purpose Calculates the exponential function.

Format $y = \exp(x)$;

Input x NxK matrix.

Output y NxK matrix containing e, the base of natural logs, raised to the powers given by the elements of x.

Example x = eye(3);

 $y = \exp(x);$

 $1.000000\ 0.000000\ 0.000000$

 $x = 0.000000 \ 1.000000 \ 0.000000$ $0.000000 \ 0.000000 \ 1.000000$

2.718282 1.000000 1.000000

 $y = 1.000000 \ 2.718282 \ 1.000000$

1.000000 1.000000 2.718282

This example creates a 3x3 identity matrix and computes the exponential function for each one of its elements. Note that exp(1) returns e, the base of natural logs.

See also ln

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extern (dataloop)

extern (dataloop)

Purpose

Allows access to matrices or strings in memory from inside a data loop.

Format

extern variable list;

Remarks

Commas in *variable_list* are optional.

extern tells the translator not to generate local code for the listed variables, and not to assume they are elements of the input data set.

extern statements should be placed before any reference to the symbols listed. The specified names should not exist in the input data set, or be used in a **make** statement.

Example

This example shows how to assign the contents of an external vector to a new variable in the data set, by iteratively assigning a range of elements to the variable. The reserved variable $\mathbf{x}_{-}\mathbf{x}$ contains the data read from the input data set on each iteration. The external vector must have at least as many rows as the data set.

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x y z

external

external

Purpose

Lets the compiler know about symbols that are referenced above or in a separate file from their definitions.

Format

```
external proc dog,cat;
external keyword dog;
external fn dog;
external matrix x,y,z;
external string mstr,cstr;
```

Remarks

See "Procedures and Keywords" in the *User's Guide*.

You may have several procedures in different files that reference the same global variable. By placing an **external** statement at the top of each file, you can let the compiler know if the symbol is a matrix, string, or procedure. If the symbol is listed and strongly typed in an active library, no **external** statement is needed.

If a matrix or string appears in an **external** statement it needs to appear once in a **declare** statement. If no declaration is found, an **Undefined symbol** error will result.

Example

The general eigenvalue procedure, eigrg, sets a global variable _eigerr if it cannot compute all of the eigenvalues.

```
external matrix _eigerr;

x = rndn(4,4);

xi = inv(x);

xev = eigrg(x);

if _eigerr;

  print "Eigenvalues not computed";
  end;
end;
```

Without the **external** statement, the compiler would assume that **_eigerr** was a procedure and incorrectly compile this program. The file containing the **eigrg** procedure also contains an external statement that

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external

defines _eigerr as a matrix, but this would not be encountered until the if statement containing the reference to _eigerr in the main program file had already been incorrectly compiled.

See also declare

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eye

Purpose Creates an identity matrix.

Format y = eye(n);

Input n scalar, size of identity matrix to be created.

Output y NxN identity matrix.

Remarks If n is not an integer, it will be truncated to an integer.

The matrix created will contain 1's down the diagonal and 0's everywhere

else.

Example x = eye(3);

1.000000 0.000000 0.000000

 $x = 0.000000 \ 1.000000 \ 0.000000$

 $0.000000 \ 0.000000 \ 1.000000$

See also zeros, ones

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fcheckerr

fcheckerr

Purpose Gets the error status of a file.

Format err = fcheckerr(f);

Input f scalar, file handle of a file opened with **fopen**.

Output *err* scalar, error status.

Remarks If there has been a read or write error on a file, **fcheckerr** returns 1, otherwise 0.

If you pass **fcheckerr** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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w x y z

fclearerr

fclearerr

Purpose Gets the error status of a file, then clears it.

Format err = fclearerr(f);

Input f scalar, file handle of a file opened with **fopen**.

Output *err* scalar, error status.

Remarks

Each file has an error flag that is set when there is an I/O error on the file. Typically, once this flag is set, you can no longer do I/O on the file, even if the error is a recoverable one. **fclearerr** clears the file's error flag, and you can attempt to continue using it.

If there has been a read or write error on a file, **fclearerr** returns 1, otherwise 0.

If you pass **fclearerr** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

The flag accessed by **fclearerr** is not the same as that accessed by **fstrerror**.

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feq, fge, fgt, fle, flt, fne

feq, fge, fgt, fle, flt, fne

Purpose

Fuzzy comparison functions. These functions use **_fcmptol** to fuzz the comparison operations to allow for roundoff error.

Format

```
y = feq(a,b);
y = fge(a,b);
y = fgt(a,b);
y = fle(a,b);
y = flt(a,b);
```

y = fne(a,b);

Input

NxK matrix, first matrix.

b LxM matrix, second matrix, ExE compatible with a.

Global Input

_fcmptol global scalar, comparison tolerance. The default value is 1.0e-15.

Output

scalar, 1 (true) or 0 (false).

Remarks

The return value is true if every comparison is true.

The statement:

```
y = feq(a,b);
```

is equivalent to:

$$y = a eq b;$$

For the sake of efficiency, these functions are not written to handle missing values. If a and b have missing values, use **missrv** to convert the missing values to something appropriate before calling a fuzzy comparison function.

The calling program can reset **_fcmptol** before calling these procedures.

```
_{\text{fcmptol}} = 1e-12;
```

Example

```
x = rndu(2,2);

y = rndu(2,2);
```

8

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W

хух

feq, fge, fgt, fle, flt, fne

t = fge(x,y);

 $x = \begin{array}{cc} 0.0382895 & 0.07253527 \\ 0.01471395 & 0.96863611 \end{array}$

 $y = \begin{array}{c} 0.25622293 & 0.70636474 \\ 0.00361912 & 0.35913385 \end{array}$

t = 0.0000000

Source fcompare.src

Globals _fcmptol

See also dotfeq-dotfne

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fflush

fflush

Purpose Flushes a file's output buffer.

Format ret = fflush(f);

Input f scalar, file handle of a file opened with **fopen**.

Output *ret* scalar, 0 if successful, -1 if not.

Remarks If fflush fails, you can call fstrerror to find out why.

If you pass **fflush** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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fft

Purpose Computes a 1- or 2-D Fast Fourier transform.

Format y = fft(x);

Input x NxK matrix.

Output y LxM matrix, where L and M are the smallest powers of 2 greater than or equal to N and K, respectively.

Remarks This computes the FFT of x, scaled by 1/N.

This uses a Temperton Fast Fourier algorithm.

If N or K is not a power of 2, x will be padded out with zeros before computing the transform.

Example $x = \{ 22 24, \\ 23 25 \};$ y = fft(x); $y = \frac{23.500000 -1.0000000}{-0.50000000 0.000000000}$

See also ffti, rfft, rffti

a

b

c d

0

f

g h

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J k

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Purpose Computes an inverse 1- or 2-D Fast Fourier transform.

Format y = ffti(x);

Input x NxK matrix.

Output y LxM matrix, where L and M are the smallest prime factor products greater than or equal to N and K, respectively.

Remarks Computes the inverse FFT of x, scaled by 1/N.

This uses a Temperton prime factor Fast Fourier algorithm.

Example $x = \{ 22 \ 24 ,$

 $23 \ 25$ }; y = fft(x);

23.500000 -1.000000

-0.500000 0.0000000

fi = ffti(y);

 $fi = \begin{array}{c} 22.000000 \ 24.000000 \\ 23.000000 \ 25.000000 \end{array}$

See also fft, rfft, rffti

fftm

Purpose Computes a multi-dimensional FFT.

Format y = fftm(x, dim);

Input x Mx1 vector, data.

dim Kx1 vector, size of each dimension.

Output y Lx1 vector, FFT of x.

Remarks

The multi-dimensional data are laid out in a recursive or heirarchical fashion in the vector x. That is to say, the elements of any given dimension are stored in sequence left to right within the vector, with each element containing a sequence of elements of the next smaller dimension. In abstract terms, a 4-dimensional 2x2x2x2 hypercubic x would consist of two cubes in sequence, each cube containing two matrices in sequence, each matrix containing two rows in sequence, and each row containing two columns in sequence. Visually, x would look something like this:

$$\begin{aligned} X_{hyper} &= X_{cube1} \mid X_{cube2} \\ X_{cube1} &= X_{mat1} \mid X_{mat2} \\ X_{mat1} &= X_{row1} \mid X_{row2} \\ X_{row1} &= X_{col1} \mid X_{col2} \end{aligned}$$

Or, in an extended GAUSS notation, *x* would be:

Xhyper =
$$x[1,...,.]$$
 | $x[2,...,.];$
Xcubel = $x[1,1,...]$ | $x[1,2,...];$
Xmatl = $x[1,1,1,...]$ | $x[1,1,2,...];$
Xrowl = $x[1,1,1,1]$ | $x[1,1,1,2];$

To be explicit, *x* would be laid out like this:

$$x[1,1,1,1]$$
 $x[1,1,1,2]$ $x[1,1,2,1]$ $x[1,1,2,2]$ $x[1,2,1,1]$ $x[1,2,2,2]$ $x[1,2,1,1]$ $x[1,2,1,2]$ $x[2,1,2,1]$ $x[2,1,2,2]$ $x[2,1,1,1]$ $x[2,1,1,2]$ $x[2,1,2,1]$ $x[2,1,2,2]$ $x[2,2,1,1]$ $x[2,2,2,2,2]$

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If you look at the last diagram for the layout of x you'll notice that each line actually constitutes the elements of an ordinary matrix in normal row-major order. This is easy to achieve with **vecr**. Further, each pair of lines or "matrices" constitutes one of the desired cubes, again with all the elements in the correct order. And finally, the two cubes combine to form the hypercube. So, the process of construction is simply a sequence of concatenations of column vectors, with a **vecr** step if necessary to get started.

Here's an example, this time working with a 2x3x2x3 hypercube.

```
let dim = 2 3 2 3;
let x1[2,3] = 1 2 3 4 5 6;
let x2[2,3] = 6 5 4 3 2 1;
let x3[2,3] = 1 2 3 5 7 11;
xc1 = vecr(x1)|vecr(x2)|vecr(x3); /* cube 1 */
let x1 = 1 1 2 3 5 8;
let x2 = 1 2 6 24 120 720;
let x3 = 13 17 19 23 29 31;
xc2 = x1|x2|x3; /* cube 2 */
xh = xc1|xc2; /* hypercube */
xhfft = fftm(xh,dim);
let dimi = 2 4 2 4;
xhffti = fftmi(xhfft,dimi);
```

We left out the **vecr** step for the 2^{nd} cube. It's not really necessary when you're constructing the matrices with **let** statements.

 \mathtt{dim} contains the dimensions of x, beginning with the highest dimension. The last element of \mathtt{dim} is the number of columns, the next to the last element of \mathtt{dim} is the number of rows, and so on. Thus

```
dim = \{ 2, 3, 3 \};
```

fftm

indicates that the data in x is a 2x3x3 three-dimensional array, i.e., two 3x3 matrices of data. Suppose that x1 is the first 3x3 matrix and x2 the second 3x3 matrix, then $x = vecr(x1) \mid vecr(x2)$.

The size of dim tells you how many dimensions x has.

The arrays have to be padded in each dimension to the nearest power of two. Thus the output array can be larger than the input array. In the 2x3x2x3 hypercube example, x would be padded from 2x3x2x3 out to 2x4x2x4. The input vector would contain 36 elements, while the output vector would contain 64 elements. You may have noticed that we used a dimi with padded values at the end of the example to check our answer.

Source fftm.src

See also fftmi, fft, ffti, fftn

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Computes a multi-dimensional inverse FFT.

Format y = fftmi(x, dim);

Input х Mx1 vector, data.

> Kx1 vector, size of each dimension. dim

Output Lx1 vector, inverse FFT of x.

Remarks

Purpose

The multi-dimensional data are laid out in a recursive or heirarchical fashion in the vector x. That is to say, the elements of any given dimension are stored in sequence left to right within the vector, with each element containing a sequence of elements of the next smaller dimension. In abstract terms, a 4-dimensional 2x2x2x2 hypercubic x would consist of two cubes in sequence, each cube containing two matrices in sequence, each matrix containing two rows in sequence, and each row containing two columns in sequence. Visually, x would look something like this:

$$\begin{aligned} \mathbf{X}_{hyper} &= \mathbf{X}_{cube1} \mid \mathbf{X}_{cube2} \\ &\mathbf{X}_{cube1} &= \mathbf{X}_{mat1} \mid \mathbf{X}_{mat2} \\ &\mathbf{X}_{mat1} &= \mathbf{X}_{row1} \mid \mathbf{X}_{row2} \\ &\mathbf{X}_{row1} &= \mathbf{X}_{col1} \mid \mathbf{X}_{col2} \end{aligned}$$

Or, in an extended GAUSS notation, x would be:

Xhyper = x[1,...] | x[2,....]; Xcube1 = x[1,1,...] | x[1,2,...1; Xmat1 = x[1,1,1,...] | x[1,1,2,...]; = x[1,1,1,1] | x[1,1,1,2]Xrow1];

To be explicit, x would be laid out like this:

x[1,1,1,1] x[1,1,1,2] x[1,1,2,1] x[1,1,2,2]x[1,2,1,1] x[1,2,1,2] x[1,2,2,1] x[1,2,2,2]x[2,1,1,1] x[2,1,1,2] x[2,1,2,1] x[2,1,2,2]x[2,2,1,1] x[2,2,1,2] x[2,2,2,1] x[2,2,2,2]

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If you look at the last diagram for the layout of x you'll notice that each line actually constitutes the elements of an ordinary matrix in normal row-major order. This is easy to achieve with **vecr**. Further, each pair of lines or "matrices" constitutes one of the desired cubes, again with all the elements in the correct order. And finally, the two cubes combine to form the hypercube. So, the process of construction is simply a sequence of concatenations of column vectors, with a **vecr** step if necessary to get started.

Here's an example, this time working with a 2x3x2x3 hypercube.

```
let dim = 2 3 2 3;

let x1[2,3] = 1 2 3 4 5 6;

let x2[2,3] = 6 5 4 3 2 1;

let x3[2,3] = 1 2 3 5 7 11;

xc1 = vecr(x1) | vecr(x2) | vecr(x3); /* cube 1 */

let x1 = 1 1 2 3 5 8;

let x2 = 1 2 6 24 120 720;

let x3 = 13 17 19 23 29 31;

xc2 = x1 | x2 | x3; /* cube 2 */

xh = xc1 | xc2; /* hypercube */
xhffti = fftmi(xh,dim);
```

We left out the **vecr** step for the 2^{nd} cube. It's not really necessary when you're constructing the matrices with **let** statements.

dim contains the dimensions of x, beginning with the highest dimension. The last element of **dim** is the number of columns, the next to the last element of **dim** is the number of rows, and so on. Thus

```
dim = \{ 2, 3, 3 \};
```

indicates that the data in x is a 2x3x3 three-dimensional array, i.e., two 3x3 matrices of data. Suppose that x1 is the first 3x3 matrix and x2 the second 3x3 matrix, then $x = vecr(x1) \mid vecr(x2)$.

The size of dim tells you how many dimensions x has.

fftmi

The arrays have to be padded in each dimension to the nearest power of two. Thus the output array can be larger than the input array. In the 2x3x2x3 hypercube example, *x* would be padded from 2x3x2x3 out to 2x4x2x4. The input vector would contain 36 elements, while the output vector would contain 64 elements.

Source fftm.src

See also fftmi, fft, ffti, fftn

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fftn

Purpose Computes a complex 1- or 2-D FFT.

Format y = fftn(x);

Input x NxK matrix.

Output y LxM matrix, where L and M are the smallest prime factor products greater than or equal to N and K, respectively.

Remarks

fftn uses the Temperton prime factor FFT algorithm. This algorithm can compute the FFT of any vector or matrix whose dimensions can be expressed as the product of selected prime number factors. GAUSS implements the Temperton algorithm for any power of 2, 3, and 5, and one factor of 7. Thus, **fftn** can handle any matrix whose dimensions can be expressed as

$$2^p \times 3^q \times 5^r \times 7^s$$
, p,q,r nonnegative integers $s = 0$ or 1

If a dimension of *x* does not meet this requirement, it will be padded with zeros to the next allowable size before the FFT is computed.

fftn pads matrices to the next allowable dimensions; however, it generally runs faster for matrices whose dimensions are highly composite numbers, i.e., products of several factors (to various powers), rather than powers of a single factor. For example, even though it is bigger, a 33600x1 vector can compute as much as 20% faster than a 32768x1 vector, because 33600 is a highly composite number, $2^6 \times 3 \times 5^2 \times 7$, whereas 32768 is a simple power of 2, 2^{15} . For this reason, you may want to hand-pad matrices to optimum dimensions before passing them to **fftn**. The Run-Time Library includes a routine, **optn**, for determining optimum dimensions.

The Run-Time Library also includes the **nextn** routine, for determining allowable dimensions for a matrix. (You can use this to see the dimensions to which **fftn** would pad a matrix.)

fftn scales the computed FFT by 1/(L*M).

See also fft, ffti, fftm, fftmi, rfft, rffti, rfftip, rfftn, rfftnp, rfftp

fgets

fgets

Purpose Reads a line of text from a file.

Format str = fgets(f, maxsize);

Input f scalar, file handle of a file opened with **fopen**.

maxsize scalar, maximum size of string to read in, including the

terminating null byte.

Output *str* string.

Remarks fgets reads text from a file into a string. It reads up to a newline, the end of the file, or *maxsize-1* characters. The result is placed in *str*, which is then terminated with a null byte. The newline, if present, is retained.

If the file is already at end-of-file when you call **fgets**, your program will terminate with an error. Use **eof** in conjunction with **fgets** to avoid this.

If the file was opened for update (see **fopen**) and you are switching from writing to reading, don't forget to call **fseek** or **fflush** first, to flush the file's buffer.

If you pass **fgets** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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fgetsa

fgetsa

Purpose Reads lines of text from a file into a string array.

Format sa = fgetsa(f, numl);

Input f scalar, file handle of a file opened with **fopen**.

numl scalar, number of lines to read.

Output sa Nx1 string array, N <= numl.

Remarks

fgetsa reads up to *numl* lines of text. If fgetsa reaches the end of the file before reading *numl* lines, *sa* will be shortened. Lines are read in the same manner as fgets, except that no limit is placed on the size of a line. Thus, fgetsa always returns complete lines of text. Newlines are retained. If *numl* is 1, fgetsa returns a string. (This is one way to read a line from a file without placing a limit on the length of the line.)

If the file is already at end-of-file when you call **fgetsa**, your program will terminate with an error. Use **eof** in conjunction with **fgetsa** to avoid this. If the file was opened for update (see **fopen**) and you are switching from writing to reading, don't forget to call **fseek** or **fflush** first, to flush the file's buffer.

If you pass **fgetsa** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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fgetsat

fgetsat

Purpose Reads lines of text from a file into a string array.

Format sa = fgetsat(f, numl);

Input f scalar, file handle of a file opened with **fopen**.

numl scalar, number of lines to read.

Output sa Nx1 string array, N <= numl.

Remarks fgetsat operates identically to fgetsa, except that newlines are not retained as text is read into sa.

In general, you don't want to use **fgetsat** on files opened in binary mode (see **fopen**). **fgetsat** drops the newlines, but it does NOT drop the carriage returns that precede them on some platforms. Printing out such a string array can produce unexpected results.

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fgetst

fgetst

Purpose Reads a line of text from a file.

Format str = fgetst(f, maxsize);

Input f scalar, file handle of a file opened with **fopen**.

maxsize scalar, maximum size of string to read in, including the null

terminating byte.

Output *str* string.

Remarks fgetst operates identically to fgets, except that the newline is not retained in the string.

In general, you don't want to use **fgetst** on files opened in binary mode (see **fopen**). **fgetst** drops the newline, but it does NOT drop the preceding carriage return used on some platforms. Printing out such a string can produce unexpected results.

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fileinfo

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fileinfo

Purpose Returns names and information for files that match a specification. **Format** { fnames, finfo } = fileinfo(fspec); Input string, file specification. Can include path. Wildcards are fspec allowed in the filename. **Output** fnames Nx1 string array of all filenames that match, null string if none are found. finfo Nx13 matrix, information about matching files. UNIX filesystem ID [N, 1] inode number [N, 2] [N, 3] mode bit mask number of links [N, 4]user ID [N, 5]group ID [N, 6] device ID (char/block special files [N, 7] only) size in bytes [N, 8] [N, 9] last access time last data modification time [N, 10][N,11]last file status change time preferred I/O block size [N, 12]number of 512-byte blocks allocated [N, 13]OS/2, Windows drive number (A = 0, B = 1, etc.) [N, 1] n/a, 0 [N, 2] mode bit mask [N, 3]number of links, always 1 [N, 4][N, 5]n/a. 0 n/a, 0 [N, 6]n/a, 0 [N, 7]size in bytes [N, 8]

fileinfo

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	[N, 9]	last access time
	[N,10]	last data modification time
	[N,11]	creation time
	[N,12]	n/a, 0
	[N,13]	n/a, 0
DOS		
	[N, 1]	drive number ($A = 0$, $B = 1$, etc.)
	[N, 2]	n/a, 0
	[N, 3]	mode bit mask
	[N, 4]	number of links, always 1
	[N, 5]	n/a, 0
	[N, 6]	n/a, 0
	[N, 7]	n/a, 0
	[N, 8]	size in bytes
	[N, 9]	n/a, 0
	[N,10]	last data modification time
	[N,11]	n/a, 0
	[N,12]	n/a, 0
	[N,13]	n/a, 0

finfo will be a scalar zero if no matches are found.

Remarks

fnames will contain file *names* only; any path information that was passed is dropped.

The time stamp fields (*finfo*[N,9]-[N,11]) are expressed as the number of seconds since midnight, Jan. 1, 1970, Coordinated Universal Time (UTC).

See also files, filesa

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filesa

filesa

a **Purpose** Returns a string array of file names. b **Format** y = filesa(n);Input string, file specification to search for. Can include path. d n Wildcards are allowed in the filename. е **Output** Nx1 string array of all filenames that match, or null string if none are found. Remarks y will contain file *names* only; any path information that was passed is h dropped. Example y = filesa("ch*"); In this example all files listed in the current directory that begin with "ch" will be returned. k proc exist(filename); retp(not filesa(filename) \$== ""); m endp; This procedure will return 1 if the file exists or 0 if not. 0 See also fileinfo, files, shell p q t

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floor

Purpose Rounds down toward $-\infty$.

Format y = floor(x);

Input x NxK matrix.

Output y NxK matrix containing the elements of x rounded down.

Remarks This rounds every element in the matrix *x* down to the nearest integer.

Example x = 100*rndn(2,2);

$$x = \begin{array}{c} 77.68 & -14.10 \\ 4.73 & -158.88 \end{array}$$

f = floor(x);

$$f = \begin{array}{cc} 77.00 & -15.00 \\ 4.00 & -159.00 \end{array}$$

See also ceil, round, trunc

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fmod

fmod

a b

Computes the floating-point remainder of x/y.

Format

Purpose

r = fmod(x, y);

Input

x NxK matrix.

y LxM matrix, ExE conformable with x.

Output

 $r = \max(N,L)$ by $\max(K,M)$ matrix.

Remarks

Returns the floating-point remainder r of x/y such that x = iy + r, where i is an integer, r has the same sign as x, and |r| < |y|.

Compare this with %, the modulo division operator. (See "Operators" in the *User's Guide*.)

Example

x = seqa(1.7, 2.3, 5)';

y = 2;

r = fmod(x,y);

 $x = 1.7 \quad 4 \quad 6.3 \quad 8.6 \quad 10.9$

 $r = 1.7 \quad 0 \quad 0.3 \quad 0.6 \quad 0.9$

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fn

Purpose Allows user to create one-line functions.

Format fn fn_name(args) = code_for_function;

Remarks Functions can be called in the same way as other procedures.

Example fn area(r) = pi*r*r;

a = area(4);

a = 50.265482

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fonts

Purpose Loads fonts to be used in the graph.

Library pgraph

Format fonts(str);

Input str string or character vector containing the names of fonts to be

used in the plot.

Simplex standard sans serif font.
Simgrma Simplex Greek, math.

Microb bold and boxy.

Complex standard font with serif.

The first font specified will be used for the axes numbers.

If *str* is a null string, or **fonts** is not called, Simplex is loaded

by default.

Remarks For information on how to select fonts within a text string, see

"Publication Quality Graphics in the User's Guide.

Source pgraph.src

See also title, xlabel, ylabel, zlabel

w x y z

fopen

fopen

Purpose Opens a file.

Format f = fopen(filename, omode);

Input filename string, name of file to open.

omode string, file I/O mode. (See Remarks, below.)

Output f scalar, file handle.

Portability UNIX

Carriage return-linefeed conversion for files opened in text mode is unnecessary, because in UNIX a newline is simply a linefeed.

Remarks *filename* can contain a path specification.

omode is a sequence of characters that specify the mode in which to open the file. The first character must be one of:

- r Open an existing file for reading. If the file does not exist, **fopen** fails.
- w Open or create a file for writing. If the file already exists, its current contents will be destroyed.
- a Open or create a file for appending. All output is appended to the end of the file.

To this can be appended a + and/or a **b**. The + indicates the file is to opened for reading and writing, or update, as follows:

- **r**+ Open an existing file for update. You can read from or write to any location in the file. If the file does not exist, **fopen** fails.
- w+ Open or create a file for update. You can read from or write to any location in the file. If the file already exists, its current contents will be destroyed.
- Open or create a file for update. You can read from any location in the file, but all output will be appended to the end of the file.

Finally, the **b** indicates whether the file is to be opened in text or binary mode. If the file is opened in binary mode, the contents of the file are read verbatim; likewise, anything output to the file is written verbatim. In text mode (the default), carriage return-linefeed sequences are converted on

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fopen

input to linefeeds, or newlines. Likewise on output, newlines are converted to carriage return-linefeeds. Also in text mode, if a CTRL-Z (char 26) is encountered during a read, it is interpreted as an end-of-file character, and reading ceases. In binary mode, CTRL-Z is read in uninterpreted.

The order of + and **b** is not significant; **rb**+ and **r**+**b** mean the same thing.

You can both read from and write to a file opened for update. However, before switching from one to the other, you must make an **fseek** or **fflush** call, to flush the file's buffer.

If **fopen** fails, it returns a 0.

Use close and closeall to close files opened with fopen.

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for

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Purpose
               Begins a for loop.
  Format
                for i (start, stop, step);
                endfor:
    Input
                i
                        literal, the name of the counter variable.
                start
                        scalar expression, the initial value of the counter.
                stop
                        scalar expression, the final value of the counter.
                        scalar expression, the increment value.
                step
Remarks
                The counter is strictly local to the loop. The expressions start, stop, and
                step are evaluated only once when the loop initializes. They are converted
                to integers and stored local to the loop.
                The for loop is optimized for speed and is much faster than a do loop.
                The commands break and continue are supported. The continue
                command steps the counter and jumps to the top of the loop. The break
                command terminates the current loop.
                The loop terminates when the value of i exceeds stop. If break is used to
                terminate the loop and you want the final value of the counter, you need to
                assign it to a variable before the break statement (see the third example,
                following).
Example
               Example 1
               x = zeros(10, 5);
                for i(1, rows(x), 1);
                   for j (1, cols(x), 1);
                       x[i,j] = i*j;
                   endfor;
```

endfor;

for

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x y z

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Example 2
x = rndn(3,3);
y = rndn(3,3);
for i(1, rows(x), 1);
   for j(1, cols(x), 1);
      if x[i,j] >= y[i,j];
        continue;
      endif;
     temp = x[i,j];
     x[i,j] = y[i,j];
     y[i,j] = temp;
   endfor;
endfor;
Example 3
li = 0;
x = rndn(100,1);
y = rndn(100,1);
for i(1, rows(x), 1);
   if x[i] /= y[i];
      li = i;
     break;
   endif;
endfor;
if li;
   print "Compare failed on row " li;
endif;
```

3-324

format

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format

Purpose Controls the format of matrices and numbers printed out with print or lprint statements. **Format** format [[/typ]] [[/fmted]] [[/mf]] [[/jnt]] [[f,p]]; Input literal, symbol type flag(s). Indicate which symbol types you are /typ setting the output format for. /mat, /sa, /str Formatting parameters are maintained separately for matrices (/mat), string arrays (/sa), and strings (/str). You can specify more than one /typ flag; the format will be set for all types indicated. If no /typ flag is listed, format assumes /mat. /fmted literal, enable formatting flag. /on, /off Enable/disable formatting. When formatting is disabled, the contents of a variable are dumped to the window in a "raw" format. /off is currently supported only for strings. Raw format for strings means that the entire string is printed, starting at the current cursor position. When formatting is enabled for strings, they are handled the same as string arrays. This shouldn't be too surprising, since a string is actually a 1x1 string array. /mf literal, matrix row format flag. no delimiters before or after rows when /m0printing out matrices. print 1 carriage return/line feed pair /ml or /mbl before each row of a matrix with more than 1 row. /m2 or /mb2print 2 carriage return/line feed pairs before each row of a matrix with more than 1 row. /m3 or /mb3print "Row 1", "Row 2"... before each row of a matrix with more than one row.

format

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/ma1 print 1 carriage return/line feed pair after each row of a matrix with more than 1 row. print 2 carriage return/line feed pairs /ma2 after each row of a matrix with more than 1 row. print 1 carriage return/line feed pair /a1 after each row of a matrix. print 2 carriage return/line feed pairs /a2 after each row of a matrix. /b1 print 1 carriage return/line feed pair before each row of a matrix. print 2 carriage return/line feed pairs /b2 before each row of a matrix. print "Row 1", "Row 2"... before each /b3 row of a matrix. literal, matrix element format flag controls justification, notation /jnt and trailing character. Right-Justified Signed decimal number in the form /rd [-]] ####.###, where #### is one or more decimal digits. The number of digits before the decimal point depends on the magnitude of the number, and the number of digits after the decimal point depends on the precision. If the precision is 0, no decimal point will be printed. Signed number in the form /re [-]#.##E±###, where # is one decimal digit, ## is one or more decimal digits depending on the precision, and ### is three decimal digits. If precision is 0, the form will be \[-\] #E±### with no decimal point printed.

w x y z

u

This will give a format like /rd or /re

depending on which is most compact for the number being printed. A format

format

like /re will be used only if the exponent value is less than -4 or greater than the precision. If a /re format is used, a decimal point will always appear. The precision signifies the number of significant digits displayed. This will give a format like /rd or /re /rz depending on which is most compact for the number being printed. A format like /re will be used only if the exponent value is less than -4 or greater than the precision. If a /re format is used, trailing zeros will be supressed and a decimal point will appear only if one or more digits follow it. The precision signifies the number of significant digits displayed. Left-Justified /ld Signed decimal number in the form [-] ####.####, where #### is one or more decimal digits. The number of digits before the decimal point depends on the magnitude of the number, and the number of digits after the decimal point depends on the precision. If the precision is 0, no decimal point will be printed. If the number is positive, a space character will replace the leading minus sign. /le Signed number in the form [-] #.##E±###, where # is one decimal digit, ## is one or more decimal digits depending on the precision, and ### is three decimal digits. If precision is 0, the form will be \[-\] #E±### with no decimal point printed. If the number is positive, a space character will replace the leading minus sign.

/ro

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format

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		/lc	This will give a format like /ld or /le depending on which is most compact for the number being printed. A format like /le will be used only if the exponent value is less than -4 or greater than the precision. If a /le format is used, a decimal point will always appear. If the number is positive, a space character will replace the leading minus sign. The precision specifies the number of significant digits displayed. This will give a format like /ld or /le depending on which is most compact for the number being printed. A format like /le will be used only if the exponent value is less than -4 or greater than the precision. If a /le format is used, trailing zeros will be supressed and a decimal point will appear only if one or more digits follow it. If the number is positive, a space character will replace the leading minus sign. The precision specifies the number of
		Trailing Character	significant digits displayed.
		The following characters can be added to the /jnt parameters above to control the trailing character if any:	
			format /rdn 1,3;
		s	The number will be followed immediately by a space character. This is the default.
		С	The number will be followed immediately by a comma.
		t	The number will be followed immediately by a tab character.
		n	No trailing character.
	f	scalar expression, controls the field width.	
	p	scalar expression, controls the precision.	
Remarks	If character elements are to be printed, the precision should be at least 8 or		

the elements will be truncated. This does not affect the string data type.

3-328

format

For numeric values in matrices, p sets the number of significant digits to be printed. For string arrays, strings, and character elements in matrices, p sets the number of characters to be printed. If a string is shorter than the specified precision, the entire string is printed. For string arrays and strings, p = -1 means print the entire string, regardless of its length. p = -1 is illegal for matrices; setting p >= 8 means the same thing for character elements.

The /xxx slash parameters are optional. Field and precision are optional also but if one is included, then both must be included.

Slash parameters, if present, must precede the field and precision parameters.

A **format** statement stays in effect until it is overridden by a new **format** statement. The slash parameters may be used in a **print** statement to override the current default.

f and p may be any legal expressions that return scalars. Nonintegers will be truncated to integers.

The total width of field will be overridden if the number is too big to fit into the space allotted. For instance, **format** /rds 1,0 can be used to print integers with a single space between them, regardless of the magnitudes of the integers.

Complex numbers are printed with the sign of the imaginary half separating them and an "i" appended to the imaginary half. Also, the field parameter refers to the width of field for each half of the number, so a complex number printed with a field of 8 will actually take (at least) 20 spaces to print. The character printed after the imaginary part can be changed (for example, to a "j") with the **sysstate** function, case 9.

The default when GAUSS is first started is:

format /mb1 /ros 16,8;

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c d

P

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Example

format

a

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хуг

```
This code:
x = rndn(3,3);
format /m1 /rd 16,8;
print x;
produces:
  -1.63533465 1.61350700
                               -1.06295179
   0.26171282 0.27972294
                               -1.38937242
   0.58891114 0.46812202
                               1.08805960
This code:
format /m1 /rzs 1,10;
print x;
produces:
-1.6353346
                1.613507
                             -1.0629518
 0.26171282
            0.27972294
                             -1.3893724
 0.58891114 0.46812202
                             1.0880596
This code:
format /m3 /rdn 16,4;
print x;
produces:
Row 1
       -1.6353 1.6135
                        -1.0630
Row 2
        0.2617 0.2797
                         -1.3894
Row 3
        0.5889 0.4681
                          1.0881
```

3-330

format

a

b

С

d

e

f

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h

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хух

```
This code:
format /m1 /ldn 16,4;
print x;
produces:
  -1.6353
            1.6135
                       -1.0630
   0.2617 0.2797
                       -1.3894
   0.5889
          0.4681
                        1.0881
This code:
format /m1 /res 12,4;
print x;
produces:
  -1.6353E+000
                  1.6135E+000
                                 -1.0630E+000
   2.6171E-001
                  2.7972E-001
                                 -1.3894E+000
   5.8891E-001
                  4.6812E-001
                                  1.0881E+000
```

See also formatcv, formatnv, print, lprint, output

formatcv

formatcv

a **Purpose** Sets the character data format used by **printfmt**. b Format oldfmt = formatcv(newfmt); Input *newfmt* 1x3 vector, the new format specification. d е Output 1x3 vector, the old format specification. oldfmt See **printfm** for details on the format vector. Remarks Example This example saves the old format, sets the format desired for printing x, h prints x, then restores the old format. This code: $x = \{ A 1, B 2, C 3 \};$ oldfmt = formatcv("*.*s" ~ 3 ~ 3); call printfmt(x,0~1); k call formatcv(oldfmt); produces: m 1 Α n В 2 0 C3 Source p gauss.src q Globals fmtcv See also formatny, printfm, printfmt

u

V

W

x y z

formatny

formatnv

Purpose Sets the numeric data format used by printfmt.

Format oldfmt = formatnv(newfmt);

Input *newfmt* 1x3 vector, the new format specification.

Output *oldfmt* 1x3 vector, the old format specification.

Remarks See printfm for details on the format vector.

Example This example saves the old format, sets the format desired for printing x, prints x, then restores the old format. This code:

x = { A 1, B 2, C 3 };
oldfmt = formatnv("*.*lf" ~ 8 ~ 4);
call printfmt(x,0~1);

call formatnv(oldfmt);
produces:

A 1

в 2

C 3

Source gauss.src

Globals __fmtnv

See also formatcv, printfm, printfmt

a

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c d

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fputs

fputs

Purpose Writes strings to a file.

Format numl = fputs(f, sa);

Input f scalar, file handle of a file opened with **fopen**.

sa string or string array.

Output *numl* scalar, the number of lines written to the file.

Portability UNIX

Carriage return-linefeed conversion for files opened in text mode is unnecessary, because in UNIX a newline is simply a linefeed.

Remarks fputs writes the contents of each string in sa, minus the null terminating

byte, to the file specified. If the file was opened in text mode (see **fopen**), any newlines present in the strings are converted to carriage return-linefeed sequences on output. If *numl* is not equal to the number of elements in *sa*, there may have been an I/O error while writing the file. You can use **fcheckerr** or **fclearerr** to check this. If there was an error, you can call **fstrerror** to find out what it was. If the file was opened for update (see **fopen**) and you are switching from reading to writing, don't forget to call **fseek** or **fflush** first, to flush the file's buffer. If you pass **fputs** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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fputst

fputst

Purpose Writes strings to a file.

Format numl = fputst (f, sa);

Input f scalar, file handle of a file opened with **fopen**.

sa string or string array.

Output *numl* scalar, the number of lines written to the file.

Portability UNIX

Carriage return-linefeed conversion for files opened in text mode is unnecessary, because in UNIX a newline is simply a linefeed.

Remarks fputst works identically to fputs, except that a newline is appended

to each string that is written to the file. If the file was opened in text mode (see **fopen**), these newlines are also converted to carriage return-

linefeed sequences on output.

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V

W

fseek

fseek

h d h k m 0 p u V

```
Purpose
                Positions the file pointer in a file.
 Format
                 ret = fseek(f, offs, base);
    Input
                        scalar, file handle of a file opened with fopen.
                offs
                        scalar, offset (in bytes).
                        scalar, base position.
                base
                                beginning of file.
                        1
                                current position of file pointer.
                        2
                                end of file.
 Output
                ret
                        scalar, 0 if successful, 1 if not.
```

Portability UNIX

Carriage return-linefeed conversion for files opened in text mode is unnecessary, because in UNIX a newline is simply a linefeed.

Remarks

fseek moves the file pointer *offs* bytes from the specified *base* position. *offs* can be positive or negative. The call may fail if the file buffer needs to be flushed (see **fflush**).

If fseek fails, you can call fstrerror to find out why.

For files opened for update (see **fopen**), the next operation can be a read or a write.

fseek is not reliable when used on files opened in text mode (see **fopen**). This has to do with the conversion of carriage return-linefeed sequences to newlines. In particular, an **fseek** that follows one of the **fgets**xx or **fputs**xx commands may not produce the expected result. For example:

```
p = ftell(f);
s = fgetsa(f,7);
call fseek(f,p,0);
```

is not reliable. The best results are obtained by **fseek**'ing to the beginning of the file and then **fseek**'ing to the desired location, as in

W

хуZ

fseek

```
p = ftell(f);
s = fgetsa(f,7);
call fseek(f,0,0);
call fseek(f,p,0);
```

If you pass **fseek** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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fstrerror

fstrerror

Purpose Returns an error message explaining the cause of the most recent file I/O error.

Format *s* = fstrerror;

Output *s* string, error message.

Remarks
Any time an I/O error occurs on a file opened with fopen, an internal error flag is updated. (This flag, unlike those accessed by fcheckerr and fclearerr, is not specific to a given file; rather, it is system-wide.) fstrerror returns an error message based on the value of this flag, clearing it in the process. If no error has occurred, a null string is returned.

Since **fstrerror** clears the error flag, if you call it twice in a row, it will always return a null string the second time.

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ftell

ftell

Purpose Gets the position of the file pointer in a file.

Format pos = ftell(f);

Input f scalar, file handle of a file opened with **fopen**.

Output pos scalar, current position of the file pointer in a file.

Remarks ftell returns the position of the file pointer in terms of bytes from the beginning of the file. The call may fail if the file buffer needs to be

flushed (see fflush).

If an error occurs, **ftell** returns -1. You can call **fstrerror** to find out what the error was.

If you pass **ftell** the handle of a file opened with **open** (i.e., a data set or matrix file), your program will terminate with a fatal error.

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ftocv

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S

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Purpose Converts a matrix containing floating point numbers into a matrix containing the decimal character representation of each element.

Format y = ftocv(x, field, prec);

Input *x* NxK matrix containing numeric data to be converted.

field scalar, minimum field width.

prec scalar, the numbers created will have prec places after the

decimal point.

Output y NxK matrix containing the decimal character equivalent of the corresponding elements in x in the format defined by *field* and *prec*.

Remarks If a number is narrower than *field*, it will be padded on the left with zeros.

If prec = 0, the decimal point will be suppressed.

Example y = seqa(6,1,5);x = 0 \$+ "cat" \$+ ftocv(y,2,0);

cat06

cat07

x = cat08

cat09

cat10

Notice that the (0 \$+) above was necessary to force the type of the result to matrix because the string constant **cat** would be of type string. The left operand in an expression containing a \$+ operator controls the type of the result.

See also ftos

ftos

Purpose

Converts a scalar into a string containing the decimal character representation of that number.

Format

$$y = ftos(x, fmat, field, prec);$$

Input

x scalar, the number to be converted.

fmat string, the format string to control the conversion.

field scalar or 2x1 vector, the minimum field width. If *field* is 2x1, it specifies separate field widths for the real and imaginary parts of *x*.

prec scalar or 2x1 vector, the number of places following the decimal point. If prec is 2x1, it specifies separate precisions for the real and imaginary parts of x.

Output

y string containing the decimal character equivalent of x in the format specified.

Remarks

The format string corresponds to the **format** /jnt (justification, notation, trailing character) slash parameter as follows:

```
/rdn "%*.*lf"
/ren "%*.*lE"
/ron "%#*.*lG"
/rzn "%*.*lG"

/ldn "%- *.*lf"
/len "%- *.*lE"
/lon "%-# *.*lG"

/lzn "%- *.*lG"
```

If *x* is complex, you can specify separate formats for the real and imaginary parts by putting two format specifications in the format string. You can also specify separate fields and precisions. You can position the sign of the imaginary part by placing a "+" between the two format specifications. If you use two formats, no "i" is appended to the imaginary

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ftos

part. This is so you can use an alternate format if you prefer, for example, prefacing the imaginary part with a "j".

The format string can be a maximum of 80 characters.

If you want special characters to be printed after *x*, include them as the last characters of the format string. For example:

```
"%*.*lf," right-justified decimal followed by a comma.
"%-*.*s" left-justified string followed by a space.
"%*.*lf" right-justified decimal followed by nothing.
```

You can embed the format specification in the middle of other text.

```
"Time: %*.*lf seconds."
```

If you want the beginning of the field padded with zeros, then put a "0" before the first "*" in the format string:

```
"%0*.*lf" right-justified decimal.
```

If prec = 0, the decimal point will be suppressed.

Example

You can create custom formats for complex numbers with **ftos**. For example,

```
let c = 24.56124+6.3224e-2i;

field = 1;
prec = 3|5;
fmat = "%lf + j%le is a complex number.";
cc = ftos(c,fmat,field,prec);

results in
    cc = "24.561 + j6.32240e-02 is a complex number."
```

Some other things you can do with ftos:

```
let x = 929.857435324123;
let y = 5.46;
let z = 5;
field = 1;
```

a

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a

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w x y z

ftos

a

b

С

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V

x y z

```
prec = 0;
              fmat = "%*.*lf";
              zz = ftos(z,fmat,field,prec);
              field = 1;
              prec = 10;
              fmat = "%*.*lE";
              xx = ftos(x,fmat,field,prec);
              field = 7;
              prec = 2;
              fmat = "%*.*lf seconds";
              s1 = ftos(x,fmat,field,prec);
              s2 = ftos(y,fmat,field,prec);
              field = 1;
              prec = 2;
              fmat = "The maximum resistance is %*.*lf
                      ohms.";
              om = ftos(x,fmat,field,prec);
           The results:
              zz = "5"
              xx = "9.2985743532E+02"
              s1 = " 929.86 seconds"
              s2 = " 5.46 seconds"
              om = "The maximum resistance is 929.86 ohms."
See also ftocv, stof, format
```

ftostrC

ftostrC

a **Purpose** Converts a matrix to a string array using a C language format h specification. **Format** sa = ftostrC(x, fmt);d Input NxK matrix, real or complex. х fmt Kx1, 1xK or 1x1 string array containing format information. Output NxK string array. sa Remarks If fmt has K elements, each column of sa can be formatted separately. If x h is complex, there must be two format specifications in each element of fmt. **Example** declare string fmtr = { "%6.31f", k "%11.81f" **}**; m declare string fmtc = { n "(%6.31f, %6.31f)", 0 "(%11.8lf, %11.8lf)" p }; q xr = rndn(4, 2);xc = sqrt(xr')';sar = ftostrC(xr, fmtr); u sac = ftostrC(xc, fmtc); V print sar; W print sac;

x y z

ftostrC

```
produces:
```

(0.92784529, 0.00000000)

See also strtof, strtofcplx

(0.888, 0.000)

a

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С

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gamma

gamma

a b

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k

m

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0 p

q

u

V

W

x y z

Purpose Returns the value of the gamma function.

Format y = gamma(x);

Input NxK matrix.

Output NxK matrix.

For each element of x, this function returns the integral Remarks

$$\int_0^\infty t^{(x-1)} e^{-t} dt$$

All elements of x must be positive and less than or equal to 169. Values of x greater than 169 will cause an overflow.

The natural log of gamma is often what is required and it can be computed without the overflow problems of gamma. Infact can be used to compute log gamma.

Example y = gamma(2.5);

y = 1.32934

See also cdfchic, cdfbeta, cdffc, cdfn, cdfnc, cdftc, erf,

erfc

gammaii

gammaii

Purpose Computes the inverse incomplete gamma function.

Format x = gammaii(a,p);

Input *a* MxN matrix, exponents.

p KxL matrix, ExE conformable with a, incomplete gamma

values.

Output $x = \max(M,K)$ by $\max(N,L)$ matrix, abscissae.

Source cdfchii.src

Globals _ginvinc, __macheps

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gausset

a

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gausset

Purpose Resets the global control variables declared in gauss.dec.

Format gausset;

Source gauss.src

Globals __altnam, __con, __ff, __fmtcv, __fmtnv,

__header, __miss, __output, __row, __rowfac,

__sort, __title, __tol, __vpad, __vtype, __weight

u

V

getarray

getarray

Purpose Gets a contiguous subarray from an N-dimensional array.

Format y = getarray(a, loc);

Input *a* N-dimensional array.

loc Mx1 vector of indices into the array to locate the subarray of

interest, where M is a value from 1 to N.

Output y [N-M]-dimensional subarray or scalar.

Remarks If N-M > 0, getarray will return an array of [N-M] dimensions, otherwise, if N-M = 0, it will return a scalar.

Example a = seqa(1,1,720);

a = areshape(a, 2|3|4|5|6);

loc = { 2,1 };

y = getarray(a,loc);

y will be a 4x5x6 array of sequential values, beginning at [1,1,1] with 361, and ending at [4,5,6] with 480.

See also getmatrix

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getdims

a

b

d

e

g

h

k

m

n

0

p

q

getdims

Purpose Gets the number of dimensions in an array.

Format y = getdims(a);

Input a N-dimensional array.

Output y scalar, the number of dimensions in the array.

Example a = arrayinit(3|4|5|6|7|2,0);
 dims = getdims(a);

dims = 6

See also getorders

w x y z

u

V

getf

Purpose Loads an ASCII or binary file into a string.

Format y = getf(filename, mode);

Input *filename* string, any valid file name.

mode scalar 1 or 0 which determines if the file is to be loaded in

ASCII mode (0) or binary mode (1).

Output y string containing the file.

Remarks

If the file is loaded in ASCII mode, it will be tested to see if it contains any end of file characters. These are ^Z (ASCII 26). The file will be truncated before the first ^Z and there will be no ^Z's in the string. This is the correct way to load most text files because the ^Z's can cause problems when trying to print the string to a printer.

If the file is loaded in binary mode, it will be loaded just like it is with no changes.

Example

Create a file examp. e containing the following program.

```
library pgraph;
```

```
graphset;
```

```
x = seqa(0,0.1,100);
```

$$y = \sin(x);$$

Then execute the following.

```
y = getf("examp.e", 0);
```

```
print y;
```

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getf

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```

```
This produces:
library pgraph;
graphset;
x = seqa(0,0.1,100);
y = sin(x);
xy(x,y);
```

See also load, save, let, con

V

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getmatrix

getmatrix

Purpose Gets a contiguous matrix from an N-dimensional array.

Format y = getmatrix(a, loc);

Input *a* N-dimensional array.

loc Mx1 vector of indices into the array to locate the matrix of interest, where M equals N, N-1 or N-2.

Output y KxL or 1xL matrix or scalar, where L is the size of the fastest moving dimension of the array and K is the size of the second fastest moving dimension.

Remarks Inputting an Nx1 locator vector will return a scalar, an (N-1)x1 locator vector will return a 1xL matrix, and an (N-2)x1 locator vector will return a KxL matrix.

Example a = seqa(1,1,120);
a = areshape(a,2|3|4|5);
loc = { 1,2 };
y = getmatrix(a,loc);

21 22 23 24 25
y = 26 27 28 29 30
31 32 33 34 35

36 37 38 39 40

See also getarray, getmatrix4D

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getmatrix4D

getmatrix4D

Purpose Gets a contiguous matrix from a 4-dimensional array. h **Format** y = getmatrix4D(a,i1,i2);Input d 4-dimensional array. *i1* scalar, index into the slowest moving dimension of the array. *i*2 scalar, index into the second slowest moving dimension of the array. **Output** KxL matrix, where L is the size of the fastest moving dimension of the array and K is the size of the second fastest moving h dimension. Remarks getmatrix4D returns the contiguous matrix that begins at the [i1,i2,1,1] position in array a and ends at the [i1,i2,K,L] position. A call to **getmatrix4D** is faster than using the more general k **getmatrix** function to get a matrix from a 4-dimensional array, especially when i1 and i2 are the counters from nested **for** loops. **Example** a = sega(1,1,120);m a = areshape(a, 2|3|4|5);y = getmatrix4D(a,2,3);0 p 101 102 103 104 105 106 107 108 109 110 v =111 112 113 114 115 116 117 118 119 120 See also getmatrix, getscalar4D, getarray u

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getname

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getname

Purpose Returns a column vector containing the names of the variables in a GAUSS data set. **Format** y = getname(dset);Input string specifying the name of the data set from which the dset function will obtain the variable names. **Output** Nx1 vector containing the names of all of the variables in the specified data set. Remarks The output, y, will have as many rows as there are variables in the data set. **Example** y = getname("olsdat"); format 8,8; print \$y; produces: TIME DIST TEMP FRICT The above example assumes the data set **olsdat** contained the variables TIME, DIST, TEMP, FRICT. Note that the extension is not included in the filename passed to the getname function. See also getnamef, indcv

getnamef

getnamef

a **Purpose** Returns a string array containing the names of the variables in a GAUSS b data set. **Format** y = getnamef(f);d Input scalar, file handle of an open data set. **Output** Nx1 string array containing the names of all of the variables in the specified data set. Remarks The output, y, will have as many rows as there are variables in the data h set. Example open f = olsdat for read; y = getnamef(f);k t = vartypef(f);print y; m produces: time 0 dist temp p frict q The above example assumes the data set **olsdat** contained the variables time, dist, temp, frict. Note the use of **vartypef** to determine the types of these variables. See also getname, indcv, vartypef u V

W

x y z

getNextTradingDay

getNextTradingDay

Purpose Returns the next trading day.

Format n = getNextTradingDay(a)

Input a scalar, date in DT scalar format.

Output n next trading day in DT scalar format

Remarks A trading day is a weekday that is not a holiday as defined by the New

York Stock Exchange from 1888 through 2004. Holidays are defined in

holidays.asc. You may edit that file to modify or add holidays.

Source finutils.src

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getNextWeekDay

getNextWeekDay

Purpose Returns the next day that is not on a weekend.

Format n = getNextWeekDay(a)

Input a scalar, date in DT scalar format.

Output n next week day in DT scalar format

Source finutils.src

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getnr

getnr

Purpose Computes number of rows to read per iteration for a program that reads data from a disk file in a loop.

Format nr = getnr(nsets,ncols);

Input *nsets* scalar, estimate of the maximum number of duplicate copies of

the data matrix read by **readr** to be kept in memory during each

iteration of the loop.

ncols scalar, columns in the data file.

Output *nr* scalar, number of rows **readr** should read per iteration of the

read loop.

Remarks If __row is greater than 0, *nr* will be set to __row.

If an insufficient memory error is encountered, change **__rowfac** to a number less than 1.0 (e.g., 0.75). The number of rows read will be

reduced in size by this factor.

Source gauss.src

Globals __row, __rowfac

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getorders

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x y z

getorders

Purpose Gets the vector of orders corresponding to an array. Format y = getorders(a); Input N-dimensional array. Output Nx1 vector of orders, the sizes of the dimensions of the array. Example a = arrayalloc(7|6|5|4|3,0);orders = getorders(a); 7 6 orders = 3 See also getdims

3-360

getpath

getpath

Purpose Returns an expanded filename including the drive and path. **Format** fname = getpath(pfname); Input string, partial filename with only partial or missing path information. Output fname string, filename with full drive and path. Remarks This function handles relative path references. Example y = getpath("temp.e"); print y; produces: /gauss/temp.e Source getpath.src

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getPreviousTradingDay

getPreviousTradingDay

Purpose Returns the previous trading day.

Format n = getPreviousTradingDay(a)

Input a scalar, date in DT scalar format.

Output n Previous trading day in DT scalar format

Remarks A trading day is a weekday that is not a holiday as defined by the New

York Stock Exchange from 1888 through 2004. Holidays are defined in holidays. asc. You may edit that file to modify or add holidays.

Source finutils.src

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getPreviousWeekDay

getPreviousWeekDay

Purpose Returns the previous day that is not on a weekend.

Format n = getPreviousWeekDay(a)

Input a scalar, date in DT scalar format.

Output n previous week day in DT scalar format

Source finutils.src

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getscalar3D

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getscalar3D

Purpose Gets a scalar from a 3-dimensional array. **Format** y = getscalar3D(a,i1,i2,i3);Input 3-dimensional array. i1scalar, index into the slowest moving dimension of the array. *i*2 scalar, index into the second slowest moving dimension of the array. iЗ scalar, index into the fastest moving dimension of the array. **Output** scalar, the element of the array indicated by the indices. Remarks **getscalar3D** returns the scalar that is located in the [i1,i2,i3] position of array a. A call to **getscalar3D** is faster than using the more general **getmatrix** function to get a scalar from a 3-dimensional array. **Example** a = seqa(1,1,24);a = areshape(a, 2|3|4);y = getscalar3D(a,1,3,2);y = 10See also getmatrix, getscalar4D, getarray

getscalar4D

getscalar4D

Purpose Gets a scalar from a 4-dimensional array.

Format y = getscalar4D(a,i1,i2,i3,i4);

Input

- *a* 4-dimensional array.
- *i1* scalar, index into the slowest moving dimension of the array.
- *i*2 scalar, index into the second slowest moving dimension of the array.
- *i3* scalar, index into the second fastest moving dimension of the array.
- *i4* scalar, index into the fastest moving dimension of the array.

Output

y scalar, the element of the array indicated by the indices.

Remarks

getscalar4D returns the scalar that is located in the [i1,i2,i3,i4] position of array a.

A call to **getscalar4D** is faster than using the more general **getmatrix** function to get a scalar from a 4-dimensional array.

Example

```
a = seqa(1,1,120);
a = areshape(a,2|3|4|5);
y = getscalar4D(a,1,3,2,5);
```

y = 50

See also

getmatrix, getscalar3D, getarray

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getwind

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getwind

Purpose Retrieves the current graphic panel number.

Library pgraph

Format n = getwind;

Output *n* scalar, graphic panel number of current graphic panel.

Remarks The current graphic panel is the graphic panel in which the next graph

will be drawn.

Source pwindow.src

See also endwind, begwind, window, setwind, nextwind

хуг

3-366

gosub

gosub

Purpose Causes a branch to a subroutine.

Format gosub label;

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.

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label:

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.

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return;

Remarks

For multi-line recursive user-defined functions, see "Procedures and Keywords" in the *User's Guide*.

When a **gosub** statement is encountered, the program will branch to the label and begin executing from there. When a **return** statement is encountered, the program will resume executing at the statement following the **gosub** statement. Labels are 1-32 characters long and are followed by a colon. The characters can be A-Z or 0-9 and they must begin with an alphabetic character. Uppercase or lowercase is allowed.

It is possible to pass parameters to subroutines and receive parameters from them when they return. See the second example, following.

The only legal way to enter a subroutine is with a **gosub** statement.

If your subroutines are at the end of your program, you should have an **end** statement before the first one to prevent the program from running into a subroutine without using a **gosub**. This will result in a "return without gosub" error message.

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gosub

The variables used in subroutines are not local to the subroutine and can be accessed from other places in your program. (See "Procedures and Keywords" in the *User's Guide*.)

Example

In the program below, the name **mysub** is a label. When the **gosub** statement is executed, the program will jump to the label **mysub** and continue executing from there. When the **return** statement is executed, the program will resume executing at the statement following the **gosub**.

```
x = rndn(3,3); z = 0;
gosub mysub;
print z;
end;
/* ----- Subroutines Follow ----- */
mysub:
   z = inv(x);
   return;
```

Parameters can be passed to subroutines in the following way (line numbers are added for clarity):

```
    gosub mysub(x,y);
    pop j; /* b will be in j */
    pop k; /* a will be in k */
    t = j*k;
    print t;
    end;
    /* ---- Subroutines Follow ----- */
    mysub:
    pop b; /* y will be in b */
```

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qosub

- 12. pop a; /* x will be in a */
- 13.
- 14. a = inv(b)*b+a;
- 15. b = a'b;
- 16. return(a,b);

In the previous example, when the **gosub** statement is executed, the following sequence of events results:

- 1. **x** and **y** are pushed on the stack and the program branches to the label **mysub** in line 10.
- 11. the second argument that was pushed, y, is pop'ped into b.
- 12. the first argument that was pushed, \mathbf{x} , is **pop**'ped into \mathbf{a} .
- 14. inv(b)*b+a is assigned to a.
- 15. **a/b** is assigned to **b**.
- 16. **a** and **b** are pushed on the stack and the program branches to the statement following the **gosub**, which is line 2.
- 2. the second argument that was pushed, **b**, is **pop**'ped into **j**.
- 3. the first argument that was pushed, **a**, is **pop**'ped into **k**.
- 4. j*k is assigned to t.
- 5. **t** is printed.
- 6. the program is terminated with the **end** statement.

Matrices are pushed on a last-in/first-out stack in the **gosub()** and **return()** statements. They must be popped off in the reverse order. No intervening statements are allowed between the label and the **pop** or the **gosub** and the **pop**. Only one matrix may be popped per **pop** statement.

See also goto, proc, pop, return

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goto

a **Purpose** h **Format** label: Remarks h m **Example** x = seqa(.1,.1,5);0 p q print x; end; fip: print n; u produces: V W

```
Causes a branch to a label.
goto label;
```

Label names can be any legal GAUSS names up to 32 alphanumeric characters, beginning with an alphabetic character or an underscore, not a reserved word.

Labels are always followed immediately by a colon.

Labels do not have to be declared before they are used. GAUSS knows they are labels by the fact that they are followed immediately by a colon.

When GAUSS encounters a goto statement, it jumps to the specified label and continues execution of the program from there.

Parameters can be passed in a goto statement the same way as they can with a gosub.

```
n = \{ 1 2 3 \};
goto fip;
```

1.0000000 2.0000000 3.0000000

x y z

goto

See also gosub, if

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gradMT

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x y z

gradMT

Purpose Computes numerical gradient. **Format** g = gradMT(&fct,parl,datal);Include optim.sdf Input &fct scalar, pointer to procedure returning either Nx1 vector or 1x1 scalar. an instance of structure of type PV containing parameter vector par1 at which gradient is to be evaluated. structure of type DS containing any data needed by fct. data1 **Output** NxK Jacobian or 1xK gradient. Remarks par1 must be created using the pvPack procedures. **Example** #include optim.sdf struct PV p1; p1 = pvCreate; p1 = pvPack(p1, 0.1 | 0.2, "P");struct DS d0; d0 = dsCreate; d0.dataMatrix = seqa(1,1,15);proc fct(struct PV p0, struct DS d0); local p,y; p = pvUnpack(p0,"P"); y = p[1] * exp(-p[2] * d0.dataMatrix);retp(y); endp;

gradMT

g = gradMT(&fct,p1,d0);

Source gradmt.src

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gradMTm

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x y z

gradMTm

Purpose Computes numerical gradient with mask. **Format** g = gradMTm(&fct,parl,datal,mask);Include optim.sdf Input &fct scalar, pointer to procedure returning either Nx1 vector or 1x1 scalar. an instance of structure of type PV containing parameter vector par1 at which gradient is to be evaluated. structure of type DS containing any data needed by fct. data1 Kx1 matrix, elements in g corresponding to elements of mask set mask to zero are not computed, otherwise are computed. **Output** NxK Jacobian or 1xK gradient. g Remarks parl must be created using the **pvPack** procedures. **Example** #include optim.sdf struct PV p1; p1 = pvCreate; p1 = pvPack(p1, 0.1 | 0.2, "P");struct DS d0; d0 = dsCreate; d0.dataMatrix = seqa(1,1,15);proc fct(struct PV p0, struct DS d0); local p,y; p = pvUnpack(p0,"P"); y = p[1] * exp(-p[2] * d0.dataMatrix);retp(y);

${\tt gradMTm}$

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```
endp;
          mask = \{ 0, 1 \};
          g = gradMTm(&fct,p1,d0,mask);
Source
         gradmt.src
```

gradp

gradp

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Purpose

Computes the gradient vector or matrix (Jacobian) of a vector-valued function that has been defined in a procedure. Single-sided (forward difference) gradients are computed.

Format

g = gradp(&f,x0);

Input

& f a pointer to a vector-valued function (f:Kx1 -> Nx1) defined as a procedure. It is acceptable for f(x) to have been defined in terms of global arguments in addition to x, and thus f can return an Nx1 vector:

```
proc f(x);
    retp( exp(x.*b) );
endp;
```

x0 Kx1 vector of points at which to compute gradient.

Output

NXK matrix containing the gradients of f with respect to the variable x at x0.

Remarks

gradp will return a row for every row that is returned by f. For instance, if f returns a scalar result, then **gradp** will return a 1xK row vector. This allows the same function to be used regardless of N, where N is the number of rows in the result returned by f. Thus, for instance, **gradp** can be used to compute the Jacobian matrix of a set of equations.

Example

gradp

It is a 3x3 matrix because we are passing it 3 arguments and **myfunc** returns 3 results when we do that; the off-diagonals are zeros because the cross-derivatives of 3 arguments are 0.

Source gradp.src

See also hessp

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graphprt

graphprt

Purpose Controls automatic printer hardcopy and conversion file output.

Library pgraph

Format graphprt(str);

Input *str* string, control string.

Portability UNIX

Not supported, use WinPrintPQG instead.

Remarks graphprt is used to create hardcopy output automatically without user intervention. The input string *str* can have any of the following items,

separated by spaces. If str is a null string, the interactive mode is entered.

This is the default.

-P print graph.

-PO=c set print orientation.

L landscape.

P portrait.

-C=n convert to another file format.

1 Encapsulated PostScript file.

3 HPGL Plotter file.

5 BMP (Windows only)

8 WMF (Windows only)

-CF=*name* set converted output file name.

-I Minimize (iconize) the graphics window.

-Q Close window after processing.

-W=n display graph, wait *n* seconds, then continue.

If you are not using graphic panels, you can call **graphprt** anytime before the call to the graphics routine. If you are using graphic panels, call **graphprt** just before the **endwind** statement.

The print option default values are set from the viewer application. Any parameters passed through **graphprt** will override the default values. (See "Publication Quality Graphics" in the *User's Guide*.)

Under DOS, this uses a utility called **vwr.exe** by default.

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graphprt

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W

x y z

Example Automatic print using a single graphics call.

```
library pgraph;
graphset;
load x,y;
graphprt("-p"); /* tell "xy" to print */
xy(x,y); /* create graph and print */
```

Automatic print using multiple graphics graphic panels. Note **graphprt** is called once just before the **endwind** call.

```
library pgraph;
graphset;
load x,y;
begwind;
window(1,2,0); /* create two windows */
setwind(1);
xy(x,y); /* first graphics call */
nextwind;
xy(x,y); /* second graphics call */
graphprt("-p");
endwind; /* print page containing all graphs */
```

The next example shows how to build a string to be used with graphprt.

```
library pgraph;
graphset;
load x,y;
cvtnam = "mycvt.eps"; /* name of output file
   /* concatenate options into one string */
```

graphprt

The above string *cmdstr* will produce:

"-
$$c = 1_n$$
 - $cf = mycvt.eps_n$ - q"

Source pgraph.src

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graphset

graphset

Purpose Resets graphics globals to default values.

Library pgraph

Format graphset;

Remarks This procedure is used to reset the defaults between graphs.

graphset may be called between each graphic panel to be displayed.

To change the default values of the global control variables, make the appropriate changes in the file pgraph.dec and to the procedure

graphset.

Source pgraph.src

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hasimag

hasimag

Purpose

Tests whether the imaginary part of a complex matrix is negligible.

Format

y = hasimag(x);

Input

x NxK matrix.

Output

scalar, 1 if the imaginary part of x has any nonzero elements, 0 if it consists entirely of 0's.

The function iscplx tests whether x is a complex matrix or not, but it does not test the contents of the imaginary part of x. hasimag tests the contents of the imaginary part of x to see if it is zero.

hasimag actually tests the imaginary part of *x* against a tolerance to determine if it is negligible. The tolerance used is the imaginary tolerance set with the **sysstate** command, case 21.

Some functions are not defined for complex matrices. iscplx can be used to determine whether a matrix has no imaginary part and so can pass through those functions. hasimag can be used to determine whether a complex matrix has a negligible imaginary part and could thus be converted to a real matrix to pass through those functions.

iscplx is useful as a preliminary check because for large matrices it is much faster than **hasimag**.

Example

```
x = \{ 1 2 3i, \\ 4-i 5 6i, \\ 7 8i 9 \};

y = hasimag(x);

y = 1.0000000
```

See also

iscplx

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header

Purpose Prints a header for a report. **Format** header(prcnm,dataset,ver); Input string, name of procedure that calls header. prcnm dataset string, name of data set. 2x1 numeric vector, the first element is the major version ver number of the program, the second element is the revision number. Normally this argument will be the version/revision global (?? ver) associated with the module within which header is called. This argument will be ignored if set to 0. **Global Input** header string, containing the letters: title is to be printed t 1 lines are to bracket the title a date and time is to be printed d v version number of program is printed file name being analyzed is printed f title string, title for header. Source gauss.src Globals header, title

hess

hess

a h

d

Purpose

Computes the Hessenberg form of a square matrix.

Format

 $\{h,z\} = hess(x);$

Input

x KxK matrix.

Output

h KxK matrix, Hessenberg form.

z KxK matrix, transformation matrix.

Remarks

hess computes the Hessenberg form of a square matrix. The Hessenberg form is an intermediate step in computing eigenvalues. It also is useful for solving certain matrix equations that occur in control theory (see Van Loan, Charles F. "Using the Hessenberg Decomposition in Control Theory," *Algorithms and Theory in FIltering and Control*. Sorenson, D.C., and R.J. Wets, eds., Mathematical Programming Study No. 18, No. Holland, Amsterdam, 1982, 102-11).

z is an orthogonal matrix that transforms x into h and vice versa. Thus:

$$h = z'x z$$

and since z is orthogonal,

$$x = z h z'$$

x is reduced to upper Hessenberg form using orthogonal similarity transformations. This preserves the Frobenious norm of the matrix and the condition numbers of the eigenvalues.

hess uses the ORTRAN and ORTHES functions from EISPACK.

Example

let
$$x[3,3] = 1 2 3$$

4 5 6
7 8 9;

$$\{h,z\} = hess(x);$$

$$h = -8.06225775$$
 14.04615385 2.83076923 0.00000000 0.83076923 -0.04615385

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hess

See also schur

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x y z

hessMT

Purpose Computes numerical Hessian. **Format** h = hessMT(&fct,parl,datal);Include optim.sdf Input &fct scalar, pointer to procedure returning either Nx1 vector or 1x1 scalar. an instance of structure of type PV containing parameter vector par1 at which Hessian is to be evaluated. structure of type DS containing any data needed by fct. data l **Output** h KxK matrix, Hessian. Remarks par1 must be created using the **pvPack** procedures. **Example** #include optim.sdf struct PV p1; p1 = pvCreate; p1 = pvPack(p1, 0.1 | 0.2, "P");struct DS d0; d0 = dsCreate; d0.dataMatrix = seqa(1,1,15);proc fct(struct PV p0, struct DS d0); local p,y; p = pvUnpack(p0,"P"); y = p[1] * exp(-p[2] * d0.dataMatrix);retp(y); endp;

hessMT

h = hessMT(&fct,p1,d0);

Source hessmt.src

a

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С

d

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h

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q

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V

W

hessMTg

b

d

h

m

n

0

p

q

u

V

W

x y z

hessMTg

```
Purpose
             Computes numerical Hessian using gradient procedure.
  Format
             h = hessMTg(\&gfct, parl, datal);
 Include
             optim.sdf
    Input
             &gfct
                    scalar, pointer to procedure computing either 1xK gradient or
                    NxK Jacobian.
                    an instance of structure of type PV containing parameter vector
             par1
                    at which Hessian is to be evaluated.
                    structure of type DS containing any data needed by gfct.
             data1
  Output
                    KxK matrix, Hessian.
Remarks
             par1 must be created using the pvPack procedures.
Example
             #include optim.sdf
             struct PV p1;
             p1 = pvCreate;
             p1 = pvPack(p1, 0.1 | 0.2, "P");
             struct DS d0;
             d0 = dsCreate;
             d0.dataMatrix = seqa(1,1,15);
             proc gfct(&fct, struct PV p0, struct DS d0);
                 local p,y,g1,g2;
                 p = pvUnpack(p0,"P");
                g1 = exp(-p[2] * d0.dataMatrix);
                y = p[1] * exp(-p[2] * d0.dataMatrix);
                 q2 = -p[1] * d0.dataMatrix .* q1;
```

hessMTg

retp(g1~g2);
endp;

h = hessMTg(&gfct,p1,d0);

Source hessmt.src

a

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V

hessMTqw

a

b

d

h

m

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0

p

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u

V

W

x y z

hessMTgw

Purpose Computes numerical Hessian using gradient procedure with weights. **Format** h = hessMTgw(&gfct,parl,datal,wgts);Include optim.sdf Input &gfct scalar, pointer to procedure computing either NxK Jacobian. an instance of structure of type PV containing parameter vector par1 at which Hessian is to be evaluated. structure of type DS containing any data needed by gfct. data1 Nx1 vector. wgts **Output** KxK matrix, Hessian. Remarks par1 must be created using the pvPack procedures. **Example** #include optim.sdf struct PV p1; p1 = pvCreate; p1 = pvPack(p1, 0.1 | 0.2, "P");struct DS d0; d0 = dsCreate; d0.dataMatrix = seqa(1,1,15); $wqts = zeros(5,1) \mid ones(10,1);$ proc gfct(&fct, struct PV p0, struct DS d0); local p,y,q1,q2; p = pvUnpack(p0,"P"); g1 = exp(-p[2] * d0.dataMatrix);

3-390

hessMTgw

```
y = p[1] * exp( -p[2] * d0.dataMatrix );
g2 = -p[1] * d0.dataMatrix .* g1;
retp(g1~g2);
endp;

h = hessMTgw(&gfct,p1,d0,wgts);

Source hessmt.src
```

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V

hessMTm

a

b

d

h

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0

p

q

u

V

W

x y z

hessMTm

Purpose Computes numerical Hessian with mask. **Format** h = hessMTm(&fct,parl,datal,mask);Include optim.sdf Input &fct scalar, pointer to procedure returning either Nx1 vector or scalar. par1 an instance of structure of type PV containing parameter vector at which Hessian is to be evaluated. data1 structure of type DS containing any data needed by fct. KxK matrix, elements in h corresponding to elements of mask set mask to zero are not computed, otherwise are computed. **Output** h KxK matrix, Hessian. Remarks parl must be created using the **pvPack** procedures. Only lower left part of mask looked at. **Example** #include optim.sdf struct PV p1; p1 = pvCreate; p1 = pvPack(p1, 0.1 | 0.2, "P");struct DS d0; d0 = dsCreate; d0.dataMatrix = seqa(1,1,15); $mask = \{ 1 1 \}$ 1 0 }; proc fct(struct PV p0, struct DS d0); local p,y;

hessMTm

```
p = pvUnpack(p0,"P");
            y = p[1] * exp(-p[2] * d0.dataMatrix);
            retp(y);
            endp;
         h = hessMTm(&fct,p1,d0,mask);
Source
         hessmt.src
```

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hessMTmw

hessMTmw

a **Purpose** Computes numerical Hessian with mask and weights. b **Format** h = hessMTmw(&fct,parl,datal,mask,wgts);Include optim.sdf d Input &fct scalar, pointer to procedure returning Nx1 vector. par1 an instance of structure of type PV containing parameter vector at which Hessian is to be evaluated. structure of type DS containing any data needed by fct. data1 KxK matrix, elements in h corresponding to elements of mask set mask h to zero are not computed, otherwise are computed. Nx1 vector, weights. wgts **Output** h KxK matrix, Hessian. Remarks fct must evaluate to an Nx1 vector conformable to the weight vector. par1 must be created using the pvPack procedures. m **Example** #include optim.sdf n struct PV p1; 0 p1 = pvCreate; p p1 = pvPack(p1,0.1|0.2,"P"); q struct DS d0; d0 = dsCreate; d0.dataMatrix = seqa(1,1,15); $wgts = zeros(5,1) \mid ones(10,1);$ u V $mask = { 1 1,}$ W 1 0 };

x y z

hessMTmw

a

b

С

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m

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q

t

V

```
proc fct(&fct, struct PV p0, struct DS d0, wgts);
            local p,y;
            p = pvUnpack(p0,"P");
            y = p[1] * exp(-p[2] * d0.dataMatrix);
            retp(y);
         endp;
         h = hessMTmw(&fct,p1,d0,mask,wgt);
Source
         hessmt.src
```

hessMTw

a

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h

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V

W

x y z

hessMTw

Purpose Computes numerical Hessian with weights. **Format** h = hessMTw(&fct, parl, datal, wgts);Include optim.sdf Input &fct scalar, pointer to procedure returning Nx1 vector. par1 an instance of structure of type PV containing parameter vector at which Hessian is to be evaluated. structure of type DS containing any data needed by fct. data1 Nx1 vector, weights. wgts **Output** h KxK matrix, Hessian. Remarks fct must evaluate to an Nx1 vector conformable to the weight vector. par1 must be created using the **pvPack** procedures. **Example** #include optim.sdf struct PV p1; p1 = pvCreate; p1 = pvPack(p1, 0.1 | 0.2, "P");struct DS d0; d0 = dsCreate;d0.dataMatrix = seqa(1,1,15); $wgt = zeros(5,1) \mid ones(10,1);$ proc fct(&fct, struct PV p0, struct DS d0, wgt); local p,y; p = pvUnpack(p0,"P"); y = p[1] * exp(-p[2] * d0.dataMatrix);

hessMTw

a

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С

d

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h

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V

```
retp(y);
         endp;
         h = hessMTw(&fct,p1,d0,wgt);
Source
         hessmt.src
```

hessp

hessp

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хуг

Purpose Computes the matrix of second partial derivatives (Hessian matrix) of a function defined as a procedure.

Format h = hessp(&f,x0);

Input

pointer to a single-valued function f(x), defined as a procedure, taking a single Kx1 vector argument (f:Kx1 -> 1x1); f(x) may be defined in terms of global arguments in addition to x.

x0 Kx1 vector specifying the point at which the Hessian of f(x) is to be computed.

Output

KxK matrix of second derivatives of f with respect to x at x0; this matrix will be symmetric.

Remarks

This procedure requires $K^*(K+1)/2$ function evaluations. Thus if K is large, it may take a long time to compute the Hessian matrix.

No more than 3-4 digit accuracy should be expected from this function, though it is possible for greater accuracy to be achieved with some functions.

It is important that the function be properly scaled, in order to obtain greatest possible accuracy. Specifically, scale it so that the first derivatives are approximately the same size. If these derivatives differ by more than a factor of 100 or so, the results can be meaningless.

Example $x = \{ 1, 2, 3 \};$

```
proc g(b);
    retp( exp(x'b) );
endp;

b0 = { 3, 2, 1 };
h = hessp(&g,b0);
```

The resulting matrix of second partial derivatives of **g(b)** evaluated at **b=b0** is:

hessp

22027.12898372 44054.87238165 66083.36762901 44054.87238165 88111.11102645 132168.66742899 66083.36762901 132168.66742899 198256.04087836

Source hessp.src

See also gradp

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hist

hist

a h

h

m

0

p

q

u V

W

x y z

Purpose

Computes and graphs a frequency histogram for a vector. The actual frequencies are plotted for each category.

Library pgraph

 $\{b,m,freq\} = hist(x,v);$ **Format**

Input \boldsymbol{x}

Mx1 vector of data.

Nx1 vector, the breakpoints to be used to compute the frequencies,

or

scalar, the number of categories.

Output

b Px1 vector, the breakpoints used for each category.

Px1 vector, the midpoints of each category. m

Px1 vector of computed frequency counts. freq

Remarks

If a vector of breakpoints is specified, a final breakpoint equal to the maximum value of x will be added if the maximum breakpoint value is smaller.

If a number of categories is specified, the data will be divided into vevenly spaced categories.

Each time an element falls into one of the categories specified in b, the corresponding element of *freq* will be incremented by one. The categories are interpreted as follows:

freq[1]b[1]freq[2]b[1]b[2]x <=

freq[3]b[2]<= b[3]

freq[P]b[P-1]b[P]<=

hist

Example library pgraph;
 x = rndn(5000,1);
 { b,m,f } = hist(x,20);

Source phist.src

See also histp, histf, bar

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histf

a

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k

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V

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histf

Purpose Graphs a histogram given a vector of frequency counts. Library

Format histf(f,c);

pgraph

Input fNx1 vector, frequencies to be graphed.

> Nx1 vector, numeric labels for categories. If this is a scalar 0, a csequence from 1 to rows(f) will be created.

Remarks The axes are not automatically labeled. Use xlabel for the category axis and **ylabel** for the frequency axis.

Source phist.src

See also hist, bar, xlabel, ylabel

histp

Purpose

Computes and graphs a percent frequency histogram of a vector. The percentages in each category are plotted.

Library

pgraph

Format

 $\{b,m,freq\} = histp(x,v);$

Input

x Mx1 vector of data.

v Nx1 vector, the breakpoints to be used to compute the frequencies,

or

scalar, the number of categories.

Output

b Px1 vector, the breakpoints used for each category.

m Px1 vector, the midpoints of each category.

freq Px1 vector of computed frequency counts. This is the vector of counts, not percentages.

Remarks

If a vector of breakpoints is specified, a final breakpoint equal to the maximum value of *x* will be added if the maximum breakpoint value is smaller.

If a number of categories is specified, the data will be divided into v evenly spaced categories.

Each time an element falls into one of the categories specified in b, the corresponding element of freq will be incremented by one. The categories are interpreted as follows:

$$freq[1] = x \leftarrow b[1]$$

$$freq[2] = b[1] < x <= b[2]$$

$$freq[3] = b[2] < x <= b[3]$$

Source

phist.src

See also

hist, histf, bar

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hsec

hsec

h

d

Purpose Returns the number of hundredths of a second since midnight.

Format

y = hsec;

Remarks

The number of hundredths of a second since midnight can also be accessed as the [4,1] element of the vector returned by the **date** function.

Example

```
x = rndu(100, 100);
```

ts = hsec;

y = x*x;

et = hsec-ts;

In this example, **hsec** is used to time a 100x100 multiplication in GAUSS. A 100x100 matrix, **x**, is created, and the current time, in hundredths of a second since midnight, is stored in the variable **ts**. Then the multiplication is carried out. Finally, **ts** is subtracted from **hsec** to give the time difference which is assigned to **et**.

See also

date, time, timestr, ethsec, etstr

V

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V

W

x y z

if

Purpose Controls program flow with conditional branching.

Format

if scalar_expression;
list of statements;

elseif scalar_expression;

list of statements;

elseif scalar_expression;

list of statements;

else;

list of statements;

endif;

Remarks

scalar_expression is any expression that returns a scalar. It is *TRUE* if it is not zero, and *FALSE* if it is zero.

A list of statements is any set of GAUSS statements.

GAUSS will test the expression after the **if** statement. If it is *TRUE* (nonzero), then the first list of statements is executed. If it is *FALSE* (zero), then GAUSS will move to the expression after the first **elseif** statement if there is one and test it. It will keep testing expressions and will execute the first list of statements that corresponds to a *TRUE* expression. If no expression is *TRUE*, then the list of statements following the **else** statement is executed. After the appropriate list of statements is executed, the program will go to the statement following the **endif** and continue on.

if statements can be nested.

One **endif** is required per **if** statement. If an **else** statement is used, there may be only one per **if** statement. There may be as many **elseif**'s as are required. There need not be any **elseif**'s or any **else** statement within an **if** statement.

Note the semicolon after the **else** statement.

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XX7

if

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Example if x < 0;
 y = -1;
 elseif x > 0;
 y = 1;
 else;
 y = 0;

endif;

See also do

imag

imag

Purpose Returns the imaginary part of a matrix.

Format zi = imag(x);

Input x NxK matrix.

Output zi NxK matrix, the imaginary part of x.

Remarks If x is real, zi will be an NxK matrix of zeros.

Example $x = \{ 4i \ 9 \ 3, \}$

2 5-6i 7i };

y = imag(x);

 $y = \begin{cases} 4.0000000 & 0.0000000 & 0.0000000 \\ 0.0000000 & -6.0000000 & 7.0000000 \end{cases}$

See also complex, real

a

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#include

#include

Purpose Inserts code from another file into a GAUSS program.

Format #include filename; #include "filename";

Remarks filename can be any legitimate file name.

> This command makes it possible to write a section of general-purpose code, and insert it into other programs.

The code from the **#include**'d file is inserted literally as if it were merged into that place in the program with a text editor.

If a path is specified for the file, then no additional searching will be attempted if the file is not found.

If a path is not specified, the current directory will be searched first, then each directory listed in **src path**. **src path** is defined in qauss.cfq.

No additional search will be #include /gauss/myprog.prc; made if the file is not found.

#include myprog.prc; The directories listed in

src path will be searched for myprog.prc if the file is not found in the current

directory.

Compile time errors will return the line number and the name of the file in which they occur. For execution time errors, if a program is compiled with **#lineson**, the line number and name of the file where the error occurred will be printed. For files that have been **#include**'d this reflects the actual line number within the **#include**'d file. See **#lineson** for a more complete discussion of the use of and the validity of line numbers when debugging.

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#include

Example #include "/gauss/inc/cond.inc";

The command will cause the code in the file <code>cond.inc</code> to be merged into the current program at the point at which this statement appears.

See also run, #lineson

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XX7

indcv

indcv

a h h m 0 p q

u V W

Remarks

Purpose

Checks one character vector against another and returns the indices of the elements of the first vector in the second vector.

Format z = indcv(what, where);

Input Nx1 character vector which contains the elements to be found in what vector where.

> where Mx1 character vector to be searched for matches to the elements of what.

Output Nx1 vector of integers containing the indices of the corresponding element of what in where.

> If no matches are found for any of the elements in what, then the corresponding elements in the returned vector are set to the GAUSS missing value code.

Both arguments will be forced to uppercase before the comparison.

If there are duplicate elements in where, the index of the first match will be returned.

Example let what = AGE PAY SEX;

let where = AGE SEX JOB "date" PAY;

z = indcv(what, where);

AGE what =PAY

SEX

AGE SEX

date

where =

PAY

IOB

indcv

$$Z = \begin{array}{c} 1 \\ 5 \\ 2 \end{array}$$

a

b

С

d e

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h i

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indexcat

indexcat

Purpose

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t

u

V W

хух

h

Returns the indices of the elements of a vector which fall into a specified category.

Format

y = indexcat(x, y);

Input

Nx1 vector.

scalar or 2x1 vector.

If scalar, the function returns the indices of all elements of x equal to v.

If 2x1, then the function returns the indices of all elements of x that fall into the range:

$$v[1] < x \le v[2].$$

If v is scalar, it can contain a single missing to specify the missing value as the category.

Output

Lx1 vector, containing the indices of the elements of x which fall into the category defined by v. It will contain error code 13 if there are no elements in this category.

Remarks

Use a loop to pull out indices of multiple categories.

Example

let $x = 1.0 \ 4.0 \ 3.3 \ 4.2 \ 6.0 \ 5.7 \ 8.1 \ 5.5$;

let v = 46;

y = indexcat(x,v);

1.0

4.0

3.3

4.2 x =

6.0

5.7

8.1

5.5

indexcat

$$v = \frac{4}{6}$$

$$y = \begin{cases} 4\\5\\6\\8 \end{cases}$$

a

b

С

d e

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g h

i

J k

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XX7

indices

indices

Purpose

Processes a set of variable names or indices and returns a vector of

variable names and a vector of indices.

Format { name,indx } = indices(dataset,vars);

Input dataset string, the name of the data set.

vars Nx1 vector, a character vector of names or a numeric vector of

column indices.

If scalar 0, all variables in the data set will be selected.

Output *name* Nx1 character vector, the names associated with *vars*.

indx Nx1 numeric vector, the column indices associated with vars.

Remarks If an error occurs, **indices** will either return a scalar error code or

terminate the program with an error message, depending on the trap state. If the low order bit of the trap flag is 0, indices will terminate with an error message. If the low order bit of the trap flag is 1, indices will return an error code. The value of the trap flag can be tested with trapchk; the return from indices can be tested with scalerr. You only need to check one argument; they will both be the same. The following error codes are possible:

1 Can't open dataset.

2 Index of variable out of range, or undefined data set variables.

Source indices.src

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indices2

indices2

Purpose Processes two sets of variable names or indices from a single file. The

first is a single variable and the second is a set of variables. The first must not occur in the second set and all must be in the file.

Format { name1,indx1,name2,indx2 } =
indices2(dataset,var1,var2);

Input dataset string, the name of the data set.

var1 string or scalar, variable name or index.

This can be either the name of the variable, or the column

index of the variable.

If null or 0, the last variable in the data set will be used.

var2 Nx1 vector, a character vector of names or a numeric vector of

column indices.

If scalar 0, all variables in the data set except the one

associated with var1 will be selected.

Output *name1* scalar character matrix containing the name of the variable

associated with *var1*.

indx1 scalar, the column index of var1.

name2 Nx1 character vector, the names associated with var2.

indx2 Nx1 numeric vector, the column indices of var2.

Remarks

If an error occurs, indices2 will either return a scalar error code or terminate the program with an error message, depending on the trap state. If the low order bit of the trap flag is 0, indices2 will terminate with an error message. If the low order bit of the trap flag is 1, indices2 will return an error code. The value of the trap flag can be tested with trapchk; the return from indices2 can be tested with scalerr. You only need to check one argument; they will all be the same. The following error codes are possible:

- **1** Can't open dataset.
- 2 Index of variable out of range, or undefined data set variables.
- 3 First variable must be a single name or index.
- First variable contained in second set.

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indices2

Source indices2.src

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indnv

indnv

Purpose Checks one numeric vector against another and returns the indices of the elements of the first vector in the second vector.

Format z = indnv(what, where);

Input what Nx1 numeric vector which contains the values to be found in vector where.

where Mx1 numeric vector to be searched for matches to the values in what.

Output z Nx1 vector of integers, the indices of the corresponding elements of what in where.

Remarks If no matches are found for any of the elements in *what*, then those elements in the returned vector are set to the GAUSS missing value code.

If there are duplicate elements in *where*, the index of the first match will be returned.

Example let what = 8 7 3;

let where = 2 7 8 4 3;

z = indnv(what, where);

$$what = \begin{cases} 8\\ 7\\ 3 \end{cases}$$

b

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е

f g

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V

$\verb"indnv"$

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z = 2

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indsav

indsav

Purpose Checks one string array against another and returns the indices of the first string array in the second string array.

Format indx = indsav(what, where);

Input what Nx1 string array which contains the values to be found in

vector where.

where Mx1 string array to be searched for the corresponding

elements of what in where.

Output *indx* Nx1 vector of indices, the values of *what* in *where*.

Remarks If no matches are found, those elements in the returned vector are set to

the GAUSS missing value code.

If there are duplicate elements in *where*, the index of the first match will

be returned.

Source indsav.src

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intgrat2

Purpose

Integrates the following double integral, using user-defined functions f, g_1 and g_2 , and scalars a and b:

$$\int_{a}^{b} \int_{g_{2}(x)}^{g_{1}(x)} f(x, y) dy dx$$

Format

y = intgrat2(&f,xl,gl);

Input

scalar, pointer to the procedure containing the function to be integrated.

xl 2x1 or 2xN matrix, the limits of x. These must be scalar limits.

gl 2x1 or 2xN matrix of function pointers, the limits of y. For xl and gl, the first row is the upper limit and the second row is the lower limit. N integrations are computed.

Global Input

_intord scalar, the order of the integration. The larger _intord,

the more precise the final result will be. **_intord** may

be set to 2, 3, 4, 6, 8, 12, 16, 20, 24, 32, 40.

Default = 12.

intrec

scalar. This variable is used to keep track of the level of

recursion of intgrat2 and may start out with a different value if your program terminated inside of the integration function on a previous run. Always set

_intrec explicitly to 0 before any call to **intgrat2**.

Output

NX1 vector of the estimated integral(s) of f(x,y) evaluated between the limits given by xl and gl.

Remarks

The user-defined functions specified by f and gl must either

Return a scalar constant, OR

Return a vector of function values. **intgrat2** will pass to user-defined functions a vector or matrix for x and y and expect a vector or matrix to be returned. Use .* and ./ instead of * and /.

W

u

V

a

b

С

d

f

g

h

i

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1

m

n

0

p

q

r

t

u

V

W

x y z

```
Example
             proc f(x,y);
                 retp(cos(x) + 1).*(sin(y) + 1));
              endp;
             proc g1(x);
                 retp(sqrt(1-x^2));
              endp;
             proc g2(x);
                 retp(0);
              endp;
             x1 = 1 | -1;
             g0 = &g1 | &g2;
              _{intord} = 40;
             _{intrec} = 0;
             y = intgrat2(&f,xl,g0);
              This will integrate the function f(x,y) = (cos(x)+1)(sin(y)+1) over the
              upper half of the unit circle. Note the use of the . * operator instead of just
             * in the definition of f(x,y). This allows f to return a vector or matrix of
             function values.
  Source
             intgrat.src
 Globals
             _intord, _intq12, _intq16, _intq2, _intq20,
             _intq24, _intq3, _intq32, _intq4, _intq40,
             _intq6, _intq8, _intrec
See also
             intgrat3, intquad1, intquad2, intquad3, intsimp
```

intgrat3

Purpose

h

h

0

p

Integrates the following triple integral, using user-defined functions and scalars for bounds:

$$\int_{a}^{b} \int_{g_{2}(x)}^{g_{1}(x)} \int_{h_{2}(x,y)}^{h_{1}(x,y)} f(x, y, z) dz dy dx$$

Format

y = intgrat3(&f,xl,gl,hl);

Input

&f scalar, pointer to the procedure containing the function to be integrated. F is a function of (x,y,z).

xl 2x1 or 2xN matrix, the limits of x. These must be scalar limits.

gl 2x1 or 2xN matrix of function pointers. These procedures

are functions of x.

hl 2x1 or 2xN matrix of function pointers. These procedures are functions of *x* and *y*.

For xl, gl, and hl, the first row is the upper limit and the second row is the lower limit. N integrations are computed.

Global Input

_intord scalar, the order of the integration. The larger **_intord**,

the more precise the final result will be. _intord may

be set to 2, 3, 4, 6, 8, 12, 16, 20, 24, 32, 40.

Default = 12.

_intrec scalar. This variable is used to keep track of the level of

recursion of intgrat3 and may start out with a different value if your program terminated inside of the integration function on a previous run. Always set

_intrec explicitly to 0 before any call to **intgrat3**.

Output

Nx1 vector of the estimated integral(s) of f(x,y,z) evaluated between the limits given by xl, gl, and hl.

u v

W

Remarks User-defined functions *f*, and those used in *gl* and *hl*, must either:

Return a scalar constant, OR

Return a vector of function values. **intgrat3** will pass to user-defined functions a vector or matrix for x and y and expect a vector or matrix to be returned. Use **.*** and **.**/ operators instead of just ***** or /.

Example

```
proc f(x,y,z);
   retp(2);
endp;
proc g1(x);
   retp(sqrt(25-x^2));
endp;
proc g2(x);
   retp(-g1(x));
endp;
proc h1(x,y);
   retp(sqrt(25 - x^2 - y^2));
endp;
proc h2(x,y);
   retp(-h1(x,y));
endp;
```

a

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c d

e f

g

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r

S

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m

n

0

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q

r

t

u

V

W

x y z

xl = 5|-5;
g0 = &g1|&g2;
h0 = &h1|&h2;
_intrec = 0;
_intord = 40;
y = intgrat3(&f,xl,gl,hl);

This will integrate the function f(x,y,z) = 2 over the sphere of radius 5. The result will be approximately twice the volume of a sphere of radius 5.

Source intgrat.src

See also intgrat2, intquad1, intquad2, intquad3, intsimp

intquad1

Purpose Integrates a specified function using Gauss-Legendre quadrature. A suite of upper and lower bounds may be calculated in one procedure call. **Format** y = intquad1(&f,xl);Input &f scalar, pointer to the procedure containing the function to be integrated. This must be a function of x. xl2xN matrix, the limits of x. The first row is the upper limit and the second row is the lower limit. N integrations are computed. intord scalar, the order of the integration. The larger intord, the more precise the final result will be. _intord may be set to 2, 3, 4, 6, 8, 12, 16, 20, 24, 32, 40. Default = 12. **Global Input** scalar, the order of the integration. The larger intord, intord the more precise the final result will be. **intord** may be set to 2, 3, 4, 6, 8, 12, 16, 20, 24, 32, 40. Default = 12.

Output

Nx1 vector of the estimated integral(s) of f(x) evaluated between the limits given by xl.

Remarks

The user-defined function f must return a vector of function values. **intquad1** will pass to the user-defined function a vector or matrix for x and expect a vector or matrix to be returned. Use the **.*** and **./** instead of ***** and **/**.

Example

```
retp(x.*sin(x));
endp;

xl = 1|0;
y = intquad1(&f,xl);
```

proc f(x);

a

b

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С

e f

> g h

i

1

k

n

m

0

q

r

t

u

V

X y Z

405

This will integrate the function f(x) = xsin(x) between 0 and 1. Note the use of the \cdot * instead of *.

Source integral.src

Globals _intord, _intq12, _intq16, _intq2, _intq20,

_intq24, _intq3, _intq32, _intq4, _intq40,

_intq6, _intq8

See also intsimp, intquad2, intquad3, intgrat2, intgrat3

a

b

c d

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1

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1

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0

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q

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t

u

V W

intquad2

Purpose

Integrates a specified function using Gauss-Legendre quadrature. A suite of upper and lower bounds may be calculated in one procedure call.

Format

```
y = intquad2(&f,xl,yl);
```

Input

scalar, pointer to the procedure containing the function to be integrated.

xl 2x1 or 2xN matrix, the limits of x.

yl 2x1 or 2xN matrix, the limits of y.

For xl and yl, the first row is the upper limit and the second row is the lower limit. N integrations are computed.

Global Input

intord

global scalar, the order of the integration. The larger _intord, the more precise the final result will be. _intord may be set to 2, 3, 4, 6, 8, 12, 16, 20, 24, 32, 40.

Default = 12.

intrec

global scalar. This variable is used to keep track of the level of recursion of **intquad2** and may start out with a different value if your program terminated inside of the integration function on a previous run. Always set **_intrec** explicitly to 0 before any calls to **intquad2**.

Output

y Nx1 vector of the estimated integral(s) of f(x,y) evaluated between the limits given by xl and yl.

Remarks

The user-defined function f must return a vector of function values. **intquad2** will pass to user-defined functions a vector or matrix for x and y and expect a vector or matrix to be returned. Use •* and •/ instead of * and /.

intquad2 will expand scalars to the appropriate size. This means that functions can be defined to return a scalar constant. If users write their functions incorrectly (using * instead of .*, for example), intquad2 may not compute the expected integral, but the integral of a constant function.

To integrate over a region which is bounded by functions, rather than just scalars, use intgrat2 or intgrat3.

a

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c d

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u v

XX7

b

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h

k

m

n

0

p

q

r

t

u

V

W

```
Example
            proc f(x,y);
                retp(x.*sin(x+y));
             endp;
             x1 = 1 | 0;
             y1 = 1 | 0;
             _{intrec} = 0;
             y = intquad2(&f,xl,yl);
             This will integrate the function \mathbf{x}.*sin(\mathbf{x}+\mathbf{y}) between \mathbf{x}=0 and 1, and
             between y = 0 and 1.
  Source
             integral.src
 Globals
             _intord, _intq12, _intq16, _intq2, _intq20,
             _intq24, _intq3, _intq32, _intq4, _intq40,
             _intq6, _intq8, _intrec
See also
             intquad1, intquad3, intsimp, intgrat2, intgrat3
```

intquad3

Purpose	Integrates	a speci	ified fu	unction	using	Gauss-	-Legend
					. 0		

dre quadrature. A suite of upper and lower bounds may be calculated in one procedure call.

Format y = intquad3(&f,xl,yl,zl);

Input scalar, pointer to the procedure containing the function to be &f integrated. f is a function of (x,y,z).

> xl2x1 or 2xN matrix, the limits of x.

> γl 2x1 or 2xN matrix, the limits of y.

> zl2x1 or 2xN matrix, the limits of z.

> > For xl, yl, and zl, the first row is the upper limit and the second row is the lower limit. N integrations are computed.

Global Input

global scalar, the order of the integration. The larger intord **_intord**, the more precise the final result will be.

_intord may be set to 2, 3, 4, 6, 8, 12, 16, 20, 24, 32,

40.

Default = 12.

intrec

global scalar. This variable is used to keep track of the level of recursion of **intguad3** and may start out with a different value if your program terminated inside of the integration function on a previous run. Always set **_intrec** explicitly to 0 before any calls to

intquad3.

Output

Nx1 vector of the estimated integral(s) of f(x,y,z) evaluated between the limits given by xl, yl, and zl.

Remarks

The user-defined function f must return a vector of function values. intquad3 will pass to the user-defined function a vector or matrix for x, y, and z and expect a vector or matrix to be returned. Use .* and ./ instead of * and /.

intguad3 will expand scalars to the appropriate size. This means that functions can be defined to return a scalar constant. If users write their functions incorrectly (using * instead of .*, for example), intquad3 may not compute the expected integral, but the integral of a constant function.

a

b

С d

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k

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n 0

q

t

u

V

To integrate over a region which is bounded by functions, rather than just scalars, use intgrat2 or intgrat3.

Example

```
proc f(x,y,z);
    retp(x.*y.*z);
endp;

xl = 1|0;
yl = 1|0;
zl = { 1 2 3, 0 0 0 };
```

_intrec = 0;
y = intquad3(&f,xl,yl,zl);

This will integrate the function f(x) = x*y*z over 3 sets of limits, since z1 is defined to be a 2x3 matrix.

Source

integral.src

Globals

```
_intord, _intq12, _intq16, _intq2, _intq20,
_intq24, _intq3, _intq32, _intq4, _intq40,
_intq6, _intq8, _intrec
```

See also

intquad1, intquad2, intsimp, intgrat2, intgrat3

a

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m n

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intrleav

intrleav

Purpose Interleaves the rows of two files that have been sorted on a common variable, to give a single file sorted on that variable.

Format intrleav(infile1, infile2, outfile, keyvar, keytyp);

Input *infile1* string, name of input file 1.

infile2 string, name of input file 2.outfile string, name of output file.

keyvar string, name of key variable, this is the column the files are

sorted on.

keytyp scalar, data type of key variable.

1 numeric key, ascending order

2 character key, ascending order

-1 numeric key, descending order

-2 character key, descending order

Remarks The two files MUST have exactly the same variables, i.e., the same

number of columns AND the same variable names. They must both already be sorted on the key column. This procedure will combine them

into one large file, sorted by the key variable.

If the inputs are null or 0, the procedure will ask for them.

Example intrleav("freq.dat", "freqata.dat",

"intfile", "AGE",1);

Source sortd.src

a

b

С

d

f

g h

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k

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m

n

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q

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S

t

u

V

intrsect

intrsect

a **Purpose** Returns the intersection of two vectors, with duplicates removed. b **Format** y = intrsect(v1, v2, flag); Input v1Nx1 vector. d v2Mx1 vector. e flag scalar; if 1, v1, and v2 are numeric, if 0, character. **Output** Lx1 vector containing all unique values that are in both v1 and v2, sorted in ascending order. h Place smaller vector first for fastest operation. Remarks If there are a lot of duplicates within a vector, it is faster to remove them with unique before calling intrsect. Source intrsect.src k **Example** let v1 = mary jane linda dawn; let v2 = mary sally lisa ruth linda; m y = intrsect(v1, v2, 0);n 0 **LINDA** y =MARY p q

x y z

u

V

W

intsimp

intsimp

Purpose Integrates a specified function using Simpson's method with end correction. A single integral is computed in one function call. **Format** y = intsimp(&f,xl,tol);Input pointer to the procedure containing the function to be integrated. &f xl2x1 vector, the limits of x. The first element is the upper limit and the second element is the lower limit. tolThe tolerance to be used in testing for convergence. **Output** The estimated integral of f(x) between xl[1] and xl[2]. **Example** proc f(x); retp(sin(x)); endp; $let xl = \{ 1,$ 0 }; y = intsimp(&f,xl,1E-8);v = 0.45969769This will integrate the function between 0 and 1. Source intsimp.src See also intquad1, intquad2, intquad3, intgrat2, intgrat3

a

b

c d

e f

g

h i

j k

1

m

n o

n

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V

inv, invpd

inv, invpd

a

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S

u

V

W

хух

Purpose inv returns the inverse of an invertible matrix.

invpd returns the inverse of a symmetric, positive definite matrix.

Format

y = inv(x);y = invpd(x);

Input

x NxN matrix.

Output

NxN matrix containing the inverse of x.

Remarks

x can be any legitimate matrix expression that returns a matrix that is legal for the function.

For **inv**, *x* must be square and invertible.

For **invpd**, *x* must be symmetric and positive definite.

If the input matrix is not invertible by these functions, they will either terminate the program with an error message or return an error code which can be tested for with the **scalerr** function. This depends on the **trap** state as follows:

trap 1, return error code

inv invpd 50 20

trap 0, terminate with error message

inv

invpd

Matrix singular

Matrix not positive definite

If the input to **invpd** is not symmetric, it is possible that the function will (erroneously) appear to operate successfully.

Positive definite matrices can be inverted by **inv**. However, for symmetric, positive definite matrices (such as moment matrices), **invpd** is about twice as fast as **inv**.

inv, invpd

Example

```
x1 = rndn(n,1);
x = ones(n,1)~x1;
btrue = { 1, 0.5 };
y = x*btrue + rndn(n,1);
bols = invpd(x'x)*x'y;
bols = 1.017201 0.484244
```

n = 4000;

This example simulates some data and computes the ols coefficient estimator using the invpd function. First, the number of observations is specified. Second, a vector x1 of standard Normal random variables is generated and is concatenated with a vector of 1's (to create a constant term). The true coefficients are specified, and the dependent variable y is created. Then the ols coefficient estimates are computed.

See also

scalerr, trap, prcsn

Technical Notes

For complex matrices, **inv** uses the ZGECO, ZGEDI path in the LINPACK routines. For real matrices, it uses the **croutp** function.

The **inv** function uses the Crout decomposition. The advantage of this routine is that on some platforms it allows most of the intermediate results to be computed in extended precision.

The **invpd** function uses the Cholesky decomposition and is based upon the LINPACK routines for positive definite matrices. On OS/2 and DOS, if **prcsn** 80 is in effect, all intermediate calculations and intermediate results will be in the 80-bit extended precision of the 80x87 temporary real format. The final results will be rounded to 64-bit double precision.

The tolerance used to determine singularity is 1.0e-14. This can be changed. See "Singularity Tolerance" in the *User's Guide*.

a

b

c d

е

g

h i

j k

1

m

n

0

p

q

S

t u

V

W

invswp

invswp

Purpose

h

d

h

m

0

p

q

Computes a generalized sweep inverse.

Format

y = invswp(x);

Input

x NxN matrix.

Output

y NxN matrix, the generalized inverse of x.

Remarks

This will invert any general matrix. That is, even matrices which will not invert using **inv** because they are singular will invert using **invswp**.

x and *y* will satisfy the four Moore-Penrose conditions:

1. xyx = x

 $2. \quad yxy = y$

3. xy is symmetric

4. yx is symmetric

The tolerance used to determine if a pivot element is zero is taken from the **crout** singularity tolerance. The corresponding row and column are zeroed out. See "Appendix C" in the *User's Guide*.

Example

let x[3,3] = 1 2 3 4 5 6 7 8 9;

y = invswp(x);

-1.6666667 0.66666667 0.0000000

y = 1.3333333 -0.33333333 0.00000000

 $0.0000000 \quad 0.0000000 \quad 0.0000000$

u

V W

хух

3-436

iscplx

iscplx

Purpose Returns whether a matrix is complex or real.

Format y = iscplx(x);

Input x NxK matrix.

Output y scalar, 1 if x is complex, 0 if it is real.

Example $x = \{ 1, 2i, 3 \};$

y = iscplx(x);

y = 1

See also hasimag, iscplxf

a

b

c d

0

f

g h

i

j

k 1

m

n

0

p

q

r

S

t

u

V

XX7

iscplxf

iscplxf

Purpose Returns whether a data set is complex or real.

Format y = iscplxf(fh);

Input *fh* scalar, file handle of an open file.

Output y scalar, 1 if the data set is complex, 0 if it is real.

See also hasimag, iscplx

f g

a

b

С

d

e

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h

k

m

n

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p

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r

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t

u

V

W

isinfnanmiss

isinfnanmiss

Purpose Returns true if the argument contains an infinity, NaN, or missing value.

Format y = isinfnanmiss(x);

Input x NxK matrix.

Output y scalar, 1 if x contains any infinities, NaNs, or missing values, else 0.

See also scalinfnanmiss, ismiss, scalmiss

a

b

c d

е

f g

h

i

j k

1

m

n

0

p

q

r

S

t

u

V

W

ismiss

ismiss

b

Purpose Returns a 1 if its matrix argument contains any missing values, otherwise returns a 0.

Format

y = ismiss(x);

Input

x NxK matrix.

Output

y scalar, 1 or 0.

Remarks

y will be a scalar 1 if the matrix x contains any missing values, otherwise it will be a 0.

An element of x is considered to be a missing if and only if it contains a missing value in the real part. Thus, for x = 1 + .i, **ismiss**(x) will return a 0.

Example

 $x = \{ 1634 \};$

y = ismiss(x);

y = 0

See also

scalmiss, miss, missrv

a

b

d

0

g

h

_

k

m

n

0

p

q

1

+

u

V

W

isSparse

isSparse

Purpose Tests whether a matrix is a sparse matrix.

Format r = isSparse(x);

Input x MxN sparse or dense matrix.

Output r scalar, 1 if x is sparse, 0 otherwise.

Source sparse.src

a

b

c d

f

g

h i

J

k

m

n

0

p

q

1

5

t

u

V

W

keep (dataloop)

b

h

k

m

0

p

u

V

W

хух

keep (dataloop)

Purpose Specifies columns (variables) to be saved to the output data set in a data

loop.

Format keep variable_list;

Remarks Commas are optional in variable_list.

Retains only the specified variables in the output data set. Any variables referenced must already exist, either as elements of the source data set, or

as the result of a previous make, vector, or code statement.

If neither **keep** nor **drop** is used, the output data set will contain all variables from the source data set, as well as any newly defined variables.

The effects of multiple **keep** and **drop** statements are cumulative.

Example keep age, pay, sex;

See also drop

3-442

a

b

С

d

e

f

g

h

k

m

n

0

q

u

x y z

key

Purpose Returns the ASCII value of the next key available in the keyboard buffer.

Format y = key;

Remarks

If you are working in terminal mode, **key** does not "see" any keystrokes until ENTER is pressed. The value returned will be zero if no key is available in the buffer or it will equal the ASCII value of the key if one is available. The key is taken from the buffer at this time and the next call to **key** will return the next key.

Here are the values returned if the key pressed is not a standard ASCII character in the range of 1-255.

1015	SHIFT+TAB
1016-1025	ALT+Q, W, E, R, T, Y, U, I, O, P
1030-1038	ALT+A, S, D, F, G, H, J, K, L
1044-1050	ALT+Z, X, C, V, B, N, M
1059-1068	F1-F10
1071	HOME
1072	CURSOR UP
1073	PAGE UP
1075	LEFT ARROW
1077	RIGHT ARROW
1079	END
1080	DOWN ARROW
1081	PAGE DOWN
1082	INSERT
1083	DELETE
1084-1093	SHIFT+F1-F10
1094-1103	CTRL+F1-F10
1104-1113	ALT+F1-F10
1114	CTRL+PRINT SCREEN
1115	CTRL+LEFT ARROW
1116	CTRL+RIGHT ARROW
1117	CTRL+END
1118	CTRL+PAGE DOWN

key

a

h

d

е

f

h

k

m

n

0

p

q

t

u

V

W

хух

1119 CTRL+HOME 1120-1131 ALT+1,2,3,4,5,6,7,8,9,0,HYPHEN, **EQUAL SIGN** CTRL+PAGE UP 1132 **Example** format /rds 1,0; kk = 0;do until kk == 27; kk = key;if kk == 0;continue; elseif kk == vals(" "); print "space \\" kk; elseif kk == vals("\r"); print "carriage return \\" kk; elseif kk >= vals("0") and kk <= vals("9");print "digit \\" kk chrs(kk); elseif vals(upper(chrs(kk))) >= vals("A") and vals(upper(chrs(kk))) <= vals("Z");</pre> print "alpha \\" kk chrs(kk); else; print "\\" kk; endif; endo; This is an example of a loop that processes keyboard input. This loop will continue until the ESC key (ASCII 27) is pressed. See also vals, chrs, upper, lower, con, cons

keyav

keyav

Purpose Check if keystroke is available.

Format x = keyav;

Output x scalar, value of key or 0 if no key is available.

See also keyw, key

a

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c d

е

f

g h

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1

m

n

Ο

p

q

1

S

t

u

V

W

keyw

a

b

d

h

k

m

n

0

p

q

u

V

W

хух

keyw

Purpose Waits for and gets a key.

Format k = keyw;

Output k scalar, ASCII value of the key pressed.

Remarks If you are working in terminal mode, GAUSS will not see any input until

you press the ENTER key. **keyw** gets the next key from the keyboard buffer. If the keyboard buffer is empty, **keyw** waits for a keystroke. For normal keys, **keyw** returns the ASCII value of the key. See **key** for a

table of return values for extended and function keys.

See also key

3-446

keyword

keyword

Purpose

Begins the definition of a keyword procedure. Keywords are user-defined functions with local or global variables.

Format

keyword name(str);

Input

name literal, name of the keyword. This name will be a global symbol.
 str string, a name to be used inside the keyword to refer to the argument that is passed to the keyword when the keyword is called. This will always be local to the keyword, and cannot be accessed from outside the keyword or from other keywords or procedures.

Remarks

A keyword definition begins with the **keyword** statement and ends with the **endp** statement. See "Procedures and Keywords" in *User's Guide*.

Keywords always have 1 string argument and 0 returns. GAUSS will take everything past *name*, excluding leading spaces, and pass it as a string argument to the keyword. Inside the keyword, the argument is a local string. The user is responsible to manipulate or parse the string.

An example of a keyword definition is:

```
keyword add(str);
  local tok,sum;
  sum = 0;
  do until str $== "";
     { tok, str } = token(str);
     sum = sum + stof(tok);
  endo;
  print "Sum is: " sum;
endp;
```

To use this keyword, type:

add 1 2 3 4 5;

а

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С

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keyword

This keyword will respond by printing:

Sum is: 15

See also proc, local, endp

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lag (dataloop)

lag (dataloop)

Purpose Lags variables a specified number of periods.

Format lag nv1 = var1:p1 [[nv2 = var2:p2...]];

Input *var* name of the variable to lag.

p scalar constant, number of periods to lag.

Output nv name of the new lagged variable.

Remarks You can specify any number of variables to lag. Each variable can be lagged a different number of periods. Both positive and negative lags are allowed.

Lagging is executed before any other transformations. If the new variable name is different from that of the variable to lag, the new variable is first created and appended to a temporary data set. This temporary data set becomes the input data set for the data loop, and is then automatically deleted.

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lag1

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lag1

Purpose Lags a matrix by one time period for time series analysis.

Format y = lag1(x);

Input x NxK matrix.

Output y NxK matrix, x lagged 1 period.

Remarks lag1 lags x by one time period, so the first observations of y are missing.

Source lag.src

See also lagn

3-450

lagn

lagn

Purpose Lags a matrix a specified number of time periods for time series analysis.

Format y = lagn(x,t);

Input x NxK matrix.

t scalar, number of time periods.

Output y NxK matrix, x lagged t periods.

Remarks If t is positive, lagn lags x back t time periods, so the first t observations

of y are missing. If t is negative, **lagn** lags x forward t time periods, so

the last *t* observations of *y* are missing.

Source lag.src

See also lag1

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lapeighb

lapeighb

Purpose

Computes eigenvalues only of a real symmetric or complex Hermitian matrix selected by bounds.

Format

ve = lapeighb(x, vl, vu);

Input

x NxN matrix, real symmetric or complex Hermitian.

vl scalar, lower bound of the interval to be searched for eigenvalues.

vu scalar, upper bound of the interval to be searched for eigenvalues; vu must be greater than vl.

abstol scalar, the absolute error tolerance for the eigenvalues. An approximate eigenvalue is accepted as converged when it is determined to lie in an interva [a,b] of width less than or equal to ABSTOL + EPS*max(|a|,|b|), where EPS is machine precision. If ABSTOL is less than or equal to zero, then EPS*||T|| will be used in its place, where T is the tridiagonal matrix obtained by reducing the input matrix to tridiagonal form.

Output

ve

Mx1 vector, eigenvalues, where M is the number of eigenvalues on the half open interval (*vl*,*vu*]. If no eigenvalues are found then *s* is a scalar missing value.

Remarks

lapeighb computes eigenvalues only which are found on on the half open interval (vl,vu]. To find eigenvalues within a specified range of indices see lapeighi. For eigenvectors see lapeighvi, or lapeighvb lapeighb is based on the LAPACK drivers DYESVX and ZHEEVX. Further documentation of these functions may be found in the LAPACK User's Guide.

Example

```
x = { 5   2   1,
        2   6   2,
        1   2   9 };
vl = 5;
vu = 10;
ve = lapeighi(x,il,iu,0);
```

9

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lapeighb

print ve;

6.0000

See also lapeighb, lapeighvi, lapeighvb

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t

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...

lapeighi

lapeighi

Purpose

Computes eigenvalues only of a real symmetric or complex Hermitian matrix selected by index.

Format

```
ve = lapeighi(x,il,iu,abstol);
```

Input

x NxN matrix, real symmetric or complex Hermitian.

il scalar, index of the smallest desired eigenvalue ranking them from smallest to largest.

iu scalar, index of the largest desired eigenvalue, *iu* must be greater than *il*.

abstol scalar, the absolute error tolerance for the eigenvalues. An approximate eigenvalue is accepted as converged when it is determined to lie in an interval [a,b] of width less than or equal to ABSTOL + EPS*max(|a|,|b|), where EPS is machine precision. If ABSTOL is less than or equal to zero, then EPS*||T|| will be used in its place, where T is the tridiagonal matrix obtained by reducing the input matrix to tridiagonal form.

Output

ve (iu-il+1)x1 vector, eigenvalues.

Remarks

lapeighi computes *iu-il+1* eigenvalues only given a range of indices, i.e., the i-th to j-th eigenvalues, ranking them from smallest to largest. To find eigenvalues within a specified range see lapeighxb. For eigenvectors see LEIGHVX, lapeighvi, or lapeighvb. lapeighi is based on the LAPACK drivers DYESVX and ZHEEVX. Further documentation of these functions may be found in the LAPACK User's Guide.

Example

```
x = { 5   2   1,
        2   6   2,
        1   2   9 };
il = 2;
iu = 3;
ve = lapeighi(x,il,iu,0);
print ve;
```

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lapeighi

6.0000 10.6056

See also lapeighb, lapeighvi, lapeighvb

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1

m

n

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r

S

t

u

V

lapeighvb

lapeighvb

Purpose

Computes eigenvalues and eigenvectors of a real symmetric or complex Hermitian matrix selected by bounds.

Format

```
\{ ve, va \} = lapeighvb(x, vl, vu, abstol);
```

Input

x NxN matrix, real symmetric or complex Hermitian.

vl scalar, lower bound of the interval to be searched for eigenvalues.

vu scalar, upper bound of the interval to be searched for eigenvalues; vu must be greater than vl.

abstol scalar, the absolute error tolerance for the eigenvalues. An approximate eigenvalue is accepted as converged when it is determined to lie in an interval [a,b] of width less than or equal to ABSTOL + EPS*max(|a|,|b|), where EPS is machine precision. If ABSTOL is less than or equal to zero, then EPS*||T|| will be used in its place, where T is the tridiagonal matrix obtained by reducing the input matrix to tridiagonal form.

Output

ve Mx1 vector, eigenvalues, where M is the number of eigenvalues on the half open interval (vl,vu]. If no eigenvalues are found then s is a scalar missing value.

va NxM matrix, eigenvectors.

Remarks

lapeighvb computes eigenvalues and eigenvectors which are found on the half open interval (*vl*,*vu*]. **lapeighvb** is based on the LAPACK drivers DYESVX and ZHEEVX. Further documentation of these functions may be found in the LAPACK User's Guide.

Example

```
x = { 5   2   1,
        2   6   2,
        1   2   9 };
vl = 5;
vu = 10;
{ ve,va } = lapeighvb(x,il,iu,0);
```

a

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W

lapeighvb

print ve;

6.0000

print va;

-0.5774 -0.5774

0.5774

See also lapeighvb

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d

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337

lapeighvi

lapeighvi

Purpose

Computes selected eigenvalues and eigenvectors of a real symmetric or complex Hermitian matrix.

Format

```
\{ ve, va \} = lapeighvi(x, il, iu, abstol);
```

Input

x NxN matrix, real symmetric or complex Hermitian.

il scalar, index of the smallest desired eigenvalue ranking them from smallest to largest.

iu calar, index of the largest desired eigenvalue, *iu* must be greater than *il*.

abstol scalar, the absolute error tolerance for the eigenvalues. An approximate eigenvalue is accepted as converged when it is determined to lie in an interval [a,b] of width less than or equal to ABSTOL + EPS*max(|a|,|b|), where EPS is machine precision. If ABSTOL is less than or equal to zero, then EPS*||T|| will be used in its place, where T is the tridiagonal matrix obtained by reducing the input matrix to tridiagonal form.

Output

ve (iu-il+1)x1 vector, eigenvalues.

va Nx(*iu-il*+1) matrix, eigenvectors.

Remarks

lapeighvi computes *iu-il+*1 eigenvalues and eigenvectors given a range of indices, i.e., the i-th to j-th eigenvalues, ranking them from smallest to largest. To find eigenvalues and eigenvectors within a specified range see **lapeighvb**. **lapeighvi** is based on the LAPACK drivers DYESVX and ZHEEVX. Further documentation of these functions may be found in the LAPACK User's Guide.

Example

```
x = { 5   2   1,
        2   6   2,
        1   2   9 };
il = 2;
iu = 3;
{ ve,va } = lapeighvi(x,il,iu,0);
```

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b

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m

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q

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u

V

W

lapeighvi

print ve;

6.0000 10.6056

print va;

-0.5774 0.3197 -0.5774 0.4908

0.5774 0.8105

See also lapeighbb, lapeighb

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u

V

W

lapgeig

lapgeig

Purpose Computes generalized eigenvalues for a pair of real or complex general matrices.

Format $\{ val, va2 \} = lapgeig(A,B);$

Input \boldsymbol{A} NxN matrix, real or complex general matrix.

> \boldsymbol{R} NxN matrix, real or complex general matrix.

Output va1 Nx1 vector, numerator of eigenvalues.

> va2 Nx1 vector, denominator of eigenvalues.

Remarks val and va2 are the vectors of the numerators and denominators

respectively of the eigenvalues of the solution of the generalized symmetric eigenproblem of the form Aw = eBw where A and B are real or complex general matrices and w = va1 / va2. The generalized eigenvalues are not computed directly because some elements of va2 may be zero, i.e., the eigenvalues may be infinite. This procedure calls the LAPACK

routines DGEGV and ZGEGV.

See also lapgeig, lapgeigh

3-460

b

d

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m

0 p

u

V

W

lapgeigh

lapgeigh

Purpose Computes generalized eigenvalues for a pair of real symmetric or Hermitian matrices.

Format ve = lapgeigh(A, B);

Input A NxN matrix, real or complex symmetric or Hermitian matrix.

B NxN matrix, real or complex positive definite symmetric or Hermitian matrix.

Output *ve* Nx1 vector, eigenvalues.

Remarks ve is the vector of eigenvalues of the solution of the generalized symmetric eigenproblem of the form $Ax = \lambda Bx$.

Example A = { 3 4 5, 2 5 2, 3 2 4 }; B = { 4 2 2,

2 6 1, 2 1 8 };

ve = lapgeigh(A,B);

print ve; -0.18577146

0.50880165 1.1335370

This procedure calls the LAPACK routines DSYGV and ZHEGV.

See also lapgeig, lapgeighv

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n m

n

0

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q

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S

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u

V

lapgeighv

lapgeighv

h

Purpose

Computes generalized eigenvalues and eigenvectors for a pair of real symmetric or Hermitian matrices.

Format

$$\{ ve, va \} = lapgeighv(A,B);$$

Input

A NXN matrix, real or complex symmetric or Hermitian matrix.

B NxN matrix, real or complex positive definite symmetric or Hermitian matrix.

Output

ve Nx1 vector, eigenvalues.

va NxN matrix, eigenvectors.

Remarks

ve and va are the eigenvalues and eigenvectors of the solution of the generalized symmetric eigenproblem of the form $Ax = \lambda Bx$. Equivalently, va diagonalizes $U^{-1}AU^{-1}$ in the following way

$$vaU'^{-1}AU^{-1}va' = e$$

where B = U'U. This procedure calls the LAPACK routines DSYGV and ZHEGV.

Example

a

d

е

g

h

j

k

m

n

0

p q

r

S

t

u

V

W

lapgeighv

print va;

0.3575 -0.0996 0.9286

-0.2594 0.9446 0.2012

-0.8972 -0.3128 0.3118

See also lapgeig, lapgeigh

a

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l m

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q

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u

V

XX7

lapgeigv

lapgeigv

Purpose h

d

h

k

1

m

0

p

Computes generalized eigenvalues, left eigenvectors, and right eigenvectors for a pair of real or complex general matrices.

Format

 $\{ val, va2, lve, rve \} = lapgeigv(A,B);$

Input

A NxN matrix, real or complex general matrix.

B NxN matrix, real or complex general matrix.

Output

val Nx1 vector, numerator of eigenvalues.

va2 Nx1 vector, denominator of eigenvalues.

lve NxN left eigenvectors.

rve NxN right eigenvectors.

Remarks

va1 and va2 are the vectors of the numerators and denominators respectively of the eigenvalues of the solution of the generalized symmetric eigenproblem of the form $Aw = \lambda Bw$ where A and B are real or complex general matrices and w = va1. va2. The generalized eigenvalues are not computed directly because some elements of va2 may be zero, i.e., the eigenvalues may be infinite.

The left and right eigenvectors diagonalize $U^{-1}A$ U^{-1} where B = U'U, that is,

$$lve U'^{-1}AU lve' = w$$

and

$$rve'U'^{-1}AU^{-1}rve = w$$

This procedure calls the LAPACK routines DGEGV and ZGEGV.

See also

lapgeig, lapgeigh

V W

u

хуZ

lapgsvds

lapgsvds

Purpose Compute the generalized singular value decomposition of a pair of real or complex general matrices.

Format $\{C,S,R\} = \text{lapgsvds}(A,B);$

Input A MxN real or complex matrix.

B PxN real or complex matrix.

Output C Lx1 vector, singular values for A.

S Lx1 vector, singular values for B.

R (K+L)x(K+L) upper triangular matrix.

Remarks (1) The generalized singular value decomposition of A and B is

$$U'AQ = D_1Z$$

$$V'BQ = D_2Z$$

where U, V, and Q are orthogonal matrices (see **lapgsvdcst** and **lapgsvdst**). Letting K+L = the rank of A|B then R is a (K+L)x(K+L) upper triangular matrix, D1 and D2 are Mx(K+L) and Px(K+L) matrices with entries on the diagonal, Z = [0 R], and if M-K-L >= 0

$$D_{1} = \begin{pmatrix} K & L \\ K & I & O \\ O & C \\ M - K - L & O & O \end{pmatrix}$$

$$D_2 = \begin{array}{c} K L \\ L \begin{bmatrix} O & S \\ O & O \end{bmatrix} \end{array}$$

a

b

С

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j

k

n m

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0

p

q

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v

XX7

lapgsvds

a

b

С

d

е

J

m

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U

p

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r

2

u

V

W

хуг

$$[O R] = K \begin{bmatrix} N - K - L & K & L \\ O & R_{11} & R_{12} \\ D & O & R_{22} \end{bmatrix}$$

$$D_1 = \begin{bmatrix} K & M-K & K+L-M \\ K & I & O & O \\ M-K & O & C & O \end{bmatrix}$$

$$D_2 = \begin{bmatrix} K & M-K & K+L-M \\ M-K & O & S & O \\ O & O & I \\ P-L & O & O & O \end{bmatrix}$$

$$[O\ R] = M - K \begin{bmatrix} N - K - L & K & M - K & K + L - M \\ K & O & R_{11} & R_{12} & R_{13} \\ O & O & R_{22} & R_{23} \\ K + L - M & O & O & O & R_{33} \end{bmatrix}$$

(2) Form the matrix

$$X = Q \begin{bmatrix} I & O \\ O & R^{-1} \end{bmatrix}$$

then

$$A = U'^{-1}E_1X$$

$$B = V^{\prime^{-1}} E_2 X^{-1}$$

where
$$E_1 = \begin{bmatrix} O & D_2 \end{bmatrix}$$
. $E_2 = \begin{bmatrix} O & A_2 \end{bmatrix}$

lapgsvds

(3) The generalized singular value decomposition of A and B implicitly produces the singular value decomposition of AB^{-1} :

$$AB^{-1} = UD_1D_2^{-1}V'$$

This procedure calls the LAPACK routines DGGSVD and ZGGSVD.

See also lapgsvdcst and lapgsvdst

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lapgsvdcst

lapgsvdcst

Purpose

Compute the generalized singular value decomposition of a pair of real or complex general matrices.

Format

 $\{C,S,R,U,V,Q\} = lapgsvdcst(A,B);$

Input

A MxN matrix.

B PxN matrix.

Output

C Lx1 vector, singular values for A.

S Lx1 vector, singular values for B.

R (K+L)x(K+L) upper triangular matrix.

U MxM matrix, orthogonal transformation matrix.

V PxP matrix, orthogonal transformation matrix.

U NxN matrix, orthogonal transformation matrix.

Remarks

(1) The generalized singular value decomposition of A and B is

$$U'AQ = D_1Z$$

$$V'BQ = D_2Z$$

where U, V, and Q are orthogonal matrices (see **lapgsvdcst** and **lapgsvdst**). Letting K+L = the rank of A|B then R is a (K+L)x(K+L) upper triangular matrix, D1 and D2 are Mx(K+L) and Px(K+L) matrices with entries on the diagonal, Z = [0 R], and if M-K-L >= 0

$$D_{1} = KL \\ K \begin{bmatrix} I & O \\ O & C \\ M - K - L \end{bmatrix}$$

$$D_2 = \begin{array}{c} K L \\ L \begin{bmatrix} O & S \\ O & O \end{bmatrix} \end{array}$$

b

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f

h

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k

1

m

0

р

q

1

,

u

V

W

x y z

lapgsvdcst

$$[O R] = K \begin{bmatrix} N - K - L & K & L \\ O & R_{11} & R_{12} \\ D & O & R_{22} \end{bmatrix}$$

$$D_1 = \begin{bmatrix} K & M-K & K+L-M \\ K & I & O & O \\ M-K & O & C & O \end{bmatrix}$$

$$D_2 = K + L - M \begin{bmatrix} K & M - K & K + L - M \\ M - K \begin{bmatrix} O & S & O \\ O & O & I \\ P - L & O & O \end{bmatrix}$$

$$[O \ R] = M - K \begin{bmatrix} N - K - L & K & M - K & K + L - M \\ K & O & R_{11} & R_{12} & R_{13} \\ O & O & R_{22} & R_{23} \\ K + L - M & O & O & R_{33} \end{bmatrix}$$

(2) Form the matrix

$$X = Q \begin{bmatrix} I & O \\ O & R^{-1} \end{bmatrix}$$

then

$$A = U'^{-1}E_1X$$

$$B = V^{\prime -1} E_2 X^{-1}$$

where
$$E_1 = \begin{bmatrix} O & D_2 \end{bmatrix}$$
. $E_2 = \begin{bmatrix} O & A_2 \end{bmatrix}$

a

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c d

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f g

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i

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q

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V

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lapgsvdcst

(3) The generalized singular value decomposition of A and B implicitly produces the singular value decomposition of AB^{-1} :

$$AB^{-1} = UD_1D_2^{-1}V'$$

This procedure calls the LAPACK routines DGGSVD and ZGGSVD.

See also lapgsvds and lapgsvdst

a

b

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m

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V

W

lapgsvdst

lapgsvdst

Purpose Compute the generalized singular value decomposition of a pair of real or complex general matrices.

Format { D1,D2,Z,U,V,Q } = lapgsvdst(A,B);

Input A MxN matrix.

B PxN matrix.

Output D1 Mx(K+L) matrix, with singular values for A on diagonal.

D2 Px(K+L) matrix, with singular values for B on diagonal.

Z (K+L)xN matrix, partitioned matrix composed of a zero matrix and upper triangular matrix.

U MxM matrix, orthogonal transformation matrix.

V PxP matrix, orthogonal transformation matrix.

U NxN matrix, orthogonal transformation matrix.

Remarks (1) The generalized singular value decomposition of A and B is

$$U'AQ = D_1Z$$

$$V'BQ = D_2Z$$

where U, V, and Q are orthogonal matrices (see **lapgsvdcst** and **lapgsvdst**). Letting K+L = the rank of A|B then R is a (K+L)x(K+L) upper triangular matrix, D1 and D2 are Mx(K+L) and Px(K+L) matrices with entries on the diagonal, Z = [0 R], and if M-K-L >= 0

$$D_{1} = KL K \begin{bmatrix} I & O \\ O & C \\ M - K - L \end{bmatrix}$$

$$D_2 = \begin{bmatrix} K & L \\ O & S \\ P - L & O & O \end{bmatrix}$$

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W

lapgsvdst

a

b

C

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1.

i

J

I

m

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9

Γ

2

u

V

W

хуг

$$[O R] = K \begin{bmatrix} N - K - L & K & L \\ O & R_{11} & R_{12} \\ O & O & R_{22} \end{bmatrix}$$

$$D_1 = \begin{bmatrix} K & M-K & K+L-M \\ K & O & O \\ M-K & O & C \end{bmatrix}$$

$$D_2 = \begin{bmatrix} K & M-K & K+L-M \\ M-K & O & S & O \\ O & O & I \\ P-L & O & O & O \end{bmatrix}$$

$$[O\ R] = M - K \begin{bmatrix} N - K - L & K & M - K & K + L - M \\ K & O & R_{11} & R_{12} & R_{13} \\ O & O & R_{22} & R_{23} \\ K + L - M & O & O & R_{33} \end{bmatrix}$$

(2) Form the matrix

$$X = Q \begin{bmatrix} I & O \\ O & R^{-1} \end{bmatrix}$$

then

$$A = U'^{-1}E_1X$$

$$B = V^{\prime^{-1}} E_2 X^{-1}$$

where
$$E_1 = \begin{bmatrix} O & D_2 \end{bmatrix}$$
. $E_2 = \begin{bmatrix} O & A_2 \end{bmatrix}$

lapgsvdst

(3) The generalized singular value decomposition of A and B implicitly produces the singular value decomposition of AB^{-1} :

$$AB^{-1} = UD_1D_2^{-1}V'$$

This procedure calls the LAPACK routines DGGSVD and ZGGSVD.

See also lapgsvds and lapgsvdcst

a

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VV

lapschur

h

h

k

m

p

lapschur

Purpose

Compute the generalized Schur form of a pair of real or complex general matrices.

Format

 $\{ sa, sb, q, z \} = lapschur(A,B);$

Input

A NxN matrix, real or complex general matrix.

B NxN matrix, real or complex general matrix.

Output

sa NxN matrix, Schur form of A.

sb NxN matrix, Schur form of B.

q NxN matrix, left Schur vectors.

z NxN matrix, right Schur vectors.

Remarks

The pair of matrices *A* and *B* are in generalized real Schur form when *B* is upper triangular with non-negative diagonal, and *A* is block upper triangular with 1x1 and 2x2 blocks. The 1x1 blocks correspond to real generalized eigenvalues and the 2x2 blocks to pairs of complex conjugate eigenvalues. The real generalized eigenvalues can be computed by dividing the diagonal element of sa by the corresponding diagonal element of *sb*. The complex generalized eigenvalues are computed by first constructing two complex conjugate numbers from 2x2 block where the real parts are on the diagonal of the block and the imaginary part on the off-diagonal. The eigenvalues are then computed by dividing the two complex conjugate values by their corresponding diagonal elements of *sb*. The generalized Schur vectors q and z are orthogonal matrices that reduce *A* and *B* to Schur form:

sb q'Bz

This procedure calls the LAPACK routines DGEGS and ZGEGS.

u

V

W

lapsvdcusv

lapsvdcusv

Purpose Computes the singular value decomposition a real or complex rectangular matrix, returns compact u and v.

Format $\{u,s,v\} = \text{lapsvdcusv}(x);$

Input x MxN matrix, real or complex rectangular matrix.

Output u Mxmin(M,N) matrix, left singular vectors.

 $s = \min(M,N)xN$ matrix, singular values.

v NxN matrix, right singular values.

Remarks lapsvdcusv computes the singular value decomposition of a real or complex rectangular matrix. The SVD is

```
x = usv'
```

where v is the matrix of right singular vectors. **lapsvdcusv** is based on the LAPACK drivers DGESVD and ZGESVD. Further documentation of these functions may be found in the LAPACK User's Guide.

```
Example
```

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lapsvdcusv

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print s;

13.895868 0.0000000 0.0000000
0.0000000 2.1893939 0.0000000
0.0000000 0.0000000 1.4344261

print v;

-0.13624432 -0.62209955 -0.77099263
0.46497296 0.64704876 -0.60425826
0.87477862 -0.44081748 0.20110275

See also lapsvds, lapsvdusv

lapsvds

lapsvds

Purpose Computes the singular values of a real or complex rectangular matrix

Format s = lapsvds(x);

Input *x* MxN matrix, real or complex rectangular matrix.

Output s MiN(M,N)x1 vector, singular values.

Remarks lapsvd computes the singular values of a real or complex rectangular matrix. The svd is

x = usv'

where v is the matrix of right singular vectors. For the computation of the singular vectors, see **lapsvdcusv** and **lapsvdusv**.

lapsvd is based on the LAPACK drivers DGESVD and ZGESVD. Further documentation of these functions may be found in the LAPACK User's Guide.

Example

10.352877 4.0190557 2.3801546

See also lapsvdcusv, lapsvdusv

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lapsvdusv

lapsvdusv

Purpose Computes the singular value decomposition a real or complex rectangular matrix.

Format $\{u,s,v\} = lapsvdusv(x);$

Input x MxN matrix, real or complex rectangular matrix.

Output u MxM matrix, left singular vectors.

s MxN matrix, singular values.

v NxN matrix, right singular values.

Remarks lapsvdusv computes the singular value decomposition of a real or complex rectangular matrix. The SVD is

x = usv'

where *v* is the matrix of right singular vectors. **lapsvdusv** is based on the LAPACK drivers DGESVD and ZGESVD. Further documentation of these functions may be found in the LAPACK User's Guide.

Example

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lapsvdusv

print v;
-0.1362 0.4650 0.8748

0.6221 0.6470 -0.4408 -0.7710 -0.6043 0.2011

See also lapsvds, lapsvdcusv

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Purpose

Creates a matrix from a list of numeric or character values. The result is always of type matrix, string or string array.

Format

let x = constant_list;

Remarks

Expressions and matrix names are not allowed in the **let** command. Expressions such as this:

let
$$x[2,1] = 3*a b$$

are illegal. To define matrices by combining matrices and expressions, use an expression containing the concatenation operators: ~ and | .

Numbers can be entered in scientific notation. The syntax is $dE\pm n$, where d is a number and n is an integer (denoting the power of 10).

let
$$x = 1e+10 \ 1.1e-4 \ 4.019e+2;$$

Complex numbers can be entered by joining the real and imaginary parts with a sign (+ or -); there should be no spaces between the numbers and the sign. Numbers with no real part can be entered by appending an "i" to the number.

let
$$x = 1.2+23 8.56i 3-2.1i -4.2e+6i$$

 $1.2e-4-4.5e+3i$

If curly braces are used, the **let** is optional. You will need the **let** for statements that you want to protect from the beautifier using the **-1** flag on the beautifier command line.

let
$$x = \{ 1 2 3, 4 5 6, 7 8 9 \};$$

 $x = \{ 1 2 3, 4 5 6, 7 8 9 \};$

If indices are given, a matrix of that size will be created:

let
$$x[2,2] = 1 2 3 4;$$

$$x = \begin{array}{c} 1 & 2 \\ 3 & 4 \end{array}$$

If indices are not given, a column vector will be created:

let
$$x = 1 2 3 4;$$

$$x = \begin{array}{c} 1 \\ 2 \\ 3 \\ 4 \end{array}$$

You can create matrices with no elements, i.e., "empty matrices". Just use a set of empty curly braces.

$$x = \{\};$$

Empty matrices are chiefly used as the starting point for building up a matrix, for example in a **do** loop. For more information on empty matrices, see "Language Fundamentals" in the *User's Guide*.

Character elements are allowed in a **let** statement:

let
$$x = age pay sex;$$

$$x = \begin{array}{c} AGE \\ PAY \\ SEX \end{array}$$

Lowercase elements can be created if quotation marks are used. Note that each element must be quoted.

$$x = \begin{cases} age \\ pay \end{cases}$$

Example

$$x = 0$$

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c d

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let $x = \{ 1 2 3, 4 5 6, 7 8 9 \};$

1 2 3

x = 456 789

let x[3,3] = 1 2 3 4 5 6 7 8 9;

1 2 3

x = 456

7 8 9

let x[3,3] = 1;

1 1 1

x = 1 1 1

1 1 1

let x[3,3];

0 0 0

x = 0 0 0

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let x = 1 2 3 4 5 6 7 8 9;1 2 3 x = 56 7 8 9 let x = dog cat;DOG x =CATlet x = "dog" "cat"; $x = \log$ cat let string x = { "Median Income" "Country"; Median Income Country See also con, cons, declare, load

lib

lib

a **Purpose** Builds and updates library files. h **Format** lib library [file] [-flag -flag...]; Input d library literal, name of library. file optional literal, name of source file to be updated or added. optional literal preceded by '-', controls operation of library flags update. To control handling of path information on source filenames: -addpath (default) add paths to entries without paths and expand relative paths. h reset all paths using a normal file -gausspath search. -leavepath leave all path information untouched. drop all path information. -nopath k To specify a library update or a complete library build: (default) update the symbol information -update for the specified file only. update the symbol information for every m -build library entry by compiling the actual source file. delete a file from the library. -delete 0 -list list files in a library. p To control the symbol type information placed in the library file: (default) use strongly typed symbol -strong entries. -weak save no type information. This should only be used to build a library compatible with a previous version of t GAUSS. u V W

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To control location of temporary files for a complete library build:

-tmp	(default) use the directory pointed to by
	the tmp_path configuration variable.
	The directory will usually be on a RAM
	disk. If tmp_path is not defined, lib
	will look for a tmp environment

variable.

-disk use the same directory listed in the lib_path configuration variable.

Remarks

The flags can be shortened to one or two letters, as long as they remain unique — for example, **-b** to **-build** a library, **-li** to list files in a library.

If the filenames include a full path, the compilation process is faster because no unnecessary directory searching is needed during the autoloading process. The default path handling adds a path to each file listed in the library and also expands any relative paths so the system will work from any drive or subdirectory.

When a path is added to a filename containing no path information, the file is searched for on the current directory and then on each subdirectory listed in **src_path**. The first path encountered that contains the file is added to the filename in the library entry.

See also library

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library

library

Purpose Sets up the list of active libraries.

Format library [-1] lib1, lib2, lib3, lib4; library;

Remarks If no arguments are given, the list of current libraries will be printed out.

The -1 option will write a file containing a listing of libraries, files, and symbols for all active libraries. This file will reside in the directory defined by the lib_path configuration variable. Under Windows and UNIX, the file will have a unique name beginning with liblst_. Under OS/2 and DOS, the file will be called gausslib.lst; if it already exists it will be overwritten.

For more information about the library system, see "Libraries" in the *User's Guide*.

The default extension for library files is .lcg.

If a list of library names is given, they will be the new set of active libraries. The two default libraries are user.lcg and gauss.lcg. Unless otherwise specified, user.lcg will be searched first and gauss.lcg will be searched last. Any other user-specified libraries will be searched after user.lcg in the order they were entered in the library statement.

If the statement:

```
y = doq(x);
```

is encountered in a program, **dog** will be searched for in the active libraries. If it is found, it will be compiled. If it cannot be found in a library, the deletion state determines how it is handled:

autodelete on search for dog.g autodelete off return Undefined symbol error message

If dog calls cat and cat calls bird and they are all in separate files, they will all be found by the autoloader.

The source browser and the help facility will search for **dog** in exactly the same sequence as the autoloader. The file containing **dog** will be displayed in the window and you can scroll up and down and look at the code and comments.

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library

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Library files are simple ASCII files that you can create with the editor. Here is an example:

```
/*
** This is a GAUSS library file.
* /
eig.src
     eig
                : proc
     eigsym
                : proc
     eigerr
                : matrix
svd.src
     cond
                : proc
     pinv
                : proc
     rank
                : proc
     svd
                : proc
     svdtol
                : matrix
```

The lines not indented are the file names. The lines that are indented are the symbols defined in that file. As you can see, a GAUSS library is a dictionary of files and the global symbols they contain.

Any line beginning with /*, ** or */ is considered a comment. Blank lines are okay.

Here is a debugging hint. If your program is acting strange and you suspect it is autoloading the wrong copy of a procedure, use the source browser or help facility to locate the suspected function. It will use the same search path that the autoloader uses.

See also declare, external, lib, proc

#lineson, #linesoff

#lineson, #linesoff

Purpose

The **#lineson** command causes GAUSS to embed line number and file name records in a program for the purpose of reporting the location where an error occurs. The **#linesoff** command causes GAUSS to stop embedding line and file records in a program.

Format

#lineson;
#linesoff;

Remarks

In the "lines on" mode, GAUSS keeps track of line numbers and file names and reports the location of an error when an execution time error occurs. In the "lines off" mode, GAUSS does not keep track of lines and files at execution time. During the compile phase, line numbers and file names will always be given when errors occur in a program stored in a disk file.

It is easier to debug a program when the locations of errors are reported, but this slows down execution. In programs with several scalar operations, the time spent tracking line numbers and file names is most significant.

These commands have no effect on interactive programs (that is, those typed in the window and run from the command line), since there are no line numbers in such programs.

Line number tracking can be turned on and off through the user interface, but the **#lineson** and **#linesoff** commands will override that.

The line numbers and file names given at run-time will reflect the last record encountered in the code. If you have a mixture of procedures that were compiled without line and file records and procedures that were compiled with line and file records, use the **trace** command to locate exactly where the error occurs.

The **Currently active call** error message will always be correct. If it states that it was executing procedure xyz at line number nnn in file ABC and xyz has no line nnn or is not in file ABC, you know that it just did not encounter any line or file records in xyz before it crashed.

When using **#include**'d files, the line number and file name will be correct for the file the error was in, within the limits stated above.

See also

trace

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listwise (dataloop)

listwise (dataloop)

Purpose Controls listwise deletion of missing values.

Format listwise [read] | [write];

Remarks If read is specified, the deletion of all rows containing missing values

happens immediately after reading the input file and before any transformations. If **write** is specified, the deletion of missing values happens after any transformations and just before writing to the output file. If no **listwise** statement is present, rows with missing values are not deleted.

The default is **read**.

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Purpose Computes the natural log of all elements of a matrix.

Format $y = \ln(x)$;

Input x NxK matrix.

Output y NxK matrix containing the natural log values of the elements of x.

Remarks In is defined for $x \neq 0$.

If x is negative, complex results are returned.

You can turn the generation of complex numbers for negative inputs on or off in the GAUSS configuration file, and with the **sysstate** function, case 8. If you turn it off, **1n** will generate an error for negative inputs.

If *x* is already complex, the complex number state doesn't matter; **1n** will compute a complex result.

x can be any expression that returns a matrix.

Example y = ln(16);

y = 2.7725887

lncdfbvn

lncdfbvn

Purpose Computes natural log of bivariate Normal cumulative distribution function.

Format y = lncdfbvn(x1, x2, r);

Input *x1* NxK matrix, abscissae.

x2 LxM matrix, abscissae.r PxQ matrix, correlations.

Output $y = \max(N,L,P) \times \max(K,M,Q) \text{ matrix, } \ln \Pr(X < x1, X < x2 \mid r).$

Remarks x1, x2, and r must be ExE conformable.

Source lncdfn.src

See also cdfbvn, lncdfmvn

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lncdfbvn2

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lncdfbvn2

Purpose Returns log of cdfbvn of a bounded rectangle. **Format** y = lncdfbvn2(h,dh,k,dk,r);Input h Nx1 vector, upper limits of integration for variable 1. dh Nx1 vector, increments for variable 1. kNx1 vector, upper limits of integration for variable 2. dkNx1 vector, increments for variable 2. Nx1 vector, correlation coefficients between the two variables. r **Output** Nx1 vector, the log of the integral from h,k to h+dh,k+dk of the y standardized bivariate Normal distribution. Remarks Scalar input arguments are okay; they will be expanded to Nx1 vectors. **lncdfbvn2** will abort if the computed integral is negative. **lncdfbvn2** computes an error estimate for each set of inputs--the real integral is exp(y)±err. The size of the error depends on the input arguments. If trap 2 is set, a warning message is displayed when err >= exp(y)/100. For an estimate of the actual error, see **cdfbvn2e**. **Example** Example 1 lncdfbvn2(1,1,1,1,0.5); produces: -3.2180110258198771e+000 Example 2 trap 0,2; lncdfbvn2(1,1e-15,1,1e-15,0.5); produces: -7.1171016046360151e+001

lncdfbvn2

```
Example 3
trap 2,2;
lncdfbvn2(1,-1e-45,1,1e-45,0.5);
WARNING: Dubious accuracy from lncdfbvn2:
0.000e+000 ± 2.8e-060
-INF
```

See also cdfbvn2, cdfbvn2e

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lncdfmvn

lncdfmvn

Purpose Computes natural log of multivariate Normal cumulative distribution function.

Format y = lncdfmvn(x,r);

Input *x* KxL matrix, abscissae.

r KxK matrix, correlation matrix.

Output y Lx1 vector, $ln Pr(X < x \mid r)$.

Remarks You can pass more than one set of abscissae at a time; each column of x is

treated separately.

Source lncdfn.src

See also cdfmvn, lncdfbvn

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lncdfn

lncdfn

Purpose Computes natural log of Normal cumulative distribution function.

Format y = lncdfn(x);

Input *x* NxK matrix, abscissae.

Output y NxK matrix, ln Pr(X < x).

Source lncdfn.src

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lncdfn2

lncdfn2

a **Purpose** h **Format** d Input **Output** h Remarks k m Example n 0 p q Source See also u

```
Computes natural log of interval of Normal cumulative distribution
function.
y = lncdfn2(x,r);
        MxN matrix, abscissae.
х
        KxL matrix, ExE conformable with x, intervals.
r
        max(M,K) \times max(N,L) matrix, the log of the integral from x to
       x+dx of the Normal distribution, i.e., ln Pr(x < X < x + dx).
The relative error is:
    |x| \le 1 and dx \le 1
                                       \pm 1e-14
    1 < |x| < 37 and |dx| < 1|x|
                                      \pm 1e-13
    min(x,x + dx) > -37 and y > -690
                                       \pm 1e-11 or better
A relative error of \pm 1e-14 implies that the answer is accurate to better
than \pm 1 in the 14th digit.
print lncdfn2(-10,29);
    -7.6198530241605269e-24
print lncdfn2(0,1);
    -1.0748623268620716e+00
print lncdfn2(5,1);
    -1.5068446096529453e+01
```

lncdfn.src

cdfn2

V

W

x y z

lncdfnc

lncdfnc

Purpose Computes natural log of complement of Normal cumulative distribution function.

Format y = lncdfnc(x);

Input *x* NxK matrix, abscissae.

Output y NxK matrix, ln(1 - Pr(X < x)).

Source lncdfn.src

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1nfact

lnfact

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Purpose Computes the natural log of the factorial function and can be used to compute log gamma.

Format y = lnfact(x);

Input x NxK matrix, all elements must be positive.

Output y NxK matrix containing the natural log of the factorial of each of the elements of x.

Remarks For integer x, this is (approximately) ln(x!). However, the computation is done using a formula, and the function is defined for noninteger x.

In most formulae in which the factorial operator appears, it is possible to avoid computing the factorial directly, and to use **lnfact** instead. The advantage of this is that **lnfact** does not have the overflow problems that the factorial (!) operator has.

For x >= 1, this function has at least 6 digit accuracy, for x > 4 it has at least 9 digit accuracy, and for x > 10 it has at least 12 digit accuracy. For 0 < x < 1, accuracy is not known completely but is probably at least 6 digits.

Sometimes log gamma is required instead of log factorial. These functions are related by:

lngamma(x) = lnfact(x-1);

Example let x = 100 500 1000;

y = lnfact(x);

363.739375560

y = 2611.33045846

5912.12817849

Source lnfact.src

See also gamma

lnfact

Technical Notes

For x > 1, Stirling's formula is used.

For 0 < x <= 1, ln(gamma(x+1)) is used.

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lnpdfn

lnpdfn

Purpose Computes standard Normal log-probabilities.

Format z = lnpdfn(x);

Input x NxK matrix, data.

Output *z* NxK matrix, log-probabilities.

Remarks This computes the log of the scalar Normal density function for each element of x. z could be computed by the following GAUSS code:

 $y = -\ln(\text{sqrt}(2*\text{pi})) - x.*x/2;$

For multivariate log-probabilities, see **lnpdfmvn**.

Example $x = \{ .2, -1, 0, 1, 2 \};$

z = lnpdfn(x);

-2.9189385

-1.4189385

z = -0.91893853

-1.4189385

-2.9189385

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x y z

lnpdfmvn

lnpdfmvn

Purpose Computes multivariate Normal log-probabilities.

Format z = lnpdfmvn(x,s);

Input x NxK matrix, data.

s KxK matrix, covariance matrix.

Output z Nx1 vector, log-probabilities.

Remarks This computes the multivariate Normal log-probability for each row of *x*.

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lnpdfmvt

lnpdfmvt

Purpose Computes multivariate Student's t log-probabilities.

Format z = lnpdfmvt(x, s, nu);

Input x NxK matrix, data.

s KxK matrix, covariance matrix.

nu scalar, degrees of freedom.

Output z Nx1 vector, log-probabilities.

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lnpdft

lnpdft

Purpose Computes Student's t log-probabilities.

Format z = lnpdft(x, nu);

Input x NxK matrix, data.

nu scalar, degrees of freedom.

Output *z* NxK matrix, log-probabilities.

Remarks This does not compute the log of the joint Student's t pdf. Instead, the

scalar Normal density function is computed element by element.

For multivariate probabilities with covariance matrix see lnpdfmvt.

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load, loadf, loadk, loadm, loadp, loads

load, loadf, loadk, loadm, loadp, loads

Purpose Loads from a disk file.

Format load [path=path] x, y[]=filename, z=filename;

Remarks All the **load***xx* commands use the same syntax — they only differ in the types of symbols you use them with.

load, loadm matrix
loads string
loadf function

loadf function (fn)

loadk keyword (keyword)
loadp procedure (proc)

If no filename is given as with *x* above, then the symbol name the file is to be loaded into is used as the filename and the proper extension is added.

If more than one item is to be loaded in a single statement, the names should be separated by commas.

The filename can be either a literal or a string. If the filename is in a string variable, then the ^ (caret) operator must precede the name of the string, as in:

```
filestr = "mydata/char";
loadm x = ^filestr;
```

If no extension is supplied, the proper extension for each type of file will be used automatically as follows:

load .fmt - matrix file or delimited ASCII file
loadm .fmt - matrix file or delimited ASCII file
loads .fst - string file
loadf .fcg - user-defined function (fn) file
loadk .fcg - user-defined keyword (keyword) file
loadp .fcg - user-defined procedure (proc) file

These commands also signal to the compiler what type of object the symbol is so that later references to it will be compiled correctly.

A dummy definition must exist in the program for each symbol that is loaded in using loadf, loadk, or loadp. This resolves the need to

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load, loadf, loadk, loadm, loadp, loads

have the symbol initialized at compile time. When the load executes, the dummy definition will be replaced with the saved definition.

```
proc corrmat; endp;
loadp corrmat;
y = corrmat;

keyword regress(x); endp;
loadk regress;
regress x on y z t from data01;

fn sqrd=;
loadf sqrd;
y = sqrd(4.5);
```

To load GAUSS files created with the **save** command, no brackets are used with the symbol name.

If you use **save** to save a scalar error code 65535 (i.e., **error**(65535)), it will be interpreted as an empty matrix when you **load** it again.

ASCII data files

To load ASCII data files, square brackets follow the name of the symbol.

Numbers in ASCII files must be delimited with spaces, commas, tabs, or newlines. If the size of the matrix to be loaded is not explicitly given, as in:

```
load x[] = data.asc;
```

GAUSS will load as many elements as possible from the file and create an Nx1 matrix. This is the preferred method of loading ASCII data from a file, especially when you want to verify if the load was successful. Your program can then see how many elements were actually loaded by testing the matrix with the **rows** command, and if that is correct, the Nx1 matrix can be reshaped to the desired form. You could, for instance, put the number of rows and columns of the matrix right in the file as the first and second elements and reshape the remainder of the vector to the desired form using those values.

load, loadf, loadk, loadm, loadp, loads

If the size of the matrix is explicitly given in the **load** command, then no checking will be done. If you use:

load
$$x[500,6] = data.asc;$$

GAUSS will still load as many elements as possible from the file into an Nx1 matrix and then automatically reshape it using the dimensions given.

If your file contains nine numbers (1 2 3 4 5 6 7 8 9), then the matrix *x* that was created would be as follows:

load
$$x[1,9] = data.asc;$$

$$x = 123456789$$

load x[3,3] = data.asc;

$$x = 456$$

7 8 9

load x[2,2] = data.asc;

$$x = \begin{array}{c} 1 & 2 \\ 3 & 4 \end{array}$$

load x[2,9] = data.asc;

$$x = \begin{array}{rrrr} 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 \\ 1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 & 9 \end{array}$$

load x[3,5] = data.asc;

$$x = 67891$$

2 3 4 5 6

load accepts pathnames. The following is legal:

$$loadm k = /gauss/x;$$

This will load /gauss/x.fmt into \mathbf{k} .

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x y z

load, loadf, loadk, loadm, loadp, loads

If the **path=** subcommand is used with **load** and **save**, the path string will be remembered until changed in a subsequent command. This path will be used whenever none is specified. There are four separate paths for:

- 1. load, loadm
- 2. loadf, loadp
- 3. loads
- 4. save

Setting any of the four paths will not affect the others. The current path settings can be obtained (and changed) with the **sysstate** function, cases 4-7.

```
loadm path = /data;
```

This will change the **loadm** path without loading anything.

```
load path = /gauss x,y,z;
```

This will load x.fmt, 0y.fmt, and z.fmt using /gauss as a path. This path will be used for the next load if none is specified.

The load path or save path can be overridden in any particular load or save by putting an explicit path on the filename given to load from or save to as follows:

```
loadm path = /miscdata;
loadm x = /data/mydata1, y, z = hisdata;
```

In the above program:

```
/data/mydata1.fmt would be loaded into a matrix called x.
/miscdata/y.fmt would be loaded into a matrix called y.
/miscdata/hisdata.fmt would be loaded into a matrix called z.
oldmpath = sysstate(5,"/data");
load x, y;
```

This will get the old **loadm** path, set it to /data, load x.fmt and y.fmt, and reset the **loadm** path to its original setting.

See also loadd, save, let, con, cons, sysstate

call sysstate(5,oldmpath);

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loadd

loadd

a b d e h k 1 m n

0 p q

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V

W

x y z

Purpose Loads a data set. **Format** y = loadd(dataset);Input string, name of data set. dataset **Output** NxK matrix of data. **Remarks** The data set must not be larger than a single GAUSS matrix. If dataset is a null string or 0, the data set temp. dat will be loaded. To load a matrix file, use an .fmt extension on dataset. Source saveload.src **Globals** maxvec

loadwind

loadwind

Purpose Loads a previously saved graphic panel configuration.

Library pgraph

Format err = loadwind(namestr);

Input *namestr* string, name of file to be loaded.

Output *err* scalar, 0 if successful, 1 if graphic panel matrix is invalid. Note

that the current graphic panel configuration will be overwritten

in either case.

Source pwindow.src

See also savewind

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local

local

Purpose Declares variables that are to exist only inside a procedure.

Format local x, y, f:proc;

Remarks The statement above would place the names x, y, and f in the local symbol table for the current procedure being compiled. This statement is legal only between the **proc** statement and the **endp** statement of a procedure definition.

These symbols cannot be accessed outside of the procedure.

The symbol *f* in the example above will be treated as a procedure whenever it is accessed in the procedure. What is actually passed in is a pointer to a procedure.

See "Procedures and Keywords" in the *User's Guide*.

See also proc

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0

p q

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u v

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x y z

3-510

locate

locate

Purpose Positions the cursor in the window.

Format locate m, n;

Portability Windows only

Locates the cursor in the current output window.

Remarks *m* and *n* denote the row and column, respectively, at which the cursor is to be located.

The origin (1,1) is the upper left corner.

m and n may be any expressions that return scalars. Nonintegers will be truncated to an integer.

traneated to an intege

Example r = csrlin;

c = csrcol;

cls;

locate r,c;

In this example the window is cleared without affecting the cursor

position.

See also csrlin, csrcol

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j k

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loess

loess

a Computes coefficients of locally weighted regression. **Purpose** b { yhat, ys, xs } = loess(depvar, indvars); **Format** Input Nx1 vector, dependent variable. d depvar indvars NxK matrix, independent variables. е **Global Input** loess Span scalar, degree of smoothing. Must be greater than 2 / N. Default = .67777. g **loess NumEval** scalar, number of points in ys and xs. Default = 50. scalar, if 2, quadratic fit, otherwise linear. Default loess Degree h = 1.loess WgtType scalar, type of weights. If 1, robust, symmetric weights, otherwise Gaussian. Default = 1. scalar, if 1, iteration information and results are output printed, otherwise nothing is printed. k 1 **Output** yhat Nx1 vector, predicted *depvar* given *indvars*. loess numEvalx1 vector, ordinate values given abscissae VS m values in xs. n loess numEvalX1 vector, equally spaced abscissae values. xs0 Remarks Based on Cleveland, William S. "Robust Locally Weighted Regression and Smoothing Scatterplots." JASA. Vol. 74, 1979, 829-36. p Source q loess.src

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W

log

Purpose Computes the \log_{10} of all elements of a matrix.

Format $y = \log(x)$;

Input x NxK matrix.

Output y NxK matrix containing the log 10 values of the elements of x.

Remarks log is defined for $x \neq 0$.

If *x* is negative, complex results are returned.

You can turn the generation of complex numbers for negative inputs on or off in the GAUSS configuration file, and with the **sysstate** function, case 8. If you turn it off, **log** will generate an error for negative inputs.

If x is already complex, the complex number state doesn't matter; \log will compute a complex result.

x can be any expression that returns a matrix.

Example

```
x = round(rndu(3,3)*10+1);
y = log(x);
```

7.0000000000 2.0000000000 6.0000000000

0.60205999 0.30103 0.30103

1.0000000000 0.60205999 0.90308999 0.8450980400 0.30103 0.77815125

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i j

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n m

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loglog

loglog

Purpose Graphs X vs. Y using log coordinates.

Library pgraph

Format loglog(x,y);

Input x Nx1 or NxM matrix. Each column contains the X values for a particular line.

y Nx1 or NxM matrix. Each column contains the Y values for a particular line.

Source ploglog.src

See also xy, logy, logx

хух

3-514

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p q

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logx

logx

Purpose Graphs X vs. Y using log coordinates for the X axis.

Library pgraph

Format logx(x,y);

Input x Nx1 or NxM matrix. Each column contains the X values for a particular line.

y Nx1 or NxM matrix. Each column contains the Y values for a particular line.

Source plogx.src

See also xy, logy, loglog

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logy

logy

Purpose Graphs X vs. Y using log coordinates for the Y axis.

Library pgraph

Format logy(x,y);

Input Nx1 or NxM matrix. Each column represents the X values for a х particular line.

> Nx1 or NxM matrix. Each column represents the Y values for a y particular line.

Source plogy.src

See also xy, logx, loglog

3-516

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loopnextindex

loopnextindex

Purpose

Increments an index vector to the next logical index and jumps to the specified label if the index did not wrap to the beginning.

Format

loopnextindex lab,i,o [,dim];

Input

label to jump to if loopnextindex succeeds.

i Mx1 vector of indices into an array, where $M \le N$.

Nx1 vector of orders of an N-dimensional array.

dim scalar [1-M], index into the vector of indices *i*, corresponding to the dimension to walk through, positive to walk the index forward, or negative to walk backward.

Remarks

If the argument *dim* is given, **loopnextindex** will walk through only the dimension indicated by *dim* in the specified direction. Otherwise, if *dim* is not given, each call to **loopnextindex** will increment *ind* to index the next element or subarray of the corresponding array.

loopnextindex will jump to the label indicated by *lab* if the index can walk further in the specified dimension and direction, otherwise it will fall out of the loop and continue through the program.

When the index matches the vector of orders, the index will be reset to the beginning and program execution will resume at the statement following the **loopnextindex** statement.

Example

```
orders = { 2,3,4,5,6,7 };
a = arrayalloc(orders,0);
ind = { 1,1,1,1 };
loopni:
    setarray a, ind, rndn(6,7);
    loopnextindex loopni, ind, orders;
```

This example sets each 6x7 subarray of array a, by incrementing the index at each call of **loopnextindex** and then going to the label *loopni*. When *ind* cannot be incremented, the program drops out of the loop and continues.

ind =
$$\{1,1,4,5\};$$

a

b

c d

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i

k

n l

n

0

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...

u v

117

loopnextindex

```
loopni2:
```

```
setarray a, ind, rndn(6,7);
loopnextindex loopni2, ind, orders, 2;
```

Using the array and vector of orders from the example above, this example increments the second value of the index vector *ind* during each call to **loopnextindex**. This loop will set the 6x7 subarrays of a that begin at [1,1,4,5,1,1], [1,2,4,5,1,1], and [1,3,4,5,1,1], and then drop out of the loop.

See also nextindex, previousindex, walkindex

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lower

lower

Purpose Converts a string or character matrix to lowercase. **Format** y = lower(x);Input x string or NxK matrix of character data to be converted to lowercase. **Output** string or NxK matrix which contains the lowercase equivalent of the data in x. Remarks If x is a numeric matrix, y will contain garbage. No error message will be generated since GAUSS does not distinguish between numeric and character data in matrices. **Example** x = "MATH 401"; y = lower(x);print y; produces: math 401 See also upper

a

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V

Purpose

Output

lowmat, lowmat1

b

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m

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0 p

q

u

W

x y z

Source V

diag.src

See also

upmat, upmat1, diag, diagrv, crout, croutp

Format L = lowmat(x);

L = lowmat1(x);

main diagonal with ones.

Input NxN matrix. \boldsymbol{x}

> LNxN matrix containing the lower elements of the matrix. The upper elements are replaced with zeros. lowmat returns the main diagonal intact. **lowmat1** replaces the main diagonal with

Returns the lower portion of a matrix. **lowmat** returns the main diagonal

and every element below. **lowmat1** is the same except it replaces the

ones.

Example $x = \{ 1 \ 2 \ -1,$

> $2 \quad 3 \quad -2$ 1 -2 1 };

L = lowmat(x);

L1 = lowmat1(x);

The resulting matrices are:

1 0 0

 $L = 2 \quad 3 \quad 0$

1 - 2 1

L1 = 2 10

1 - 2 1

lpos

Purpose

Returns the current position of the print head within the printer buffer for the printer.

Format

y = lpos;

Remarks

This function is basically equivalent to function **csrcol** but this returns the current column position for the standard printer.

The value returned is the column position of the next character to be printed to the printer buffer. This does not necessarily reflect the actual physical position of the print head at the time of the call.

If this function returns a number greater than 1, there are characters in the buffer for the standard printer which have not yet been sent to the printer. This buffer can be flushed at any time by lprint'ing a carriage return/line feed sequence, or a form feed character.

Example

```
if lpos > 60;
    lprint;
endif;
```

In this example, if the print buffer contains 60 characters or more, a carriage return/line feed sequence will be printed.

See also

lprint, lpwidth

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lprint

lprint

Remarks

1

Purpose Controls printing to the line printer.

Format lprint [/typ] [/fmted] [/mf] [/jnt] [list of expressions separated by spaces] [;];

This function was originally written for line printers. It is still supported for backwards compatibility purposes, but if you're using a page-oriented printer (such as a laser or inkjet printer), it may not give you the results you're expecting.

lprint statements work in essentially the same way that **print** statements work. The main difference is that **lprint** statements cannot be directed to the auxiliary output. Also, the **locate** statement has no meaning with **lprint**.

Two semicolons following an **lprint** statement will suppress the final line feed.

See **print** for information on /typ, /fmted, /mf, and /jnt.

A list of expressions is a list of GAUSS expressions, separated by spaces. In lprint statements, because a space is the delimiter between expressions, no spaces are allowed inside expressions unless they are within index brackets, they are in quotes, or the whole expression is in parentheses.

Printer width can be specified by the **lpwidth** statement:

```
lpwidth 132;
```

This statement remains in effect until cancelled. The default printer width is 80. That is, GAUSS automatically sends a line feed to the printer after printing 80 characters.

lpos can be used to determine the (column) position of the next character that will be printed in the buffer.

An lprint statement by itself will cause a blank line to be printed:

```
lprint;
```

The printing of special characters is accomplished by the use of the backslash (\) within double quotes. The options are:

"\b"	backspace (ASCII 8)
"\e"	escape (ASCII 27)

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lprint

"\f" form feed (ASCII 12)
"\1" line feed (ASCII 10)
"\r" carriage return (ASCII 13)
" t" tab (ASCII 9)
"\###" the character whose ASCII value is "###" (decimal)

GAUSS also has an automatic line printer mode which causes the results of all global assignment statements to be printed out on the printer. This is controlled by the lprint on and lprint off commands. (See lprint on, lprint off.)

Example lprint 3*4 5+2;

See also print, lprint on, lpos, lpwidth, format

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lpwidth

lpwidth

Purpose Specifies the width of the printer.

Format lpwidth n;

Remarks *n* is a scalar which specifies the width of the printer in columns (characters). That is, after printing *n* characters on a line, GAUSS will send a carriage return and a line feed, so that the print head will move to the beginning of the next line.

If a matrix is being printed, the line feed sequence will always be inserted between separate elements of the matrix rather than being inserted between digits of a single element.

n may be any scalar-valued expression. Nonintegers will be truncated to an integer.

The default is 80 columns.

Note: This does not send control characters to the printer to automatically switch the mode of the printer to a different character pitch because each printer is different. This only controls the frequency of carriage return/line feed sequences.

Example lpwidth 132;

This statement will change the printer width to 132 columns.

See also lprint, lpos, outwidth

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ltrisol

ltrisol

Purpose Computes the solution of Lx = b where L is a lower triangular matrix.

Format x = ltrisol(b, L);

Input b PxK matrix.

L PxP lower triangular matrix.

Output x PxK matrix.

ltrisol applies a forward solve to Lx = b to solve for x. If b has more than one column, each column will be solved for separately, i.e.,

ltrisol will apply a forward solve to L * x[.,i] = b[.,i].

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lu

a h d h k 1 m 0 p q t u V W

```
Purpose
              Computes the LU decomposition of a square matrix with partial (row)
              pivoting, such that X = L\bar{U}.
  Format
              \{l,u\}=lu(x);
    Input
                     NxN square nonsingular matrix.
  Output
              1
                     NxN "scrambled" lower triangular matrix. This is a lower
                     triangular matrix that has been reordered based on the row
                     pivoting.
                     NxN upper triangular matrix.
              и
Example
              rndseed 13;
              format /rd 10,4;
              x = complex(rndn(3,3),rndn(3,3));
              \{ 1,u \} = lu(x);
              x2 = 1*11;
                      0.1523 + 0.7685i -0.8957 + 0.0342i 2.4353 + 2.7736i
              x = -1.1953 + 1.2187i 1.2118 + 0.2571i - 0.0446 - 1.7768i
                                         1.2950 - 1.6929i 1.6267 + 0.2844i
                      0.8038 + 1.3668i
                   0.2589 - 0.3789i -1.2417 + 0.5225i
                                                          1.0000
              1 =
                             1.0000
                                                          0.0000
                                                0.0000
                   0.2419 - 0.8968i
                                                 1.0000
                                                          0.0000
                     -1.1953 + 1.2187i \ 1.2118 + 0.2571i
                                                         -0.0446 - 1.7768i
               u =
                                0.0000 \ 0.7713 - 0.6683i \ 3.2309 + 0.6742i
                                0.0000
                                                 0.0000
                                                           6.7795 + 5.7420i
```

lu

$$x^{2} = -1.1953 + 1.2187i - 1.2118 + 0.2571i - 0.0446 - 1.7768i$$

$$0.8038 + 1.3668i - 1.2950 - 1.6929i - 1.6267 + 0.2844i$$

See also crout, croutp, chol

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> : 1

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lusol

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V

W

lusol

Purpose Computes the solution of LUx = b where L is a lower triangular matrix and U is an upper triangular matrix.

Format x = lusol(b, L, U);

Input b PxK matrix.

L PxP lower triangular matrix.

U PxP upper triangular matrix.

Output x PxK matrix.

Remarks If *b* has more than one column, each column is solved for separately, i.e., lusol solves LUx[.,i] = b[.,i].

machEpsilon

machEpsilon

Purpose Returns the smallest number such that 1 + eps > 1.

Format eps = machEpsilon;

Output *eps* scalar, machine epsilon.

Source machconst.src

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make (dataloop)

a

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d

h

k

m

0

p

make (dataloop)

Purpose Specifies the creation of a new variable within a data loop.

Format make [#] numvar = numeric_expression;
make \$ charvar = character expression;

Remarks A *numeric_expression* is any valid expression returning a numeric vector. A *character_expression* is any valid expression returning a character

vector. If neither '\$' nor '#' is specified, '#' is assumed.

The expression may contain explicit variable names and/or GAUSS commands. Any variables referenced must already exist, either as elements of the source data set, as **externs**, or as the result of a previous **make**, **vector**, or **code** statement. The variable name must be unique. A variable cannot be made more than once, or an error is generated.

Example make sqvpt = sqrt(velocity * pressure * temp);

make \$ sex = lower(sex);

See also vector

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V

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3-530

makevars

makevars

Purpose Creates separate global vectors from the columns of a matrix.

Format makevars(x, vnames, xnames);

Input x NxK matrix whose columns will be converted into individual vectors.

vnames string or Mx1 character vector containing names of global vectors to create. If 0, all names in xnames will be used.

xnames string or Kx1 character vector containing names to be

associated with the columns of the matrix x.

Remarks

If xnames = 0, the prefix X will be used to create names. Therefore, if there are 9 columns in x, the names will be X1-X9, if there are 10, they will be X01-X10, and so on.

If *xnames* or *vnames* is a string, the individual names must be separated by spaces or commas.

vnames = "age pay sex";

Since these new vectors are created at execution time, the compiler will not know they exist until after **makevars** has executed once. This means that you cannot access them by name unless you previously **clear** them or otherwise add them to the symbol table. (See **setvars** for a quick interactive solution to this.)

This function is the opposite of mergevar.

Example

Two global vectors, called age and pay, are created from the columns of x.

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makevars

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V

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хух

This is the same as the example above, except that strings are used for the variable names.

Source vars.src

Globals __vpad

See also mergevar, setvars

makewind

makewind

Purpose Creates a graphic panel of specific size and position and add it to the list

of graphic panels.

Library pgraph

Format makewind(xsize, ysize, xshft, yshft, typ);

Input *xsize* scalar, horizontal size of the graphic panel in inches.

ysize scalar, vertical size of the graphic panel in inches.

xshft scalar, horizontal distance from left edge of window in inches.

yshft scalar, vertical distance from bottom edge of window in inches.

typ scalar, graphic panel attribute type. If this value is 1, the graphic

panels will be transparent. If 0, the graphic panels will be

nontransparent.

Remarks Note that if this procedure is used when rotating the page, the passed

parameters are scaled appropriately to the newly oriented page. The size and shift values will not be true inches when printed, but the graphic panel size to page size ratio remains the same. The result of this implementation automates the rotation and eliminates the required

graphic panel recalculations by the user.

See the **window** command for creating tiled graphic panels. For more information on using graphic panels, see "Publication Quality Graphics"

in the *User's Guide*.

Source pwindow.src

See also window, endwind, setwind, getwind, begwind,

nextwind

3-533

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margin

margin

a **Purpose** Sets the margins for the current graph graphic panel. b Library pgraph Format margin(l,r,t,b); d Input scalar, the left margin in inches. scalar, the right margin in inches. rt scalar, the top margin in inches. scalar, the bottom margin in inches. h h Remarks By default, the dimensions of the graph are the same as the graphic panel dimensions. With this function the graph dimensions may be decreased. The result will be a smaller plot area surrounded by the specified margin. This procedure takes into consideration the axes labels and numbers for correct placement. k All input inch values for this procedure are based on a full size window of 9 x 6.855 inches. If this procedure is used with a graphic panel, the values will be scaled to window inches automatically. m If the axes must be placed an exact distance from the edge of the page, axmargin should be used. 0 Source pgraph.src p See also axmargin

u

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matalloc

matalloc

Purpose Allocates a matrix with unspecified contents.

Format y = matalloc(r,c);

Input r scalar, rows.

c scalar, columns.

Output y = rxc matrix.

Remarks The contents are unspecified. This function is used to allocate a matrix

that will be written to in sections using indexing or used with the Foreign Language Interface as an output matrix for a function called with

dllcall().

See also matinit, ones, zeros, eye

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matinit

matinit

Purpose Allocates a matrix with a specified fill value.

Format y = matinit(r,c,v);

Input r scalar, rows.

c scalar, columns.

v scalar, value to initialize.

Output y rxc matrix with each element equal to the value of v.

See also matalloc, ones, zeros, eye

3-536

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mattoarray

mattoarray

Purpose Changes a matrix to a type array. **Format** y = mattoarray(x);Input matrix. X Output 1-or-2-dimensional array. If the argument *x* is a scalar, **arraytomat** will simply return the scalar, Remarks without changing it to a type array. Example x = 5*ones(2,3);y = mattoarray(x);y will be a 2x3 array of fives. See also arraytomat

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maxc

maxc

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Purpose

Returns a column vector containing the largest element in each column of a matrix.

Format

$$y = \max(x)$$
;

Input

x NxK matrix.

Output

Kx1 matrix containing the largest element in each column of x.

Remarks

If x is complex, **maxc** uses the complex modulus (**abs**(x)) to determine the largest elements.

To find the maximum elements in each row of a matrix, transpose the matrix before applying the **maxc** function.

To find the maximum value in the whole matrix if the matrix has more than one column, nest two calls to maxe:

$$y = maxc(maxc(x));$$

Example

$$x = rndn(4,2);$$

$$y = maxc(x);$$

$$x = \begin{array}{c} 0.348110 & 1.172391 \\ -0.027064 & 0.796867 \end{array}$$

$$1.421940 - 0.351313$$

$$y = \frac{1.421940}{1.376765}$$

See also

minc, maxindc, minindc

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maxindc

maxindc

Purpose

Returns a column vector containing the index (i.e., row number) of the maximum element in each column in a matrix.

Format

$$y = maxindc(x);$$

Input

x NxK matrix.

Output

y Kx1 matrix containing the index of the maximum element in each column of x.

Remarks

If x is complex, maxc uses the complex modulus (abs(x)) to determine the largest elements.

To find the index of the maximum element in each row of a matrix, transpose the matrix before applying **maxindc**.

If there are two or more "largest" elements in a column (i.e., two or more elements equal to each other and greater than all other elements), then **maxindc** returns the index of the first one found, which will be the smallest index.

Example

$$x = round(rndn(4,4)*5);$$

$$y = maxc(x);$$

$$z = maxindc(x);$$

$$x = \begin{array}{rrrr} 1 & -11 & 0 & 5 \\ 0 & 0 & -2 & -6 \\ -8 & 0 & 3 & 2 \\ -11 & 5 & -4 & 5 \end{array}$$

$$y = \begin{array}{c} 1 \\ 5 \\ 3 \\ 5 \end{array}$$

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maxindc

 $z = \frac{2}{3}$

See also maxc, minindc, minc

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maxvec

maxvec

Purpose Returns maximum vector length allowed.

Format y = maxvec;

Global Input __maxvec scalar, maximum vector length allowed.

Output y scalar, maximum vector length.

Remarks maxvec returns the value in the global scalar __maxvec, which can be reset in the calling program. This must never be set to 8190.

maxvec is called by Run-Time Library functions and applications when determining how many rows can be read from a data set in one call to **readr**.

On systems without virtual memory you can use 536870910. Otherwise a smaller value like 20000-30000 is necessary to prevent excessive disk thrashing. The trick is to allow the algorithm making the disk reads to execute entirely in RAM.

Example y = maxvec;

print y;

produces:

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Source system.src

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Purpose

Computes modified and exponentially scaled modified Bessels of the first kind of the nth order.

Format

```
y = mbesseli(x,n,alpha);
y = mbesseli0(x);
y = mbesseli1(x);

y = mbesselei(x,n,alpha);
y = mbesselei0(x);
y = mbesselei1(x);
```

Input

x Kx1 vector, abscissae.

n scalar, highest order. alpha scalar, $0 \le alpha < 1$.

Output

KxN matrix, evaluations of the modified Bessel or the exponentially scaled modified Bessel of the first kind of the nth orders.

Remarks

For the functions that permit you to specify the order, the returned matrix contains a sequence of modified or exponentially scaled modified Bessel values of different orders. For the ith row of y:

$$y[i,.] = I_{\alpha}(x[i]) I_{\alpha+1}(x[i]) ... I_{\alpha+n-1}(x[i])$$

The remaining functions generate modified Bessels of only the specified order.

The exponentially scaled modified Bessels are related to the unscaled modifed Bessels in the following way:

$$mbesselei0(x) = exp(-x) * mbesseli0(x)$$

The use of the scaled versions of the modified Bessel can improve the numerical properties of some calculations by keeping the intermediate numbers small in size.

Example

This example produces estimates for the "circular" response regression model (Fisher, N.I. *Statistical Analysis of Circular Data*. NY: Cambridge

mbesseli

University Press, 1993.), where the dependent variable varies between $-\pi$ and π in a circular manner. The model is

$$y = \mu + G(XB)$$

where B is a vector of regression coefficients, X a matrix of independent variables with a column of 1's included for a constant, and y a vector of "circular" dependent variables, and where G() is a function mapping XB onto the $[-\pi, \pi]$ interval.

The log-likelihood for this model is from Fisher, N.I. ... 1993, 159;

$$logL = -N x ln((I_0(\kappa)) + \kappa) \sum_{i}^{N} cos(y_i - \mu - G(X_i B))$$

To generate estimates it is necessary to maximize this function using an iterative method. **QNewton** is used here.

 κ is required to be nonnegative and therefore in the example below, the exponential of this parameter is estimated instead. Also, the exponentially scaled modified Bessel is used to improve numerical properties of the calculations.

The **arctan** function is used in G()to map XB to the $[-\pi, \pi]$ interval as suggested by Fisher, N.I. ... 1993, 158.

```
proc G(u);
    retp(2*atan(u));
endp;

proc lpr(b);
    local dev;
```

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```
/*
   ** b[1] - kappa
   ** b[2] - mu
   ** b[3] - constant
   ** b[4:rows(b)] - coefficients
   * /
     dev = y - b[2] - G(b[3] + x * b[4:rows(b)]);
     retp(rows(dev)*ln(mbesselei0(exp(b[1])) -
     sumc(exp(b[1])*(cos(dev)-1)));
   endp;
   loadm data;
  y0 = data[.,1];
  x0 = data[.,2:cols(data)];
  b0 = 2*ones(cols(x),1);
   { b,fct,grd,ret } = QNewton(&lpr,b0);
   cov = invpd(hessp(&lpr,b));
  print "estimates standard errors";
  print;
  print b~sqrt(diag(cov));
ribesl.src
```

хух

Source

meanc

meanc

Purpose Computes the mean of every column of a matrix.

Format y = meanc(x);

Input x NxK matrix.

Output y Kx1 matrix containing the mean of every column of x.

Example x = meanc(rndu(2000,4));

0.492446

 $x = \begin{array}{c} 0.503543 \\ 0.502905 \end{array}$

0.509283

In this example, 4 columns of uniform random numbers are generated in a matrix, and the mean is computed for each column.

See also stdc

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median

median

Purpose

Computes the medians of the columns of a matrix.

Format m = median(x);

Input x NxK matrix.

Output m Kx1 vector containing the medians of the respective columns of x.

Example $x = \{ 84, 68, 37 \};$ y = median(x);

 $y = \begin{array}{c} 6.0000000 \\ 7.0000000 \end{array}$

Source median.src

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mergeby

mergeby

Purpose Merges two sorted files by a common variable.

Format mergeby(infile1, infile2, outfile, keytyp);

Input *infile1* string, name of input file 1.

infile2 string, name of input file 2. outfile string, name of output file.

keytyp scalar, data type of key variable.

1 numeric

2 character

Remarks

This will combine the variables in the two files to create a single large file. The following assumptions hold:

- 1. Both files have the key variable in the first column.
- 2. All of the values of the key variable within a file are unique.
- 3. Each file is already sorted on that variable.

The output file will contain the key variable in its first column.

It is not necessary for the two files to have the same number of rows. For each row for which the key variables match, a row will be created in the output file. *outfile* will contain the columns from *infile1* followed by the columns of *infile2* minus the key column from the second file.

If the inputs are null or 0, the procedure will ask for them.

Example mergeby("freq", "freqdata", "mergex",1);

Source sortd.src

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mergevar

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x y z

mergevar

Accepts a list of names of global matrices, and concatenates the **Purpose** corresponding matrices horizontally to form a single matrix. **Format** x = mergevar(vnames);Input string or Kx1 column vector containing the names of K global vnames matrices. **Output** NXM matrix that contains the concatenated matrices, where M is the sum of the columns in the K matrices specified in *vnames*. Remarks The matrices specified in *vnames* must be globals and they must all have the same number of rows. This function is the opposite of **makevars**. **Example** let vnames = age pay sex; x = mergevar(vnames);The matrices age, pay, and sex will be concatenated horizontally to create x. Source vars.src See also makevars

minc

Purpose Returns a column vector containing the smallest element in each column in a matrix.

Format $y = \min(x)$;

Input x NxK matrix.

Output y Kx1 matrix containing the smallest element in each column of x.

Remarks If x is complex, minc uses the complex modulus (abs(x)) to determine the smallest elements.

To find the minimum element in each row, transpose the matrix before applying the **minc** function.

To find the minimum value in the whole matrix, nest two calls to minc:

$$y = minc(minc(x));$$

Example x = rndn(4,2);

y = minc(x);

-1.061321 -0.729026

 $x = \begin{array}{rrr} -0.021965 & 0.184246 \\ 1.843242 & -1.847015 \end{array}$

1.977621 -0.532307

 $y = \begin{array}{r} -1.061321 \\ -1.847015 \end{array}$

See also maxc, minindc, maxindc

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Purpose

Returns a column vector containing the index (i.e., row number) of the smallest element in each column in a matrix.

Format

y = minindc(x);

Input

NxK matrix.

Output

Kx1 matrix containing the index of the smallest element in each column of x.

Remarks

If x is complex, minindc uses the complex modulus (abs(x)) to determine the smallest elements.

To find the index of the smallest element in each row, transpose the matrix before applying minindc.

If there are two or more "smallest" elements in a column (i.e., two or more elements equal to each other and less than all other elements), then minindc returns the index of the first one found, which will be the smallest index.

Example

x = round(rndn(5,4)*5);

y = minc(x);

z = minindc(x);

$$-5$$
 6 -4 -1

$$x = \begin{bmatrix} 2 & -2 & 1 & 3 \\ 6 & 0 & 1 & -7 \end{bmatrix}$$

$$-6 \quad 0 \quad 8 \quad -4$$

$$-6$$

$$y =$$

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minindc

$$x = \begin{cases} 4 \\ 5 \\ 1 \\ 3 \end{cases}$$

See also maxindc, minc, maxc

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miss, missrv

miss, missrv

Purpose

miss converts specified elements in a matrix to GAUSS's missing value code. missrv is the reverse of this, and converts missing values into specified values.

Format

y = miss(x,v);y = missrv(x,v);

Input

x NxK matrix.

v LxM matrix, ExE conformable with x.

Output

 $\max(N,L)$ by $\max(K,M)$ matrix.

Remarks

For **miss**, elements in x that are equal to the corresponding elements in v will be replaced with the GAUSS missing value code.

For **missrv**, elements in *x* that are equal to the GAUSS missing value code will be replaced with the corresponding element of *v*.

For complex matrices, the missing value code is defined as a missing value entry in the real part of the matrix. For complex x, then, **miss** replaces elements with a ". + 0i" value, and **missrv** examines only the real part of x for missing values. If, for example, an element of x = 1 + i, **missrv** will not replace it.

These functions act like element-by-element operators. If v is a scalar, for instance -1, then all -1's in x are converted to missing. If v is a row (column) vector with the same number of columns (rows) as x, then each column (row) in x is transformed to missings according to the corresponding element in v. If v is a matrix of the same size as x, then the transformation is done corresponding element by corresponding element.

Missing values are given special treatment in the following functions and operators: b/a (matrix division when a is not square and neither a nor b is scalar), counts, ismiss, maxc, maxindc, minc, minindc, miss, missex, missry, moment, packr, scalmiss, sortc.

As long as you know a matrix contains no missings to begin with, **miss** and **missrv** can be used to convert one set of numbers into another. For example:

```
y=missrv(miss(x,0),1);
```

will convert 0's to 1's.

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miss, missrv

Example $v = -1 \sim 4 \sim 5;$

y = miss(x,v);

If \mathbf{x} has 3 columns, all -1's in the first column will be changed to missings, along with all 4's in the second column and 5's in the third column.

See also counts, ismiss, maxc, maxindc, minc, minindc, missex, moment, packr, scalmiss, sortc

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Purpose

Converts numeric values to the missing value code according to the values given in a logical expression.

Format

y = missex(x,e);

Input

x NxK matrix.

e NxK logical matrix (matrix of 0's and 1's) that serves as a "mask" for x; the 1's in e correspond to the values in x that are to be converted into missing values.

Output

NXK matrix that equals x, but with those elements that correspond to the 1's in e converted to missing.

Remarks

The matrix e will usually be created by a logical expression. For instance, to convert all numbers between 10 and 15 in x to missing, the following code could be used:

```
y = missex(x, (x .> 10) .and (x .< 15));
```

Note that "dot" operators MUST be used in constructing the logical expressions.

For complex matrices, the missing value code is defined as a missing value entry in the real part of the matrix. For complex x, then, **missex** replaces elements with a ". + 0i" value.

This function is like **miss**, but is more general in that a range of values can be converted into missings.

Example

```
x = rndu(3,2);
/* logical expression */
e = (x .> .10) .and (x .< .20);
y = missex(x,e);</pre>
```

A 3x2 matrix of uniform random numbers is created. All values in the interval (0.10, 0.20) are converted to missing.

Source

datatran.src

See also

miss, missrv

moment

moment

Purpose Computes a cross-product matrix. This is the same as x'x.

Format y = moment(x,d);

Input x NxK matrix.

d scalar, controls handling of missing values.

- 0 missing values will not be checked for. This is the fastest option.
- "listwise deletion" is used. Any row that contains a missing value in any of its elements is excluded from the computation of the moment matrix. If every row in x contains missing values, then moment (x,1) will return a scalar zero.
- 2 "pairwise deletion" is used. Any element of *x* that is missing is excluded from the computation of the moment matrix. Note that this is seldom a satisfactory method of handling missing values, and special care must be taken in computing the relevant number of observations and degrees of freedom.

Output y KxK matrix which equals x'x.

Remarks

The fact that the moment matrix is symmetric is taken into account to cut execution time almost in half.

If there is no missing data then d = 0 should be used because it will be faster.

The / operator (matrix division) will automatically form a moment matrix (performing pairwise deletions if trap 2 is set) and will compute the ols coefficients of a regression. However, it can only be used for data sets that are small enough to fit into a single matrix. In addition, the moment matrix and its inverse cannot be recovered if the / operator is used.

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moment

Example

```
xx = moment(x,2);
ixx = invpd(xx);
b = ixx*missrv(x,0)'y;
```

In this example, the regression of y on x is computed. The moment matrix **xx** is formed using the **moment** command (with pairwise deletion, since the second parameter is 2). Then **xx** is inverted using the **invpd** function. Finally, the **ols** coefficients are computed. **missrv** is used to emulate pairwise deletion by setting missing values to 0.

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momentd

Purpose	Computes a moment (X'X) matrix from a GAUSS data set.				
Format	<pre>m = momentd(dataset, vars);</pre>				
Input	dataset vars	string, name of data set. Kx1 character vector, names of variables. or Kx1 numeric vector, indices of columns.			
	any order. Defaults ar be ignored	These can be any size subset of the variables in the data set, and can be in any order. If a scalar 0 is passed, all columns of the data set will be used. Defaults are provided for the following global input variables so they can be ignored unless you need control over the other options provided by this procedure.			
Global Input	conmissrow	scalar, default 1. 1 a constant term will be added. 0 no constant term will be added. scalar, default 0. 0 there are no missing values (fastest). 1 do listwise deletion, drop an observation if any missings occur in it. 2 do pairwise deletion. This is equivalent to setting missings to 0 when calculating m. scalar, default 0, the number of rows to read per iteration of the read loop.			
Output	m	If 0, the number of rows will be calculated internally. If you get an Insufficient memory error message, or you want the rounding to be exactly the same between runs, you can set the number of rows to read before calling momentd . MxM matrix, where M = K +con, the moment matrix constructed by calculating X'X where X is the data, with or without a constant vector of ones. Error handling is controlled by the low order bit of the trap flag.			

momentd

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terminate with error message trap 0 trap 1 return scalar error code in m33 too many missings file not found 34 Example $z = \{ age, pay, sex \};$ m = momentd("freq",z); Source momentd.src Globals __con, __miss, __row

msym

Purpose Allows the user to set the symbol that GAUSS uses when missing values are converted to ASCII and vice versa.

Format msym str;

Input str literal or ^string (up to 8 letters) which, if not surrounded by

quotes, is forced to uppercase. This is the string to be printed for

missing values. The default is '.'.

Remarks The entire string will be printed out when converting to ASCII in print, lprint, and printfm statements.

When converting ASCII to binary in **loadm** and **let** statements, only the first character is significant. In other words,

msym HAT;

will cause 'H' to be converted to missing on input.

This does not affect writer which outputs data in binary format.

See also print, lprint, printfm

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nametype

nametype

Purpose Provides support for programs following the upper/lowercase convention in GAUSS data sets. (See "File I/O" in the User's Guide.) Returns a h vector of names of the correct case and a 1/0 vector of type information. **Format** { vname, vtype } = nametype(vname, vtype); d Input vname Nx1 character vector of variable names. vtype scalar or Nx1 vector of 1's and 0's to determine the type and therefore the case of the output vname. If this is scalar 0 or 1 it will be expanded to Nx1. If -1, nametype will assume that vname follows the upper/lowercase convention. h **Output** *vname* Nx1 character vector of variable names of the correct case. uppercase if numeric, lowercase if character. vtype Nx1 vector of ones and zeros, 1 if variable is numeric, 0 if character. Example vn = { age, pay, sex }; $vt = \{ 1, 1, 0 \};$ m { vn, vt } = nametype(vn, vt); print \$vn; 0 **AGE** p vn =PAY sex Source nametype.src Globals vartype u

V

W

x y z

new

Purpose

Erases everything in memory including the symbol table; closes all open files, the auxiliary output, and turns the window on if it was off; also allows the size of the new symbol table and the main program space to be specified.

Format

new [[nos [[, mps]]];

Input

nos scalar, which indicates the maximum number of global symbols allowed. See your platform supplement for the maximum number of globals allowed in this implementation.

mps scalar, which indicates the number of bytes of main program space to be allocated. See your platform supplement for the maximum amount allowed in this implementation.

Remarks

Procedures, user-defined functions, and global matrices strings and string arrays are all global symbols.

The main program space is the amount of memory available for nonprocedure, nonfunction program code.

This command can be used with arguments as the first statement in a program to clear the symbol table and to allocate only as much space for program code as your program actually needs. When used in this manner, the auxiliary output will not be closed. This will allow you to open the auxiliary output from command level and run a program without having to remove the **new** at the beginning of the program. If this command is not the first statement in your program, it will cause the program to terminate.

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new

```
Example
                                    /*
                                        clear global symbols. */
                      new;
                                    /*
                      new 300;
                                        clear global symbols, set */
                                    /*
                                        maximum number of global */
                                     /*
                                        symbols to 300, and leave */
h
                                    /*
                                        program space unchanged. */
d
                      new
                                     /*
                                        clear global symbols, set */
                      200,100000;
                                    /*
                                        maximum number of global */
                                    /*
                                        symbols to 200, and allocate */
                                    /*
                                        100000 bytes for main */
                                        program code. */
                                     /*
                                        clear global symbols, */
                       new ,100000;
h
                                        allocate 100000 bytes for */
                                    /*
                                        main program code, and leave */
                                    /*
                                    /*
                                        maximum number of globals */
                                    /*
                                        unchanged. */
k
           See also clear, delete, output
m
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nextindex

nextindex

Purpose Returns the index of the next element or subarray in an array.

Format ni = nextindex(i, o);

Input i Mx1 vector of indices into an array, where M<=N.

o Nx1 vector of orders of an N-dimensional array.

Output ni Mx1 vector of indices, the index of the next element or subarray in the array corresponding to o.

Remarks nextindex will return a scalar error code if the index cannot be incremented.

 $ind = \begin{pmatrix} 2 \\ 4 \\ 1 \end{pmatrix}$

In this example, **nextindex** incremented *ind* to index the next 6x7 subarray in array a.

See also previousindex, loopnextindex, walkindex

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nextn, nextnevn

nextn, nextnevn

a

Purpose Returns allowable matrix dimensions for computing FFT's.

Format

n = nextn(n0);

Input

n0 scalar, the length of a vector or the number of rows or columns in

a matrix.

 $n = nextnevn(n\theta);$

Output

scalar, the next allowable size for the given dimension for computing an FFT or RFFT. $n \ge n0$.

Remarks

nextn and **nextnevn** determine allowable matrix dimensions for computing FFT's. The Temperton FFT routines (see table below) can handle any matrix whose dimensions can be expressed as:

$$2^p \times 3^q \times 5^r \times 7^s$$
, p,q,r nonnegative integers $s = 0$ or 1

with one restriction: the vector length or matrix column size must be even (*p* must be positive) when computing RFFT's.

fftn, etc., automatically pad matrices (with zeros) to the next allowable dimensions; **nextn** and **nextnevn** are provided in case you want to check or fix matrix sizes yourself.

Use the following table to determine what to call for a given function and matrix:

FFT	Vector	Matrix	Matrix
Function	Length	Rows	Columns
fftn	nextn	nextn	nextn
rfftn	nextnevn	nextn	nextnevn
rfftnp	nextnevn	nextn	nextnevn

Example

n = nextn(456);

n = 480

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nextn, nextnevn

Source optim.src

See also fftn, optn, optnevn, rfftn, rfftnp

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nextwind

nextwind

Purpose Sets the current graphic panel to the next available graphic panel.

Library pgraph

Format nextwind;

Remarks This function selects the next available graphic panel to be the current

graphic panel. This is the graphic panel in which the next graph will be

drawn.

See the discussion on using graphic panels in "Publication Quality

Graphics" in the *User's Guide*.

Source pwindow.src

See also endwind, begwind, setwind, getwind, makewind,

window

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null

null

Purpose Computes an orthonormal basis for the (right) null space of a matrix.

Format b = null(x);

Input x NxM matrix.

Output b MxK matrix, where K is the nullity of X, such that:

$$x*b = 0$$
 (NxK matrix of zeros)

and

$$b'b = I$$
 (MXM identity matrix)

The error returns are returned in *b*:

error code reason

1 there is no null space

b is too large to return in a single matrix

Use **scalerr** to test for error returns.

Remarks The orthogonal complement of the column space of x' is computed using

the QR decomposition. This provides an orthonormal basis for the null

space of x.

Example let $x[2,4] = 2 \ 1 \ 3 \ -1$

$$b = null(x);$$

$$z = x*b;$$

$$i = b'b;$$

Source null.src

Globals _qrdc, _qrsl

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null1

Purpose Computes an orthonormal basis for the (right) null space of a matrix.

b

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d

NxM matrix.

dataset string, the name of a data set **null1** will write.

Output nu scalar, the nullity of x.

Remarks null1 computes an MxK matrix b, where K is the nullity of x, such that:

x*b = 0 (NxK matrix of zeros) and b'b = I (MxM identity matrix)

The transpose of b is written to the data set named by *dataset*, unless the nullity of x is zero. If nu is zero, the data set is not written.

Source null.src

Globals _qrdc, _qrsl

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ols

Purpose	Computes a least squares regression.						
Format	{ vnam,m,b,stb,vc,stderr,sigma,cx,rsq,resid,dwstat } = ols(dataset,depvar,indvars);						
Input	dataset	string, name of data set or null string.					
		If <i>dataset</i> is a null string, the procedure assumes that the actual data has been passed in the next two arguments.					
	depvar	If dataset contains a string:					
		string, name of dependent variable.					
		scalar, index of dependent variable. If scalar 0, the last column of the data set will be used.					
		If <i>dataset</i> is a null string or 0:					
		Nx1 vector, the dependent variable.					
	indvars	If dataset contains a string:					
		Kx1 character vector, names of independent variables.					
		Kx1 numeric vector, indices of independent variables.					
		These can be any size subset of the variables in the data set and can be in any order. If a scalar 0 is passed, all columns of the data set will be used except for the one used for the dependent variable.					
		If dataset is a null string or 0:					
		NxK matrix, the independent variables.					
Global Input	Defaults are provided for the following global input variables, so be ignored unless you need control over the other options provide this procedure.						
	altnam	global vector, default 0.					
	This can be a (K+1)x1 or (K+2)x1 character vector of alternate variable names for the output. Ifcon is 1, to must be (K+2)x1. The name of the dependent variable in the last element. con global scalar, default 1.						
		1 a constant term will be added, $D = K+1$.					
	0 no constant term will be added, $D = K$.						

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			A constant term will always be used in constructing the moment matrix m.					
	mis	ss	global scalar, default 0.					
				0	there are no miss	ing values (fastest).		
				1	listwise deletion, missings occur.	drop any cases in which		
				2	setting missings The number of c	to 0 when calculating <i>m</i> . ases computed is equal to of cases in the data set.		
	out	put	globa	bal scalar, default 1.				
			1		print the statistics.			
			0 0		do not print statistics.			
	rov	v	global scalar, the number of rows to read per iteration of the read loop. Default 0.					
		If 0, the number of rows will be you get an Insufficient while executing ols , you can that works on your system.				memory error message		
			differ iterati	ences which ion. You	nen a different num can userow to	htly due to rounding error at number of rows is read per ow to control this if you want to ang effects between several runs.		
	_olsı	res global scalar, default 0.						
			1 compute residuals (<i>resid</i>) and Durbin-Watson statistic(<i>dwstat</i>).					
				0	resid = 0, $dwstat$	=0.		
Output	vnam	the re(K+2	$(K+2)x1$ or $(K+1)x1$ character vector, the variable names used in the regression. If a constant term is used, this vector will be $(K+2)x1$, and the first name will be "CONSTANT". The last name will be the name of the dependent variable. MxM matrix, where $M = K+2$, the moment matrix constructed by calculating $\mathbf{X}'\mathbf{X}$ where \mathbf{X} is a matrix containing all useable observations and having columns in the order:					
	m	by ca						
			1.0		indvars	depvar		
		(con	stant)	(indepe	ndent variables)	(dependent variable)		
		A co	onstant term is always used in computing <i>m</i> .					
	b	Dx1	vector, the least squares estimates of parameters.					

Error handling is controlled by the low order bit of the trap flag.

trap 0 terminate with error message

trap 1 return scalar error code in b

- 30 system singular
- 31 system underdetermined
- 32 same number of columns as rows
- 33 too many missings
- 34 file not found
- 35 no variance in an independent variable

The system can become underdetermined if you use listwise deletion and have missing values. In that case, it is possible to skip so many cases that there are fewer useable rows than columns in the data set.

stb Kx1 vector, the standardized coefficients.

vc DxD matrix, the variance-covariance matrix of estimates.

stderr Dx1 vector, the standard errors of the estimated parameters.

sigma scalar, standard deviation of residual.

cx (K+1)x(K+1) matrix, correlation matrix of variables with the dependent variable as the last column.

rsq scalar, R square, coefficient of determination.

resid residuals, resid = y - x * b.

If **_olsres** = 1, the residuals will be computed.

If the data is taken from a data set, a new data set will be created for the residuals, using the name in the global string variable **_olsrnam**. The residuals will be saved in this data set as an Nx1 column. The *resid* return value will be a string containing the name of the new data set containing the residuals.

If the data is passed in as a matrix, the *resid* return value will be the Nx1 vector of residuals.

dwstat scalar, Durbin-Watson statistic.

Remarks

No output file is modified, opened, or closed by this procedure. If you want output to be placed in a file, you need to open an output file before calling ols.

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Example $y = \{ 2, \}$

In this example, the output from **ols** was put into a file called ols.out as well as being printed in the window. This example will compute a least squares regression of \mathbf{y} on \mathbf{x} . The return values were discarded by using a **call** statement.

```
data = "olsdat";
depvar = { score };
indvars = { region,age,marstat };
_olsres = 1;
output file = lpt1 on;
{ nam,m,b,stb,vc,std,sig,cx,rsq,resid,dbw } =
   ols(data,depvar,indvars);
output off;
```

In this example, the data set olsdat.dat was used to compute a regression. The dependent variable is **score**. The independent variables are **region**, **age**, and **marstat**. The residuals and Durbin-Watson statistic will be computed. The output will be sent to the printer as well as the window and the returned values are assigned to variables.

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Source ols.src

Globals _olsres, _olsrnam, __altnam, __con, __miss,

__output, __row, __vpad

See also olsqr

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olsqr

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olsqr

Purpose Computes OLS coefficients using QR decomposition.

Format b = olsqr(y,x);

Input y Nx1 vector containing dependent variable.

x NxP matrix containing independent variables.

Global Input _olsqtol global scalar, the tolerance for testing if diagonal elements are approaching zero. The default value is 10*e*-14.

Output b Px1 vector of least squares estimates of regression of y on x. If x does not have full rank, then the coefficients that cannot be estimated will be zero.

Remarks This provides an alternative to y/x for computing least squares coefficients.

This procedure is slower than the / operator. However, for near singular matrices it may produce better results.

olsqr handles matrices that do not have full rank by returning zeros for the coefficients that cannot be estimated.

Source olsqr.src

Globals _olsqtol

See also ols, olsqr2, orth, qqr

olsqr2

Purpose Computes OLS coefficients, residuals, and predicted values using the QR decomposition.

Format $\{b,r,p\} = olsqr2(y,x);$

Input y Nx1 vector containing dependent variable.

x NxP matrix containing independent variables.

Global Input _olsqtol global scalar, the tolerance for testing if diagonal elements are approaching zero. The default value is 10*e*-14.

Output b Px1 vector of least squares estimates of regression of y on x. If x does not have full rank, then the coefficients that cannot be estimated will be zero.

r Px1 vector of residuals. (r = y - x*b)

p Px1 vector of predicted values. (p = x*b)

Remarks This provides an alternative to y/x for computing least squares coefficients.

This procedure is slower than the / operator. However, for near singular matrices, it may produce better results.

olsqr2 handles matrices that do not have full rank by returning zeros for the coefficients that cannot be estimated.

Source olsqr.src

Globals _olsqtol

See also olsqr, orth, qqr

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Purpose Creates a matrix of ones.

Format

mat y = ones(r,c);

Input

r scalar, number of rows.

c scalar, number of columns.

Output

y RxC matrix of ones.

g Remarks

Noninteger arguments will be truncated to an integer.

Example

x = ones(3,2);

$$x = \begin{array}{c} 1 & 1 \\ 1 & 1 \\ 1 & 1 \end{array}$$

See also

zeros, eye

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Purpose Opens an existing GAUSS data file.

Format open fh=filename [for mode] [varindxi [offs]];

Input *filename* literal or ^string.

filename is the name of the file on the disk. The name can include a path if the directory to be used is not the current directory. This filename will automatically be given the extension .dat. If an extension is specified, the .dat will be overridden. If the file is an .fmt matrix file, the extension must be explicitly given. If the name of the file is to be taken from a string variable, the name of the string must be preceded by the ^ (caret) operator.

mode [[read]], append, update

The modes supported with the optional **for** subcommand are:

This is the default file opening mode and will be the one used if none is specified. Files opened in this mode cannot be written to. The pointer is set to the beginning of the file and the writer function is disabled for files opened in this way. This is the only mode available for matrix files (.fmt), which are always written in one piece

with the save command.

append Files opened in this mode cannot be read. The pointer will be set to the end of the file so that a subsequent write to the file with the writer function will add data to the end of the file without overwriting any of the existing data in the file. The readr function is disabled for files opened in this way. This mode is used to add additional rows to the end of a file.

update Files opened in this mode can be read from and written to. The pointer will be set to the beginning of the file. This mode is used to make changes in a file.

offs scalar.

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The optional **varindxi** subcommand tells GAUSS to create a set of global scalars that contain the index (column position) of the variables in a GAUSS data file. These "index variables" will have the same names as the corresponding variables in the data file but with "i" added as a prefix. They can be used inside index brackets, and with functions like **submat** to access specific columns of a matrix without having to remember the column position.

The optional *offs* is an offset that will be added to the index variables. This is useful if data from multiple files are concatenated horizontally in one matrix. It can be any scalar expression. The default is 0.

The index variables are useful for creating submatrices of specific variables without requiring that the positions of the variables be known. For instance, if there are two variables, **xvar** and **yvar** in the data set, the index variables will have the names **ixvar**, **iyvar**. If **xvar** is the first column in the data file, and **yvar** is the second, and if no offset, offs, has been specified, then **ixvar** and **iyvar** will equal 1 and 2, respectively. If an offset of 3 had been specified, then these variables would be assigned the values 4 and 5, respectively.

The **varindxi** and **varindx** options cannot be used with . fmt matrix files because no column names are stored with them.

If varindxi is used, GAUSS will ignore the Undefined symbol error message for global symbols that start with "i". This makes it much more convenient to use index variables because they don't have to be cleared before they are accessed in the program. Clearing is otherwise necessary because the index variables do not exist until execution time when the data file is actually opened and the names are read in from the header of the file. At compile time a statement like: y=x[.,ixvar]; will be illegal if the compiler has never heard of ixvar. If varindxi is used, this error will be ignored for symbols beginning with "i". Any symbols that are accessed before they have been initialized with a real value will be trapped at execution time with a Variable not initialized error message.

Output *fh* scalar.

fh is the file handle which will be used by most commands to refer to the file within GAUSS. This file handle is actually a scalar containing an integer value that uniquely identifies each file. This value is assigned by GAUSS when the **open** command is executed. If the file was not successfully opened, the file handle will be set to -1.

Remarks

The file must exist before it can be opened with the **open** command. (To create a new file, see **create** or **save**.)

A file can be opened simultaneously under more than one handle. See the second example following.

If the value that is in the file handle when the **open** command begins to execute matches that of an already open file, the process will be aborted and a **File already open** error message will be given. This gives you some protection against opening a second file with the same handle as a currently open file. If this happens, you would no longer be able to access the first file.

It is important to set unused file handles to zero because both **open** and **create** check the value that is in a file handle to see if it matches that of an open file before they proceed with the process of opening a file. This should be done with **close** or **closeall**.

Example

```
fname = "/data/rawdat";
open dt = ^fname for append;
if dt == -1;
    print "File not found";
    end;
end;
endif;
y = writer(dt,x);
if y /= rows(x);
    print "Disk Full";
    end;
endif;
dt = close(dt);
```

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In the example above, the existing data set /data/rawdat.dat is opened for appending new data. The name of the file was in the string variable **fname**. In this example the file handle is tested to see if the file was opened successfully. The matrix **x** is written to this data set. The number of columns in **x** must be the same as the number of columns in the existing data set. The first row in **x** will be placed after the last row in the existing data set. The **writer** function will return the number of rows actually written. If this does not equal the number of rows that were attempted, then the disk is probably full.

```
open fin = mydata for read;
open fout = mydata for update;
do until eof(fin);
    x = readr(fin,100);
    x[.,1 3] = ln(x[.,1 3]);
    call writer(fout,x);
endo;
closeall fin,fout;
```

In the above example, the same file, mydata.dat, is opened twice with two different file handles. It is opened for read with the handle **fin**, and it is opened for update with the handle **fout**. This will allow the file to be transformed in place without taking up the extra space necessary for a separate output file. Notice that **fin** is used as the input handle and **fout** is used as the output handle. The loop will terminate as soon as the input handle has reached the end of the file. Inside the loop the file is read into a matrix called **x** using the input handle, the data are transformed (columns 1 and 3 are replaced with their natural logs), and the transformed data is written back out using the output handle. This type of operation works well as long as the total number of rows and columns does not change.

The following example assumes a data file named dat1.dat that has the variables: visc, temp, lub, rpm.

```
open f1 = dat1 varindxi;
dtx = readr(f1,100);
x = dtx[.,irpm ilub ivisc];
y = dtx[.,itemp];
call seekr(f1,1);
```

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In this example, the data set dat1.dat is opened for reading (the .dat and the **for read** are implicit). **varindxi** is specified with no constant. Thus, index variables are created that give the positions of the variables in the data set. The first 100 rows of the data set are read into the matrix **dtx**. Then, specified variables in a specified order are assigned to the matrices **x** and **y** using the index variables. The last line uses the **seekr** function to reset the pointer to the beginning of the file.

```
open q1 = dat1 varindx;
open q2 = dat2 varindx colsf(q1);
nr = 100;
y = readr(q1,nr)~readr(q2,nr);
closeall q1,q2;
```

In this example, two data sets are opened for reading and index variables are created for each. A constant is added to the indices for the second data set (q2), equal to the number of variables (columns) in the first data set (q1). Thus, if there are three variables x1, x2, x3 in q1, and three variables y1, y2, y3 in q2, the index variables that were created when the files were opened would be ix1, ix2, ix3, iy1, iy2, iy3. The values of these index variables would be 1, 2, 3, 4, 5, 6, respectively. The first 100 rows of the two data sets are read in and concatenated to give the matrix y. The index variables will thus give the correct positions of the variables in y.

```
open fx = x.fmt;
i = 1; rf = rowsf(fx);
sampsize = round(rf*0.1);
rndsmpx = zeros(sampsize,colsf(fx));
do until i > sampsize;
  r = ceil(rndu(1,1)*rf);
  call seekr(fx,r);
  rndsmpx[i,.] = readr(fx,1);
  i = i+1;
endo;
fx = close(fx);
```

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w x y z

In this example, a 10% random sample of rows is drawn from the matrix file x.fmt and put into the matrix rndsmpx. Note that the extension .fmt must be specified explicitly in the open statement. The rowsf command is used to obtain the number of rows in x.fmt. This number is multiplied by 0.10 and the result is rounded to the nearest integer; this yields desired sample size. Then random integers (r) in the range 1 to rf are generated. seekr is used to locate to the appropriate row in the matrix, and the row is read with readr and placed in the matrix rndsmpx. This is continued until the complete sample has been obtained.

See also

create, close, closeall, readr, writer, seekr, eof

optn, optnevn

optn, optnevn

Purpose Returns optimal matrix dimensions for computing FFT's.

Format $n = optn(n\theta)$;

n = optnevn(n0);

Input n0 scalar, the length of a vector or the number of rows or columns in

a matrix.

Output n scalar, the next optimal size for the given dimension for

computing an FFT or RFFT. $n \ge n0$.

Remarks optn and optnevn determine optimal matrix dimensions for computing FFT's. The Temperton FFT routines (see table following) can handle any matrix whose dimensions can be expressed as:

$$2^p \times 3^q \times 5^r \times 7^s$$
, p, q, r nonnegative integers $s = 0$ or 1

with one restriction: the vector length or matrix column size must be even (*p* must be positive) when computing RFFT's.

fftn, etc., pad matrices to the next allowable dimensions; however, they generally run faster for matrices whose dimensions are highly composite numbers, that is, products of several factors (to various powers), rather than powers of a single factor. For example, even though it is bigger, a 33600x1 vector can compute as much as 20 percent faster than a 32768x1 vector, because 33600 is a highly composite number, $2^6 \times 3 \times 5^2 \times 7$, whereas 32768 is a simple power of 2, 2^{15} . **optn** and **optnevn** are provided so you can take advantage of this fact by hand-sizing matrices to optimal dimensions before computing the FFT.

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optn, optnevn

Use the following table to determine what to call for a given function and matrix:

FFT	Vector	Matrix	Matrix
Function	Length	Rows	Columns
fftn	optn	optn	optn
rfftn	optnevn	optn	optnevn
rfftnp	optnevn	optn	optnevn

Example n = optn(231);

n = 240.00000

See also fftn, nextn, nextnevn, rfftn, rfftnp

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orth

Purpose Computes an orthonormal basis for the column space of a matrix.

Format y = orth(x);

Input x NxK matrix.

Global Input _orthtol global scalar, the tolerance for testing if diagonal elements are approaching zero. The default is 1.0*e*-14.

Output y NxL matrix such that y'y = eye(L) and whose columns span the same space as the columns of x; L is the rank of x.

Example $x = \{ 654, 275 \};$ y = orth(x); $y = \begin{cases} -0.58123819 -0.81373347 \\ -0.81373347 & 0.58123819 \end{cases}$

Source qqr.src

Globals _orthtol

See also qqr, olsqr

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output

output

Purpose

This command makes it possible to direct the output of **print** statements to two different places simultaneously. One output device is always the window or standard output. The other can be selected by the user to be any disk file or other suitable output device such as a printer.

Format output [file=filename] [[on|off|reset]];

Input *filename* literal or ^string.

The **file**=*filename* subcommand selects the file or device to which output is to be sent.

If the name of the file is to be taken from a string variable, the name of the string must be preceded by the ^ (caret) operator.

The default file name is output.out.

on, off, literal, mode flag reset

on opens the auxiliary output file or device and causes the results of all print statements to be sent to that file or device. If the file already exists, it will be opened for appending. If the file does not already exist, it will be created.

off closes the auxiliary output file and turns off the auxiliary output.

reset similar to the on subcommand, except that it always creates a new file. If the file already exists, it will be destroyed and a new file by that name will be created. If it does not exist, it will be created.

Remarks After you have written to an output file you have to close the file before you can print it or edit it with the GAUSS editor. Use **output off**.

The selection of the auxiliary output file or device remains in effect until a new selection is made, or until you exit GAUSS. Thus, if a file is named as the output device in one program, it will remain the output device in subsequent programs until a new **file**=*filename* subcommand is encountered.

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The command output file=filename; will select the file or device but will not open it. A subsequent output on; or output reset; will open it and turn on the auxiliary output.

The command **output off** will close the file and turn off the auxiliary output. The filename will remain the same. A subsequent **output on** will cause the file to be opened again for appending. A subsequent **output reset** will cause the existing file to be destroyed and then recreated and will turn on the auxiliary output.

The command **output** by itself will cause the name and status (i.e., open or closed) of the current auxiliary output file to be printed in the window.

The output to the console can be turned off and on using the screen off and screen on commands. Output to the auxiliary file or device can be turned off or on using the output off or output on command. The defaults are screen on and output off.

The auxiliary file or device can be closed by an explicit **output off** statement, by an **end** statement, or by an interactive **new** statement. However, a **new** statement at the beginning of a program will not close the file. This allows programs with **new** statements in them to be run without reopening the auxiliary output file.

If a program sends data to a disk file, it will execute much faster if the window is off.

The **outwidth** command will set the line width of the output file. The default is 80.

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output

Example

```
output file = out1.out on;
```

This statement will open the file out1.out and will cause the results of all subsequent **print** statements to be sent to that file. If out1.out already exists, the new output will be appended.

```
output file = out2.out;
output on;
```

This is equivalent to the previous example.

```
output reset;
```

This statement will create a new output file using the current filename. If the file already exists, any data in it will be lost.

```
output file = mydata.asc reset;
screen off;
format /m1/rz 1,8;
open fp = mydata;
do until eof(fp);
   print readr(fp,200);;
endo;
fp = close(fp);
end;
```

The program above will write the contents of the GAUSS file mydata.dat into an ASCII file called mydata.asc. If there had been an existing file by the name of mydata.asc, it would have been overwritten.

The /ml parameter in the format statement in combination with the ;; at the end of the print statement will cause one carriage return/line feed pair to be written at the beginning of each row of the output file. There will not be an extra line feed added at the end of each 200 row block.

The end statement above will automatically perform output off and screen on.

See also

outwidth, screen, end, new

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outtyp (dataloop)

outtyp (dataloop)

Purpose Specifies the precision of the output data set.

Format outtyp num_constant;

Remarks *num_constant* must be 2, 4, or 8, to specify integer, single precision, or

double precision, respectively.

If **outtyp** is not specified, the precison of the output data set will be that of the input data set. If character data is present in the data set, the

precision will be forced to double.

Example outtyp 8;

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outwidth

outwidth

Purpose Specifies the width of the auxiliary output.

Format outwidth *n*;

Remarks n specifies the width of the auxiliary output in columns (characters). After

printing n characters on a line, GAUSS will output a line feed.

If a matrix is being printed, the line feed sequence will always be inserted between separate elements of the matrix rather than being inserted between digits of a single element.

n may be any scalar-valued expression in the range of 2-256. Nonintegers will be truncated to an integer. If 256 is used, no additional lines will be inserted.

The default is 80 columns.

Example outwidth 132;

This statement will change the auxiliary output width to 132 columns.

See also lpwidth, output, print

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w x y z

3-590

pacf

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V

x y z

pacf

```
Purpose
             Computes sample partial autocorrelations.
 Format
             rkk = pacf(y,k,d);
    Input
             v
                    Nx1 vector, data.
             k
                   scalar, maximum number of partial autocorrelations to compute.
                   scalar, order of differencing.
             d
  Output
                   Kx1 vector, sample partial autocorrelations.
             rkk
Example
             proc pacf(y,k,d);
                local a,l,j,r,t;
                r = acf(y,k,d);
                a = zeros(k,k);
                a[1,1] = r[1];
                t = 1;
                1 = 2;
                do while 1 le k;
                    a[1,1] = (r[1]-a[1-1,1:t]*rev(r[1:1-1]))/
                    (1-a[1-1,1:t]*r[1:t]);
                    j = 1;
                   do while j <= t;</pre>
                       a[l,j] = a[l-1,j] - a[l,l]*a[l-1,l-j];
                       j = j+1;
                    endo;
                    t = t+1;
                    1 = 1+1;
                endo;
```

pacf

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retp(diag(a));
endp;

Source tsutil.src

w x y z

packr

packr

Purpose Deletes the rows of a matrix that contain any missing values.

Format y = packr(x);

Input x NxK matrix.

Output y LxK submatrix of x containing only those rows that do not have missing values in any of their elements.

Remarks This function is useful for handling missing values by "listwise deletion," particularly prior to using the / operator to compute least squares coefficients.

If all rows of a matrix contain missing values, **packr** returns a scalar missing value. This can be tested for quickly with the **scalmiss** function.

Example x = miss(ceil(rndu(3,3)*10),1);
y = packr(x);

$$\begin{array}{r}
 . 9 10 \\
 x = 4 2 . \\
 3 4 9
 \end{array}$$

$$y = 349$$

In this example, the matrix **x** is formed with random integers and missing values. **packr** is used to delete rows with missing values.

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packr

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```
open fp = mydata;
obs = 0;
sum = 0;
do until eof(fp);
  x = packr(readr(fp,100));
if not scalmiss(x);
  obs = obs+rows(x);
  sum = sum+sumc(x);
  endif;
endo;
mean = sum/obs;
In this example, the sums of each column well as a count of the rows that do not con
```

In this example, the sums of each column in a data file are computed as well as a count of the rows that do not contain any missing values. **packr** is used to delete rows that contain missings and **scalmiss** is used to skip the two sum steps if all the rows are deleted for a particular iteration of the read loop. Then the sums are divided by the number of observations to obtain the means.

See also scalmiss, miss, missrv

w x y z

u

V

parse

parse

Purpose Parses a string, returning a character vector of tokens.

Format tok = parse(str,delim);

Input *str* string consisting of a series of tokens and/or delimiters.

delim NxK character matrix of delimiters that might be found in str.

Output *tok* Mx1 character vector consisting of the tokens contained in *str*.

All tokens are returned; any delimiters found in str are ignored.

Remarks The tokens in *str* must be 8 characters or less in size. If they are longer,

the contents of tok is unpredictable.

See also token

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pause

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pause

Purpose Pauses for a specified number of seconds.

Format pause(sec);

Input *sec* seconds to pause.

Source pause.src

See also wait

w x y z

u

V

pdfn

pdfn

Purpose Computes the standard Normal (scalar) probability density function.

Format y = pdfn(x);

Input x NxK matrix.

Output y NxK matrix containing the standard Normal probability density function of x.

Remarks This does not compute the joint Normal density function. Instead, the scalar Normal density function is computed element-by-element. *y* could

be computed by the following GAUSS code:

$$y = (1/sqrt(2*pi))*exp(-(x.*x)/2);$$

Example x = rndn(2,2);

y = pdfn(x);

 $= \begin{array}{rrr} -1.828915 & 0.514485 \\ -0.550219 & -0.275229 \end{array}$

 $y = \begin{array}{c} 0.074915 \ 0.349488 \\ 0.342903 \ 0.384115 \end{array}$

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pi

a b d е g h m n 0 p q u

```
Purpose Returns the mathematical constant π.

Format y = pi;

Example format /rdn 16,14;
    print pi;

produces:
    3.14159265358979
```

V

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pinv

Purpose Computes the Moore-Penrose pseudo-inverse of a matrix, using the singular value decomposition.

This pseudo-inverse is one particular type of generalized inverse.

Format y = pinv(x);

Input x NxM matrix.

Global Input_svdtol global scalar, any singular values less than _svdtol are treated as zero in determining the rank of the input matrix.

The default value for svdtol is 1.0e-13.

Output y MxN matrix that satisfies the 4 Moore-Penrose conditions:

XYX = X

YXY = Y

XY is symmetric

YX is symmetric

Global _svderr global scalar, if not all of the singular values can be computed _svderr will be nonzero.

Example $x = \{ 654, 275 \} ;$

y = pinv(x);

0.22017139 -0.16348055

 $y = -0.05207647 \quad 0.13447594$

 $-0.0151615 \quad 0.07712591$

Source svd.src

Globals _svdtol, _svderr

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polar

Purpose Graphs data using polar coordinates.

Library pgraph

Format polar(radius, theta);

Input radius Nx1 or NxM matrix. Each column contains the magnitude for

a particular line.

theta Nx1 or NxM matrix. Each column represents the angle values

for a particular line.

Source polar.src

See also xy, logx, logy, loglog, scale, xtics, ytics

polychar

polychar

Purpose Computes the characteristic polynomial of a square matrix.

Format c = polychar(x);

Input x NxN matrix.

Output c (N+1)x1 vector of coefficients of the Nth order characteristic polynomial of x:

 $p(z)=c[1]*z^n + c[2]*z^{(n-1)} + ... + c[n]*z + c[n+1];$

Remarks The coefficient of z^n is set to unity (c[1]=1).

Source poly.src

See also polymake, polymult, polyroot, polyeval

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polyeval

polyeval

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Purpose Evaluates polynomials. Can either be 1 or more scalar polynomials or a single matrix polynomial.

Format y = polyeval(x,c);

Input

1xK or NxN; that is, x can either represent K separate scalar values at which to evaluate the (scalar) polynomial(s), or it can represent a single NxN matrix.

c (P+1)xK or (P+1)x1 matrix of coefficients of polynomials to evaluate. If x is 1xK, then c must be (P+1)xK. If x is NxN, c must be (P+1)x1. That is, if x is a matrix, it can only be evaluated at a single set of coefficients.

Output

Kx1 vector (if c is (P+1)xK) or NxN matrix (if c is (P+1)x1 and x is NxN):

$$y = (c[1,.].*x^p + c[2,.].*x^{(p-1)} + ... + c[p+1,.])';$$

Remarks

In both the scalar and the matrix case, Horner's rule is used to do the evaluation. In the scalar case, the function **recsercp** is called (this implements an elaboration of Horner's rule).

Example x = 2i

let c = 1 1 0 1 1;
y = polyeval(x,c);

The result is 27. Note that this is the decimal value of the binary number 11011.

y = polyeval(x,1|zeros(n,1));

This will raise the matrix x to the n^{th} power (e.g. x*x*x*x*...*x).

Source poly.src

See also polymake, polychar, polymult, polyroot

polyint

polyint

Purpose Calculates an N^{th} order polynomial interpolation.

Format y = polyint(xa, ya, x);

Input xa Nx1 vector, X values.

ya Nx1 vector, Y values.

x scalar, X value to solve for.

Global Input _poldeg global scalar, the degree of polynomial required, default 6.

Output y result of interpolation or extrapolation.

Global Output **_polerr** global scalar, interpolation error.

Remarks Calculates an N^{th} order polynomial interpolation or extrapolation of X on

Y given the vectors xa and ya and the scalar x. The procedure uses Neville's algorithm to determine an up to N^{th} order polynomial and an error estimate.

ciroi estimate.

Polynomials above degree 6 are not likely to increase the accuracy for most data. Test _polerr to determine the required _poldeg for your problem.

problem.

Source polyint.src

Technical Notes

Press, W.P., B.P. Flannery, S.A. Tevkolsky, and W.T. Vettering. *Numerical Recipes: The Art of Scientific Computing*. NY: Cambridge Press, 1986.

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polymake

polymake

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хуг

Purpose Computes the coefficients of a polynomial given the roots.

Format c = polymake(r);

Input r Nx1 vector containing roots of the desired polynomial.

Output c (N+1)x1 vector containing the coefficients of the N^{th} order polynomial with roots r:

 $p(z)=c[1]*z^n+c[2]*z^{(n-1)}+...+c[n]*z+c[n+1];$

Remarks The coefficient of z^n is set to unity (c[1]=1).

Example $r = \{ 2, 1, 3 \};$

c = polymake(r);

 $c = \begin{array}{r} -1.0000000 \\ -6.0000000 \\ 11.000000 \\ -6.0000000 \end{array}$

Source poly.src

See also polychar, polymult, polyroot, polyeval

polymat

polymat

```
Purpose
              Returns a matrix containing the powers of the elements of x from 1 to p.
  Format
              y = polymat(x,p);
    Input
                     NxK matrix.
              x
                     scalar, positive integer.
              p
  Output
                     Nx(p*K) matrix containing powers of the elements of x from 1 to
                     p. The first K columns will contain first powers, the second K
                     columns contain the second powers, and so on.
Remarks
              To do polynomial regression use ols:
                  { vnam,m,b,stb,vc,stderr,sigma,cx,rsq,resid,
                  dwstat } = ols(0,y,polymat(x,p));
  Source
              polymat.src
```

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polymroot

polymroot

a b d h m n 0

p q u

V W x y z

```
Purpose
              Computes the roots of the determinant of a matrix polynomial
  Format
              r = polymroot(c);
    Input
                     (N+1)*KxK matrix of coefficients of an Nth order
                     polynomial of rank K.
  Output
                     K*N vector containing the roots of the determinantal equation.
Remarks
              c is constructed of N+1 KxK coefficient matrices stacked vertically with
              the coefficient matrix of the t^n at the top, t^(n-1) next, down to the t^0
              matrix at the bottom.
              Note that this procedure solves the scalar problem as well, that is, the one
              that POLYROOT solves.
Example
              Solve det(A2*t^2 + A1*t + A0) = 0 where:
              A2 = [1 2]
                     [21]
              A1 = [58]
                     [10
                         7 1
              A0 = [3 \ 4]
                     [65]
              a2 = \{ 1 2, 2 1 \};
              a1 = \{ 5 8, 10 7 \};
              a0 = \{ 34, 65 \};
              print polymroot(a2|a1|a0);
                  -4.3027756
                  -.69722436
```

-2.6180340

-.38196601

polymult

polymult

Purpose Multiplies polynomials.

Format c = polymult(c1, c2);

Input *c1* (D1+1)x1 vector containing the coefficients of the first polynomial.

c2 (D2+1)x1 vector containing the coefficients of the second polynomial.

Output c (D1+D2)x1 vector containing the coefficients of the product of the two polynomials.

Example $c1 = \{ 2, 1 \};$ $c2 = \{ 2, 0, 1 \};$

c = polymult(c1,c2);

4.0000000 2.0000000

c = 2.0000000

1.0000000

Source poly.src

See also polymake, polychar, polyroot, polyeval

Technical Notes If the degree of cI is DI (e.g., if DI=3, then the polynomial corresponding to cI is cubic), then there must be DI+I elements in cI (e.g., 4 elements for a cubic). Thus, for instance the coefficients for the polynomial $5*x^3 + 6*x + 3$ would be: cI=5|0|6|3. (Note that zeros must be explicitly given if there are powers of x missing.)

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polyroot

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хух

polyroot

Purpose Computes the roots of a polynomial given the coefficients.

Format y = polyroot(c);

Input c (N+1)x1 vector of coefficients of an N^{th} order polynomial:

 $p(z)=c[1]*z^n + c[2]*z^{(n-1)} + ... + c[n]*z + c[n+1]$

Zero leading terms will be stripped from c. When that occurs the order of y will be the order of the polynomial after the leading zeros have been stripped.

c[1] need not be normalized to unity.

Output y Nx1 vector, the roots of c.

Source poly.src

See also polymake, polychar, polymult, polyeval

pop

Purpose

Provides access to a last-in, first-out stack for matrices.

Format

pop b; pop a;

Remarks

This is used with **gosub**, **goto**, and **return** statements with parameters. It permits passing parameters to subroutines or labels, and returning parameters from subroutines.

The **gosub** syntax allows an implicit **push** statement. This syntax is almost the same as that of a standard **gosub**, except that the matrices to be **push**'ed "into the subroutine" are in parentheses following the label name. The matrices to be **push**'ed back to the main body of the program are in parentheses following the **return** statement. The only limit on the number of matrices that can be passed to and from subroutines in this way is the amount of room on the stack.

No matrix expressions can be executed between the (implicit) **push** and the **pop**. Execution of such expressions will alter what is on the stack.

Matrices must be **pop**'ped in the reverse order that they are **push**'ed, therefore the statements:

Note that matrices are **pop**'ped in reverse order, and that there is a separate **pop** statement for each matrix popped.

See also

gosub, goto, return

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W

pqgwin

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pqgwin

Purpose Sets the graphics viewer mode.

Library pgraph

Format pagwin arg;

Input *arg* string literal.

"one" Use only one viewer.

"many" Use a new viewer for each graph.

Remarks If "one" is set, the viewer executable will be vwr.exe.

"manual" and "auto" are supported for backwards compatibility,

manual=one, auto=many.

Example: pqgwin one;

pqgwin many;

Source pgraph.src

See also setvwrmode

w x y z

3-610

setvwrmode

Purpose Sets the computational precision of some of the matrix operators.

Format prcsn n;

Input n scalar, 64 or 80.

Portability UNIX, Windows

This function has no effect under UNIX or Windows. All computations are done in 64-bit precision.

Remarks n is a scalar containing either 64 or 80. The operators affected by this command are **chol**, **solpd**, **invpd**, and b/a (when neither a nor b is

scalar and *a* is not square).

prcsn 80 is the default. Precision is set to 80 bits (10 bytes), which corresponds to about 19 digits of precision.

prcsn 64 sets the precision to 64 bits (8 bytes), which is standard IEEE double precision. This corresponds to 15-16 digits of precision. 80-bit precision is still maintained within the 80x87 math coprocessor so that actual precision is better than double precision.

When **prcsn** 80 is in effect, all temporary storage and all computations for the operators listed above are done in 80 bits. When the operator is finished, the final result is rounded to 64-bit double precision.

See also chol, solpd, invpd

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previousindex

previousindex

a Purpose Returns the index of the previous element or subarray in an array. h Format pi = previousindex(i, o);Input i d Mx1 vector of indices into an array, where $M \le N$. Nx1 vector of orders of an N-dimensional array. 0 **Output** рi Mx1 vector of indices, the index of the previous element or subarray in the array corresponding to o. Remarks **previousindex** will return a scalar error code if the index cannot be h decremented. orders = $\{3,4,5,6,7\};$ Example a = areshape(1,orders); orders = getorders(a); ind = $\{2,3,1\};$ ind = previousindex(ind, orders); m ind = 20 5 p

In this example, **previousindex** decremented *ind* to index the previous 6x7 subarray in array *a*.

See also nextindex, loopnextindex, walkindex

w x y z

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princomp

princomp

Purpose Computes principal components of a data matrix.

Format $\{p, v, a\} = princomp(x, j);$

Input x NxK data matrix, N > K, full rank.

j scalar, number of principal components to be computed (J \leq K).

Output p NxJ matrix of the first j principal components of x in descending order of amount of variance explained.

v Jx1 vector of fractions of variance explained.

a JxK matrix of factor loadings, such that x = p*a +error.

Remarks Adapted from a program written by Mico Loretan.

The algorithm is based on Theil, Henri "Principles of Econometrics." Wiley, NY, 1971, 46-56.

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	print								
a	Purpose	Prints matrices or strings to the window and/or auxiliary output.							
b	Format	<pre>print [[/flush]] [[/typ]] [[/fmted]] [[/mf]] [[/jnt]]</pre>							
C			_express						
d	Input	/typ literal, symbol type flag.							
e			/mat /str	, /	sa,	Indicate which symbol types you are setting the output format for: matrices			
f g			/ 501			(/mat), string arrays (/sa), and/or strings (/str). You can specify more			
h						than one /typ flag; the format will be set			
i						for all types indicated. If no /typ flag is listed, print assumes /mat.			
		/fmted literal, enable formatting flag.							
j			/on,	/o	ff	Enable/disable formatting. When			
k 1						formatting is disabled, the contents of a variable are dumped to the window in a "raw" format.			
m		/mf literal, matrix format. It controls the way rows of a matrix are separated from one another. The possibilities are:							
n			/m0			no delimiters before or after rows when printing out matrices.			
0 n			/m1 (or	/mb1	print 1 carriage return/line feed pair before each row of a matrix with more			
p						than 1 row.			
q			/m2 d	or	/mb2	print 2 carriage return/line feed pairs			
r						before each row of a matrix with more than 1 row.			
S			/m3 d	or	/mb3	print "Row 1", "Row 2" before each			
t			/ 1			row of a matrix with more than one row.			
u			/ma1			print 1 carriage return/line feed pair after each row of a matrix with more than 1 row.			
V			/ma2			print 2 carriage return/line feed pairs after			
W						each row of a matrix with more than 1 row.			
хух						IOW.			

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		-					
	/a1	print 1 carriage return/line feed pair after each row of a matrix.					
	/a2	print 2 carriage return/line feed pairs after each row of a matrix.					
	/b1	print 1 carriage return/line feed pair before each row of a matrix.					
	/b2	print 2 carriage return/line feed pairs before each row of a matrix.					
	/b3	print "Row 1", "Row 2" before each row of a matrix.					
/jnt	literal, controls justifi	literal, controls justification, notation, and the trailing character.					
. .	Right-Justified						
	/rd	Signed decimal number in the form					
		[-]] ####.#### where #### is one or more decimal digits. The number of digits					
		before the decimal point depends on the					
		magnitude of the number, and the number					
		of digits after the decimal point depends					
		on the precision. If the precision is 0, no decimal point will be printed.					
	/re	Signed number in the form [] #.##E±###,					
	/16	where # is one decimal digit, ## is one or					
		more decimal digits depending on the					
		precision, and ### is three decimal digits.					
		If precision is 0, the form will be #### with no decimal point printed.					
	/ro	This will give a format like /rd or /re					
	,	depending on which is most compact for	П				
		the number being printed. A format like /re will be used only if the exponent					
		value is less than -4 or greater than the					
		precision. If a /re format is used a					
		decimal point will always appear. The precision signifies the number of					
		significant digits displayed.					

/rz

This will give a format like /rd or /re depending on which is most compact for the number being printed. A format like /re will be used only if the exponent value is less than -4 or greater than the precision. If a /re format is used, trailing zeros will be supressed and a decimal point will appear only if one or more digits follow it. The precision signifies the number of significant digits displayed.

Left-Justified

/ld

Signed decimal number in the form [] ####.###, where #### is one or more decimal digits. The number of digits before the decimal point depends on the magnitude of the number, and the number of digits after the decimal point depends on the precision. If the precision is 0, no decimal point will be printed. If the number is positive, a space character will replace the leading minus sign.

/le

/lo

This will give a format like /ld or /le depending on which is most compact for the number being printed. A format like /le will be used only if the exponent value is less than -4 or greater than the precision. If a /le format is used a decimal point will always appear. If the number is positive, a space character will replace the leading minus sign. The precision specifies the number of significant digits displayed.

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/1z This will give a format like /ld or /le depending on which is most compact for the number being printed. A format like /le will be used only if the exponent value is less than -4 or greater than the precision. If a /le format is used, trailing zeros will be supressed and a decimal point will appear only if one or more digits follow it. If the number is positive, a space character will replace the leading minus sign. The precision specifies the number of significant digits displayed.

Trailing Character

The following characters can be added to the /int parameters above to control the trailing character if any:

format /rdn 1,3;

The number will be followed immediately s

by a space character. This is the default.

The number will be followed immediately C

with a comma.

The number will be followed immediately t

with a tab character.

n No trailing character.

The default when GAUSS is first started is:

format /m1 /r0 16,8;

;;

Double semicolons following a print

statement will suppress the final carriage

return/line feed.

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Remarks

The list of expressions MUST be separated by spaces. In **print** statements, because a space is the delimiter between expressions, NO SPACES are allowed inside expressions unless they are within index brackets, quotes, or parentheses.

The printing of special characters is accomplished by the use of the backslash (\) within double quotes. The options are:

\b	backspace (ASCII 8)
\e	escape (ASCII 27)
\ f	form feed (ASCII 12)
\g	beep (ASCII 7)
\1	line feed (ASCII 10)
\r	carriage return (ASCII 13)
\t	tab (ASCII 9)
\###	the character whose ASCII value is "###" (decimal).

Thus, \13\10 is a carriage return/line feed sequence. The first three digits will be picked up here. So if the character to follow a special character is a digit, be sure to use three digits in the escape sequence. For example: \0074 will be interpreted as 2 characters (ASCII 7 followed by the character "4").

An expression with no assignment operator is an implicit **print** statement.

If output on has been specified, then all subsequent print statements will be directed to the auxiliary output as well as the window. (See output.) The locate statement has no effect on what will be sent to the auxiliary output, so all formatting must be accomplished using tab characters or some other form of serial output.

If the name of the symbol to be printed is prefixed with a '\$', it is assumed that the symbol is a matrix of characters.

```
print $x;
```

Note that GAUSS makes no distinction between matrices containing character data and those containing numeric data, so it is the responsibility of the user to use functions which operate on character matrices only on those matrices containing character data.

These matrices of character strings have a maximum of 8 characters per element. A precision of 8 or more should be set when printing out character matrices or the elements will be truncated.

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Complex numbers are printed with the sign of the imaginary half separating them and an "i" appended to the imaginary half. Also, the current field width setting (see **format**) refers to the width of field for each half of the number, so a complex number printed with a field of 8 will actually take (at least) 20 spaces to print.

A **print** statement by itself will cause a blank line to be printed:

```
print;
```

GAUSS also has an *automatic print mode* which causes the results of all global assignment statements to be printed out. This is controlled by the **print on** and **print off** commands. (See **print on**.)

Example

```
x = rndn(3,3);
format /rd 16,8;
   print x;
format /re 12,2;
   print x;
   print /rd/m3 x;
    0.14357994 -1.39272762 -0.91942414
    0.51061645 \quad -0.02332207 \quad -0.02511298
   -1.54675893 -1.04988540
                              0.07992059
   1.44E-001 -1.39E+000 -9.19E-001
   5.11E-001
              -2.33E-002
                          -2.51E-002
 -1.55E+000 -1.05E+000
                           7.99E-002
Row 1
          0.14
                  -1.39
                            -0.92
Row 2
          0.51
                  -0.02
                            -0.03
Row 3
         -1.55
                  -1.05
                             0.08
```

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In this example, a 3x3 random matrix is printed using 3 different formats. Notice that in the last statement the format is overridden in the **print** statement itself but the field and precision remain the same.

```
let x = AGE PAY SEX;
format /ml 8,8;
print $x;
produces:
   AGE
   PAY
   SEX
```

See also lprint, print on, lprint on, printfm, printdos

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printdos

printdos

Purpose Prints a string to the standard output.

Format printdos s;

Input s string, containing the string to be printed to the standard output.

Remarks This function is useful for printing messages to the window when

screen off is in effect. The output of this function will not go to the

auxiliary output.

This function can also be used to send escape sequences to the

ansi.sys device driver.

Example printdos "\27[7m"; /* set for reverse video */

printdos "\27[0m"; /* set for normal text */

See the DOS manuals for more complete information.

See also print, lprint, printfm, screen

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printfm

printfm

Purpose

Prints a matrix using a different format for each column of the matrix.

Format

y = printfm(x, mask, fmt);

Input

x NxK matrix which is to be printed and which may contain both character and numeric data.

mask LxM matrix, ExE conformable with x, containing ones and zeros which is used to specify whether the particular row, column, or element is to be printed as a string (0) or numeric (1) value.

fmt Kx3 or 1x3 matrix where each row specifies the format for the respective column of x.

Output

scalar, 1 if the function is successful and 0 if it fails.

Remarks

The mask is applied to the matrix x following the rules of standard element-by-element operations. If the corresponding element of mask is 0, then that element of x is printed as a character string of up to 8 characters. If mask contains a 1, then that element of x is assumed to be a double precision floating point number.

The contents of *fmt* are as follows:

[K,1] format string, a string 8 characters maximum.

[K,2] field width, a number < 80. [K,3] precision, a number < 17.

The format strings correspond to the **format** slash commands as follows:

/rdn "*.*lf" /ren "*.*1E" "#*,*1G" /ron "*,*1G" /rzn "-*.*lf" /ldn "-*,*1E" /len "-#*.*1G" /lon "-*.*1G" /lzn

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x y z

Complex numbers are printed with the sign of the imaginary half separating them and an "i" appended to the imaginary half. The field width refers to the width of field for each half of the number, so a complex number printed with a field of 8 will actually take (at least) 20 spaces to print.

If the precision = 0, the decimal point will be suppressed.

The format string can be a maximum of 8 characters and is appended to a % sign and passed directly to the **fprintf** function in the standard C language I/O library. The **lf**, etc., are case sensitive. If you know C, you will easily be able to use this.

If you want special characters to be printed after x, then include them as the last characters of the format string. For example:

```
"*.*1f," right-justified decimal followed by a comma.

"-*.*s" left-justified string followed by a space.

"*.*1f" right-justified decimal followed by nothing.
```

If you want the beginning of the field padded with zeros, then put a "0" before the first "*" in the format string:

Example

Here is an example of **printfm** being used to print a mixed numeric and character matrix:

printfm

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```
"*.*le" 12 4; /* third column format */
d = printfm(x,mask,fmt);
```

The output looks like this:

```
AGE 5.123, 2.2346E+000
PAY 1.235, 1.2346E+000
SEX 1.145, 3.4471E+000
JOB 4.114, 8.5564E+000
```

When the column of x to be printed contains all string elements, use a format string of "*.*s" if you want it right-justified, or "-*.*s" if you want it left-justified. If the column is mixed string and numeric elements, then use the correct numeric format and **printfm** will substitute a default format string for those elements in the column that are strings.

Remember, the mask value controls whether an element will be printed as a number or a string.

See also print, lprint, printdos

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printfmt

printfmt

Purpose Prints character, numeric, or mixed matrix using a default format controlled by the functions **formatcv** and **formatnv**. **Format** y = printfmt(x, mask);Input NxK matrix which is to be printed. scalar. 1 if x is numeric or 0 if x is character. mask 1xK vector of 1's and 0's. The corresponding column of x will be printed as numeric where mask = 1 and as character where mask = 0. Output scalar, 1 if the function is successful and 0 if it fails. Remarks Default format for numeric data is: "*.*lg" 16 8 Default format for character data is: "*.*s" 8 8 Example x = rndn(5,4);call printfmt(x,1); Source gauss.src Globals fmtcv, fmtnv See also formatcv, formatnv

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Purpose

Begins the definition of a multi-line recursive procedure. Procedures are user-defined functions with local or global variables.

Format

proc [(nrets) =] name(arglist);

Input

nrets constant, number of objects returned by the procedure. If nrets is not explicitly given, the default is 1. Legal values are 0 to 1023. The retp statement is used to return values from a procedure.

name literal, name of the procedure. This name will be a global symbol.

arglist a list of names, separated by commas, to be used inside the procedure to refer to the arguments that are passed to the procedure when the procedure is called. These will always be local to the procedure, and cannot be accessed from outside the procedure or from other procedures.

Remarks

A procedure definition begins with the **proc** statement and ends with the **endp** statement.

An example of a procedure definition is:

proc

Procedures can be used just as if they were functions intrinsic to the language. Below are the possible variations depending on the number of items the procedure returns.

Returns 1 item:

$$y = dog(i,j,k);$$

Returns multiple items:

$$\{x,y,z\} = cat(i,j,k);$$

Returns no items:

If the procedure does not return any items or you want to discard the returned items:

Procedure definitions may not be nested.

For more details on writing procedures, see "Procedures and Keywords" in the *User's Guide*.

See also keyword, call, endp, local, retp

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Purpose Computes the products of all elements in each column of a matrix.

Format y = prodc(x);

Input x NxK matrix.

Output y Kx1 matrix containing the products of all elements in each column of x.

Remarks To find the products of the elements in each row of a matrix, transpose before applying **prodc**. If *x* is complex, use the bookkeeping transpose (.').

To find the products of all of the elements in a matrix, use the **vecr** function before applying **prodc**.

Example let x[3,3] = 1 2 3 4 5 6 7 8 9; y = prodc(x); 28y = 80

162 **See also** sumc, meanc, stdc

putarray

putarray

Purpose

Puts a contiguous subarray into an N-dimensional array and returns the resulting array.

Format

```
y = putarray(a, loc, src);
```

Input

a N-dimensional array.

loc Mx1 vector of indices into the array to locate the subarray of interest, where M is a value from 1 to N.

src [N-M]-dimensional array, matrix, or scalar.

Output

y N-dimensional array.

Remarks

If *loc* is an Nx1 vector, then *src* must be a scalar. If *loc* is an [N-1]x1 vector, then *src* must be a 1-dimensional array or a 1xL vector, where L is the size of the fastest moving dimension of the array. If *loc* is an [N-2]x1 vector, then *src* must be a KxL matrix, or a KxL 2-dimensional array, where K is the size of the second fastest moving dimension.

Otherwise, if *loc* is an Mx1 vector, then *src* must be an [N-M]-dimensional array, whose dimensions are the same size as the corresponding dimensions of array *a*.

Example

```
a = arrayalloc(2|3|4|5|6,0);
src = arrayinit(4|5|6,5);
loc = { 2,1 };
a = putarray(a,loc,src);
```

This example sets the contiguous 4x5x6 subarray of a beginning at [2,1,1,1,1] to the array src, in which each element is set to the specified value 5.

See also

setarray

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putf

putf

Purpose

Purpose Writes the contents of a string to a file.

Format ret = putf(filename, str, start, len, mode, append);

Input *filename* string, name of output file.

str string to be written to *filename*. All or part of *str* may be

written out.

start scalar, beginning position in str of output string.

len scalar, length of output string.

mode scalar, output mode, (0) ASCII or (1) binary.

append scalar, file write mode, (0) overwrite or (1) append.

Output *ret* scalar, return code.

Remarks If *mode* is set to (1) binary, a string of length *len* will be written to *filename*. If *mode* is set to (0) ASCII, the string will be output up to length *len* or until **putf** encounters a ^Z (ASCII 26) in *str*. The ^Z will not be written to *filename*.

If *append* is set to (0) overwrite, the current contents of *filename* will be destroyed. If *append* is set to (1) append, *filename* will be created if it does not already exist.

If an error occurs, **putf** will either return an error code or terminate the program with an error message, depending on the **trap** state. If bit 2 (the 4's bit) of the trap flag is 0, **putf** will terminate with an error message. If bit 2 of the trap flag is 1, **putf** will return an error code. The value of the trap flag can be tested with **trapchk**.

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putf

ret can have the following values:

- 0 normal return
- 1 null file name
- 2 file open error
- 3 file write error
- 4 output string too long
- 5 null output string, or illegal *mode* value
- 6 illegal *append* value
- append specified but file did not exist; file was created (warning only)

Source putf.src

See also getf

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pvCreate

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pvCreate

Purpose Returns an initialized an instance of structure of type PV.

Format p1 = pvCreate;

Output p1 an instance of structure of type PV

Example struct PV p1;

p1 = pvCreate;

Source pv.src

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pvGetIndex

pvGetIndex

Purpose Gets row indices of a matrix in a parameter vector.

Format id = pvGetIndex(p1,nm1);

Input p1 an instance of structure of type PV.

nm1 name or row number of matrix.

Output *id* Kx1 vector, row indices of matrix described by *nm1* in parameter

vector.

Source pv.src

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pvGetParNames

pvGetParNames

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Purpose Generates names for parameter vector stored in structure of type PV. **Format** s = pvGetParNames(p1);Include pv.sdf Input p1an instance of structure of type PV. **Output** Kx1 string array, names of parameters. Remarks If the vector in the structure of type PV was generated with matrix names, the parameter names will be concatenations of the matrix name with row and column numbers of the parameters in the matrix. Otherwise the names will have a generic prefix with concatenated row and column numbers. **Example** #include pv.sdf struct PV p1; p1 = pvCreate; $x = \{ 1 2,$ 3 4 }; $mask = \{ 10,$ 0 1 };

p1 = pvPackm(p1,x,"P",mask);

print pvGetParNames(p1); P[1,1] P[2,2] x y z

pvGetParNames

Source pv.src

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pvGetParVector

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pvGetParVector

Purpose Retrieves parameter vector from structure of type PV. **Format** p = pvGetParVector(p1); Include pv.sdf Input p1 an instance of structure of type PV. **Output** Kx1 vector, parameter vector. Remarks Matrices or portions of matrices (stored using a mask) are stored in the structure of type PV as a vector in the p member. **Example** #include pv.sdf struct PV p1; p1 = pvCreate; $x = \{ 1 2,$ 3 4 }; $mask = \{ 10,$ 0 1 }; p1 = pvPackm(p1,x,"X",mask); print pvUnpack(p1,1); 1.000 2.000 3.000 4.000 print pvGetParVector(p1);

pvGetParVector

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Source pv.src

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pvLength

pvLength

Purpose Returns length of vector p.

Format n = pvLength(p1);

Input p1 an instance of structure of type PV.

Output n scalar, length of parameter vector in p1.

Source pv.src

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pvList

pvList

Purpose Retrieves names of packed matrices in structure of type PV.

Format n = pvList(p1);

Input p1 an instance of structure of type PV.

Output n Kx1 string vector, names of packed matrices.

Source pv.src

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pvPack

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x y z

pvPack

Purpose Packs general matrix into a structure of type PV with matrix name. **Format** p1 = pvPack(p1,x,nm);Include pv.sdf Input p1an instance of structure of type PV. MxN matrix or N-dimensional array. \boldsymbol{x} string, name of matrix/array. nmOutput an instance of structure of type PV. p1Example #include pv.sdf y = rndn(100,1);x = rndn(100,5);struct PV p1; p1 = pvCreate; p1 = pvPack(p1,x,"Y");p1 = pvPack(p1, y, "X");These matrices can be extracted using the unpack command: y = pvUnpack(p1,"Y"); x = pvUnpack(p1, "X");Source pv.src See also pvPackm, pvPacks, pvUnpack

pvPacki

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pvPacki

```
Purpose
              Packs general matrix or array into a PV instance with name and index.
  Format
              p1 = pvPacki(p1,x,nm,i);
 Include
              pv.sdf
    Input
              p1
                     an instance of structure of type PV.
                     MxN matrix or N-dimensional array.
              \boldsymbol{x}
                     string, name of matrix or array, or null string.
              nm
                     scalar, index of matrix or array in lookup table.
              i
  Output
              p1
                     an instance of structure of type PV.
Example
              #include pv.sdf
              y = rndn(100,1);
              x = rndn(100,5);
              struct PV p1;
              p1 = pvCreate;
              p1 = pvPacki(p1,y,"Y",1);
              p1 = pvPacki(p1,x,"X",2);
              These matrices can be extracted using the pvUnpack command.
              y = pvUnpack(p1,1);
              x = pvUnpack(p1,2);
See also
              pvPack, pvUnpack
```

pvPackm

pvPackm

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```
Packs general matrix into a structure of type PV with a mask and matrix
Purpose
               name.
  Format
               p1 = pvPackm(p1, x, nm, mask);
 Include
               pv.sdf
    Input
                      an instance of structure of type PV.
               p1
               x
                      MxN matrix or N-dimensional array.
               nm
                      string, name of matrix/array or N-dimensional array.
                      MxN matrix, mask matrix of zeros and ones.
               mask
  Output
               p1
                      an instance of structure of type PV.
Remarks
               The mask allows storing a selected portion of a matrix into the packed
               vector. The 1's in the mask matrix indicate an element to be stored in the
               packed matrix. When the matrix is unpacked (using pvUnpack) the
               elements corresponding to the zeros are restored. Elements corresponding
               to the ones come from the packed vector which may have been changed.
Example
               #include pv.sdf
               struct PV p1;
               p1 = pvCreate;
               x = \{ 1 2,
                       3 4 };
               mask = \{ 10,
                           0 1 };
```

p1 = pvPackm(p1,x,"X",mask);

print pvUnpack(p1,1);

pvPackm

```
1.000 2.000
3.000 4.000

pl = pvPutParVector(pl,5|6);

print pvUnpack(pl,"X");

5.000 2.000
3.000 6.000

Source pv.src
```

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pvPackmi

pvPackmi

h h m 0 p u V W

```
Purpose
               Packs general matrix or array into a PV instance with a mask, name, and
               index.
  Format
               pl = pvPackmi(pl,x,nm,mask,i);
 Include
               pv.sdf
    Input
                      an instance of structure of type PV.
               p1
               x
                      MxN matrix or N-dimensional array.
               nm
                      string, matrix or array name.
                      MxN matrix or N-dimensional array, mask of zeros and
               mask
                      ones.
               i
                      scalar, index of matrix or array in lookup table.
  Output
                      an instance of structure of type PV.
               p1
Remarks
               The mask allows storing a selected portion of a matrix into the parameter
               vector. The 1's in the mask matrix indicate an element to be stored in the
               parameter matrix. When the matrix is unpacked (using pvUnpackm) the
               elements corresponding to the zeros are restored. Elements
               corresponding to the ones come from the parameter vector.
Example
               #include pv.sdf
               struct PV p1;
               p1 = pvCreate;
               x = \{ 1 2,
                       3 4 };
               mask = \{ 10,
                           0 1 };
               p1 = pvPackmi(p1, x, "X", mask, 1);
```

print pvUnpack(p1,1);

хуZ

pvPackmi

```
1.000 2.000
3.000 4.000

p1 = pvPutParVector(p1,5|6);

print pvUnpack(p1,1);

5.000 2.000
3.000 6.000
```

See also pvPackm, pvUnpack

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pvPacks

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pvPacks

```
Purpose
             Packs symmetric matrix into a structure of type PV.
  Format
             p1 = pvPacks(p1,x,nm);
 Include
             pv.sdf
    Input
             p1
                    an instance of structure of type PV.
                    MxM symmetric matrix.
             х
                    string, matrix name.
             nm
  Output
                    an instance of structure of type PV.
             p1
Remarks
             pvPacks does not support the packing of arrays.
Example
             #include pv.sdf
             struct PV p1;
             p1 = pvCreate;
             x = \{ 1 2,
                    2 1 };
             p1 = pvPacks(p1,x,"A");
             p1 = pvPacks(p1,eye(2),"I");
             These matrices can be extracted using the pvUnpack command:
             print pvUnpack(p1,"A");
                    1.000
                            2.000
                    2.000
                            1.000
             print pvUnpack(p1,"I");
```

pvPacks

1.000 0.000

0.000 1.000

Source pv.src

See also pvPacksm, pvUnpack

a

b

С

d

e

f g

h

i

k

1

m

n

Ο

p

q

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S

t

u

V

pvPacksi

b

d

h

m

n

0

p

q

u

V

W

хуz

pvPacksi

Purpose Packs symmetric matrix into a PV instance with matrix name and index. **Format** pl = pvPacksi(pl,x,nm,i);Include pv.sdf Input p1an instance of structure of type PV. MxM symmetric matrix. \boldsymbol{x} string, matrix name. nmscalar, index of matrix in lookup table. i**Output** p1an instance of structure of type PV. pvPacksi does not support the packing of arrays. Remarks **Example** #include pv.sdf struct PV p1; p1 = pvCreate; $x = \{ 1 2, 2 1 \};$ p1 = pvPacksi(p1,x,"A",1);p1 = pvPacksi(p1,eye(2),"I",2); These matrices can be extracted using the pvUnpack command. print pvUnpack(p1,1); 1.000 2.000 2.000 1.000

pvPacksi

print pvUnpack(p1,2);

1.000 0.000

0.000 1.000

See also pvPacks, pvUnpack

a

b

С

d e

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h .

j

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m

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q

r

S

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u

V

XX7

pvPacksm

pvPacksm

b d h m n 0 p q

u V

```
Purpose
               Packs symmetric matrix into a structure of type PV with a mask.
  Format
               pl = pvPacksm(pl,x,nm,mask);
  Include
               pv.sdf
     Input
               p1
                       an instance of structure of type PV.
               x
                       MxM symmetric matrix.
                       string, matrix name.
               nm
               mask
                       MxM matrix, mask matrix of zeros and ones.
  Output
                       an instance of structure of type PV.
               p1
Remarks
               pvPacksm does not support the packing of arrays.
               The mask allows storing a selected portion of a matrix into the packed
               vector. The 1's in the mask matrix indicate an element to be stored in the
               packed matrix. When the matrix is unpacked (using pvUnpack) the
               elements corresponding to the zeros are restored. Elements corresponding
               to the ones come from the packed vector which may have been changed.
               Only the lower left portion of the mask matrix is used, and only the lower
               left portion of the x matrix is stored in the packed vector.
Example
               #include pv.sdf
               struct PV p1;
               p1 = pvCreate;
               x = \{ 1 2 4,
                        2 3 5,
                        4 5 6};
               mask = \{ 1 0 1, \}
```

0 1 0,

W

x y z

pvPacksm

```
1 0 1 };
p1 = pvPacksm(p1,x,"A",mask);
print pvUnpack(p1, "A");
      1.000
             2.000
                    4.000
      2.000
             3.000
                    5.000
             5.000
      4.000
                    6.000
p2 = pvGetParVector(p1);
print p2;
      1.000
      2.000
      3.000
      4.000
      5.000
      6.000
p3 = \{ 10, 11, 12, 13 \};
p1 = pvPutParVector(p1,p3);
print pvUnpack(p1, "A");
```

Source pv.src

10.000

2.000

12.000

2.000

5.000

11.000 5.000

4.000

13.000

a

b

c d

e

f g

h

j

k

1 m

n

0

p

q

r

S

t

u

V

W

pvPacksmi

pvPacksmi

Format

Input

a h

Purpose

Packs symmetric matrix into a PV instance with a mask, matrix name, and index.

pl = pvPacksmi(pl,x,nm,mask,i);

Include

pv.sdf

.

h

m

0

p1 an instance of structure of type PV.

х

MxM symmetric matrix. string, matrix name.

nm

mask MxM matrix, symmetric mask matrix of zeros and ones.

i scalar, index of matrix in lookup table.

Output

p1 an instance of structure of type PV.

Remarks

pvPacksmi does not support the packing of arrays.

The mask allows storing a selected portion of a matrix into the parameter vector. The 1's in the mask matrix indicate an element to be stored in the parameter vector. When the matrix is unpacked (using **pvUnpackm**) the elements corresponding to the zeros are restored. Elements corresponding to the ones come from the parameter vector.

Only the lower left portion of the mask matrix is used, and only the lower left portion of the *x* matrix is stored in the packed vector.

p

Example #include pv.sdf

struct PV p1;

p1 = pvCreate;

 $x = \{ 1 2 4,$

2 3 5,

4 5 6};

u

V

W

pvPacksmi

a

b

С

d

f

g

h

1

m

n

0

p

q

t

u

V

w xyz

```
mask = \{ 1 0 1, 
          0 1 0,
          1 0 1 };
 p1 = pvPacksmi(p1,x,"A",mask,1);
 print pvUnpack(p1,1);
       1.000
              2.000
                     4.000
       2.000 3.000
                     5.000
       4.000 5.000
                     6.000
 p2 = pvGetParVector(p1);
 print p2;
       1.000
       3.000
       4.000
       6.000
 p3 = \{ 10, 11, 12, 13 \};
 p1 = pvPutParVector(p1,p3);
 print pvUnpack(p1,1);
              2.000
       10.000
                      12.000
       2.000
              11.000
                      5.000
       12.000
              5.000
                      13.000
pvPacksm, pvUnpack
```

See also

pvPutParVector

b

d

h

m

n

0

p

q

u

V

W

pvPutParVector

Purpose Inserts parameter vector into structure of type PV. **Format** p1 = pvPutParVector(p); Include pv.sdf Input Kx1 vector, parameter vector. **Output** an instance of structure of type PV. p1Remarks Matrices or portions of matrices (stored using a mask) are stored in the structure of type PV as a vector in the p member. **Example** #include pv.sdf struct PV p1; p1 = pvCreate; $x = \{ 1 2 4,$ 2 3 5, 4 5 6}; $mask = \{ 1 0 1,$ 0 1 0, 1 0 1 }; p1 = pvPackm(p1,x,"A",mask);print pvUnpack(p1, "A"); 1.000 2.000 4.000 2.000 3.000 5.000

хух

pvPutParVector

```
4.000 5.000 6.000

p3 = { 10, 11, 12, 13 };

p1 = pvPutParVector(p1,p3);

print pvUnpack(p1,"A");

10.000 2.000 4.000
2.000 11.000 5.000
12.000 5.000 13.000
```

Source

pv.src

a

b

С

d

е

f

g

h

.

k

1

m

n

О

p

q

r

S

t

u

V

W

хух

pvTest

a

b

d

е

g

h

k

m

n

0

p

q

pvTest

Purpose Tests an instance of structure of type PV to determine if it is a proper structure of type PV.

Format i = pvTest(p1);

Input p1 an instance of structure of type PV.

Output i scalar, if 0 p1 is a proper structure of type PV, else if 1 an improper or unitialized structure of type PV.

Source pv.src

x y z

u

V

W

3-656

pvUnpack

pvUnpack

Purpose Unpacks matrices stored in a structure of type PV.

Format x = pvUnpack(pl, m);

Input p1 an instance of structure of type PV.

m string, name of matrix, or integer, index of matrix.

Output x MxN general matrix or MxM symmetric matrix or

N-dimensional array.

Source pv.src

a

b

c d

e

f

g

h

.

1

m

n

Ο

p

q

r

S

t

u

V

QNewton

h

h

m

0

p

u

V

W

хух

QNewton

Purpose Optimizes a function using the BFGS descent algorithm. **Format** $\{x,f,g,ret\} =$ QNewton(&fct,start); Input &fct pointer to a procedure that computes the function to be minimized. This procedure must have one input argument, a vector of parameter values, and one output argument, the value of the function evaluated at the input vector of parameter values. Kx1 vector, start values. start **Global Input** qn RelGradTol scalar, convergence tolerance for relative gradient of estimated coefficients. Default = 1e-5. scalar, pointer to a procedure that computes the gn GradProc gradient of the function with respect to the parameters. This procedure must have a single input argument, a Kx1 vector of parameter values, and a single output argument, a Kx1 vector of gradients of the function with respect to the parameters evaluated at the vector of parameter values. If **gn GradProc** is 0, ONewton uses gradp. scalar, maximum number of iterations. Default on MaxIters = 1e+5. Termination can be forced by pressing C on the keyboard. an PrintIters scalar, if 1, print iteration information. Default = 0. Can be toggled during iterations by pressing P on the keyboard. Kx1 vector, labels for parameters. qn ParNames gn PrintResults scalar, if 1, results are printed. **Output** Kx1 vector, coefficients at the minimum of the function. х f scalar, value of function at minimum. Kx1 vector, gradient at the minimum of the function. g scalar, return code. ret normal convergence

ONewton

a

b

С

d

е

f

g

h

m

n

0

p

q

u

V

x y z

- 1 forced termination
- 2 max iterations exceeded
- 3 function calculation failed
- 4 gradient calculation failed
- 5 step length calculation failed
- 6 function cannot be evaluated at initial parameter values

Remarks

If you are running in terminal mode, GAUSS will not see any input until you press ENTER. Pressing C on the keyboard will terminate iterations, and pressing P will toggle iteration output.

To reset global variables for this function to their default values, call **qnewtonset**.

Example

This example computes maximum likelihood coefficients and standard errors for a Tobit model:

```
/*
    ** qnewton.e - a Tobit model
    */

z = loadd("tobit"); /* get data */
b0 = { 1, 1, 1, 1 };
{b,f,g,retcode} = qnewton(&lpr,b0);

/*
    ** covariance matrix of parameters
    */
h = hessp(&lpr,b);
output file = qnewton.out reset;
print "Tobit Model";
print;
```

QNewton

b

d

h

m

n

0

p

q

u

V

W

x y z

```
print "coefficients standard errors";
print b~sqrt(diag(invpd(h)));
output off;
/*
** log-likelihood proc
* /
proc lpr(b);
   local s,m,u;
   s = b[4];
   if s \le 1e-4;
      retp(error(0));
   endif;
   m = z[.,2:4]*b[1:3,.];
   u = z[.,1] ./= 0;
   retp(-sumc(u.*lnpdfn2(z[.,1]-m,s) +
                    (1-u).*(ln(cdfnc(m/sqrt(s)))));
endp;
produces:
Tobit Model
coefficients standard errors
    0.010417884
                 0.080220019
    -0.20805753
                 0.094551107
   -0.099749592 0.080006676
     0.65223067 0.099827309
qnewton.src
```

3-660

Source

QProg

a

b

С

d

e

f

g

h

k

 ${\rm m}$

n o

p

q

r

S

t

V

хух

QProg

Purpose	Solves	s the quadratic programming problem.
Format	{ <i>x,u1</i> ,	$\{u2,u3,u4,ret\}$ = QProg($start,q,r,a,b,c,d,bnds$);
Input	start	Kx1 vector, start values.
	q	KxK matrix, symmetric model matrix.
	r	Kx1 vector, model constant vector.
	а	MxK matrix, equality constraint coefficient matrix, or scalar 0, no equality constraints.
	b	Mx1 vector, equality constraint constant vector, or scalar 0, will be expanded to Mx1 vector of zeros.
	С	NxK matrix, inequality constraint coefficient matrix, or scalar 0, no inequality constraints.
	d	Nx1 vector, inequality constraint constant vector, or scalar 0, will be expanded to Nx1 vector of zeros.
	bnds	Kx2 matrix, bounds on x , the first column contains the lower bounds on x , and the second column the upper bounds. If scalar 0, the bounds for all elements will default to $\pm 1e200$.
Global Input	_qpr	og_maxit scalar, maximum number of iterations. Default = 1000.
Output	x	Kx1 vector, coefficients at the minimum of the function.
	u1	Mx1 vector, Lagrangian coefficients of equality constraints.
	<i>u</i> 2	Nx1 vector, Lagrangian coefficients of inequality constraints.
	иЗ	Kx1 vector, Lagrangian coefficients of lower bounds.
	и4	Kx1 vector, Lagrangian coefficients of upper bounds.
	ret	scalar, return code.
		0 successful termination
		1 max iterations exceeded
		2 machine accuracy is insufficient to maintain decreasing function values
		3 model matrices not conformable
		<0 active constraints inconsistent

QProg

Remarks

QProg solves the standard quadratic programming problem:

 $min\frac{1}{2}x'Qx - x'R$

subject to constraints,

$$Ax = B$$

$$Cx \ge D$$

and bounds,

$$x_{low} \le x \le x_{up}$$

Source qprog.src

8

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C

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C

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хух

qqr

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X, such that:

$$X = Q_1 R$$

Format

$$\{ q1,r \} = qqr(x);$$

Input

x NxP matrix.

Output

q1 NxK unitary matrix, K = min(N,P).

r KxP upper triangular matrix.

Remarks

Given X, there is an orthogonal matrix Q such that Q'X is zero below its diagonal, i.e.,

$$Q'X = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X = Q_1 R$$

is the QR decomposition of X. If X has linearly independent columns, R is also the Cholesky factorization of the moment matrix of X, i.e., of X'X.

If you want only the R matrix, see the function \mathbf{qr} . Not computing Q_1 can produce significant improvements in computing time and memory usage.

An unpivoted *R* matrix can also be generated using **cholup**:

For linear equation or least squares problems, which require Q_2 for computing residuals and residual sums of squares, see **olsqr** and **qtyr**.

a

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w x y z qqr

For most problems an explicit copy of Q_1 or Q_2 is not required. Instead one of the following, Q'Y, QY, Q_1Y , Q_1Y , Q_2Y , or Q_2Y , for some Y, is required. These cases are all handled by **qtyr** and **qyr**. These functions are available because Q and Q_1 are typically very large matrices while their products with Y are more manageable.

If N < P the factorization assumes the form:

$$Q'X = \left[R_1 R_2\right]$$

where R_1 is a PxP upper triangular matrix and R_2 is Px(N – P). Thus Q is a PxP matrix and R is a PxN matrix containing R_1 and R_2 . This type of factorization is useful for the solution of underdetermined systems. However, unless the linearly independent columns happen to be the initial rows, such an analysis also requires pivoting (see **qre** and **qrep**).

Source qqr.src

See also qre, qrep, qtyr, qtyre, qtyrep, qyr, qyre, qyrep, olsqr

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р **q**

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qqre

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X, such that:

$$X[., E] = Q_1 R$$

Format
$$\{q1,r,e\} = qqre(x);$$

Input x NxP matrix.

Output q1 NxK unitary matrix, K = min(N,P).

r KxP upper triangular matrix.

e Px1 permutation vector.

Remarks

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X[.,E] = Q_1 R$$

is the QR decomposition of X[.,E].

If you want only the R matrix, see **qre**. Not computing Q_1 can produce significant improvements in computing time and memory usage.

If X has rank P, then the columns of X will not be permuted. If X has rank M < P, then the M linearly independent columns are permuted to the front of X by E. Partition the permuted X in the following way:

$$X[.,E] = \left[X_1 \ X_2\right]$$

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W

qqre

where X_1 is NxM and X_2 is Nx(P – M). Further partition R in the following way:

$$R = \begin{bmatrix} R_{11} & R_{12} \\ 0 & 0 \end{bmatrix}$$

where R_{11} is MxM and R_{12} is Mx(P – M). Then

$$A = R_{11}^{-1} R_{12}$$

and

$$X_2 = X_1 A$$

that is, A is an Mx(P - N) matrix defining the linear combinations of X_2 with respect to X_1 .

If N < P the factorization assumes the form:

$$Q'X = \begin{bmatrix} R_1 & R_2 \end{bmatrix}$$

where R_1 is a PXP upper triangular matrix and R_2 is PX (N – P). Thus Q is a PXP matrix and R is a PXN matrix containing R_1 and R_2 . This type of factorization is useful for the solution of underdetermined systems. For the solution of

$$X[.,E]b = Y$$

it can be shown that

$$b = qrsol(Q'Y,R1) \mid zeros(N-P,1);$$

The explicit formation here of Q, which can be a very large matrix, can be avoided by using the function **qtyre**.

For further discussion of QR factorizations see the remarks under qqr.

Source qqr.src

See also qqr, qtyre, olsqr

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W

x y z

qqrep

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X, such that:

$$X[., E] = Q_1 R$$

Format

$$\{q1,r,e\} = qqrep(x,pvt);$$

Input

x NxP matrix.

pvt Px1 vector, controls the selection of the pivot columns:

if pvt[i] > 0, x[i] is an initial column

if pvt[i] = 0, x[i] is a free column

if pvt[i] < 0, x[i] is a final column

The initial columns are placed at the beginning of the matrix and the final columns are placed at the end. Only the free columns will be moved during the decomposition.

Output

q1 NxK unitary matrix, K = min(N,P).

r KxP upper triangular matrix.

e Px1 permutation vector.

Remarks

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \begin{bmatrix} Q_1 & Q_2 \end{bmatrix}$$

where Q_1 has P columns, then

$$X[.,E] = \left[Q_1 R\right]$$

is the QR decomposition of X[.,E].

a

b

c d

е

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i

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n

0

р **q**

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qqrep

b

d

h

m

n

0

p

q

qqrep allows you to control the pivoting. For example, suppose that X is a data set with a column of ones in the first column. If there are linear dependencies among the columns of X, the column of ones for the constant may get pivoted away. This column can be forced to be included among the linearly independent columns using pvt.

If you want only the R matrix, see **qrep**. Not computing Q_1 can produce significant improvements in computing time and memory usage.

Source qqr.src

See also qqr, qre, olsqr

u

V

qr

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X, such that:

$$X = Q_1 R$$

Format
$$r = qr(x)$$
;

Input x NxP matrix.

Output r KxP upper triangular matrix, K = min(N,P).

Remarks

qr is the same as qqr but doesn't return the Q_1 matrix. If Q_1 is not wanted, qr will save a significant amount of time and memory usage, especially for large problems.

Given X, there is an orthogonal matrix Q such that Q'X is zero below its diagonal, i.e.,

$$Q'X = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \begin{bmatrix} Q_1 & Q_2 \end{bmatrix}$$

where Q_1 has P columns, then

$$X = Q_1 R$$

is the QR decomposition of X. If X has linearly independent columns, R is also the Cholesky factorization of the moment matrix of X, i.e., of X'X.

 \mathbf{qr} does not return the Q_1 matrix because in most cases it is not required and can be very large. If you need the Q_1 matrix see the function \mathbf{qqr} . If you need the entire Q matrix call \mathbf{qyr} with Y set to a conformable identity matrix.

For most problems Q'Y, Q_1Y , or QY, Q_1Y , for some Y, are required. For these cases see **qtyr** and **qyr**.

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qr

For linear equation or least squares problems, which require Q_2 for computing residuals and residual sums of squares, see **olsqr**.

If N < P the factorization assumes the form:

$$Q'X = \begin{bmatrix} R_1 & R_2 \end{bmatrix}$$

where R_1 is a PxP upper triangular matrix and R_2 is Px(N – P). Thus Q is a PxP matrix and R is a PxN matrix containing R_1 and R_2 . This type of factorization is useful for the solution of underdetermined systems. However, unless the linearly independent columns happen to be the initial rows, such an analysis also requires pivoting (see **qre** and **qrep**).

Source qr.src

See also qqr, qrep, qtyre

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Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X, such that:

$$X[., E] = Q_1 R$$

Format
$$\{r,e\} = qre(x);$$

Input
$$x$$
 NxP matrix.

Output
$$r$$
 KxP upper triangular matrix, $K = \min(N,P)$.

Remarks

qre is the same as **qqre** but doesn't return the Q_1 matrix. If Q_1 is not wanted, **qre** will save a significant amount of time and memory usage, especially for large problems.

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \begin{bmatrix} Q_1 & Q_2 \end{bmatrix}$$

where Q_1 has P columns, then

$$X[., E] = Q_1 R$$

is the QR decomposition of X[.,E].

qre does not return the Q_1 matrix because in most cases it is not required and can be very large. If you need the Q_1 matrix see the function **qqre**. If you need the entire Q matrix call **qyre** with Y set to a conformable identity matrix. For most problems Q'Y, Q_1Y , or QY, Q_1Y , for some Y, are required. For these cases see **qtyre** and **qyre**.

a

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С

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qre

If X has rank P, then the columns of X will not be permuted. If X has rank M < P, then the M linearly independent columns are permuted to the front of X by E. Partition the permuted X in the following way:

$$X[., E] = \begin{bmatrix} X_1 & X_2 \end{bmatrix}$$

where X_1 is NxM and X_2 is Nx(P – M). Further partition R in the following way:

$$R = \begin{bmatrix} R_{11} & R_{12} \\ 0 & 0 \end{bmatrix}$$

where R_{11} is MxM and R_{12} is Mx(P – M). Then

$$A = R_{11}^{-1} R_{12}$$

and

$$X_2 = X_1 A$$

that is, A is an Mx(P - N) matrix defining the linear combinations of X_2 with respect to X_1 .

If N < P the factorization assumes the form:

$$Q'X = \left[R_1 \ R_2\right]$$

where R_1 is a PXP upper triangular matrix and R_2 is PX(N – P). Thus Q is a PXP matrix and R is a PXN matrix containing R_1 and R_2 . This type of factorization is useful for the solution of underdetermined systems. For the solution of

$$X[.,E]b = Y$$

it can be shown that

$$b = qrsol(Q'Y,R1) \mid zeros(N-P,1);$$

The explicit formation here of Q, which can be a very large matrix, can be avoided by using the function **qtyre**.

For further discussion of QR factorizations see the remarks under qqr.

Source qr.src

See also qqr, olsqr

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grep

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X, such that:

$$X[., E] = Q_1 R$$

Format

$$\{r,e\} = qrep(x,pvt);$$

Input

x NxP matrix.

pvt Px1 vector, controls the selection of the pivot columns:

if pvt[i] > 0, x[i] is an initial column

if pvt[i] = 0, x[i] is a free column

if pvt[i] < 0, x[i] is a final column

The initial columns are placed at the beginning of the matrix and the final columns are placed at the end. Only the free columns will be moved during the decomposition.

Output

r KXP upper triangular matrix, K = min(N,P).

e Px1 permutation vector.

Remarks

grep is the same as **ggrep** but doesn't return the Q_1 matrix. If Q_1 is not wanted, **grep** will save a significant amount of time and memory usage, especially for large problems.

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \begin{bmatrix} Q_1 & Q_2 \end{bmatrix}$$

where Q_1 has P columns, then

$$X[., E] = Q_1 R$$

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is the QR decomposition of X[.,E].

qrep does not return the Q_1 matrix because in most cases it is not required and can be very large. If you need the Q_1 matrix see the function **qqrep**. If you need the entire Q matrix call **qyrep** with Y set to a conformable identity matrix. For most problems Q'Y, Q_1Y , or QY, Q_1Y , for some Y, are required. For these cases see **qtyrep** and **qyrep**.

grep allows you to control the pivoting. For example, suppose that X is a data set with a column of ones in the first column. If there are linear dependencies among the columns of X, the column of ones for the constant may get pivoted away. This column can be forced to be included among the linearly independent columns using pvt.

Source qr.src

See also qr, qre, qqrep

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qrsol

qrsol

Purpose Computes the solution of Rx = b where R is an upper triangular matrix.

Format x = qrsol(b,R);

Input b PxL matrix.

R PxP upper triangular matrix.

Output x PxL matrix.

Remarks qrsol applies a backsolve to Rx = b to solve for x. Generally R will be

the R matrix from a QR factorization. **qrsol** may be used, however, in

any situation where R is upper triangular.

Source qrsol.src

See also qqr, qr, qtyr, qrtsol

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qrtsol

Purpose Computes the solution of R'x = b where R is an upper triangular matrix.

Format x = qrtsol(b,R);

Input b PxL matrix.

R PxP upper triangular matrix.

Output x PxL matrix.

Remarks qrtsol applies a forward solve to R'x = b to solve for x. Generally R

will be the R matrix from a QR factorization. **qrtsol** may be used, however, in any situation where R is upper triangular. If R is lower

triangular, transpose before calling qrtsol.

If R is not transposed, use **qrsol**.

Source qrsol.src

See also qqr, qr, qtyr, qrsol

qtyr

Purpose Computes the orthogonal-triangular (QR) decomposition of a matrix X and returns O'Y and R.

Format $\{qty,r\} = qtyr(y,x);$

Input y NxL matrix.

x NxP matrix.

Output qty NxL unitary matrix.

r KxP upper triangular matrix, K = min(N,P).

Remarks Given X, there is an orthogonal matrix Q such that Q'X is zero below its diagonal, i.e.,

$$Q'X = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X = Q_1 R$$

is the QR decomposition of X. If X has linearly independent columns, R is also the Cholesky factorization of the moment matrix of X, i.e., of X'X. For most problems Q or Q_1 is not what is required. Rather, we require Q'Y or Q_1Y where Y is an NxL matrix (if either QY or Q_1Y are required, see qyr). Since Q can be a very large matrix, qtyr has been provided for the calculation of Q'Y which will be a much smaller matrix. Q_1Y will be a submatrix of Q'Y. In particular,

$$G = Q_1'Y = qty[1:P,.]$$

and Q_2Y is the remaining submatrix:

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qtyr

 $H = Q_2' Y = qty[P+1:N,.]$

Suppose that X is an NxK data set of independent variables, and v is an Nx1 vector of dependent variables. Then it can be shown that

$$b = R^{-1}G$$

and

$$s_j = \sum_{i=1}^{N-P} H_{i,j}, j = 1, 2, ...L$$

where b is a PXL matrix of least squares coefficients and s is a 1XL vector of residual sums of squares. Rather than invert R directly, however, it is better to apply **qrsol** to

$$Rb = Q_1'Y$$

For rank deficient least squares problems, see **qtyre** and **qtyrep**.

Example The QR algorithm is the superior numerical method for the solution of least squares problems:

Source qtyr.src

See also qqr, qtyre, qtyrep, olsqr

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qtyre

Purpose Computes the orthogonal-triangular (QR) decomposition of a matrix X and returns O'Y and R.

Format $\{qty,r,e\} = qtyre(y,x);$

Input y NxL matrix.

x NxP matrix.

Output *qty* NxL unitary matrix.

r KxP upper triangular matrix, K = min(N,P).

e Px1 permutation vector.

Remarks Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X[., E] = Q_1 R$$

is the QR decomposition of X[.,E].

If X has rank P, then the columns of X will not be permuted. If X has rank M < P, then the M linearly independent columns are permuted to the front of X by E. Partition the permuted X in the following way:

$$X[., E] = \begin{bmatrix} X_1 & X_2 \end{bmatrix}$$

where X_1 is NxM and X_2 is Nx(P – M). Further partition R in the following way:

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qtyre

 $R = \begin{bmatrix} R_{11} & R_{12} \\ 0 & 0 \end{bmatrix}$

where R_{11} is MxM and R_{12} is Mx(P – M). Then

$$A = R_{11}^{-1} R_{12}$$

and

$$X_2 = X_1 A$$

that is, A is an Mx(P - N) matrix defining the linear combinations of X_2 with respect to X_1 .

For most problems Q or Q_1 is not what is required. Rather, we require Q'Y or Q_1Y where Y is an NxL matrix. Since Q can be a very large matrix, **qtyre** has been provided for the calculation of Q'Y which will be a much smaller matrix. Q_1Y will be a submatrix of Q'Y. In particular,

$$Q_1'Y = qty[1:P, .]$$

and Q_2Y is the remaining submatrix:

$$Q_2'Y = qty[P+1:N,.]$$

Suppose that X is an NxK data set of independent variables and Y is an Nx1 vector of dependent variables. Suppose further that X contains linearly dependent columns, i.e., X has rank M < P. Then define

$$C = Q_2'Y [1:M,.]$$

$$A = R[1:M,1:M]$$

and the vector (or matrix of L > 1) of least squares coefficients of the reduced, linearly independent problem is the solution of

$$Ab = C$$

To solve for *b* use **qrsol**:

$$b = qrsol(C,A);$$

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qtyre

If N < P the factorization assumes the form:

$$Q'X[.,E] = \begin{bmatrix} R_1 & R_2 \end{bmatrix}$$

where R_1 is a PxP upper triangular matrix and R_2 is Px(N – P). Thus Q is a PxP matrix and R is a PxN matrix containing R_1 and R_2 . This type of factorization is useful for the solution of underdetermined systems. For the solution of

$$X[.,E]b = Y$$

it can be shown that

$$b = qrsol(Q'Y,R1) \mid zeros(N-P,1);$$

Source qtyr.src

See also qqr, qre, qtyr

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qtyrep

qtyrep

a

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X using a pivot vector and returns Q'Y and R.

Format

 $\{ qty,r,e \} = qtyrep(y,x,pvt);$

Input

y NxL matrix.

x NxP matrix.

pvt Px1 vector, controls the selection of the pivot columns:

if pvt[i] > 0, x[i] is an initial column

if pvt[i] = 0, x[i] is a free column

if pvt[i] < 0, x[i] is a final column

The initial columns are placed at the beginning of the matrix and the final columns are placed at the end. Only the free columns will be moved during the decomposition.

Output

qty NxL unitary matrix.

r KxP upper triangular matrix, K = min(N,P).

e Px1 permutation vector.

Remarks

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X[., E] = Q_1 R$$

is the QR decomposition of X[.,E].

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qtyrep

qtyrep allows you to control the pivoting. For example, suppose that X is a data set with a column of ones in the first column. If there are linear dependencies among the columns of X, the column of ones for the constant may get pivoted away. This column can be forced to be included among the linearly independent columns using pvt.

Example

```
y = \{ 472,
       5 9 1,
       6 3 3 };
x = \{ 1295,
       4 3 5,
       4 2 7 };
pvt = { 11, 10, 3 };
\{ qty, r, e \} = qtyrep(y,x,pvt);
       -6.9347609
                      -9.9498744 -3.0151134
qty =
        4.0998891 3.5527137e – 15
                                   2.1929640
        3.4785054
                       6.3245553  0.31622777
     -13.266499
                -9.6483630 -8.1408063
r =
      0.0000000 -0.95346259
                              4.7673129
                              3.1622777
      0.0000000
                  0.0000000
     1.0000000
e =
     2.0000000
     3.0000000
```

Source qt

qtyr.src

See also qrep, qtyre

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quantile

Purpose Computes quantiles from data in a matrix, given specified probabilities. Format y = quantile(x,e)Input NxK matrix of data. х Lx1 vector, quantile levels or probabilities. e**Output** LxK matrix, quantiles. Remarks quantile will not succeed if N*minc(e) is less than 1, or N*maxc(e)is greater than N - 1. In other words, to produce a quantile for a level of .001, the input matrix must have more than 1000 rows. Example rndseed 345567; x = rndn(1000,4); /* data */e = { .025, .5, .975 }; /* quantile levels */ y = quantile(x,e);print "medians"; print y[2,.]; print; print "95 percentiles"; print y[1,.]; print y[3,.]; produces: medians -0.0020 -0.0408 -0.0380-0.024795 percentiles -2.1474-1.8677-1.9894-1.87471.9687 2.0899 1.8576 2.0545

quantile

Source quantile.src

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quantiled

Purpose Computes quantiles from data in a data set, given specified probabilities. **Format** y = quantiled(dataset, e, var); Input dataset string, data set name, or NxM matrix of data. Lx1 vector, quantile levels or probabilities. Kx1 vector or scalar zero. If Kx1, character vector of labels var selected for analysis, or numeric vector of column numbers in data set of variables selected for analysis. If scalar zero, all columns are selected. If *dataset* is a matrix *var* cannot be a character vector. **Output** LxK matrix, quantiles. Remarks quantiled will not succeed if N*minc(e) is less than 1, or N*maxc(e)is greater than N - 1. In other words, to produce a quantile for a level of .001, the input matrix must have more than 1000 rows. **Example** y = quantiled("tobit",e,0); print "medians"; print y[2,.]; print; print "95 percentiles"; print y[1,.]; print y[3,.]; produces: medians 0.0000 1.0000 -0.0021 -0.122895 percentiles

1.0000

-1.8139

-2.3143

-1.1198

quantiled

2.3066 1.0000 1.4590 1.6954

Source quantile.src

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x y z

Purpose Computes the orthogonal-triangular (QR) decomposition of a matrix X and returns QY and R.

Format $\{qy, q\}$

 $\{qy,r\} = qyr(y,x);$

Input

y NxL matrix.

x NxP matrix.

Output

qy NxL unitary matrix.

r KxP upper triangular matrix, K = min(N,P).

Remarks

Given X, there is an orthogonal matrix Q such that Q'X is zero below its diagonal, i.e.,

$$Q'X = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X = Q_1 R$$

is the QR decomposition of X. If X has linearly independent columns, R is also the Cholesky factorization of the moment matrix of X, i.e., of X'X.

For most problems Q or Q_1 is not what is required. Since Q can be a very large matrix, \mathbf{qyr} has been provided for the calculation of QY, where Y is some NxL matrix, which will be a much smaller matrix.

If either Q'Y or Q_1Y are required, see **qtyr**.

Example

 $x = \{ 1 11, 7 3, 2 1 \};$ $y = \{ 2 6, 5 10, 4 3 \};$ $\{ qy, r \} = qyr(y,x);$

qyr

 $qy = \begin{array}{r} 4.6288991 & 9.0506281 \\ -3.6692823 & -7.8788202 \\ 3.1795692 & 1.0051489 \end{array}$

Source qyr.src

See also qqr, qyre, qyrep, olsqr

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qyre

qyre

Purpose

Computes the orthogonal-triangular (QR) decomposition of a matrix X and returns OY and R.

Format

 $\{qy,r,e\} = qyre(y,x);$

Input

y NxL matrix.

x NxP matrix.

Output

qy NxL unitary matrix.

r KxP upper triangular matrix, K = min(N,P).

e Px1 permutation vector.

Remarks

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X[., E] = Q_1 R$$

is the QR decomposition of X[.,E].

For most problems Q or Q_1 is not what is required. Since Q can be a very large matrix, **qyre** has been provided for the calculation of QY, where Y is some NxL matrix, which will be a much smaller matrix.

If either Q'Y or $Q_{I}Y$ are required, see **qtyre**.

If N < P the factorization assumes the form:

$$Q'X[.,E] = \left[R_1 \ R_2\right]$$

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x y z

qyre

where R_1 is a PxP upper triangular matrix and R_2 is Px(N – P). Thus Q is a PxP matrix and R is a PxN matrix containing R_1 and R_2 .

Example

$$qy = \begin{array}{rrr} -0.5942276 & -3.0456088 \\ -6.2442636 & -11.647846 \\ 2.3782485 & -0.22790230 \end{array}$$

$$r = -11.445523 - 2.9705938$$
$$0.0000000 - 6.7212776$$

$$e = \begin{array}{c} 2.0000000 \\ 1.0000000 \end{array}$$

Source

qyr.src

See also

qqr, qre, qyr

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qyrep

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Purpose Computes the orthogonal-triangular (QR) decomposition of a matrix X using a pivot vector and returns QY and R.

Format

 $\{qy,r,e\} = qyrep(y,x,pvt);$

Input

y NxL matrix.

x NxP matrix.

pvt Px1 vector, controls the selection of the pivot columns:

if pvt[i] > 0, x[i] is an initial column

if pvt[i] = 0, x[i] is a free column

if pvt[i] < 0, x[i] is a final column

The initial columns are placed at the beginning of the matrix and the final columns are placed at the end. Only the free columns will be moved during the decomposition.

Output

qy NxL unitary matrix.

r KxP upper triangular matrix, K = min(N,P).

e Px1 permutation vector.

Remarks

Given X[.,E], where E is a permutation vector that permutes the columns of X, there is an orthogonal matrix Q such that Q'X[.,E] is zero below its diagonal, i.e.,

$$Q'X[.,E] = \begin{bmatrix} R \\ 0 \end{bmatrix}$$

where R is upper triangular. If we partition

$$Q = \left[Q_1 \ Q_2 \right]$$

where Q_1 has P columns, then

$$X[., E] = Q_1 R$$

is the QR decomposition of X[.,E].

gyrep

qyrep allows you to control the pivoting. For example, suppose that X is a data set with a column of ones in the first column. If there are linear dependencies among the columns of X, the column of ones for the constant may get pivoted away. This column can be forced to be included among the linearly independent columns using pvt.

For most problems Q or Q_1 is not what is required. Since Q can be a very large matrix, **qyrep** has been provided for the calculation of QY, where Y is some NxL matrix, which will be a much smaller matrix.

If either Q'Y or Q_1Y are required, see **qtyrep**.

If N < P the factorization assumes the form:

$$Q'X[.,E] = \begin{bmatrix} R_1 & R_2 \end{bmatrix}$$

where R_1 is a PxP upper triangular matrix and R_2 is Px(N – P). Thus Q is a PxP matrix and R is a PxN matrix containing R_1 and R_2 .

Source qyr.src

See also qr, qqrep, qrep, qtyrep

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rank

Purpose Computes the rank of a matrix, using the singular value decomposition. **Format** k = rank(x);Input NxP matrix. **Global Input _svdtol** global scalar, the tolerance used in determining if any of the singular values are effectively 0. The default value is 10*e*-13. This can be changed before calling the procedure. **Output** an estimate of the rank of x. This equals the number of singular values of x that exceed a prespecified tolerance in absolute value. Global **_svderr** global scalar, if not all of the singular values can be **Output** computed _svderr will be nonzero. Source svd.src

s t u v

rankindx

rankindx

Purpose Returns the vector of ranks of a vector.

Format y = rankindx(x,flag);

Input x Nx1 vector.

flag scalar, 1 for numeric data or 0 for character data.

Output y Nx1 vector containing the ranks of x. That is, the rank of the largest element is N and the rank of the smallest is 1. (To get ranks in descending order, subtract y from N+1).

Remarks rankindx assigns different ranks to elements that have equal values (ties). Missing values are assigned the lowest ranks.

Example let $x = 12 \ 4 \ 15 \ 7 \ 8;$

r = rankindx(x,1);

4

r = 5

2

3

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```
Purpose
```

Reads a specified number of rows of data from a GAUSS data set (.dat) file or a GAUSS matrix (.fmt) file.

```
Format y = readr(fl,r);

Input fl scalar, file ha
```

fl scalar, file handle of an open file.

r scalar, number of rows to read.

Output

NxK matrix, the data read from the file.

Remarks

The first time a **readr** statement is encountered, the first r rows will be read. The next time it is encountered, the next r rows will be read in, and so on. If the end of the data set is reached before r rows can be read, then only those rows remaining will be read.

After the last row has been read, the pointer is placed immediately after the end of the file. An attempt to read the file in these circumstances will cause an error message.

To move the pointer to a specific place in the file use **seekr**.

Example

```
open dt = dat1.dat;
m = 0;
do until eof(dt);
   x = readr(dt,400);
   m = m+moment(x,0);
endo;
dt = close(dt);
```

This code reads data from a data set 400 rows at a time. The moment matrix for each set of rows is computed and added to the sum of the previous moment matrices. The result is the moment matrix for the entire data set. **eof(dt)** returns 1 when the end of the data set is encountered.

See also

open, create, writer, seekr, eof

real

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real

```
Purpose
             Returns the real part of a matrix.
 Format
             zr = real(x);
    Input
                    NxK matrix.
  Output
                    NxK matrix, the real part of x.
             z,r
Remarks
             If x is not complex, zr will be equal to x.
Example
             x = \{ 1 11,
                     7i 3,
                     2+i 1 };
             zr = real(x);
                    1.0000000 11.0000000
                    0.0000000 3.0000000
                    2.0000000 1.0000000
See also
             complex, imag
```

recode

recode

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```
Purpose Changes the values of an existing vector from a vector of new values. Used in data transformations.
```

Format y = recode(x, e, v);

Input

Nx1 vector to be recoded (changed).

e NxK matrix of 1's and 0's.

v Kx1 vector containing the new values to be assigned to the recoded variable.

Output

Nx1 vector containing the recoded values of x.

Remarks

There should be no more than a single 1 in any row of e.

For any given row N of x and e, if the K^{th} column of e is 1, the K^{th} element of v will replace the original element of x.

If every column of e contains a 0, the original value of x will be unchanged.

Example

recode

Source datatran.src

See also code, substute

b

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recode (dataloop)

recode (dataloop)

Purpose

Changes the value of a variable with different values based on a set of logical expressions.

Format

```
recode [#] [[$]] var with
val_1 for expression_1,
val 2 for expression 2,
val_n for expression_n;
```

Input

var

literal, the new variable name. scalar, value to be used if corresponding expression is true. val expression logical scalar-returning expression that returns nonzero

TRUE or zero FALSE.

Remarks

If '\$' is specified, the variable will be considered a character variable. If '#' is specified, the variable will be considered numeric. If neither is specified, the type of the variable will be left unchanged.

The logical expressions must be mutually exclusive, that is only one may return TRUE for a given row (observation).

If none of the expressions is TRUE for a given row (observation), its value will remain unchanged.

Any variables referenced must already exist, either as elements of the source data set, as externs, or as the result of a previous make, vector, or code statement.

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x y z

recode (dataloop)

See also code

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recserar

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recserar

Purpose Computes a vector of autoregressive recursive series.

Format y = recserar(x, y0, a);

Input x NxK matrix

y0 PxK matrix.

a PxK matrix.

Output y NxK matrix containing the series.

Remarks recserar is particularly useful in dealing with time series.

Typically, the result would be thought of as *K* vectors of length *N*.

y0 contains the first P values of each of these vectors (thus, these are prespecified). The remaining elements are constructed by computing a P^{th} order "autoregressive" recursion, with weights given by a, and then by adding the result to the corresponding elements of x. That is, the t^{th} row of y is given by:

$$y[t, .] = x[t, .] + a[1, .] * y[t - 1, .] + ... + a[P, .] * y[t - P, .], t = P + 1, ..., N$$

and

$$y[t,.] = y0[t,.], t = 1,...,P$$

b = 1|2|3|4;

Note that the first *P* rows of *x* are not used.

Example n = 10;

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x y z

3-702

recserar

$$y = recserar(x*b+e,y0,rho);$$

In this example, two autoregressive series are formed using simulated data. The general form of the series can be written:

$$y[1,t] = rho[1,1]*y[1,t-1] + x[t,.]*b + e[1,t]$$

 $y[2,t] = rho[2,1]*y[2,t-1] + x[t,.]*b + e[2,t]$

The error terms (e[1,t] and e[2,t]) are not individually serially correlated, but they are contemporaneously correlated with each other. The variance-covariance matrix is sig.

See also recsercp, recserrc

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recsercp

recsercp

Purpose

Computes a recursive series involving products. Can be used to compute cumulative products, to evaluate polynomials using Horner's rule, and to convert from base b representations of numbers to decimal representations among other things.

Format

y = recsercp(x,z);

Input

NxK or 1xK matrix х

NxK or 1xK matrix. Z.

Output

NxK matrix in which each column is a series generated by a recursion of the form:

$$y(1) = x(1) + z(1)$$

$$y(t) = y(t-1) \times x(t) + z(t), t = 2, ...N$$

Remarks

The following GAUSS code could be used to emulate **recsercp** when the number of rows in x and z is the same:

```
n = rows(x); /* assume here that rows(z) */
              /* is also n */
y = zeros(n,1);
y[1,.] = x[1,.] + z[1,.];
i = 2i
do until i > n;
```

y[i,.] = y[i-1,.] .* x[i,.] + z[i,.];i = i + 1;

endo;

Note that K series can be computed simultaneously, since x and z can have K columns (they must both have the same number of columns).

recsercp allows either x or z to have only 1 row.

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x y z

recsercp

recsercp(x, θ) will produce the cumulative products of the elements in x.

Example

If \mathbf{x} is a scalar and \mathbf{c} is an (N+1)x1 vector, the result \mathbf{p} will contain the value of the polynomial whose coefficients are given in \mathbf{c} . That is:

$$p = c[1, .]. \times x^{n} + c[2, .]. \times x^{(n-1)} + ... + c[n+1, .]$$

Note that both \mathbf{x} and \mathbf{c} could contain more than 1 column, and then this code would evaluate the entire set of polynomials at the same time. Note also that if $\mathbf{x} = 2$, and if \mathbf{c} contains the digits of the binary representation of a number, then \mathbf{p} will be the decimal representation of that number.

See also

recserar, recserrc

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recserrc

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```
Purpose
               Computes a recursive series involving division.
 Format
               y = recserrc(x,z);
    Input
                       1xK or Kx1 vector.
               х
                       NxK matrix.
               Z.
 Output
                       NxK matrix in which each column is a series generated by
                       a recursion of the form:
                           y[1] = x \mod z[1], x = trunc(x / z[1])
                           y[2] = x \mod z[2], x = trunc(x / z[2])
                           y[3] = x \mod z[3], x = trunc(x / z[3])
```

Remarks Can be used to convert from decimal to other number systems (radix conversion).

 $y[n] = x \mod z[n]$

```
Example
           x = 2|8|10;
           b = 2i
           n = \max(\log(x)./\log(b)) + 1;
           z = reshape(b, n, rows(x));
           y = rev(recserrc(x, z))';
```

The result, y, will contain in its rows (note that it is transposed in the last step) the digits representing the decimal numbers 2, 8, and 10 in base 2:

```
0 0 1 0
1000
1 0 1 0
```

Source recserrc.src

See also recserar, recsercp

x y z

rerun

rerun

Purpose Displays the most recently created graphics file.

Library pgraph

Format rerun;

Portability DOS only

rerun invokes the graphics utility pagrun.exe.

Remarks rerun is used by the endwind function.

Source pcart.src

Globals _pcmdlin, _pnotify, _psilent, _ptek, _pzoom

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Purpose Reshapes a matrix.

Format y = reshape(x,r,c);

Input x NxK matrix.

r scalar, new row dimension.

c scalar, new column dimension.

Output y RxC matrix created from the elements of x.

Remarks Matrices are stored in row major order.

The first c elements are put into the first row of y, the second in the second row, and so on. If there are more elements in x than in y, the remaining elements are discarded. If there are not enough elements in x to fill y, then when **reshape** runs out of elements, it goes back to the first element of x and starts getting additional elements from there.

Example y = reshape(x, 2, 6);

If
$$x = \begin{pmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \end{pmatrix}$$
 then $y = \begin{pmatrix} 1 & 2 & 3 & 4 & 5 & 6 \\ 7 & 8 & 9 & 10 & 11 & 12 \end{pmatrix}$

If
$$x = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{pmatrix}$$
 then $y = \begin{pmatrix} 1 & 2 & 3 & 4 & 5 & 6 \\ 7 & 8 & 9 & 1 & 2 & 3 \end{pmatrix}$

If
$$x = \begin{pmatrix} 1 & 2 & 3 & 4 & 5 \\ 6 & 7 & 8 & 9 & 10 \end{pmatrix}$$
 then $y = \begin{pmatrix} 1 & 2 & 3 & 4 & 5 & 6 \\ 7 & 8 & 9 & 10 & 11 & 12 \end{pmatrix}$

If
$$x = \begin{pmatrix} 1 & 2 \\ 3 & 4 \end{pmatrix}$$
 then $y = \begin{pmatrix} 1 & 2 & 3 & 4 & 1 & 2 \\ 3 & 4 & 1 & 2 & 3 & 4 \end{pmatrix}$

If
$$x = 1$$
 then $y = \begin{bmatrix} 1 & 1 & 1 & 1 & 1 \\ 1 & 1 & 1 & 1 & 1 \end{bmatrix}$

reshape

See also submat, vec

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retp

Purpose Returns from a procedure or keyword.

Format retp;

retp(*x,y,...*);

Remarks For more details, see "Procedures and Keywords" in the *User's Guide*.

In a **retp** statement 0-1023 items may be returned. The items may be expressions. Items are separated by commas.

It is legal to return with no arguments, as long as the procedure is defined

to return 0 arguments.

See also proc, keyword, endp

return

return

Purpose Returns from a subroutine.

Format return;

return(*x*,*y*,...**)**;

Remarks The number of items that may be returned from a subroutine in a return

statement is limited only by stack space. The items may be expressions.

Items are separated by commas.

It is legal to return with no arguments and therefore return nothing.

See also gosub, pop

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Purpose Reverses the order of the rows in a matrix.

Format y = rev(x);

Input x NxK matrix.

Output y NxK matrix containing the reversed rows of x.

Remarks The first row of y will be where the last row of x was and the last row will be where the first was and so on. This can be used to put a sorted matrix in descending order.

Example x = round(rndn(5,3)*10);

y = rev(x);

10 7 8 7 4 –9

 $x = _{-11} \quad 0 \quad -3$

3 18 0

9 - 1 20

9 –1 20

3 18 0

 $y = -11 \quad 0 \quad -3$

7 4 –9

10 7 8

See also sortc

rfft

rfft

Purpose Computes a real 1- or 2-D Fast Fourier transform.

Format y = rfft(x);

Input x NxK real matrix.

Output y LxM matrix, where L and M are the smallest powers of 2 greater than or equal to N and K, respectively.

Remarks Computes the RFFT of x, scaled by 1/(L*M).

This uses a Temperton Fast Fourier algorithm.

If N or K is not a power of 2, x will be padded out with zeros before computing the transform.

Example $x = \{ 69, 81 \};$ y = rfft(x);y = 6.0000000 1.00000000

y = 0.5000000 - 1.50000000 1.50000000 - 2.50000000

See also rffti, fft, ffti, fftm, fftmi

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rffti

Purpose Computes inverse real 1- or 2-D Fast Fourier transform.

Format y = rffti(x);

Input x NxK matrix.

Output y LxM real matrix, where L and M are the smallest prime factor products greater than or equal to N and K.

Remarks It is up to the user to guarantee that the input will return a real result. If in doubt, use ffti.

Example $x = \{ 61, 1.5-2.5 \};$ y = rffti(x); $y = \frac{6.00000009.00000000}{8.00000001.00000000}$

See also rfft, fft, ffti, fftm, fftmi

w x y z

3-714

rfftip

Purpose Computes a

Computes an inverse real 1- or 2-D FFT. Takes a packed format FFT as input.

Format y = rfftip(x);

Input x NxK matrix or K-length vector.

Output y LxM real matrix or M-length vector.

Remarks rfftip assumes that its input is of the same form as that output by rfftp and rfftnp.

rfftip uses the Temperton prime factor FFT algorithm. This algorithm can compute the inverse FFT of any vector or matrix whose dimensions can be expressed as the product of selected prime number factors. GAUSS implements the Temperton algorithm for any integer power of 2, 3, and 5, and one factor of 7. Thus, **rfftip** can handle any matrix whose dimensions can be expressed as:

$$2^p \times 3^q \times 5^r \times 7^s$$
, $p, q, r \ge 0$
 $s = 0 \text{ or } 1$

If a dimension of x does not meet this requirement, it will be padded with zeros to the next allowable size before the inverse FFT is computed. Note that **rfftip** assumes the length (for vectors) or column dimension (for matrices) of x is K-1 rather than K, since the last element or column does not hold FFT information, but the Nyquist frequencies.

The sizes of *x* and *y* are related as follows: L will be the smallest prime factor product greater than or equal to N, and M will be twice the smallest prime factor product greater than or equal to K-1. This takes into account the fact that *x* contains both positive and negative frequencies in the row dimension (matrices only), but only positive frequencies, and those only in the first K-1 elements or columns, in the length or column dimension.

It is up to the user to guarantee that the input will return a real result. If in doubt, use **ffti**. Note, however, that **ffti** expects a full FFT, including negative frequency information, for input.

Do not pass **rfftip** the output from **rfft** or **rfftn**— it will return incorrect results. Use **rffti** with those routines.

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rfftip

See also fft, ffti, fftm, fftmi, fftn, rfft, rffti, rfftn, rfftnp, rfftp

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rfftn

Purpose Computes a real 1- or 2-D FFT.

Format y = rfftn(x);

Input x NxK real matrix.

can be expressed as:

Output y LxM matrix, where L and M are the smallest prime factor products greater than or equal to N and K, respectively.

Remarksrfftn uses the Temperton prime factor FFT algorithm. This algorithm can compute the FFT of any vector or matrix whose dimensions can be expressed as the product of selected prime number factors. GAUSS implements the Temperton algorithm for any power of 2, 3, and 5, and one factor of 7. Thus, rfftn can handle any matrix whose dimensions

 $p, q, r \ge 0$ for rows of matrix $p > 0, q, r \ge 0$ for columns of matrix $p > 0, q, r \ge 0$ for length of vector s = 0 or 1 for all dimensions

If a dimension of *x* does not meet these requirements, it will be padded with zeros to the next allowable size before the FFT is computed.

rfftn pads matrices to the next allowable size; however, it generally runs faster for matrices whose dimensions are highly composite numbers, i.e., products of several factors (to various powers), rather than powers of a single factor. For example, even though it is bigger, a 33600x1 vector can compute as much as 20 percent faster than a 32768x1 vector, because 33600 is a highly composite number, $2^6 \times 3 \times 5^2 \times 7$, whereas 32768 is a simple power of 2, 2^{15} . For this reason, you may want to hand-pad matrices to optimum dimensions before passing them to rfftn. The Run-Time Library includes two routines, optn and optnevn, for determining optimum dimensions. Use optn to determine optimum rows for matrices, and optnevn to determine optimum columns for matrices and optimum lengths for vectors.

The Run-Time Library also includes the **nextn** and **nextnevn** routines, for determining allowable dimensions for matrices and vectors.

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(You can use these to see the dimensions to which **rfftn** would pad a matrix or vector.)

rfftn scales the computed FFT by 1/(L*M).

See also fft, ffti, fftm, fftmi, fftn, rfft, rffti, rfftip, rfftnp, rfftp

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rfftnp

rfftnp

Purpose Computes a real 1- or 2-D FFT. Returns the results in a packed format.

Format y = rfftnp(x);

Input *x* NxK real matrix or K-length real vector.

Output y Lx(M/2+1) matrix or (M/2+1)-length vector, where L and M are the smallest prime factor products greater than or equal to N and K, respectively.

Remarks

For 1-D FFT's, **rfftnp** returns the positive frequencies in ascending order in the first M/2 elements, and the Nyquist frequency in the last element. For 2-D FFT's, **rfftnp** returns the positive and negative frequencies for the row dimension, and for the column dimension it returns the positive frequencies in ascending order in the first M/2 columns, and the Nyquist frequencies in the last column. Usually the FFT of a real function is calculated to find the power density spectrum or to perform filtering on the waveform. In both these cases only the positive frequencies are required. (See also **rfft** and **rfftn** for routines that return the negative frequencies as well.)

rfftnp uses the Temperton prime factor FFT algorithm. This algorithm can compute the FFT of any vector or matrix whose dimensions can be expressed as the product of selected prime number factors. GAUSS implements the Temperton algorithm for any power of 2, 3, and 5, and one factor of 7. Thus, **rfftnp** can handle any matrix whose dimensions can be expressed as:

$$2^{p} \times 3^{q} \times 5^{r} \times 7^{s}$$
, $p, q, r \ge 0$ for rows of matrix $p > 0, q, r \ge 0$ for columns of matrix $p > 0, q, r \ge 0$ for length of vector $s = 0$ or 1 for all dimensions

If a dimension of x does not meet these requirements, it will be padded with zeros to the next allowable size before the FFT is computed.

rfftnp pads matrices to the next allowable size; however, it generally runs faster for matrices whose dimensions are highly composite numbers, i.e., products of several factors (to various powers), rather than powers of

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a single factor. For example, even though it is bigger, a 33600x1 vector can compute as much as 20 percent faster than a 32768x1 vector, because 33600 is a highly composite number, $2^6 \times 3 \times 5^2 \times 7$, whereas 32768 is a simple power of 2, 2^{15} . For this reason, you may want to hand-pad matrices to optimum dimensions before passing them to **rfftnp**. The Run-Time Library includes two routines, **optn** and **optnevn**, for determining optimum dimensions. Use **optn** to determine optimum rows for matrices, and **optnevn** to determine optimum columns for matrices and optimum lengths for vectors.

The Run-Time Library also includes the **nextn** and **nextnevn** routines, for determining allowable dimensions for matrices and vectors. (You can use these to see the dimensions to which **rfftnp** would pad a matrix or vector.)

rfftnp scales the computed FFT by 1/(L*M).

See also fft, ffti, fftm, fftmi, fftn, rfft, rffti, rfftip, rfftn, rfftp

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rfftp

Purpose Computes a real 1- or 2-D FFT. Returns the results in a packed format.

Format y = rfftp(x);

Input *x* NxK real matrix or K-length real vector.

Output y Lx(M/2+1) matrix or (M/2+1)-length vector, where L and M are the smallest powers of 2 greater than or equal to N and K, respectively.

Remarks If a dimension of x is not a power of 2, it will be padded with zeros to the next allowable size before the FFT is computed.

For 1-D FFT's, rfftp returns the positive frequencies in ascending order in the first M/2 elements, and the Nyquist frequency in the last element. For 2-D FFT's, rfftp returns the positive and negative frequencies for the row dimension, and for the column dimension it returns the positive frequencies in ascending order in the first M/2 columns, and the Nyquist frequencies in the last column. Usually the FFT of a real function is calculated to find the power density spectrum or to perform filtering on the waveform. In both these cases only the positive frequencies are required. (See also rfft and rfftn for routines that return the negative frequencies as well.)

rfftp scales the computed FFT by 1/(L*M). rfftp uses the Temperton FFT algorithm.

See also fft, ffti, fftm, fftmi, fftn, rfft, rffti, rfftip, rfftn, rfftnp

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rndbeta

Purpose Computes pseudo-random numbers with beta distribution.

Format x = rndbeta(r,c,a,b);

Input *r* scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

a MxN matrix, ExE conformable with RxC resulting matrix, shape parameters for beta distribution.

b KxL matrix, ExE conformable with RxC resulting matrix, shape parameters for beta distribution.

Output *x* RxC matrix, beta distributed pseudo-random numbers.

Remarks The properties of the pseudo-random numbers in *x* are:

$$E(x) = a / (a + b)$$

$$Var(x) = a \times b / (a + b + 1) \times (a + b)^{2}$$

$$x > 0$$

$$x < 1$$

$$a > 0$$

$$b > 0$$

Source random.src

rndcon, rndmult, rndseed

rndcon, rndmult, rndseed

Purpose

Resets the parameters of the linear congruential random number generator that is the basis for **rndu**, **rndi** and **rndn**.

Format

Portability

Windows, UNIX, OS/2

Parameter default values and ranges:

seedtime(0),
$$0 < \text{seed} < 2^32$$
a 1664525 $0 < \text{a} < 2^32$ c 1013904223 $0 < \text{c} < 2^32$

Remarks

A linear congruential uniform random number generator is used by **rndu**, and is also called by **rndn**. These statements allow the parameters of this generator to be changed.

The procedure used to generate the uniform random numbers is as follows. First, the current "seed" is used to generate a new seed:

$$new_seed = (((a * seed) % 2^{32}) + c) % 2^{32}$$

(where % is the mod operator). Then a number between 0 and 1 is created by dividing the new seed by 2^{32} :

$$x = \text{new_seed} / 2^{32}$$

rndcon resets c.

rndmult resets a.

rndseed resets *seed*. This is the initial seed for the generator. The default is that GAUSS uses the clock to generate an initial seed when GAUSS is invoked.

GAUSS goes to the clock to seed the generator only when it is first started up. Therefore, if GAUSS is allowed to run for a long time, and if large numbers of random numbers are generated, there is a possibility of recycling (that is, the sequence of "random numbers" will repeat itself). However, the generator used has an extremely long cycle, and so that should not usually be a problem.

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rndcon, rndmult, rndseed

The parameters set by these commands remain in effect until new commands are encountered, or until GAUSS is restarted.

See also rndu, rndn, rndi, rndLCi, rndKMi

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rndgam

rndgam

Purpose Computes pseudo-random numbers with gamma distribution.

Format x = rndgam(r,c,alpha);

Input r scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

alpha MxN matrix, ExE conformable with RxC resulting matrix, shape parameters for gamma distribution.

Output *x* RxC matrix, gamma distributed pseudo-random numbers.

Remarks The properties of the pseudo-random numbers in x are:

$$E(x) = alpha$$

$$Var(x) = alpha$$

$$x > 0$$

$$alpha > 0$$

To generate **gamma** (*alpha*, *theta*) pseudo-random numbers where *theta* is a scale parameter, multiply the result of **rndgam** by *theta*. Thus:

$$z = \text{theta} * \text{rndgam(1,1,} alpha)$$

has the properties

$$E(z) = alpha \times theta$$

 $Var(z) = alpha \times theta^2$
 $z > 0$
 $alpha > 0$
 $theta > 0$

Source random.src

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Purpose Returns a matrix of random integers, $0 \le y \le 2^32$.

Format y = rndi(r,c);

Input r scalar, row dimension.

c scalar, column dimension.

Output y rxc matrix of random integers between 0 and $2^32 - 1$, inclusive.

Remarks *r* and *c* will be truncated to integers if necessary.

This generator is automatically seeded using the system clock when GAUSS first starts. However, that can be overridden using the **rndseed** statement or using **rndus**.

Each seed is generated from the preceding seed, using the formula

new_seed = (((
$$a * seed$$
) % 2^{32})+ c) % 2^{32}

where % is the mod operator. The new seeds are the values returned. The muliplicative constant and the additive constant may be changed using **rndmult** and **rndcon** respectively.

See also rndu, rndus, rndn, rndcon, rndmult

rndKMbeta

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rndKMbeta

Purpose Computes beta pseudo-random numbers.

Format $\{x, newstate\} = rndKMbeta(r,c,a,b,state);$

Input r scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

a rxc matrix, or rx1 vector, or 1xc vector, or scalar, first shape argument for beta distribution.

b rxc matrix, or rx1 vector, or 1xc vector, or scalar, second shape argument for beta distribution.

state scalar or 500x1 vector.

Scalar case:

state = starting seed value only. If -1, GAUSS computes the starting seed based on the system clock.

500x1 vector case:

state = the state vector returned from a previous call to one of the **rndKM** random number functions.

Output *x r*x*c* matrix, beta distributed random numbers.

*newstate 500x1 vector, the updated state.

Remarks The properties of the pseudo-random numbers in *x* are:

$$E(x) = \frac{a}{a+b}, Var(x) = \frac{(a*b)}{(a+b+1)*(a+b)^2}$$

r and c will be truncated to integers if necessary.

Source randkm.src

Technical NotesrndKMbeta uses the recur-with-carry KISS+Monster algorithm described in the rndKMi Technical Notes.

rndKMgam

rndKMgam

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Purpose Computes Gamma pseudo-random numbers. $\{x, newstate\} = rndKMgam(r,c,alpha,state);$ **Format** Input scalar, number of rows of resulting matrix. scalar, number of columns of resulting matrix. alpha rxc matrix, or rx1 vector, or 1xc vector, or scalar, shape argument for gamma distribution. scalar or 500x1 vector. state Scalar case: state = starting seed value only. If -1, GAUSS computes the starting seed based on the system clock. 500x1 vector case: state = the state vector returned from a previous call to one of the **rndKM** random number functions. **Output** rxc matrix, gamma distributed random numbers. newstate 500x1 vector, the updated state. Remarks The properties of the pseudo-random numbers in x are: E(x) = alpha, Var(x) = alphax > 0, alpha > 0. To generate gamma(alpha, theta) pseudo-random numbers where theta is a scale parameter, multiply the result of **rndgam** by *theta*. Thus z = theta * rndgam(1,1,alpha);has the properties $E(z) = alpha * theta, Var(z) = alpha * theta ^ 2$ z > 0, alpha > 0, theta > 0. r and c will be truncated to integers if necessary.

Source

randkm.src

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${\tt rndKMgam}$

Technical Notes

rndKMgam uses the recur-with-carry KISS+Monster algorithm described in the rndKMi Technical Notes.

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Purpose Returns a matrix of random integers, 0 \le y \le 2^32, and the state of the random number generator.
```

Format $\{ y, newstate \} = rndKMi(r,c,state);$

Input

r scalar, row dimension.

c scalar, column dimension.

state scalar or 500x1 vector.

Scalar case:

state = starting seed value. If -1, GAUSS computes the starting seed based on the system clock.

500x1 vector case:

state = the state vector returned from a previous call to one of the **rndKM** random number generators.

Output

rxc matrix of random integers between 0 and 2^32 - 1, inclusive.

newstate 500x1 vector, the updated state.

Remarks *r* and *c* will be truncated to integers if necessary.

Example

This example generates two thousand vectors of random integers, each with one million elements. The state of the random number generator after each iteration is used as an input to the next generation of random numbers.

```
state = 13;
n = 2000;
k = 1000000;
c = 0;
min = 2^32+1;
max = -1;
```

rndKMi

```
do while c < n;
    { y,state } = rndKMi(k,1,state);
    min = minc(min | minc(y));
    max = maxc(max | maxc(y));
    c = c + k;
endo;

print "min " min;
print "max " max;</pre>
```

See also rndKMn, rndKMu

Technical Notes

rndKMi generates random integers using a KISS+Monster algorithm developed by George Marsaglia. KISS initializes the sequence used in the recur-with-carry Monster random number generator. For more information on this generator see http://www.Aptech.com/random.

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rndKMn

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Purpose Returns a matrix of standard normal (pseudo) random variables and the state of the random number generator.

Format { y, newstate } = rndKMn(r,c,state);
Input r scalar, row dimension.

c scalar, column dimension. state scalar or 500x1 vector.

Scalar case:

state = starting seed value. If -1, GAUSS computes the starting seed based on the system clock.

500x1 vector case:

state = the *state* vector returned from a previous call to one of the **rndKM** random number generators.

Output y rxc matrix of standard normal random numbers.

newstate 500x1 vector, the updated state.

Remarks r and c will be truncated to integers if necessary.

This example generates two thousand vectors of standard normal random numbers, each with one million elements. The state of the random number generator after each iteration is used as an input to the next generation of random numbers.

```
state = 13;
n = 2000;
k = 1000000;
c = 0;
submean = {};
```

rndKMn

```
do while c < n;
    { y,state } = rndKMn(k,1,state);
    submean = submean | meanc(y);
    c = c + k;
endo;

mean = meanc(submean);
print mean;</pre>
```

See also

rndKMu, rndKMi

Technical Notes

rndKMn calls the uniform random number generator that is the basis for rndKMu multiple times for each normal random number generated. This is the recur-with-carry KISS+Monster algorithm described in the rndKMi Technical Notes. Potential normal random numbers are filtered using the fast acceptance-rejection algorithm proposed by Kinderman, A.J. and J.G. Ramage, "Computer Generation of Normal Random Numbers," Journal of the American Statistical Association, December 1976, Volume 71, Number 356, pp. 893-896.

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rndKMnb

rndKMnb

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Purpose Computes negative binomial pseudo-random numbers.

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Format $\{x, newstate\} = rndKMnb(r,c,k,p,state);$

d

r scalar, number of rows of resulting matrix.

Input

c scalar, number of columns of resulting matrix.

k

*r*X*c* matrix, or *r*X1 vector, or 1X*c* vector, or scalar, "event" argument for negative binomial distribution.

p

rxc matrix, or rx1 vector, or 1xc vector, or scalar, "probability"

_

argument for negative binomial distribution.

state

scalar or 500x1 vector.

Scalar case:

state = starting seed value only. If -1, GAUSS computes the starting seed based on the system clock.

500x1 vector case:

state = the state vector returned from a previous call to one of
the rndKM random number functions.

Output

x rxc matrix, negative binomial distributed random numbers.

newstate 500x1 vector, the updated state.

Remarks

The properties of the pseudo-random numbers in x are:

$$E(x) = \frac{k^*p}{(1-p)}, Var(x) = \frac{k^*p}{(1-p)^2}$$

$$x = 0, 1, ..., k > 0, 0$$

r and c will be truncated to integers if necessary.

Source

randkm.src

Technical Notes rndKMnb uses the recur-with-carry KISS+Monster algorithm described in the rndKMi Technical Notes.

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rndKMp

Purpose Computes Poisson pseudo-random numbers.

Format $\{x, newstate\} = rndKMp(r,c,lambda,state);$

Input r scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

lambda rxc matrix, or rx1 vector, or 1xc vector, or scalar, shape

argument for Poisson distribution.

state scalar or 500x1 vector.

Scalar case:

state = starting seed value only. If -1, GAUSS computes the

starting seed based on the system clock.

500x1 vector case:

state = the state vector returned from a previous call to one of

the **rndKM** random number functions.

Output x rxc matrix, Poisson distributed random numbers.

newstate 500x1 vector, the updated state.

Remarks The properties of the pseudo-random numbers in *x* are:

E(x) = lambda, Var(x) = lambda

x = 0, 1,, lambda > 0.

r and c will be truncated to integers if necessary.

Source randkm.src

Technical notes rndKMp uses the recur-with-carry KISS+Monster algorithm described in the rndKMi. Technical Notes.

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Purpose Returns a matrix of uniform (pseudo) random variables and the state of the random number generator.

Format

$$\{ y, newstate \} = rndKMu(r,c,state);$$

Input

r scalar, row dimension.

c scalar, column dimension.

scalar, 2x1 vector, or 500x1 vector.

Scalar case:

state = starting seed value. If -1, GAUSS computes the starting seed based on the system clock.

2x1 vector case:

[1] the starting seed, uses the system clock if -1

[2] 0 for $0 \le y < 1$ 1 for $0 \le y \le 1$

500x1 vector case:

state = the state vector returned from a previous call to one ofthe rndKM random number generators.

Output

rxc matrix of uniform random numbers, $0 \le y < 1$.

newstate 500x1 vector, the updated state.

Remarks

r and c will be truncated to integers if necessary.

Example

This example generates two thousand vectors of uniform random numbers, each with one million elements. The state of the random number generator after each iteration is used as an input to the next generation of random numbers.

```
state = 13;
n = 2000;
k = 1000000;
c = 0;
submean = {};
```

rndKMu

```
do while c < n;
    { y,state } = rndKMu(k,1,state);
    submean = submean | meanc(y);
    c = c + k;
endo;

mean = meanc(submean);
print 0.5-mean;</pre>
```

See also

rndKMn, rndKMi

Technical Notes

rndKMu uses the recur-with-carry KISS-Monster algorithm described in the **rndKMi** Technical Notes. Random integer seeds from 0 to 2^32-1 are generated. Each integer is divided by 2^32 or 2^32-1.

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rndKMvm

Purpose	Computes von Mises pseudo-random numbers.				
Format	$\{x, newstate\} = rndKMvm(r,c,m,k,state);$				
Input	r c m k state	scalar, number of rows of resulting matrix. scalar, number of columns of resulting matrix. rxc matrix, or rx1 vector, or 1xc vector, or scalar, means for vm distribution. rxc matrix, or rx1 vector, or 1xc vector, or scalar, shape argument for vm distribution. scalar or 500x1 vector. Scalar case: state = starting seed value only. If -1, GAUSS computes the starting seed based on the system clock. 500x1 vector case:			
		state = the state vector returned from a previous call to one of the rndkm random number functions.			
Output	x newstate	rxc matrix, von Mises distributed random numbers.500x1 vector, the updated state.			
Remarks	r and c will be truncated to integers if necessary.				
Source	randkm.src				
Technical Notes	<pre>rndKMvm uses the recur-with-carry KISS+Monster algorithm described in the rndKMi Technical Notes.</pre>				

rndLCbeta

rndLCbeta

Purpose Computes beta pseudo-random numbers.

Format $\{x, newstate\} = rndLCbeta(r, c, a, b, state);$

Input

- scalar, number of rows of resulting matrix.
- c scalar, number of columns of resulting matrix.
- a rxc matrix, or rx1 vector, or 1xc vector, or scalar, first shape argument for beta distribution.
- b rxc matrix, or rx1 vector, or 1xc vector, or scalar, second shape argument for beta distribution.

state scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function.

Scalar case:

state = starting seed value only. System default values are used for the additive and multiplicative constants.

The defaults are 1013904223, and 1664525, respectively. These may be changed with **rndcon** and **rndmult**.

if *state* = -1, GAUSS computes the starting seed based on the system clock.

3x1 vector case:

- [1] the starting seed, uses the system clock if -1
- [2] the multiplicative constant
- [3] the additive constant

Output

rxc matrix, beta distributed random numbers.

newstate 4x1 vector:

- [1] the updated seed
- [2] the multiplicative constant
- [3] the additive constant
- [4] the original initialization seed

Remarks: The properties of the pseudo-random numbers in *x* are:

$$E(x) = \frac{a}{a+b}, Var(x) = \frac{(a*b)}{(a+b+1)*(a+b)^2}$$

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rndLCbeta

r and c will be truncated to integers if necessary.

Source

randlc.src

Technical Notes

This function uses a linear congruential method, discussed in Kennedy, W.J. Jr., and J.E. Gentle, Statistical Computing, Marcel Dekker, Inc. 1980, pp. 136-147. Each seed is generated from the preceding seed using the formula

new_seed = (((
$$a * seed$$
) % 2^{32})+ c) % 2^{32}

where % is the mod operator and where a is the multiplicative constant and c is the additive constant.

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rndLCgam

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Purpose	Computes Gamma pseudo-random numbers.						
Format	$\{x, newstate\} = rndLCgam(r,c,alpha,state);$						
Input	r	scalar n	umber of rows of resulting matrix.				
	c		umber of columns of resulting matrix.				
	alpha	rxc matrix, or rx1 vector, or 1xc vector, or scalar, shape argument for gamma distribution.					
	state	scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function.					
		Scalar case:					
		<i>state</i> = starting seed value only. System default values are used for the additive and multiplicative constants.					
		The defaults are 1013904223, and 1664525, respectively. These may be changed with rndcon and rndmult .					
		if <i>state</i> = -1, GAUSS computes the starting seed based on the system clock.					
		3x1 vector case:					
		[1]	the starting seed, uses the system clock if -1				
		[2]	the multiplicative constant				
		[3]	the additive constant				
Output	x	rxc mat	rix, gamma distributed random numbers.				
	newstate						
		[1]	the updated seed				
		[2]	the multiplicative constant				
		[3]	the additive constant				
		[4]	the original initialization seed				
Remarks	The prop	the pseudo-random numbers in x are:					
	E(x) = alpha, Var(x) = alpha						
	x > 0,	x > 0, $alpha > 0$.					

To generate gamma (*alpha*, *theta*) pseudo-random numbers where *theta* is a scale parameter, multiply the result of **rndgam** by *theta*.

rndLCgam

Thus

$$z = theta * rndgam(1,1,alpha);$$

has the properties

$$E(z) = alpha * theta, Var(z) = alpha * theta ^ 2$$

$$z > 0$$
, $alpha > 0$, $theta > 0$.

r and c will be truncated to integers if necessary.

Source

randlc.src

Technical Notes

This function uses a linear congruential method, discussed in Kennedy, W.J. Jr., and J.E. Gentle, Statistical Computing, Marcel Dekker, Inc. 1980, pp. 136-147. Each seed is generated from the preceding seed using the formula

$$new_seed = (((a * seed) % 2^{32}) + c) % 2^{32}$$

where % is the mod operator and where a is the multiplicative constant and c is the additive constant.

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rndLCi

rndLCi

Purpose Returns a matrix of random integers, $0 \le y \le 2^32$, and the state of the random number generator.

Format { y, newstate } = rndLCi(r,c,state);

Input r scalar, row dimension.

c scalar, column dimension.

state scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function.

Scalar case:

state = starting seed value only. System default values are used for the additive and multiplicative constants.

The defaults are 1013904223, and 1664525, respectively. These may be changed with **rndcon** and **rndmult**.

if *state* < 0, GAUSS computes the starting seed based on the system clock.

3x1 vector case:

- [1] the starting seed, uses the system clock if < 0
- [2] the multiplicative constant
- [3] the additive constant

Output y rxc matrix of random integers between 0 and 2^32 - 1, inclusive.

newstate 4x1 vector:

- [1] the updated seed
- [2] the multiplicative constant
- [3] the additive constant
- [4] the original initialization seed

Remarks r and c will be truncated to integers if necessary.

Each seed is generated from the preceding seed, using the formula

$$new_seed = (((a * seed) % 2^{32}) + c) % 2^{32}$$

where % is the mod operator and where a is the multiplicative constant and c is the additive constant. The new seeds are the values returned.

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Example state = 13;n = 2000000000;k = 1000000;c = 0; $min = 2^32+1;$ $\max = -1;$ do while c < n; { y,state } = rndLCi(k,1,state); min = minc(min | minc(y)); $max = maxc(max \mid maxc(y));$ c = c + k;endo; print "min " min; print "max " max; See also rndLCn, rndLCu, rndcon, rndmult

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rndLCn

Purpose	Returns a matrix of standard normal (pseudo) random variables and the state of the random number generator.					
Format	$\{ y, newstate \} = rndLCn(r,c,state);$					
Input	r	scalar, ro	ow dimension.			
_	c	scalar, column dimension.				
	state	scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function.				
		Scalar case:				
		<i>state</i> = starting seed value only. System default values are used for the additive and multiplicative constants.				
		The defaults are 1013904223, and 1664525, respectively. These may be changed with rndcon and rndmult .				
		if <i>state</i> < 0, GAUSS computes the starting seed based on the system clock.				
		3x1 vector case:				
		[1]	the starting seed, uses the system clock if < 0			
		[2]	the multiplicative constant			
		[3]	the additive constant			
Output	y	<i>rxc</i> matrix of standard normal random numbers.				
	newstate	e 4x1 vector:				
		[1]	the updated seed			
		[2]	the multiplicative constant			
		[3]	the additive constant			
		[4]	the original initialization seed			
Remarks	r and c will be truncated to integers if necessary.					
Example	state	state = 13; n = 200000000;				
	n = 20					
	k = 1000000;					
	c = 0;					

rndLCn

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submean = {};

do while c < n;
    { y,state } = rndLCn(k,1,state);
    submean = submean | meanc(y);
    c = c + k;
endo;

mean = meanc(submean);
print mean;</pre>
```

See also

rndLCu, rndLCi, rndcon, rndmult

Technical Notes

The normal random number generator is based on the uniform random number generator, using the fast acceptance-rejection algorithm proposed by Kinderman, A.J. and J.G. Ramage, "Computer Generation of Normal Random Numbers," Journal of the American Statistical Association, December 1976, Volume 71, Number 356, pp. 893-896. This algorithm calls the linear congruential uniform random number generator multiple times for each normal random number generated. See **rndlCu** for a description of the uniform random number generator algorithm.

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rndLCnb

rndLCnb

Purpose Computes negative binomial pseudo-random numbers.

Format $\{x, newstate\} = rndLCnb(r,c,k,p,state);$

Input *r* scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

k rxc matrix, or rx1 vector, or 1xc vector, or scalar, "event" argument for negative binomial distribution.

*p r*X*c* matrix, or *r*X1 vector, or 1X*c* vector, or scalar, "probability" argument for negative binomial distribution.

state scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function.

Scalar case:

state = starting seed value only. System default values are used for the additive and multiplicative constants.

The defaults are 1013904223, and 1664525, respectively. These may be changed with **rndcon** and **rndmult**.

if *state* = -1, GAUSS computes the starting seed based on the system clock.

3x1 vector case:

- [1] the starting seed, uses the system clock if -1
- [2] the multiplicative constant
- [3] the additive constant

Output x rxc matrix, negative binomial distributed random numbers.

newstate 4x1 vector:

- [1] the updated seed
- [2] the multiplicative constant
- [3] the additive constant
- [4] the original initialization seed

Remarks The properties of the pseudo-random numbers in *x* are:

$$E(x) = \frac{k^*p}{(1-p)}, Var(x) = \frac{k^*p}{(1-p)^2}$$

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rndLCnb

$$x = 0, 1, ..., k > 0, 0$$

r and c will be truncated to integers if necessary.

Source

randlc.src

Technical Notes

This function uses a linear congruential method, discussed in Kennedy, W.J. Jr., and J.E. Gentle, Statistical Computing, Marcel Dekker, Inc. 1980, pp. 136-147. Each seed is generated from the preceding seed using the formula

new_seed = (((
$$a * seed$$
) % 2^{32})+ c) % 2^{32}

where % is the mod operator and where a is the multiplicative constant and c is the additive constant.

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rndLCp

Purpose Computes Poisson pseudo-random numbers. **Format** $\{x, newstate\} = rndLCp(r,c,lambda,state);$ Input scalar, number of rows of resulting matrix. scalar, number of columns of resulting matrix. c lambda rxc matrix, or rx1 vector, or 1xc vector, or scalar, shape argument for Poisson distribution. scalar, 3x1 vector, or a 4x1 state vector from a previous call to state the function. Scalar case: state = starting seed value only. System default values are used for the additive and multiplicative constants. The defaults are 1013904223, and 1664525, respectively. These may be changed with **rndcon** and **rndmult**. if *state* = -1, GAUSS computes the starting seed based on the system clock. 3x1 vector case: [1] the starting seed, uses the system clock if -1 [2] the multiplicative constant [3] the additive constant **Output** rxc matrix, Poisson distributed random numbers. newstate 4x1 vector: [1] the updated seed [2] the multiplicative constant [3] the additive constant [4] the original initialization seed Remarks The properties of the pseudo-random numbers in x are: E(x) = lambda, Var(x) = lambdax = 0, 1,, lambda > 0.r and c will be truncated to integers if necessary.

rndLCp

Source

randlc.src

Technical Notes

This function uses a linear congruential method, discussed in Kennedy, W.J. Jr., and J.E. Gentle, Statistical Computing, Marcel Dekker, Inc. 1980, pp. 136-147. Each seed is generated from the preceding seed using the formula

new_seed = (((
$$a * seed$$
) % 2^{32})+ c) % 2^{32}

where % is the mod operator and where a is the multiplicative constant and c is the additive constant.

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rndLCu

rndLCu

Purpose Returns a matrix of uniform (pseudo) random variables and the state of the random number generator.

Format { y, newstate } = rndLCu(r,c,state);

Input

- r scalar, row dimension.
- c scalar, column dimension.

scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function.

Scalar case:

state = starting seed value only. System default values are used for the additive and multiplicative constants.

The defaults are 1013904223, and 1664525, respectively. These may be changed with **rndcon** and **rndmult**.

if *state* < 0, GAUSS computes the starting seed based on the system clock.

3x1 vector case:

- [1] the starting seed, uses the system clock if < 0
- [2] the multiplicative constant
- [3] the additive constant

Output

y rxc matrix of uniform random numbers, $0 \le y < 1$.

newstate 4x1 vector:

- [1] the updated seed
- [2] the multiplicative constant
- [3] the additive constant
- [4] the original initialization seed

Remarks

r and c will be truncated to integers if necessary.

Each seed is generated from the preceding seed, using the formula

new_seed = (((
$$a * seed$$
) % 2^{32})+ c) % 2^{32}

where % is the mod operator and where a is the multiplicative constant and c is the additive constant. A number between 0 and 1 is created by dividing new_seed by 2^32.

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rndLCu

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Example state = 13;n = 2000000000;k = 1000000;c = 0;submean = {}; do while c < n; { y,state } = rndLCu(k,1,state); submean = submean | meanc(y); c = c + k;endo; mean = meanc(submean); print 0.5-mean; See also rndLCn, rndLCi, rndcon, rndmult This function uses a linear congruential method, discussed in Kennedy, **Technical** W. J. Jr., and J. E. Gentle, Statistical Computing, Marcel Dekker, Inc., **Notes** 1980, pp. 136-147.

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rndLCvm

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w x y z

rndLCvm

Purpose	Computes von Mises pseudo-random numbers.				
Format	$\{x, newstate\} = rndLCvm(r,c,m,k,state);$				
Input	r c m k state	scalar, number of rows of resulting matrix. scalar, number of columns of resulting matrix. rxc matrix, or rx1 vector, or 1xc vector, or scalar, means for vm distribution. rxc matrix, or rx1 vector, or 1xc vector, or scalar, shape argument for vm distribution. scalar, 3x1 vector, or a 4x1 state vector from a previous call to the function. Scalar case: state = starting seed value only. System default values are used for the additive and multiplicative constants. The defaults are 1013904223, and 1664525, respectively. These may be changed with rndcon and rndmult.			
		 if state = -1, GAUSS computes the starting seed based on the system clock. 3x1 vector case: [1] the starting seed, uses the system clock if -1 [2] the multiplicative constant [3] the additive constant 	ie		
Output	x newstate	 rxc matrix, von Mises distributed random numbers. 4x1 vector: [1] the updated seed [2] the multiplicative constant [3] the additive constant [4] the original initialization seed 			
Remarks Source	r and c w	ll be truncated to integers if necessary.	i		

rndLCvm

Technical Notes

This function uses a linear congruential method, discussed in Kennedy, W.J. Jr., and J.E. Gentle, Statistical Computing, Marcel Dekker, Inc. 1980, pp. 136-147. Each seed is generated from the preceding seed using the formula

$$new_seed = (((a * seed) % 2^{32}) + c) % 2^{32}$$

where % is the mod operator and where a is the multiplicative constant and c is the additive constant.

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rndn

rndn

Purpose Creates a matrix of standard Normal (pseudo) random numbers.

Format y = rndn(r,c);

Input r scalar, row dimension.

c scalar, column dimension.

Output y RxC matrix of Normal random numbers having a mean of 0 and standard deviation of 1.

Remarks r and c will be truncated to integers if necessary.

The Normal random number generator is based upon the uniform random number generator. To reseed them both, use the **rndseed** statement. The other parameters of the uniform generator can be changed using **rndcon**, **rndmod**, and **rndmult**.

Example x = rndn(8100,1);

m = meanc(x);
s = stdc(x);

m = 0.002810s = 0.997087

In this example, a sample of 8100 Normal random numbers is drawn, and the mean and standard deviation are computed for the sample.

See also rndu, rndcon

Technical Notes This function uses the fast acceptance-rejection algorithm proposed by Kinderman, A. J., and J. G. Ramage. "Computer Generation of Normal Random Numbers." *Journal of the American Statistical Association*. Vol. 71 No. 356, Dec. 1976, 893-96.

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Purpose Computes pseudo-random numbers with negative binomial distribution.

Format

x = rndnb(r,c,k,p);

Input

r scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

k MxN matrix, ExE conformable with RxC resulting matrix, "event" parameters for negative binomial distribution.

p KxL matrix, ExE conformable with RxC resulting matrix, probability parameters for negative binomial distribution.

Output

x RxC matrix, negative binomial distributed pseudo-random numbers.

Remarks

The properties of the pseudo-random numbers in x are:

$$E(x) = k \times p / (1-p)$$

$$Var(x) = k \times p / (1-p)^2$$

$$x = 0, 1, 2, ..., k$$

Source

random.src

rndp

rndp

Purpose Computes pseudo-random numbers with Poisson distribution.

Format x = rndp(r,c,lambda);

Input r scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

lambda MxN matrix, ExE conformable with RxC resulting matrix,

shape parameters for Poisson distribution.

Output x RxC matrix, Poisson distributed pseudo-random numbers.

Remarks The properties of the pseudo-random numbers in *x* are:

Ex = lambda

Var(x) = lambda

x = 0, 1, 2, ...

lambda > 0

Source random.src

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rndu

rndu

Purpose Creates a matrix of uniform (pseudo) random variables. h **Format** y = rndu(r,c);Input d scalar, row dimension. cscalar, column dimension. **Output** RxC matrix of uniform random variables between 0 and 1. Remarks r and c will be truncated to integers if necessary. This generator is automatically seeded using the clock when GAUSS is first started. However, that can be overridden using the rndseed statement or by using rndus. The seed is automatically updated as a random number is generated (see above under **rndcon**). Thus, if GAUSS is allowed to run for a long time, and if large numbers of random numbers are generated, there is a possibility of recycling. This is a 32-bit generator, though, so the range is sufficient for most applications. **Example** x = rndu(8100,1);y = meanc(x);0 z = stdc(x);p q y = 0.500205z = 0.289197In this example, a sample of 8100 uniform random numbers is generated, and the mean and standard deviation are computed for the sample. See also rndn, rndcon, rndmod, rndmult, rndseed u Technical This function uses a multiplicative-congruential method. This method is V discussed in Kennedy, W.J., Jr., and J.E. Gentle. Statistical Computing. **Notes** Marcel Dekker, Inc., NY, 1980, 136-147. W

x y z

rndvm

rndvm

Purpose Computes von Mises pseudo-random numbers.

Format x = rndvm(r,c,m,k);

Input *r* scalar, number of rows of resulting matrix.

c scalar, number of columns of resulting matrix.

m NxK matrix, ExE conformable with rxc, means for von Mises distribution.

k LxM matrix, ExE conformable with *r*x*c*, shape arrgument for von Mises distribution.

Output x rxc matrix, von Mises distributed random numbers.

Source random.src

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rotater

rotater

Purpose

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Rotates the rows of a matrix.

Format

y = rotater(x,r);

Input

x NxK matrix to be rotated.

r Nx1 or 1x1 matrix specifying the amount of rotation.

Output

y NxK rotated matrix.

Remarks

The rotation is performed horizontally within each row of the matrix. A positive rotation value will cause the elements to move to the right. A negative rotation value will cause the elements to move to the left. In either case, the elements that are pushed off the end of the row will wrap around to the opposite end of the same row.

If the rotation value is greater than or equal to the number of columns in x, then the rotation value will be calculated using (r % cols(x)).

Example

y = rotater(x,r);

If
$$x = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \end{pmatrix}$$
 and $r = \begin{pmatrix} 1 \\ -1 \end{pmatrix}$ Then $y = \begin{pmatrix} 3 & 1 & 2 \\ 5 & 6 & 4 \end{pmatrix}$

If
$$x = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{pmatrix}$$
 and $r = \begin{pmatrix} 1 \\ 2 \end{pmatrix}$ Then $y = \begin{pmatrix} 6 & 4 & 5 \\ 8 & 9 & 7 \\ 10 & 11 & 12 \end{pmatrix}$

See also

shiftr

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round

round

Purpose Rounds to the nearest integer.

Format y = round(x);

Input x NxK matrix.

Output y NxK matrix containing the rounded elements of x.

Example let $x = \{ 77.68 -14.10, 4.73 -158.88 \};$ y = round(x);

$$y = \begin{array}{ccc} 78 & -14 \\ 5 & -159 \end{array}$$

See also trunc, floor, ceil

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rows

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Returns the number of rows in a matrix. Purpose **Format** y = rows(x);Input NxK matrix. Output scalar, number of rows in the specified matrix. Remarks If x is an empty matrix, rows(x) and cols(x) return 0. Example x = ones(3,5);y = rows(x);1 1 1 1 1 x =11111 1 1 1 1 1 y = 3See also cols, show

rowsf

rowsf

```
Returns the number of rows in a GAUSS data set (.dat) file or GAUSS
Purpose
             matrix (.fmt) file.
 Format
             y = rowsf(f);
    Input
                    file handle of an open file.
  Output
                    scalar, number of rows in the specified file.
Example
            open fp = myfile;
             r = rowsf(fp);
             c = colsf(fp);
See also
             colsf, open, typef
```

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Purpose Computes the reduced row echelon form of a matrix.

Format y = rref(x);

Input x MxN matrix.

Output y MxN matrix containing reduced row echelon form of x.

Remarks The tolerance used for zeroing elements is computed inside the procedure using:

tol =
$$maxc(m|n)$$
 * eps * $maxc(abs(sumc(x')))$;

where **eps** = 2.24e-16;

This procedure can be used to find the rank of a matrix. It is not as stable numerically as the singular value decomposition (which is used in the rank function), but it is faster for large matrices.

There is some speed advantage in having the number of rows be greater than the number of columns, so you may want to transpose if all you care about is the rank.

The following code can be used to compute the rank of a matrix:

where *y* is the output from **rref**, and **tol** is the tolerance used. This finds the number of rows with any nonzero elements, which gives the rank of the matrix, disregarding numeric problems.

Example

let
$$x[3,3] = 1 2 3$$

 $4 5 6$
 $7 8 9;$

y = rref(x);

$$y = \begin{array}{ccc} 1 & 0 & -1 \\ 0 & 1 & 2 \\ 0 & 0 & 0 \end{array}$$

Source rref.src

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x y z

run

Purpose Runs a source code or compiled code program.

Format run filename;

Input *filename* literal or ^string, name of file to run.

Remarks

The filename can be any legal file name. Filename extensions can be whatever you want, except for the compiled file extension, .gcg. Pathnames are okay. If the name is to be taken from a string variable, then the name of the string variable must be preceded by the ^ (caret) operator.

The **run** statement can be used both from the command line and within a program. If used in a program, once control is given to another program through the **run** statement there is no return to the original program.

If you specify a filename without an extension, GAUSS will first look for a compiled code program (i.e., a .gcg file) by that name, then a source code program by that name. For example, if you enter

run dog;

GAUSS will first look for the compiled code file dog.gcg, and run that if it finds it. If GAUSS cannot find dog.gcg, it will then look for the source code file dog with no extension.

If a path is specified for the file, then no additional searching will be attempted if the file is not found.

If a path is not specified the current directory will be searched first, then each directory listed in **src_path**. The first instance found is run. **src_path** is defined in gauss.cfg.

run /gauss/myprog.prc; No additional search will be made if the file is not found.

run myprog.prc;
The directories listed in **src_path**will be searched for myprog.prc if
the file is not found in the current

directory.

Programs can also be run by typing the filename on the OS command line when starting GAUSS.

Example Example 1

run myprog.prg;

run

Example 2

name = "myprog.prg";
run ^name;

See also #include

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save

ave					
Purpose	Saves matrices, strings, or procedures to a disk file.				
Format	<pre>save [[vflag]] [[path=path]] x, [[lpath=]]y;</pre>				
Input	vflag	version flag.			
		-v89 supported on DOS, OS/2, Windows			
		-v92 supported on UNIX, Windows			
		-v96 supported on all platforms			
	See also "File I/O" in the <i>User's Guide</i> for details various versions. The default format can be specigauss.cfg by setting the dat_fmt_versio configuration variable. If dat_fmt_version idefault is v96 .				
	path	literal or 'string, a default path to use for this and subsequent save s.			
	x	a symbol name, the name of the file the symbol will be saved in is the same as this with the proper extension added for the type of the symbol.			
	lpath	literal or 'string, a local path and filename to be used for a particular symbol. This path will override the path previously set and the filename will override the name of the symbol being saved. The extension cannot be overridden.			
	у	the symbol to be saved to <i>lpath</i> .			
Remarks	save can be used to save matrices, strings, procedures, and functions. Procedures and functions must be compiled and resident in memory before they can be save 'd.				
	The following extensions will be given to files that are saved:				
	matrix string procedure function	.fmt .fst e .fcg .fcg			
	keyword	.fcg			

if the path= subcommand is used with save, the path string will be remembered until changed in a subsequent command. This path will be

save

used whenever none is specified. The save path can be overridden in any particular save by specifying an explicit path and filename.

Example

```
spath = "/gauss";
save path = ^spath x,y,z;
```

Save **x**, **y**, and **z** using /gauss as a path. This path will be used for the next save if none is specified.

```
svp = "/gauss/data";
save path = ^svp n, k, /gauss/quad1=quad;
```

n and k will be saved using /gauss/data as the save path, quad will be saved in /gauss with the name quad1.fmt. On platforms that use the backslash as the path separator, the double backslash is required inside double quotes to get a backslash, because it is the escape character in quoted strings. It is not required when specifying literals.

```
save path=/procs;
```

Changes save path to /procs.

```
save path = /miscdata;
save /data/mydata1 = x, y, hisdata = z;
```

In the above program:

```
x would be saved in /data/mydata1.fmt
```

y would be saved in /miscdata/y.fmt

z would be saved in /miscdata/hisdata.fmt

See also

load, saveall, saved

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saveall

saveall

Purpose

Saves the current state of the machine to a compiled file. All procedures, global matrices and strings will be saved.

Format

saveall fname;

Input

fname literal or ^string, the path and filename of the compiled file to be created.

Remarks

The file extension will be .gcg.

A file will be created containing all your matrices, strings, and procedures. No main code segment will be saved. This just means it will be a .gcg file with no main program code (see compile). The rest of the contents of memory will be saved including all global matrices, strings, functions and procedures. Local variables are not saved. This can be used inside a program to take a snapshot of the state of your global variables and procedures. To reload the compiled image use run or use.

```
library pgraph;
external proc xy,logx,logy,loglog,hist;
saveall pgraph;
```

This would create a file called pgraph.gcg containing all the procedures, strings and matrices needed to run Publication Quality Graphics programs. Other programs could be compiled very quickly with the following statement at the top of each:

```
use pgraph;
```

See also

compile, run, use

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a **Purpose** Writes a matrix in memory to a GAUSS data set on disk. h **Format** y = saved(x, dataset, vnames);Input х NxK matrix to save in .dat file. d dataset string, name of data set. е vnames string or Kx1 character vector, names for the columns of the data set. **Output** scalar, 1 if successful, 0 if fail. Remarks If *dataset* is null or 0, the data set name will be temp.dat. h if *vnames* is a null or 0, the variable names will begin with "X" and be numbered 1-K. If *vnames* is a string or has fewer elements than x has columns, it will be expanded as explained under create. k The output data type is double precision. Example x = rndn(100,3);m dataset = "mydata"; vnames = { height, weight, age }; 0 if not saved(x,dataset,vnames); p errorlog "Write error"; q end; endif; Source saveload.src t See also loadd, writer, create u V

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savewind

savewind

Purpose Saves the current graphic panel configuration to a file.

Library pgraph

Format err = savewind(filename);

Input *filename* Name of file.

Output *err* scalar, 0 if successful, 1 if graphic panel matrix is invalid.

Note that the file is written in either case.

Remarks See the discussion on using graphic panels in "Publication Quality

Graphics" in the *User's Guide*.

Source pwindow.src

See also loadwind

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scale

scale

a **Purpose** h Library **Format** Input Remarks h m 0 p Source q See also u V

Fixes the scaling for subsequent graphs. The axes endpoints and increments are computed as a best guess based on the data passed to it.

pgraph

scale(x,y);

matrix, the X axis data.

matrix, the Y axis data. y

x and y must each have at least 2 elements. Only the minimum and maximum values are necessary.

This routine fixes the scaling for all subsequent graphs until graphset is called. This also clears **xtics** and **ytics** whenever it is called.

If either of the arguments is a scalar missing, the main graphics function will set the scaling for that axis using the actual data.

If an argument has 2 elements, the first will be used for the minimum and the last will be used for the maximum.

If an argument has 2 elements, and contains a missing value, that end of the axis will be scaled from the data by the main graphics function.

If you want direct control over the axes endpoints and tick marks, use xtics or ytics. If xtics or ytics have been called after scale, they will override scale.

pscale.src

xtics, ytics, ztics, scale3d

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scale3d

scale3d

Purpose Fixes the scaling for subsequent graphs. The axes endpoints and increments are computed as a best guess based on the data passed to it.

merements are compared as a best guess based on the a

Library pgraph

Format scale3d(x,y,z);

Input x matrix, the X axis data.

y matrix, the Y axis data.

z matrix, the Z axis data.

Remarks x, y and z must each have at least 2 elements. Only the minimum and maximum values are necessary.

This routine fixes the scaling for all subsequent graphs until **graphset** is called. This also clears **xtics**, **ytics** and **ztics** whenever it is called.

If any of the arguments is a scalar missing, the main graphics function will set the scaling for that axis using the actual data.

If an argument has 2 elements, the first will be used for the minimum and the last will be used for the maximum.

If an argument has 2 elements, and contains a missing value, that end of the axis will be scaled from the data by the main graphics function.

If you want direct control over the axes endpoints and tick marks, use **xtics**, **ytics**, or **ztics**. If one of these functions have been called, they will override **scale3d**.

Source pscale.src

See also scale, xtics, ytics, ztics

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scalerr

scalerr

Purpose

Tests for a scalar error code.

Format

y = scalerr(c);

Input

NXK matrix, generally the return argument of a function or procedure call.

Output

scalar, which is returned as a 0 if its argument is not a scalar error code. If the argument is an error code, then scalerr returns the value of the error code as an integer.

Remarks

Error codes in GAUSS are NaN's (Not A Number). These are not just scalar integer values. They are special floating point encodings that the math chip recognizes as not representing a valid number. See also **error**.

scalerr can be used to test for either those error codes which are predefined in GAUSS or an error code which the user has defined using **error**.

If c is an empty matrix, **scalerr** will return 65535.

Certain functions will either return an error code or terminate a program with an error message, depending on the trap state. The **trap** command is used to set the trap state. The error code that will be returned will appear to most commands as a missing value code, but the **scalerr** function can distinguish between missing values and error codes and will return the value of the error code.

Following are some of the functions affected by the trap state:

function	error code	error message
chol	10	Matrix not positive definite
invpd	20	Matrix not positive definite
solpd	30	Matrix not positive definite
/	40	Matrix not positive definite (second argument not square)
	41	Matrix singular (second argument is square)
inv	50	Matrix singular

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scalerr

```
trap 1;
cm = invpd(x);
trap 0;
if scalerr(cm);
cm = inv(x);
endif;
```

In this example **invpd** will return a scalar error code if the matrix **x** is not positive definite. If **scalerr** returns with a nonzero value, the program will use the **inv** function, which is slower, to compute the inverse. Since the trap state has been turned off, if **inv** fails the program will terminate with a **Matrix singular** error message.

See also error, trap, trapchk

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scalinfnanmiss

scalinfnanmiss

Purpose Returns true if the argument is a scalar infinity, NaN, or missing value.

Format y = scalinfnanmiss(x);

Input x NxK matrix.

Output y scalar, 1 if x is a scalar, infinity, NaN, or missing value, else 0.

See also isinfnanmiss, ismiss, scalmiss

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scalmiss

scalmiss

Purpose Tests to see if its argument is a scalar missing value.

Format y = scalmiss(x);

Input x NxK matrix.

clear s;

Output y scalar, 1 if argument is a scalar missing value, 0 if not.

Remarks

scalmiss first tests to see if the argument is a scalar. If it is not scalar, **scalmiss** returns a 0 without testing any of the elements.

The **ismiss** function will test each element of the matrix and return 1 if it encounters any missing values. **scalmiss** will execute much faster if the argument is a large matrix since it will not test each element of the matrix but will simply return a 0.

An element of x is considered to be a missing if and only if it contains a missing value in the real part. Thus, **scalmiss** and **ismiss** would return a 1 for complex x = . + 1i, a 0 for x = 1 + .i.

Example

```
do until eof(fp);
  y = readr(fp,nr);
  y = packr(y);
  if scalmiss(y);
     continue;
  endif;
  s = s+sumc(y);
endo;
```

In this example the **packr** function will return a scalar missing if every row of its argument contains missing values, otherwise it will return a matrix that contains no missing values. **scalmiss** is used here to test for a scalar missing returned from **packr**. If that is true, then the sum step will be skipped for that iteration of the read loop because there were no rows left after the rows containing missings were packed out.

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schtoc

schtoc

a **Purpose** To reduce any 2x2 blocks on the diagional of the real Schur matrix h returned from **schur**. The transformation matrix is also updated. **Format** { schc, transc } = schtoc(sch,trans); Input sch real NxN matrix in Real Schur form, i.e., upper triangular except for possibly 2x2 blocks on the diagonal. real NxN matrix, the associated transformation matrix. trans **Output** schc NxN matrix, possibly complex, strictly upper triangular. The diagonal entries are the eigenvalues. h transc NxN matrix, possibly complex, the associated transformation matrix. Remarks Other than checking that the inputs are strictly real matrices, no other checks are made. If the input matrix sch is already upper triangular it is not changed. Small off-diagional elements are considered to be zero. See the source code for the test used. **Example** { schc, transc } = schtoc(schur(a)); m This example calculates the complex Schur form for a real matrix **a**. 0 Source schtoc.src p See also schur

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schur

schur

Purpose Computes the Schur form of a square matrix.

Format $\{ s,z \} = schur(x)$

Input x KxK matrix.

Output *s* KxK matrix, Schur form.

z KxK matrix, transformation matrix.

Remarks

schur computes the real Schur form of a square matrix. The real Schur form is an upper quasi-triangular matrix, that is, it is block triangular where the blocks are 2x2 submatrices which correspond to complex eigenvalues of x. If x has no complex eigenvalues, s will be strictly upper triangular. To convert s to the complex Schur form, use the Run-Time Library function **schtoc**.

x is first reduced to upper Hessenberg form using orthogonal similarity transformations, then reduced to Schur form through a sequence of QR decompositions.

schur uses the ORTRAN, ORTHES and HQR2 functions from EISPACK.

z is an orthogonal matrix that transforms x into s and vice versa. Thus

$$s = z'xz$$

and since z is orthogonal,

$$x = zsz'$$

Example

let
$$x[3,3] = 1 2 3$$

 $4 5 6$
 $7 8 9;$
{ s, z } = schur(x);

$$s = \begin{bmatrix} 16.11684397 & 4.89897949 & 0.00000000 \\ -0.0000000 & -1.11684397 & -0.0000000 \\ 0.0000000 & 0.0000000 & 0.0000000 \end{bmatrix}$$

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schur

z = 0.23197069 0.88290596 0.40824829 0.52532209 0.23952042 -0.81649658 0.81867350 -0.40386512 0.40824829

See also hess

3-780

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screen

screen

Purpose Controls output to the screen.

Format screen on;

screen off;

screen;

Remarks

When this is **on**, the results of all print statements will be directed to the window. When this is **off**, print statements will not be sent to the window. This is independent of the statement **output on**, which will cause the results of all print statements to be routed to the current auxiliary output file.

If you are sending a lot of output to the auxiliary output file on a disk drive, turning the window off will speed things up.

The **end** statement will automatically do **output off** and **screen on**.

screen with no arguments will print "Screen is on" or "Screen is off" on the console.

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screen **Example** See also

```
output file = mydata.asc reset;
screen off;
format /m1/rz 1,8;
open fp = mydata;
do until eof(fp);
   print readr(fp,200);;
endo;
fp = close(fp);
end;
```

The program above will write the contents of the GAUSS file mydata.dat into an ASCII file called mydata.asc. If mydata.asc already exists, it will be overwritten.

Turning the window off will speed up execution. The **end** statement above will automatically perform **output off** and **screen on**.

See also output, end, new

scroll

scroll

Purpose Scrolls a section of the window.

Format scroll *v*;

Input v 6x1 vector

Portability Windows

Remarks This command is intended to be used in the DOS compatibility window to support legacy programs.

The elements of *v* are defined as:

- [1] coordinate of upper left row.
- [2] coordinate of upper left column.
- [3] coordinate of lower right row.
- [4] coordinate of lower right column.
- [5] number of lines to scroll.
- [61] value of attribute.

This assumes the origin at (1,1) in the upper left just like the **locate** command. The window will be scrolled the number of lines up or down (positive or negative 5^{th} element) and the value of the 6^{th} element will be used as the attribute as follows:

- 7 regular text
- **112** reverse video
 - 0 graphics black

If the number of lines (element 5) is 0, the entire window will be blanked.

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scroll

Example let v = 1 1 12 80 5 7;

scroll v;

This call would scroll a graphic panel 80 columns wide covering the upper twelve rows of the window. The graphic panel would be scrolled up 5 lines and the new lines would be displayed in regular text mode.

See also locate, printdos

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w x y z

seekr

seekr

Purpose Moves the pointer in a .dat or .fmt file to a particular row.

Format $y = \operatorname{seekr}(fh,r)$;

Input *fh* scalar, file handle of an open file.

r scalar, the row number to which the pointer is to be moved.

Output y scalar, the row number to which the pointer has been moved.

Remarks If r = -1, the current row number will be returned.

If r = 0, the pointer will be moved to the end of the file, just past the end of the last row.

rowsf returns the number of rows in a file.

seekr(fh,0) == rowsf(fh) + 1;

Do NOT try to seek beyond the end of a file.

See also open, readr, rowsf

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V

select (dataloop)

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x y z

select (dataloop)

Purpose Selects specific rows (observations) in a data loop based on a logical

expression.

Format select *logical_expression*;

Remarks Selects only those rows for which *logical_expression* is *TRUE*. Any

variables referenced must already exist, either as elements of the source data set, as **externs**, or as the result of a previous **make**, **vector**, or

code statement.

Example select age > 40 AND sex \$== 'MALE';

See also delete

3-786

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x y z

selif

Purpose Selects rows from a matrix. Those selected are the rows for which there is a 1 in the corresponding row of *e*.

Format y = selif(x,e);

Input x NxK matrix.

e Nx1 vector of 1's and 0's.

Output y MxK matrix consisting of the rows of x for which there is a 1 in the corresponding row of e.

Remarks The argument e will usually be generated by a logical expression using "dot" operators.

y will be a scalar missing if no rows are selected.

Example y = selif(x,x[.,2] .gt 100);

selects all rows of x in which the second column is greater than 100.

let
$$x[3,3] = 0$$
 10 20 30 40 50 60 70 80;

$$e = (x[.,1] .gt 0) .and (x[.,3] .lt 100);$$

 $y = selif(x,e);$

The resulting matrix *y* is:

30 40 50 60 70 80

All rows for which the element in column 1 is greater than 0 and the element in column 3 is less than 100 are placed into the matrix \mathbf{y} .

Source datatran.src

See also delif, scalmiss

seqa, seqm

seqa, seqm

Purpose seqa creates an additive sequence. **seqm** creates a multiplicative h sequence. **Format** y = sega(start, inc, n);y = segm(start, inc, n);Input scalar specifying the first element. start scalar specifying increment. incscalar specifying the number of elements in the sequence. n **Output** Nx1 vector containing the specified sequence. y h Remarks For **sega**, y will contain a first element equal to *start*, the second equal to start+inc, and the last equal to start+inc*(n-1). For instance. sega(1,1,10) will create a column vector containing the numbers 1, 2, ... 10. For **segm**, y will contain a first element equal to *start*, the second equal to start*inc, and the last equal to $start*inc^{(n-1)}$. For instance, 0 seqm(10,10,10)p will create a column vector containing the numbers $10, 100, \dots 10^{10}$. **Example** a = sega(2,2,10)';m = segm(2, 2, 10)';*a* = 2 4 6 8 10 12 14 16 18 20 m = 2481632641282565121024u Note that the results have been transposed in this example. Both functions return Nx1 (column) vectors. V See also recserar, recsercp W

x y z

setarray

setarray

Purpose Sets a contiguous subarray of an N-dimensional array.

Format setarray *a*, *loc*, *src*;

Input *a* N-dimensional array.

loc Mx1 vector of indices into the array to locate the subarray of

interest, where M is a value from 1 to N.

src [N-M]-dimensional array, matrix, or scalar.

Remarks setarray resets the specified subarray of *a* in place, without making a copy of the entire array. Therefore, it is faster than putarray.

If *loc* is an Nx1 vector, then *src* must be a scalar. If *loc* is an [N-1]x1 vector, then *src* must be a 1-dimensional array or a 1xL vector, where L is the size of the fastest moving dimension of the array. If *loc* is an [N-2]x1 vector, then *src* must be a KxL matrix, or a KxL 2-dimensional array, where K is the size of the second fastest moving dimension.

Otherwise, if *loc* is an Mx1 vector, then *src* must be an [N-M]-dimensional array, whose dimensions are the same size as the corresponding dimensions of array *a*.

Example a = arrayalloc(2|3|4|5|6,0);

src = arrayinit(4|5|6,5);

loc = { 2,1 };

setarray a,loc,src;

This example sets the contiguous 4x5x6 subarray of a beginning at [2,1,1,1,1] to the array src, in which each element is set to the specified value 5.

See also putarray

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setdif

setdif

a **Purpose** Returns the unique elements in one vector that are not present in a second b vector. Format y = setdif(v1, v2, flag);d Input Nx1 vector. v1 v^2 Mx1 vector. if 0, case-sensitive character comparison. flag scalar. if 1, numeric comparison. if 2, case-insensitive character comparison. h **Output** Lx1 sorted vector containing all unique values that are in v1 and are not in v2, or a scalar missing. Example let v1 = mary jane linda john; k let v2 = mary sally; flag = 0;y = setdif(v1, v2, flag);m **JANE** 0 y = **JOHN** LINDA p q setdif.src Source t

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x y z

setvars

setvars

Purpose Reads the variable names from a data set header and creates global

matrices with the same names.

Format nvec = setvars(dataset);

Input dataset string, the name of the GAUSS data set. Do not use a file

extension.

Output *nvec* Nx1 character vector, containing the variable names defined

in the data set.

Remarks setvars is designed to be used interactively.

Example nvec = setvars("freq");

Source vars.src

See also makevars

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setvwrmode

setvwrmode

a **Purpose** Sets the graphics viewer mode. b Library pgraph Format oldmode = setvwrmode(mode); d Input mode string, new mode or null string. "one" Use only one viewer. "many" Use a new viewer for each graph. Output oldmode string, previous mode. h Remarks If mode is a null string, the current mode will be returned with no changes made. If "one" is set, the viewer executable will be vwr.exe. k **Example** oldmode = setvwrmode("one"); call setvwrmode(oldmode); m Source pgraph.src See also pqgwin 0 p q

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setwind

setwind

Purpose Sets the current graphic panel to a previously created graphic panel number.

number.

Library pgraph

Format setwind(n);

Input n scalar, graphic panel number.

Remarks This function selects the specified graphic panel to be the current graphic

panel. This is the graphic panel in which the next graph will be drawn.

See the discussion on using graphic panels in "Publication Quality

Graphics in the User's Guide.

Source pwindow.src

See also begwind, endwind, getwind, nextwind, makewind,

window

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shell

shell **Purpose** Executes an operating system command. h **Format** shell [s]; Input d literal or 'string, the command to be executed. Remarks **shell** lets you run shell commands and programs from inside GAUSS. If a command is specified, it is executed; when it finishes, you automatically return to GAUSS. If no command is specified, the shell is executed and control passes to it, so you can issue commands interactively. You have to type **exit** to get back to GAUSS in that case. h If you specify a command in a string variable, precede it with the (caret) ^ . **Example** comstr = "ls ./src"; shell ^comstr; This lists the contents of the . / src subdirectory, then returns to GAUSS. m shell cmp n1.fmt n1.fmt.old; This compares the matrix file n1. fmt to an older version of itself, 0 n1.fmt.old, to see if it has changed. When cmp finishes, control is returned to GAUSS. p shell; This executes an interactive shell. The OS prompt will appear and OS commands or other programs can be executed. To return to GAUSS, type exit. See also exec u V

W

x y z

shiftr

shiftr

Purpose Shifts the rows of a matrix.

Format y = shiftr(x, s, f);

Input x NxK matrix to be shifted.

s scalar or Nx1 vector specifying the amount of shift.

f scalar or Nx1 vector specifying the value to fill in.

Output y NxK shifted matrix.

Remarks The shift is performed within each row of the matrix, horizontally. If the shift value is positive, the elements in the row will be moved to the right. A negative shift value causes the elements to be moved to the left. The elements that are pushed off the end of the row are lost, and the fill value will be used for the new elements on the other end.

Example y = shiftr(x,s,f);

If
$$x = \begin{pmatrix} 1 & 2 \\ 3 & 4 \end{pmatrix}$$
 and $s = \begin{pmatrix} 1 \\ -1 \end{pmatrix}$ and $f = \begin{pmatrix} 99 \\ 999 \end{pmatrix}$

Then
$$y = \begin{array}{ccc} 99 & 1 \\ 4 & 999 \end{array}$$

If
$$x = \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \end{bmatrix}$$
 and $s = \begin{bmatrix} 0 \\ 1 \end{bmatrix}$ and $f = 0$

Then
$$y = \begin{pmatrix} 1 & 2 & 3 \\ 0 & 4 & 5 \\ 0 & 0 & 7 \end{pmatrix}$$

See also rotater

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show, 1show

show, lshow

h

Purpose Displays the global symbol table. The output from **lshow** is sent to the printer.

Format

show [[-flags]] [[symbol]];
lshow [[-flags]] [[symbol]];

Input

flags flags to specify the symbol type that is shown.

k keywords

p procedures

f fn functions

m matrices

s strings

g show only symbols with global references

1 show only symbols with all local references

n no pause

symbol

the name of the symbol to be shown. If the last character is an asterisk (*), all symbols beginning with the supplied characters will be shown.

Remarks

If there are no arguments, the entire symbol table will be displayed.

show is directed to the auxiliary output if it is open.

Here is an example listing with an explanation of the columns:

Memory used	Address	Name	Info	Cplx	Type	References
32 bytes at	[00081b74]	AREA	1=1		FUNCTION	local refs
32 bytes at	[00081a14]	dotfeq	1=2		PROCEDURE	global refs
1144 bytes at	[0007f1b4]	indices2	4=3		PROCEDURE	local refs
144 bytes at	[0007f874]	X	3,3	C	MATRIX	
352 bytes at	[0007f6ec]	_IXCAT	44,1		MATRIX	
8 bytes at	[0007f6dc]	_olsrnam	7 char		STRING	

32000 bytes program space, 0% used 4336375 bytes workspace, 4325655 bytes free 53 global symbols, 1500 maximum, 6 shown

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show, 1show

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x y z

The "Memory used" column is the amount of memory used by the item.

The "Address" column is the address where the item is stored (hexadecimal format). This will change as matrices and strings change in size.

The "Name" column is the name of the symbol.

The "Info" column depends on the type of the symbol. If the symbol is a procedure or a function, it gives the number of values that the function or procedure returns and the number of arguments that need to be passed to it when it is called. If the symbol is a matrix, then the Info column gives the number of rows and columns. If the symbol is a string, then the Info column gives the number of characters in the string. As follows:

Rets=Args if procedure or function

Row,Col if matrix Length if string

The "Cplx" column contains a "C" if the symbol is a complex matrix.

The "Type" column specifies the symbol table type of the symbol. It can be function, keyword, matrix, procedure, or string.

If the symbol is a procedure, keyword or function, the "References" column will show if it makes any global references. If it makes only local references, the procedure or function can be saved to disk in an .fcg file with the **save** command. If the function or procedure makes any global references, it cannot be saved in an .fcg file.

The program space is the area of space reserved for all nonprocedure, nonfunction program code. It can be changed in size with the **new** command. The workspace is the memory used to store matrices, strings, procedures, and functions.

Example

show /fpg eig*;

This command will show all functions and procedures that have global references and begin with eig.

show /mn;

This command will show all matrices without pausing when the window is full.

See also new, delete

3-797

sin

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x y z

Purpose Returns the sine of its argument.

Format $y = \sin(x)$;

Input x NxK matrix.

Output y NxK matrix containing sine of x.

Remarks For real matrices, *x* should contain angles measured in radians.

To convert degrees to radians, multiply the degrees by $\frac{\pi}{180}$.

Example let $x = \{ 0, .5, 1, 1.5 \};$

 $y = \sin(x);$

0.00000000

 $y = \begin{array}{c} 0.47942554 \\ 0.84147098 \end{array}$

0.99749499

See also atan, cos, sinh, pi

singleindex

singleindex

Purpose Converts a vector of indices for an N-dimensional array to a scalar vector index.

Format si = singleindex(i, o);

Input *i* Nx1 vector of indices into an N-dimensional array.

o Nx1 vector of orders of an N-dimensional array.

Output si scalar, index of corresponding element in 1-dimensional array or vector.

Remarks This function and its opposite, arrayindex, allow you to convert between an N-dimensional index and its corresponding location in a 1-

dimensional object of the same size.

print getarray(a,ai);

$$ai = 1$$
 3

$$vi = 15$$

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singleindex

$$getarray(a, ai) = 49$$

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x y z

v[vi] = 49

This example allocates a 3-dimensional array a and sets the element corresponding to the index vector ai to 49. It then creates a vector, v, with the same data. The element in the array a that is indexed by ai corresponds to the element of the vector v that is indexed by vi.

See also arrayindex

sinh

sinh

Purpose Computes the hyperbolic sine. **Format** y = sinh(x);Input NxK matrix. **Output** NxK matrix containing the hyperbolic sines of the elements of x. let $x = \{ -0.5, -0.25, 0, 0.25, 0.5, 1 \};$ Example x = x * pi;y = sinh(x);-1.570796-0.7853980.000000 x =0.785398 1.570796 3.141593 -2.301299-0.8686710.000000 y =0.868671 2.301299 11.548739 Source trig.src

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sleep

sleep

Purpose Sleeps for a specified number of seconds.

Format unslept = sleep(secs);

Input *secs* scalar, number of seconds to sleep.

Output *unslept* scalar, number of seconds not slept.

Remarks secs does not have to be an integer. If your system does not permit sleeping for a fractional number of seconds, secs will be rounded to the nearest integer, with a minimum value of 1.

If a program sleeps for the full number of secs specified, **sleep** returns 0; otherwise, if the program is awakened early (e.g., by a signal), **sleep** returns the amount of time not slept. The DOS version always sleeps the full number of seconds, so it always returns 0.

A program may sleep for longer than *secs* seconds, due to system scheduling.

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solpd

solpd

Purpose Solves a set of positive definite linear equations.

Format x = solpd(b,A);

Input b NxK matrix.

A NxN symmetric positive definite matrix.

Output x NxK matrix, the solutions for the system of equations, Ax=b.

Remarks b can have more than one column. If so, the system of equations is solved for each column, i.e., A*x[.,i] = b[.,i].

This function uses the Cholesky decomposition to solve the system directly. Therefore it is more efficient than using inv(A)*b.

solpd does not check to see that the matrix *A* is symmetric. **solpd** will look only at the upper half of the matrix including the principal diagonal.

If the *A* matrix is not positive definite:

trap 1 return scalar error code 30.

trap 0 terminate with an error message.

One obvious use for this function is to solve for least squares coefficients. The effect of this function is thus similar to that of the / operator.

If *X* is a matrix of independent variables, and *Y* is a vector containing the dependent variable, then the following code will compute the least squares coefficients of the regression of *Y* on *X*:

b = solpd(X'Y,X'X);

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x y z

Example n = 5; format 20,8;A = rndn(n,n);A = A'A;x = rndn(n,1);b = A*x;x2 = solpd(b,A);print " X solpd(b,A) Difference"; print x~x2~x-x2; Produces: Χ solpd(b,A) Difference -0.36334089 -0.36334089 0.00000000 0.19683330 0.19683330 8.32667268E-017 0.99361330 0.99361330 2.22044605E-016 0.0000000 -1.84167681 -1.84167681 -0.88455829 -0.88455829 1.11022302E-016 See also scalerr, chol, invpd, trap

sortc, sortcc

sortc, sortcc

Purpose Sorts a matrix of numeric or character data.

Format y = sortc(x,c); y = sortcc(x,c);

Input x NxK matrix.

c scalar specifying one column of x to sort on.

Output y NxK matrix equal to x and sorted on the column c.

Remarks These functions will sort the rows of a matrix with respect to a specified column. That is, they will sort the elements of a column and will arrange all rows of the matrix in the same order as the sorted column.

sortc assumes the column to sort on is numeric. **sortcc** assumes that the column to sort on contains character data.

The matrix may contain both character and numeric data, but the sort column must be all of one type. Missing values will sort as if their value is below $-\infty$.

The sort will be in ascending order. This function uses the Quicksort algorithm.

If you need to obtain the matrix sorted in descending order, you can use:

Example let
$$x[3,3] = 4 7 3$$

1 3 2

3 4 8;

y = sortc(x,1);

 $x = \begin{array}{c} 4 & 7 & 3 \\ 1 & 3 & 2 \\ 3 & 4 & 8 \end{array}$

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sortc, sortcc

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 $y = \begin{array}{c} 1 & 3 & 2 \\ 3 & 4 & 8 \\ 4 & 7 & 3 \end{array}$

See also rev

w x y z

sortd

sortd

Purpose Sorts a data file on disk with respect to a specified variable.

Format sortd(infile,outfile,keyvar,keytyp);

Input *infile* string, name of input file.

outfile string, name of output file, must be different.

keyvar string, name of key variable. *keytyp* scalar, type of key variable.

1 numeric key, ascending order.

2 character key, ascending order.

-1 numeric key, descending order.

-2 character key, descending order.

Remarks The data set *infile* will be sorted on the variable *keyvar*, and will be placed

in *outfile*.

If the inputs are null or 0, the procedure will ask for them.

Source sortd.src

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sorthc, sorthcc

sorthc, sorthcc

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Purpose Sorts a matrix of numeric or character data.

Format

y = sorthc(x,c);

y = sorthcc(x,c);

Input

x NxK matrix.

c scalar specifying one column of x to sort on.

Output

NxK matrix equal to x and sorted on the column c.

Remarks

These functions will sort the rows of a matrix with respect to a specified column. That is, they will sort the elements of a column and will arrange all rows of the matrix in the same order as the sorted column.

sorthc assumes that the column to sort on is numeric. **sorthcc** assumes that the column to sort on contains character data.

The matrix may contain both character and numeric data, but the sort column must be all of one type. Missing values will sort as if their value is below $-\infty$.

The sort is in ascending order. This function uses the heap sort algorithm.

If you need to obtain the matrix sorted in descending order, you can use:

rev(sorthc(x,c))

Example

let x[3,3] = 4 7 3

1 3 2

3 4 8;

y = sorthc(x,1);

4 7 3

x = 132

3 4 8

sorthc, sorthcc

$$y = \begin{array}{c} 1 & 3 & 2 \\ 3 & 4 & 8 \\ 4 & 7 & 3 \end{array}$$

See also sortc, rev

a

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sortind, sortindc

sortind, sortindc

Purpose Returns the sorted index of x.

Format ind = sortind(x);
ind = sortindc(x);

Input x Nx1 column vector.

Output *ind* Nx1 vector representing sorted index of x.

Remarks sortind assumes x contains numeric data. sortindc assumes x contains character data.

This function can be used to sort several matrices in the same way that some other reference matrix is sorted. To do this, create the index of the reference matrix, then use **submat** to rearrange the other matrices in the same way.

Example let x = 5 4 4 3 3 2 1;
ind = sortind(x);

y = x[ind];

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sortmc

sortmc

Purpose Sorts a matrix on multiple columns.

Format y = sortmc(x,v);

Input x NxK matrix to be sorted.

V Lx1 vector containing integers specifying the columns, in order, that are to be sorted. If an element is negative, that column will be interpreted as character data.

Output y NxK sorted matrix.

Remarks The function works recursively and the number of sort columns is limited by the available workspace.

by the available workspace

Source sortmc.src

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sparseCols

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sparseCols

Purpose Returns the number of columns in a sparse matrix.

Format c = sparseCols(x);

Input x MxN sparse matrix.

Output c scalar, number of columns.

Source sparse.src

W

sparseEye

sparseEye

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sparseFD

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sparseFD

Purpose Converts dense matrix gto sparse matrix.

Format y = sparseFD(x, eps);

Input x MxN dense matrix.

eps scalar, elements of x less than eps will be treated as zero.

Output y MxN sparse matrix.

Remarks A dense matrix is just a normal format matrix.

Source sparse.src

sparseFP

sparseFP

Purpose Converts packed matrix to sparse matrix.

Format y = sparseFP(x,r,c);

Input x Mx3 packed matrix, see remarks for format.

r scalar, rows of output matrix.

c scalar, columns of output matrix.

Output y RxC sparse matrix.

Remarks x contains the nonzero elements of the sparse matrix. The first column of

x contains the element value, the second column the row number, and the

third column the column number.

Source sparse.src

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sparseHConcat

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sparseHConcat

Purpose Horizontally concatenates two sparse matrices.

Format z = sparseHConcat(y,x);

Input y MxN sparse matrix, left hand matrix.

x MxL sparse matrix, right hand matrix.

Output z Mx(N+L) sparse matrix.

Source sparse.src

sparseNZE

sparseNZE

Purpose Returns the number of nonzero elements in a sparse matrix.

Format r = sparseNZE(x);

Input x MxN sparse matrix.

Output r scalar, number of nonzero elements in x.

Source sparse.src

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sparseOnes

Purpose Generates sparse matrix of ones and zeros

Format y = sparseOnes(x,r,c);

Input x Mx2 matrix, first column contains row numbers of the ones, and the second column contains column numbers.

r scalar, rows of full matrix.

c scalar, columns of full matrix.

Output y sparse matrix of ones.

Source sparse.src

sparseRows

sparseRows

Purpose Returns the number of rows in a sparse matrix.

Format r = sparseRows(x);

Input x MxN sparse matrix.

Output r scalar, number of rows.

Source sparse.src

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sparseSet

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sparseSet

Purpose Resets sparse library global matrices to default values.

Format sparseSet;

Globals _sparse_ARnorm, _sparse_Acond, _sparse_Anorm,

_sparse_Atol, _sparse_Btol, _sparse_CondLimit,

_sparse_Damping, _sparse_NumIters,

_sparse_RetCode, _sparse_Rnorm, _sparse_Xnorm

Source sparse.src

sparseSolve

sparseSolve

Purpose Solves Ax = B for x when A is a sparse matrix.

Format x = sparseSolve(A,B);

Input A MxN sparse matrix.

B Nx1 vector.

Global Input _sparse_Damping

scalar, if nonzero, damping coefficient for

damped least squares solve, i.e.,

$$\begin{bmatrix} A \\ dI \end{bmatrix} = \begin{bmatrix} B \\ 0 \end{bmatrix}$$

is solved for *X* where

d = **_sparse_Damping**, *I* is a conformable identity matrix, and 0 a conformable matrix

of zeros.

_sparse_Atol scalar, an estimate of the relative error in A.

If zero, _sparse_Atol is assumed to be

machine precision. Default = 0.

_sparse_Btol an estimate of the relative error in *B*. If zero,

_sparse_Btol is assumed to be machine

precision. Default = 0.

_sparse_CondLimit upper limit on condition of A. Iterations will

be terminated if a computed estimate of the

condition of A exceeds

_sparse_CondLimit. If zero, set to 1 /

machine precision.

sparse NumIters maximum number of iterations.

Output x Nx1 vector, solution of Ax = B.

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sparseSolve

Globa
Output

_sparse_RetCode

scalar, termination condition.

- 0 *x* is the exact solution, no iterations performed.
- solution is nearly exact with accuracy on the order of _sparse_Atol and_sparse_Btol.
- solution is not exact and a least squares solution has been found with accuracy on the order of _sparse_Atol.
- 3 the estimate of the condition of A has exceeded _sparse_CondLimit.

 The system appears to be ill-conditioned.
- 4 solution is nearly exact with reasonable accuracy.
- 5 solution is not exact and a least squares solution has been found with reasonable accuracy.
- 6 iterations halted due to poor condition given machine precision.
- 7 _sparse_NumIters exceeded.

_sparse_Anorm

scalar, estimate of Frobenius norm of

$$A$$
 dI

_sparse_Acond

estimate of condition of A.

_sparse_Rnorm estimate of norm of

$$\begin{bmatrix} A \\ dI \end{bmatrix} x - \begin{bmatrix} B \\ 0 \end{bmatrix}$$

_sparse_ARnorm

estimate of norm of

$$\begin{bmatrix} A \\ dI \end{bmatrix}' \begin{bmatrix} A \\ dI \end{bmatrix}$$

_sparse_XAnorm

estimate of norm of x.

Source

sparse.src

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sparseSubmat

sparseSubmat

Purpose Returns (sparse) submatrix of sparse matrix.

Format e = sparseSubmat(x,r,c);

Input x MxN sparse matrix.

r Kx1 vector, row indices.

c Lx1 vector, column indices.

Output e KxL sparse matrix.

Remarks If *r* or *c* are scalar zeros, all rows or columns will be returned.

Source sparse.src

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sparseTD

Purpose Multiplies sparse matrix by dense matrix.

Format z = sparseTD(s,d);

Input *s* MxN sparse matrix.

d NxL dense matrix.

Output z MxL dense matrix, the result of $s \times d$.

Source sparse.src

хух

3-824

sparseTrTD

sparseTrTD

Purpose Multiplies sparse matrix transposed by dense matrix.

Format z = sparseTrTD(s,d);

Input *s* NxM sparse matrix.

d NxL dense matrix.

Output z MxL dense matrix, the result of s'd.

Source sparse.src

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sparseVConcat

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sparseVConcat

Purpose Vertically concatenates two sparse matrices.

Format z = sparseVConcat(y,x);

Input y MxN sparse matrix, top matrix.

x LxN sparse matrix, bottom matrix.

Output z (M+L)xN sparse matrix.

Source sparse.src

W

spline

Purpose Computes a two-dimensional interpolatory spline.

Format { u,v,w } = spline(x,y,z,sigma,g);

Input

- x 1xK vector, x-abscissae (x-axis values).
- y Nx1 vector, y-abscissae (y-axis values).
- z KxN matrix, ordinates (z-axis values).

sigma scalar, tension factor.

g scalar, grid size factor.

Output

- u 1xK*G vector, x-abscissae, regularly spaced.
- *v* N*Gx1 vector, y-abscissae, regularly spaced.
- w K*G x N*G matrix, interpolated ordinates.

Remarks

sigma contains the tension factor. This value indicates the curviness desired. If sigma is nearly zero (e.g., .001), the resulting surface is approximately the tensor product of cubic splines. If sigma is large (e.g., 50.0), the resulting surface is approximately bi-linear. If sigma equals zero, tensor products of cubic splines result. A standard value for sigma is approximately 1.

g is the grid size factor. It determines the fineness of the output grid. For g = 1, the output matrices are identical to the input matrices. For g = 2, the output grid is twice as fine as the input grid, i.e., u will have twice as many columns as x, v will have twice as many rows as y, and w will have twice as many rows and columns as z.

Source

spline.src

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spline1D

spline1D

tes a smoothing spline for a curve.

Format { u,v } = spline1D(x,y,d,s,sigma,g);

Input

- *x* Kx1 vector, x-abscissae (x-axis values).
- y Kx1 vector, y-ordinates (y-axis values).
- d Kx1 vector or scalar, observation weights.
- s scalar, smoothing parameter, if s = 0, curve performs an interpolation. If d contains standard deviation estimates, a reasonable value for s is K.
- sigma scalar, tension factor.G scalar, grid size factor.

Output

- *u* K*Gx1vector, x-abscissae, regularly spaced.
- v N*Gx1 vector, interpolated ordinates, regularly spaced.

Remarks

sigma contains the tension factor. This value indicates the curviness desired. If sigma is nearly zero (e.g. 0.001), the resulting curve is approximately the tensor product of cubic splines. If sigma is large (e.g. 50), the resulting curve is approximately bi-linear. If sigma equals zero, tensor products of cubic splines result. A standard value for sigma is approximately 1.

G is the grid size factor. It determines the fineness of the output grid. For G = 1, the output matrices are identical to the input matrices. For G = 2, the output grid is twice as fine as the input grid, i.e., u will have twice as many columns as x, and v will have twice as many rows as v.

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spline2D

Purpose Computes a smoothing spline for a surface.

Format { u,v,w } = spline2D(x,y,z,sigma,g);

Input

- x Kx1 vector, x-abscissae (x-axis values).
- y Nx1 vector, y-abscissae (y-axis values).
- z KxN matrix, ordinates (z-axis values).

sigma scalar, tension factor.

G scalar, grid size factor.

Output

- *u* 1xK*G vector, x-abscissae, regularly spaced.
- v N*Gx1 vector, y-abscissae, regularly spaced.
- w K*GxN*G matrix, interpolated ordinates.

Remarks

sigma contains the tension factor. This value indicates the curviness desired. If sigma is nearly zero (e.g. 0.001), the resulting surface is approximately the tensor product of cubic splines. If sigma is large (e.g. 50.), the resulting surface is approximately bi-linear. If sigma equals zero, tensor products of cubic splines result. A standard value for sigma is approximately 1.

G is the grid size factor. It determines the fineness of the output grid. For G = 1, the output matrices are identical to the input matrices. For G = 2, the output grid is twice as fine as the input grid, i.e., u will have twice as many columns as x, and v will have twice as many rows as y.

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SpreadsheetReadM

SpreadsheetReadM

Purpose Reads and writes Excel files.

Format xlsmat = SpreadsheetReadM(file, range, sheet);

Input *file* string, name of .xls file.

range string, range to read or write; e.g., "a1:b20".

sheet scalar, sheet number.

Output *xlsmat* matrix of numbers read from Excel.

Remarks If the read functions fail, they will return a scalar error code which can be

decoded with **scalerr**. If the write function fails, it returns a non-zero

error number.

See also scalerr, error

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SpreadsheetReadSA

SpreadsheetReadSA

Purpose Reads and writes Excel files.

Format xlssa = SpreadsheetReadSA(file, range, sheet);

Input *file* string, name of .xls file.

range string, range to read or write; e.g., "a1:b20".

sheet scalar, sheet number.

Output *xlssa* string array read from Excel.

Remarks If the read functions fail, they will return a scalar error code which can be

decoded with **scalerr**. If the write function fails, it returns a non-zero

error number.

See also scalerr, error

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SpreadsheetWrite

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x y z

SpreadsheetWrite

Purpose Reads and writes Excel files.

Format xlsret = SpreadsheetWrite(data, file, range, sheet);

Input *data* matrix, string or string array, data to write.

file string, name of .xls file.

range string, range to read or write; e.g., "a1:b20".

sheet scalar, sheet number.

Output *xlsret* success code, 0 if successful, else error code.

Remarks If the read functions fail, they will return a scalar error code which can be

decoded with scalerr. If the write function fails, it returns a non-zero

error number.

See also scalerr, error

sqpSolve

Purpose Solves the nonlinear programming problem using a sequential quadratic programming method.

Format { x,f,lagr,retcode } = sqpSolve(&fct,start);

pointer to a procedure that computes the function to be minimized. This procedure must have one input argument, a vector of parameter values, and one output argument, the

value of the function evaluated at the input vector of parameter values.

start Kx1 vector of start values.

Global _sqp_A MxK matrix, linear equality constraint coefficients.

_sqp_B Mx1 vector, linear equality constraint

constants.

These globals are used to specify linear equality constraints of the following type:

 $_sqp_A * X = _sqp_B$

where X is the Kx1 unknown parameter vector.

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_sqp_EqProc

sqp_C

_sqp_D

scalar, pointer to a procedure that computes the nonlinear equality constraints. For example, the statement:

tells **sqpSolve** that nonlinear equality constraints are to be placed on the parameters and where the procedure computing them is to be found. The procedure must have one input argument, the Kx1 vector of parameters, and one output argument, the Rx1 vector of computed constraints that are to be equal to zero. For example, suppose that you wish to place the following constraint:

$$P[1] * P[2] = P[3]$$

The procedure for this is:

MxK matrix, linear inequality constraint coefficients.

Mx1 vector, linear inequality constraint constants.

These globals are used to specify linear inequality constraints of the following type:

$$_sqp_C * X >= _sqp_D$$

where X is the Kx1 unknown parameter vector.

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_sqp_IneqProc

sqp Bounds

scalar, pointer to a procedure that computes the nonlinear inequality constraints. For example the statement:

tells **sqpSolve** that nonlinear equality constraints are to be placed on the parameters and where the procedure computing them is to be found. The procedure must have one input argument, the Kx1 vector of parameters, and one output argument, the Rx1 vector of computed constraints that are to be equal to zero. For example, suppose that you wish to place the following constraint:

$$P[1] * P[2] >= P[3]$$

The procedure for this is:

endp;

Kx2 matrix, bounds on parameters. The first column contains the lower bounds, and the second column the upper bounds. If the bounds for all the coefficients are the same, a 1x2 matrix may be used. Default is:

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_sqp_G	radProc	scalar, pointer to a procedure that computes the gradient of the function with respect to the parameters. For example, the statement:
		_sqp_GradProc = &gradproc
		tells sqpSolve that a gradient procedure exists and where to find it. The user-provided procedure has two input arguments, a Kx1 vector of parameter values and an NxP matrix of data. The procedure returns a single output argument, an NxK matrix of gradients of the log-likelihood function with respect to the parameters evaluated at the vector of parameter values.
		Default = 0, i.e., no gradient procedure has been provided.
_sqp_He	essProc	scalar, pointer to a procedure that computes the Hessian, i.e., the matrix of second order partial derivatives of the function with respect to the parameters. For example, the instruction:
		_sqp_HessProc = &hessproc
		will tell sqpSolve that a procedure has been provided for the computation of the Hessian and where to find it. The procedure that is provided by the user must have two input arguments, a Px1 vector of parameter values and an NxK data matrix. The procedure returns a single output argument, the PxP symmetric matrix of second order derivatives of the function evaluated at the parameter values.
_sqp_Ma	axIters	scalar, maximum number of iterations. Default = 1e+5. Termination can be forced by pressing C on the keyboard.
_sqp_D:	irTol	scalar, convergence tolerance for gradient of estimated coefficients. Default = 1e-5. When this criterion has been satisifed sqpsolve will exit the iterations.
_sqp_Pa	arNames	Kx1 character vector, parameter names.

_sqp_EqProc

_sqp_C

_sqp_D

scalar, pointer to a procedure that computes the nonlinear equality constraints. For example, the statement:

tells **sqpSolve** that nonlinear equality constraints are to be placed on the parameters and where the procedure computing them is to be found. The procedure must have one input argument, the Kx1 vector of parameters, and one output argument, the Rx1 vector of computed constraints that are to be equal to zero. For example, suppose that you wish to place the following constraint:

$$P[1] * P[2] = P[3]$$

The procedure for this is:

endp;

MxK matrix, linear inequality constraint coefficients.

Mx1 vector, linear inequality constraint constants.

These globals are used to specify linear inequality constraints of the following type:

$$_sqp_C * X >= _sqp_D$$

where X is the Kx1 unknown parameter vector.

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_sqp_IneqProc

sqp Bounds

scalar, pointer to a procedure that computes the nonlinear inequality constraints. For example the statement:

```
_sqp_EqProc = &ineqproc;
```

tells **sqpSolve** that nonlinear equality constraints are to be placed on the parameters and where the procedure computing them is to be found. The procedure must have one input argument, the Kx1 vector of parameters, and one output argument, the Rx1 vector of computed constraints that are to be equal to zero. For example, suppose that you wish to place the following constraint:

$$P[1] * P[2] >= P[3]$$

The procedure for this is:

```
proc ineqproc(p);
  retp(p[1]*[2]-p[3]);
endp;
```

Kx2 matrix, bounds on parameters. The first column contains the lower bounds, and the second column the upper bounds. If the bounds for all the coefficients are the same, a 1x2 matrix may be used. Default is:

```
[1] -1e256
[2] 1e256
```

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_sqp_GradProc

scalar, pointer to a procedure that computes the gradient of the function with respect to the parameters. For example, the statement:

_sqp_GradProc = &gradproc;

tells **sqpSolve** that a gradient procedure exists and where to find it. The user-provided procedure has two input arguments, a Kx1 vector of parameter values and an NxP matrix of data. The procedure returns a single output argument, an NxK matrix of gradients of the log-likelihood function with respect to the parameters evaluated at the vector of parameter values.

Default = 0, i.e., no gradient procedure has been provided.

_sqp_HessProc

scalar, pointer to a procedure that computes the Hessian, i.e., the matrix of second order partial derivatives of the function with respect to the parameters. For example, the instruction:

_sqp_HessProc = &hessproc;

will tell **sqpSolve** that a procedure has been provided for the computation of the Hessian and where to find it. The procedure that is provided by the user must have two input arguments, a Px1 vector of parameter values and an NxK data matrix. The procedure returns a single output argument, the PxP symmetric matrix of second order derivatives of the function evaluated at the parameter values.

_sqp_MaxIters

scalar, maximum number of iterations. Default = 1e+5. Termination can be forced by pressing C on the keyboard.

 ${\tt _sqp_DirTol}$

scalar, convergence tolerance for gradient of estimated coefficients. Default = 1e-5. When this criterion has been satisifed **sqpSolve** will exit the iterations.

sqp ParNames

Kx1 character vector, parameter names.

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_sqp_PrintIters _sqp_FeasibleTest _sqp_RandRadius		scalar, if nonzero, prints iteration information. Default = 0. Can be toggled during iterations by pressing P on the keyboard.		
		scalar, if nonzero, parameters are tested for feasibility before computing function in line search. If function is defined outside inequality boundaries, then this test can be turned off.		
		scalar, If zero, no random search is attempted. If nonzero it is the radius of random search which is invoked whenever the usual line search fails. Default = .01.		
outp	ut	scalar, if nonzero, results are printed. Default = 0.		
x	Kx1 vector of	f parameters at minimum.		
f	scalar, function	on evaluated at x.		
lagr	constraints. T	ed using vput . Contains the Lagrangean for the They may be extracted with the vread ing the following strings:		
	"lineq"	Lagrangeans of linear equality constraints		
	"nlineq"	Lagrangeans of nonlinear equality constraints		
	"linineq"	Lagrangeans of linear inequality constraints		
	"nlinineq"	Lagrangeans of nonlinear inequality constraints		
	"bounds"	Lagrangeans of bounds		
	Whenever a c will be nonze	constraint is active, its associated Lagrangean ro.		
retcode	return code:			
	0	normal convergence		
	1	forced exit		
	2	maximum number of iterations exceeded		
	3	function calculation failed		
	4	gradient calculation failed		
	5	Hessian calculation failed		
	6	line search failed		
	7	error with constraints		

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x y z

Remarks

Pressing C on the keyboard will terminate iterations, and pressing P will toggle iteration output.

sqpSolve is recursive, that is, it can call itself with another function and set of global variables.

To reset global variables for this function to their default values, call sqpSolveSet.

Example

Source

sqpsolve.src

sqpSolveSet;

```
proc fct(x);
   retp( (x[1] + 3*x[2] + x[3])^2 + 4*(x[1] -
  x[2])^2);
endp;
proc ineqp(x);
   retp(6*x[2] + 4*x[3] - x[1]^3 - 3);
endp;
proc eqp(x);
   retp(1-sumc(x));
endp;
_{sqp}Bounds = { 0 1e256 };
start = \{ .1, .7, .2 \};
_sqp_IneqProc = &ineqp;
_sqp_EqProc = &eqp;
{ x,f,lagr,ret } = sqpSolve( &fct,start );
```

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W

x y z

sqpSolveMT

Solves the nonlinear programming problem. **Purpose Format** outl = sqpSolveMT(&fct,parl,datal,cl); Include sqpSolveMT.sdf Input &fct pointer to a procedure that computes the function to be minimized. This procedure must have two input arguments, an instance of structure of type PV and an instance of a structure of type DS, and one output argument, either a 1x1 scalar or an Nx1 vector of function values evaluated at the parameters stored in the PV instance using data stored in the DS instance. par1 an instance of structure of type PV. The par1 instance is passed to the user-provided procedure pointed to by &fct. par1 is constructed using the "pack" functions. an array of instances of a DS structure. This array is passed to the data1 user-provided pointed by &fct to be used in the objective function. **sqpSolveMT** does not look at this structure. Each instance contains the the following members which can be set in whatever way that is convenient for computing the objective function: data1[i].dataMatrix NxK matrix, data matrix. data1[i].dataArray NxKxL.. array, data array. data1[i].vnames string array, variable names (optional). data1[i].dsname string, data name (optional). data1[i].type scalar, type of data (optional). c1an instance of an sqpSolveMTControl structure. Normally an instance is initialized by calling sgpSolveMTControlCreate and members of this instance can be set to other values by the user. For an instance named cI, the members are: c1.AMxK matrix, linear equality constraint coefficients: c1.A * p = c1.B where p is a vector of the parameters. c1.BMx1 vector, linear equality constraint constants: c1.A * p = c1.B where p is a vector of the parameters.

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c1.C	MxK matrix, linear inequality constraint coefficients: $c1.C * p >= c1.D$ where p is a vector of the parameters.
c1.D	Mx1 vector, linear inequality constraint constants: $c1.C * p >= c1.D$ where p is a vector of the parameters.
c1.eqProc	scalar, pointer to a procedure that computes the nonlinear equality constraints. When such a procedure has been provided, it has one input argument, a structure of type SQPdata, and one output argument, a vector of computed equality constraints. For more details see Remarks below. Default = {.}, i.e., no equality procedure.
c1.weights	vector, weights for objective function returning a vector. Default = 1.
c1.ineqProc	scalar, pointer to a procedure that computes the nonlinear inequality constraints. When such a procedure has been provided, it has one input argument, a structure of type SQPdata, and one output argument, a vector of computed inequality constraints. For more details see Remarks below. Default = {.}, i.e., no inequality procedure.
c1.bounds	1x2 or Kx2 matrix, bounds on parameters. If 1x2 all parameters have same bounds. Default = { -1e256 1e256 }.
c1.covType	scalar, if 2, QML covariance matrix, else if 0, no covariance matrix is computed, else ML covariance matrix is computed.
c1.gradProc	scalar, pointer to a procedure that computes the gradient of the function with respect to the parameters. Default = {.}, i.e., no gradient procedure has been provided.

a b			c1.hessProc		second o function Default =	ointer to a procedure that s the Hessian, i.e., the matrix of rder partial derivatives of the with respect to the parameters. = {.}, i.e., no Hessian procedure
С			c1.maxIters			provided. aximum number of iterations.
	 				Default =	= 1e+5.
d			c1.dirTol			onvergence tolerance for gradient ated coefficients. Default = 1e-5.
e f	 				When th	is criterion has been satisifed ve exits the iterations.
	 		c1.feasibleTe	est		nonzero, parameters are tested
g h					in line se outside in	collity before computing function earch. If function is defined nequality boundaries, then this
i			1 ID 1:			be turned off. Default = 1.
j			c1.randRadii	us		zero, no random search is d. If nonzero, it is the radius of
k					random s	search which is invoked or the usual line search fails.
1 m			c1.output		scalar, if Default =	nonzero, results are printed. = 0.
n			c1.printIters			nonzero, prints iteration ion. Default = 0 .
0						
p	Output	out1	an instance o named <i>out1</i> ,			out structure. For an instance
q			out1.par			ucture of type PV containing the tes will be placed in the member
r			1.0		out1.par.	
S			out1.fct			valuated at x.
t			out1.lagr	the Lag	grangeans	QPLagrange structure containing for the constraints. For an
u				instanc	e named l	agr, the members are:
V				out1 la	gr.lineq	Mx1 vector, Lagrangeans of
W				0 1	······································	linear equality constraints.

sapSolveMT

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Nx1 vector, Lagrangeans of nonlinear equality constraints.
Px1 vector, Lagrangeans of linear inequality constraints.
Qx1 vector, Lagrangeans of nonlinear inequality constraints.
Kx2 matrix, Lagrangeans of bounds.
raint is active, its associated be nonzero. For any constraint roughout the iterations as well as the corresponding Lagrangean to a scalar missing value.

out1.retcode return code:

0	normal convergence.
1	forced exit.
2	maximum number of iterations exceeded.
3	function calculation failed.
4	gradient calculation failed.
5	Hessian calculation failed.
6	line search failed.
7	error with constraints.
8	function complex.

Remarks

There is one required user-provided procedure, the one computing the objective function to be minimized, and four other optional functions, one each for computing the equality constraints, the inequality constraints, the gradient of the objective function, and the Hessian of the objective function.

All of these functions have one input argument that is an instance of a structure of type struct PV and a second argument that is an instance of a structure of type struct DS. On input to the call to **sqpSolveMT**, the first argument contains starting values for the parameters and the second argument any required data. The data are passed in a separate argument because the structure in the first argument will be copied as it is passed through procedure calls which would be very costly if it contained large data matrices. Since **sqpSolveMT** makes no changes to the second argument it will be passed by pointer thus saving time because its contents aren't copied.

Both of the structures of type PV are set up using the PV pack procedures, **pvPack**, **pvPackm**, **pvPacks**, and **pvPacksm**. These procedures allow for setting up a parameter vector in a variety of ways.

For example, we might have the following objective function for fitting a nonlinear curve to data:

```
proc Micherlitz(struct PV par1, struct DS data1);
  local p0,e,s2,x,y;
  p0 = pvUnpack(par1,"parameters");
  y = data1.dataMatrix[.,1];
  x = data1.dataMatrix[.,2];
  e = y - p0[1] - p0[2]*exp(-p0[3] * x);
  retp(e'*e);
endp;
```

In this example the dependent and independent variables are passed to the procedure as the first and second columns of a data matrix stored in a single DS structure. Alternatively these two columns of data can be entered into a vector of DS structures one for each column of data:

proc Micherlitz(struct PV par1, struct DS data1);

```
local p0,e,s2,x,y;

p0 = pvUnpack(par1,"parameters");

y = data1[1].dataMatrix;

x = data1[2].dataMatrix;

e = y - p0[1] - p0[2]*exp(-p0[3]*x);

retp(e'*e);
endp;
```

The syntax is similar for the optional user-provided procedures. For example, to constrain the squared sum of the first two parameters to be greater than one in the above problem, provide the following procedure:

```
proc ineqConst(struct PV par1, struct DS data1);
```

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```
local p0;
   p0 = pvUnpack(p0, "parameters");
   retp( (p0[2]+p0[1])^2 - 1);
endp;
The following is a complete example for estimating the parameters of the
Micherlitz equation in data with bounds constraints on the parameters and
where an optional gradient procedure has been provided:
#include sqpSolveMT.sdf
   struct DS d0;
   d0 = dsCreate;
         3.183
   y =
         3.059
         2.871
         2.622
         2.541
         2.184
         2.110
         2.075
         2.018
         1.903
         1.770|
         1.762
         1.550;
   x = seqa(1,1,13);
```

d0.dataMatrix = y~x;

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```
struct sqpSolveMTControl c0;
c0 = sqpSolveMTControlCreate;
c0.bounds = 0 \sim 100; /* constrains parameters */
                   /* to be positive */
struct PV par1;
par1 = pvCreate;
pvPack(par1,.92|2.62|.114,"parameters");
struct sqpSolveMTout out1;
out1 = sqpSolveMT(&Micherlitz,par1,d0,c0);
print " parameter estimates ";
print pvUnPack(out1.par, "parameters");
proc Micherlitz(struct PV par1, struct DS
data1);
   local p0,e,s2,x,y;
  p0 = pvUnpack(par1, "parameters");
  y = data1.dataMatrix[.,1];
  x = data1.dataMatrix[.,2];
   e = y - p0[1] - p0[2]*exp(-p0[3] * x);
  retp(e'*e);
endp;
proc grad(struct PV par1, struct DS data1);
  local p0,e,w,g,r,x,y;
  p0 = pvUnpack(par1, "parameters");
  y = data1.dataMatrix[.,1];
```

3-848

```
x = data1.dataMatrix[.,2];
g = zeros(3,1);
w = exp(-p0[3] * x);
e = y - p0[1] - p0[2]*w;
r = e'*w;
g[1] = -2*sumc(e);
g[2] = -2*r;
g[3] = 2*p0[1]*p0[2]*r;
retp(g);
endp;
```

Source sqpsolvemt.src

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sqpSolveMTcontrolCreate

sqpSolveMTcontrolCreate

Purpose Creates an instance of a structure of type sqpSolveMTcontrol set to default values.

Format s = sqpSolveMTcontrolCreate;

Output *s* instance of structure of type sqpSolveMTcontrol.

Source sqpsolvemt.src

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sqpSolveMTlagrangeCreate

sqpSolveMTlagrangeCreate

Purpose Creates an instance of a structure of type sqpSolveMTlagrange set to

default values.

Format s = sqpSolveMTlagrangeCreate;

Output *s* instance of structure of type sqpSolveMTlagrange.

Source sqpsolvemt.src

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sqpSolveMToutCreate

sqpSolveMToutCreate

Purpose Creates an instance of a structure of type sqpSolveMTout set to default values.

Format s = sqpSolveMToutCreate;

Output *s* instance of structure of type sqpSolveMTout.

Source sqpsolvemt.src

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sqrt

Purpose Computes the square root of every element in a matrix.

Format y = sqrt(x);

Input x NxK matrix.

Output y NxK matrix, the square roots of each element of x.

Remarks If *x* is negative, complex results are returned.

You can turn the generation of complex numbers for negative inputs on or off in the GAUSS configuration file, and with the **sysstate** function, case 8. If you turn it off, **sqrt** will generate an error for negative inputs.

If *x* is already complex, the complex number state doesn't matter; **sqrt** will compute a complex result.

Example let x[2,2] = 1 2 3 4;

y = sqrt(x);

x = 1.00000000 2.0000000003.00000000 4.00000000

 $y = \begin{array}{r} 1.000000000 \ 1.41421356 \\ 1.73205081 \ 2.000000000 \end{array}$

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stdc

stdc

a

Purpose Computes the standard deviation of the elements in each column of a matrix.

b

Format

y = stdc(x);

d

x NxK matrix.

f

Output

Input

Kx1 vector, the standard deviation of each column of x.

g

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k

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q

Remarks

This function essentially computes:

 $sqrt(1/(N-1)*sumc((x-meanc(x)')^2))$

Thus, the divisor is N-1 rather than N, where N is the number of elements being summed. To convert to the alternate definition, multiply by

Example

y = rndn(8100,1);

std = stdc(y);

std = 1.008377

In this example, 8100 standard Normal random variables are generated, and their standard deviation is computed.

See also

meanc

w x y z

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3-854

stocv

stocv

```
Purpose
              Converts a string to a character vector.
  Format
              v = stocv(s);
    Input
                     string, to be converted to character vector.
  Output
                     Nx1 character vector, contains the contents of s.
Remarks
              stocv breaks s up into a vector of 8-character length matrix elements.
              Note that the character information in the vector is not guaranteed to be
              null-terminated.
Example
              s = "Now is the time for all good men";
              v = stocv(s);
                     "Now is t"
                     "he time "
                    "for all "
                    "good men"
See also
              cvtos, vget, vlist, vput, vread
```

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x y z

stof

Purpose Converts a string to floating point.

Format y = stof(x);

Input x string, or NxK matrix containing character elements to be converted.

Output y matrix, the floating point equivalents of the ASCII numbers in x.

Remarks If x is a string containing "1 2 3", then **stof** will return a 3x1 matrix containing the numbers 1, 2, and 3.

If *x* is a null string, **stof** will return a 0.

This uses the same input conversion routine as **loadm** and **let**. It will convert character elements and missing values. **stof** also converts complex numbers in the same manner as **let**.

See also ftos, ftocv, chrs

stop

stop

Purpose Stops a program and returns to the command prompt. Does not close files.

Format stop;

Remarks This command has the same effect as **end**, except it does not close files

or the auxiliary output.

It is not necessary to put a stop or an end statement at the end of a

program. If neither is found, an implicit stop is executed.

See also end, new, system

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strindx

strindx

a **Purpose** Finds the index of one string within another string. h **Format** y = strindx(where, what, start); Input d where string or scalar, the data to be searched. what string or scalar, the substring to be searched for in where. scalar, the starting point of the search in where for an occurrence start of what. The index of the first character in a string is 1. **Output** scalar containing the index of the first occurrence of what, within where, which is greater than or equal to start. If no occurrence is h found, it will be 0. Remarks An example of the use of this function is the location of a name within a string of names: z = "Whatchmacallit"; x = "call";y = strindx(z,x,1);m v = 90 This function is used with **strsect** for extracting substrings. p See also strrindx, strlen, strsect, strput

3-858

u

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W

x y z

strlen

strlen

Purpose Returns the length of a string.

Format y = strlen(x);

Input x string or NxK matrix of character data.

Output y scalar containing the exact length of the string x or NxK matrix of the lengths of the elements in the matrix x.

Remarks The null character (ASCII 0) is a legal character within strings and so embedded nulls will be counted in the length of strings. The final

terminating null byte is not counted, though.

For character matrices, the length is computed by counting the characters (maximum of 8) up to the first null in each element of the matrix. The null character, therefore, is not a valid character in matrices containing character data and is not counted in the lengths of the elements of those matrices.

Example x = "How long?";

y = strlen(x);

y = 9

See also strsect, strindx, strrindx

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strput

Purpose Lays a substring over a string. **Format** y = strput(substr,str,off); Input substr string, the substring to be laid over the other string. string, the string to receive the substring. str off scalar, the offset in str to place substr. The offset of the first byte is 1. **Output** string, the new string. **Example** str = "max"; sub = "imum"; f = 4;y = strput(sub, str, f);print y; Produces: maximum Source strput.src

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strrindx

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strrindx

Purpose Finds the index of one string within another string. Searches from the end of the string to the beginning.

Format y = strrindx(where, what, start);

Input where string or scalar, the data to be searched.

what string or scalar, the substring to be searched for in where.

scalar, the starting point of the search in *where* for an occurrence of *what. where* will be searched from this point backward for *what.*

Output y scalar containing the index of the last occurrence of *what*, within *where*, which is less than or equal to *start*. If no occurrence is found, it will be 0.

Remarks A negative value for *start* causes the search to begin at the end of the string. An example of the use of **strrindx** is extracting a file name from a complete path specification:

```
path = "/gauss/src/ols.src";
ps = "/";
pos = strrindx(path,ps,-1);
if pos;
  name = strsect(path,pos+1,strlen(path));
else;
  name = "";
endif;

pos = 11
name = "ols.src"
```

strrindx can be used with strsect for extracting substrings.

See also strindx, strlen, strsect, strput

strsect

strsect

a **Purpose** Extracts a substring of a string. b **Format** y = strsect(str,start,len); Input d str string or scalar from which the segment is to be obtained. scalar, the index of the substring in str. start The index of the first character is 1. scalar, the length of the substring. len **Output** string, the extracted substring, or a null string if *start* is y greater than the length of str. h Remarks If there are not enough characters in a string for the defined substring to be extracted, then a short string or a null string will be returned. If str is a matrix containing character data, it must be scalar. k **Example** strng = "This is an example string." y = strsect(strng, 12, 7);m n y = example0 See also strlen, strindx, strrindx p q

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x y z

strsplit

strsplit

Purpose Splits an Nx1 string vector into an NxK string array of the individual tokens.

Format *sa* = strsplit(*sv*);

Input *sv* Nx1 string array.

Output *sa* NxK string array.

Remarks Each row of *sv* must contain the same number of tokens. The following characters are considered delimiters between tokens:

space	ASCII 32
tab	ASCII 9
comma	ASCII 44
newline	ASCII 10
carriage return	ASCII 13

Tokens containing delimiters must be enclosed in single or double quotes or parentheses. Tokens enclosed in single or double quotes will NOT retain the quotes upon translation. Tokens enclosed in parentheses WILL retain the parentheses after translation. Parentheses cannot be nested.

```
Example let string sv = {
```

```
"dog 'cat fish' moose",
"lion, zebra, elk",
"seal owl whale"
};
sa = strsplit(sv);
```

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strsplit

'dog' 'cat fish' 'moose'
sa = 'lion' 'zebra' 'elk'
'seal' 'owl' 'whale'

See also strsplitPad

3-864

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strsplitPad

strsplitPad

Purpose

Splits a string vector into a string array of the individual tokens. Pads on the right with null strings.

Format

```
sa = strsplitPad(sv, cols);
```

Input

Nx1 string array. sv

scalar, number of columns of output string array. cols

Output

Nxcols string array. sa

Remarks

Rows containing more than *cols* tokens are truncated and rows containing fewer than *cols* tokens are padded on the right with null strings. The following characters are considered delimiters between tokens:

space	ASCII 32
tab	ASCII 9
comma	ASCII 44
newline	ASCII 10
carriage return	ASCII 13

Tokens containing delimiters must be enclosed in single or double quotes or parentheses. Tokens enclosed in single or double quotes will NOT retain the quotes upon translation. Tokens enclosed in parentheses WILL retain the parentheses after translation. Parentheses cannot be nested.

Example

```
let string sv = {
"dog 'cat fish' moose",
"lion, zebra, elk, bird",
"seal owl whale"
};
sa = strsplitPad(sv, 4);
```

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strsplitPad

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'dog' 'cat fish' 'moose' ''
sa = 'lion' 'zebra' 'elk' 'bird'
'seal' 'owl' 'whale' ''

See also strsplit

3-866

strtodt

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V

strtodt

Purpose Converts a string array of dates to a matrix in DT scalar format. **Format** x = strtodt(sa, fmt);Input NxK string array containing dates. sa string containing date/time format characters. fmt **Output** NxK matrix of dates in DT scalar format. **Remarks** The DT scalar format is a double precision representation of the date and time. In the DT scalar format, the number 20010421183207 represents 18:32:07 or 6:32:07 PM on April 21, 2001. The following formats are supported: YYYY 4 digit year Last two digits of year YR MO Number of month, 01-12 DD Day of month, 01-31 HH Hour of day, 00-23 ΜI Minute of hour, 00-59 SS Second of minute, 00-59 **Example** x = strtodt("2001-03-25 14:58:49","YYYY-MO-DD HH:MI:SS"); print x; produces: 20010325145849.0

3-867

strtodt

```
x = strtodt("2001-03-25 14:58:49", "YYYY-MO-DD");
                       print x;
                       produces:
b
                       20010325000000.0
d
                       x = strtodt("14:58:49", "HH:MI:SS");
                       print x;
                       produces:
h
                       145849.0
                       x = strtodt("04-15-00", "MO-DD-YR");
                       print x;
                       produces:
m
                       20000415000000.0
n
           See also
                       dttostr, dttoutc, utctodt
0
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q
```

3-868

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strtof

strtof

Purpose Converts a string array to a numeric matrix.

Format x = strtof(sa);

Input *sa* NxK string array containing numeric data.

Output x NxK matrix.

Remarks This function supports real matrices only. Use **strtofcplx** for

complex data.

See also strtofcplx, ftostrC

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strtofcplx

Purpose Converts a string array to a complex numeric matrix.

Format x = strtofcplx(sa);

Input *sa* NxK string array containing numeric data.

Output x NxK complex matrix.

Remarks strtofcplx supports both real and complex data. It is slower than

strtof for real matrices. **strtofcplx** requires the presence of the

real part. The imaginary part can be absent.

See also strtof, ftostrC

x y z

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3-870

submat

submat

Purpose Extracts a submatrix of a matrix, with the appropriate rows and columns given by the elements of vectors.

Format y = submat(x,r,c);

Input x NxK matrix.

r LxM matrix of row indices.

c PxQ matrix of column indices.

Output y (L*M)x(P*Q) submatrix of x, y may be larger than x.

Remarks If r = 0, then all rows of x will be used. If c = 0, then all columns of x will be used.

Example let x[3,4] = 1 2 3 4 5 6 7 8 9 10 11 12;

let v1 = 1 3;let v2 = 2 4;

y = submat(x,v1,v2);

z = submat(x, 0, v2);

 $y = \begin{array}{cc} 2 & 4 \\ 10 & 12 \end{array}$

 $z = \begin{array}{c} 2 & 4 \\ 6 & 8 \\ 10 & 12 \end{array}$

See also diag, vec, reshape

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subscat

subscat

Purpose

Changes the values in a vector depending on the category a particular element falls in.

Format

y = subscat(x, y, s);

Input

Nx1 vector.

Px1 numeric vector, containing breakpoints specifying the ranges within which substitution is to be made. This MUST be sorted in ascending order.

> v can contain a missing value as a separate category if the missing value is the first element in v.

If *v* is a scalar, all matches must be exact for a substitution to be made.

Px1 vector, containing values to be substituted. S

Output

Nx1 vector, with the elements in s substituted for the original elements of x according to which of the regions the elements of x fall into:

$$v[1] < x \le v[1] \to s[1]$$

$$x \le v[2] \to s[2]$$

$$v[p-1] < \begin{array}{c} x \le v[p] \to & s[p] \\ x > v[p] \to & \text{the original value of } x \end{array}$$

If missing is not a category specified in v, missings in x are passed through without change.

a

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W

x y z

subscat

```
let x = 1 2 3 4 5 6 7 8 9 10;
let v = 4 5 8;
let s = 10 5 0;
y = subscat(x,v,s);
```

 $y = \begin{array}{c} 10 \\ 10 \\ 10 \\ 5 \\ 0 \\ 0 \\ 0 \\ 9 \\ 10 \end{array}$

10

a

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c d

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...

substute

substute

Purpose b

Substitutes new values for old values in a matrix, depending on the outcome of a logical expression.

Format

```
y = substute(x,e,v);
```

Input

- NxK matrix containing the data to be changed. x
- LxM matrix, ExE conformable with x containing 1's and e0's.

Elements of x will be changed if the corresponding element of e is 1.

PxQ matrix, ExE conformable with x and e, containing the v values to be substituted for the original values of x when the corresponding element of e is 1.

Output

max(N,L,P) by max(K,M,Q) matrix.

Remarks

The *e* matrix is usually the result of an expression or set of expressions using dot conditional and boolean operators.

Example

```
x = \{ Y
          55
              30,
      N
          57
              18,
      Υ
          24
             3,
      N
          63
             38,
          55
              32,
      Υ
          37
              11 };
      N
e = x[.,1] .$== "Y" .and x[.,2] .>= 55 .and x[.,3]
      .>= 30;
x[.,1] = substute(x[.,1],e,0$+"R");
```

a

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substute

$$e = \begin{array}{c} 1 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \end{array}$$

Here is what *x* looks like after substitution:

$$y = \begin{cases} R & 55 & 30 \\ N & 57 & 18 \end{cases}$$

$$y = \begin{cases} Y & 24 & 3 \\ N & 63 & 38 \\ R & 55 & 32 \\ N & 37 & 11 \end{cases}$$

Source datatran.src

See also code, recode

b

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XX7

sumc

sumc

Purpose

Computes the sum of each column of a matrix.

Format y = sumc(x);

Input x NxK matrix.

Output y Kx1 vector, the sums of each column of x.

Remarks To find the sums of the elements in each row of a matrix, transpose before applying **sumc**. If x is complex, use the bookkeeping transpose (\cdot ').

To find the sums of all of the elements in a matrix, use the **vecr** function before applying **sumc**.

Example x = round(rndu(5,3)*5);

y = sumc(x);

2 4 3

 $x = \begin{array}{cc} 2 & 1 & 2 \\ 5 & 1 & 3 \end{array}$

5 1 1

3 3 4

17

y = 10

13

See also

cumsumc, meanc, stdc

a

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11

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p

q

S

u

V

W

x y z

surface

surface

Purpose Graphs a 3-D surface.

Library pgraph

Format surface(x,y,z);

Input x 1xK vector, the X axis data.

y Nx1 vector, the Y axis data.

z NxK matrix, the matrix of height data to be plotted.

Global Input psurf 2x1 vector, co

2x1 vector, controls 3-D surface characteristics.
[1] if 1, show hidden lines. Default 0.

[2] color for base (default 7). The base is an outline of the X-Y plane with a line connecting each corner to

the surface. If 0, no base is drawn.

_pticout scalar, if 0 (default), tick marks point inward, if 1, tick marks point outward.

_pzclr Z level color control.

There are 3 ways to set colors for the Z levels of a surface graph.

1. To specify a single color for the entire surface plot, set the color control variable to a scalar value 1–15. Example:

$$_pzclr = 15;$$

2. To specify multiple colors distributed evenly over the entire Z range, set the color control variable to a vector containing the desired colors only. GAUSS will automatically calculate the required corresponding Z values for you. The following example will produce a three color surface plot, the Z ranges being lowest=blue, middle=light blue, highest=white:

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3. To specify multiple colors distributed over selected ranges, the Z ranges as well as the colors must be manually input by the user. The following example assumes -0.2 to be the minimum value in the z matrix:

Since a Z level is required for each selected color, the user must be responsible to compute the minimum value of the z matrix as the first Z range element. This may be most easily accomplished by setting the **_pzclr** matrix as shown above (the first element being an arbitrary value), then reset the first element to the minimum z value as follows:

```
_{pzclr} = \{ -0.0 1,
             0.0 10,
             0.2 15 };
pzclr [1,1] = minc(minc(z));
    0 Black
               8 Dark Grey
    1 Blue
               9 Light Blue
    2 Green
              10 Light Green
    3 Cyan
              11 Light Cyan
    4 Red
              12 Light Red
    5 Magenta 13 Light Magenta
    6 Brown
              14 Yellow
    7 Grey
              15 White
```

Remarks

surface uses only the minimum and maximum of the X axis data in generating the graph and tic marks.

Source

psurface.src

surface

See also volume, view

a

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svd

svd

a **Purpose** Computes the singular values of a matrix. b Format s = svd(x);Input NxP matrix whose singular values are to be computed. d **Output** Mx1 vector, where M = min(N,P), containing the singular S values of x arranged in descending order. Global **_svderr** global scalar, if not all of the singular values can be **Output** computed _svderr will be nonzero. The singular values h in $s[_svderr+1]$, ... s[M] will be correct. Remarks Error handling is controlled with the low bit of the trap flag. trap 0 set _svderr and terminate with message k trap 1 set _svderr and continue execution **Example** $x = \{ 4 \ 3 \ 6 \ 7,$ 8 2 9 5 }, m y = svd(x);n 0 v = 16.5217873.3212254 p Source svd.src q See also svd1, svd2, svds

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svd1

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хух

svd1

```
Computes the singular value decomposition of a matrix so that:
Purpose
              x = u * s * v'.
  Format
              \{u,s,v\} = svd1(x);
    Input
                           NxP matrix whose singular values are to be computed.
              x
  Output
                           NxN matrix, the left singular vectors of x.
                           NxP diagonal matrix, containing the singular values of x
               S
                           arranged in descending order on the principal diagonal.
                           PxP matrix, the right singular vectors of x.
               ν
   Global
               svderr
                           global scalar, if all of the singular values are correct,
  Output
                           svderr is 0. If not all of the singular values can be
                           computed, svderr is set and the diagonal elements of s
                           with indices greater than _svderr are correct.
Remarks
               Error handling is controlled with the low bit of the trap flag.
               trap 0
                           set svderr and terminate with message
               trap 1
                           set svderr and continue execution
Example
               x = rndu(3,3);
               \{ u, s, v \} = svd1(x);
                     x = 0.85474208 \quad 0.79673540 \quad 0.22482095
                     0.33340653  0.74443792  0.75698778
                     -0.57955818 0.65204491 1.48882486
               u = -0.61005618 \quad 0.05056673 \quad -0.79074298
                     -0.54031821 -0.75649219 0.36847767
```

svd1

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хух

 $s = \begin{cases} 1.84994646 & 0.00000000 & 0.000000000 \\ 0.00000000 & 0.60370542 & 0.00000000 \\ 0.00000000 & 0.00000000 & 0.47539239 \end{cases}$

 $v = \begin{array}{rrr} -0.68578561 & 0.71062560 & -0.15719208 \\ -0.54451302 & -0.64427479 & -0.53704336 \\ -0.48291165 & -0.28270348 & 0.82877927 \end{array}$

Source svd.src

See also svd, svd2, svdusv

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svd2

Computes the singular value decomposition of a matrix so that: **Purpose** x = u * s * v' (compact u). Format $\{u,s,v\} = svd2(x);$ Input NxP matrix whose singular values are to be computed. x **Output** NxN or NxP matrix, the left singular vectors of x. If N > P, then u will be NxP containing only the P left singular vectors of x. NxP or PxP diagonal matrix, containing the singular S values of x arranged in descending order on the principal diagonal. If N > P, then s will be PxP. PxP matrix, the right singular vectors of x. v Global svderr global scalar, if all of the singular values are correct, **Output** _svderr is 0. If not all of the singular values can be computed, **svderr** is set and the diagonal elements of s with indices greater than **svderr** are correct. Remarks Error handling is controlled with the low bit of the trap flag. set **svderr** and terminate with message trap 0 trap 1 set **svderr** and continue execution Source svd.src Globals svderr See also svd, svd1, svdcusv

хух

svdcusv

svdcusv

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```
Computes the singular value decomposition of a matrix so that:
Purpose
                x = u * s * v' (compact u).
  Format
                \{ u, s, v \} = svdcusv(x);
    Input
                        NxP matrix whose singular values are to be computed.
  Output
                        NxN or NxP matrix, the left singular vectors of x. If N > P,
                        u is NxP containing only the P left singular vectors of x.
                        NxP or PxP diagonal matrix, the singular values of x
                S
                        arranged in descending order on the principal diagonal.
                        If N > P. s is PxP.
                        PxP matrix, the right singular vectors of x.
Remarks
                If not all the singular values can be computed, s[1,1] is set to a scalar
                error code. Use scalerr to convert this to an integer. The diagonal
                elements of s with indices greater than scalerr (s[1,1]) are correct. If
                scalerr (s[1,1]) returns a 0, all the singular values have been
                computed.
See also
                svd2, svds, svdusv
```

W

svds

svds

Purpose Computes the singular values of a matrix.

Format s = svds(x);

Input x NxP matrix whose singular values are to be computed.

Output s min(N,P)x1 vector, the singular values of x arranged in descending order.

Remarks If not all the singular values can be computed, s[1] is set to a scalar error

code. Use **scalerr** to convert this to an integer. The elements of s with indices greater than **scalerr**(s[1]) are correct. If **scalerr**(s[1])

returns a 0, all the singular values have been computed.

See also svd, svdcusv, svdusv

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хух

svdusv

svdusv

a b

d

h

Purpose

Computes the singular value decomposition of a matrix so that:

x = u * s * v'.

Format

 $\{ u, s, v \} = svdusv(x);$

Input

x NxP matrix whose singular values are to be computed.

Output

u NxN matrix, the left singular vectors of x.

s NxP diagonal matrix, the singular values of x arranged in descending order on the principal diagonal.

 ν PxP matrix, the right singular vectors of x.

Remarks

If not all the singular values can be computed, s[1,1] is set to a scalar error code. Use **scalerr** to convert this to an integer. The diagonal elements of s with indices greater than **scalerr**(s[1,1]) are correct. If **scalerr**(s[1,1]) returns a 0, all the singular values have been computed.

See also

svd1, svdcusv, svds

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sysstate

Purpose	Gets or sets general system parameters.			
Format	{ rets } = sysstate(case,y);			
Case 1 Case 2 through 7 Case 8	Case 1	Version Information		
		Returns the current GAUSS version information in an 8-element numeric vector.		
	- cases	GAUSS System Paths		
	through 7	Get or set GAUSS system path.		
	Case 8	Complex Number Toggle		
		Controls automatic generation of complex numbers in sqrt, ln, and log for negative arguments.		
	Case 9	Complex Trailing Character		
Case 10 Case 11 Case 12 Case 13 Case 14 Case 15	Get and set trailing character for the imaginary part of a complex number.			
	Case 10	Printer Width		
	Case 11	Get and set lprint width. Auxiliary Output Width		
		Get and set the auxiliary output width.		
	Case 12	Precision		
	Case 13	Get and set precision for positive definite matrix routines. LU Tolerance		
		Get and set singularity tolerance for LU decomposition.		
	Case 14	Cholesky Tolerance		
	Get and set singularity tolerance for Cholesky decomposition.			
	Screen State			
		Get and set window state as controlled by screen command.		
	Case 16	Automatic print Mode		
		Get and set automatic print mode.		

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x y z

Case 17 Automatic lprint Mode Get and set automatic lprint mode. **Auxiliary Output** Case 18 Get auxiliary output parameters. Case 19 **Get/Set Format** Get and set format parameters. Case 21 **Imaginary Tolerance** Get and set the imaginary tolerance. Case 22 **Source Path** Get and set the path the compiler will search for source files. **Dynamic Library Directory** Case 24 Get and set the path for the default dynamic library directory. Case 25 **Temporary File Path** Case 26 Interface Mode Returns the current interface mode. Case 28 **Random Number Generator Parameters** Case 30 **Base Year Toggle** Specifies whether year value returned by date is to include base year (1900) or not. Case 1: **Version Information** Returns the current GAUSS version information in an 8-element numeric vector. **Format** vi = sysstate(1,0);**Output** 8x1 numeric vector containing version information: vi [1] Major version number. Minor version number. [2] [3] Revision. [4] Machine type. Operating system. [5] Runtime module. [6] [7] Light version. [8] Always 0.

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	vi[4] indicates the type of machine on which GAUSS is running:			
	1 Intel x86			
	2	2	Sun SPARC	
	3	3	IBM RS/6000	
	۷	1	HP 9000	
	4	5	SGI MIPS	
	(5	DEC Alpha	
	<i>vi</i> [5] indicates the operating system on which GAUSS is running:			
	1 DOS			
	2	2	SunOS 4.1.x	
	3	3	Solaris 2.x	
	۷	1	AIX	
	4	5	HP-UX	
	6	5	IRIX	
	7	7	OSF/1	
	8	3	OS/2	
	Ģ)	Windows	
Cases 2 through 7:	GAUSS System Paths			
	Get or set GAUSS system path.			
Format	oldpath = sysstate(case,path);			
Input	case scalar 2-7, path to set.			
	2	2	. exe file location.	
	3	3	loadexe path.	
	2	4	save path.	
	4	5	load, loadm path.	
	(5	loadf, loadp path.	
	-	7	loads path.	
	path s	scalar	0 to get path, or string containing the new path.	
Output	oldpath	S	tring, original path.	
Remarks	If path is	of ty	pe matrix, the path will be returned but not modified.	

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Case 8: Complex Number Toggle

Controls automatic generation of complex numbers in sqrt, ln and log for positive arguments

for negative arguments.

Format oldstate = sysstate(8,state);

Input state scalar, 1, 0, or -1.

Output *oldstate* scalar, the original state.

Remarks If *state* = 1, log, ln, and sqrt will return complex numbers for

negative arguments. If state = 0, the program will terminate with an error message when negative numbers are passed to log, ln, and sqrt. If state = -1, the current state is returned and left unchanged. The default

state is 1.

Case 9: Complex Trailing Character

Get and set trailing character for the imaginary part of a complex number.

Format oldtrail = sysstate(9,trail);

Input trail scalar 0 to get character, or string containing the new

trailing character.

Output *oldtrail* string, the original trailing character.

Remarks The default character is "i".

Case 10: Printer Width

Get and set **lprint** width.

Format oldwidth = sysstate(10,width);

Input width scalar, new printer width.

Output oldwidth scalar, the current original width.

Remarks If *width* is 0, the printer width will not be changed.

This may also be set with the **lpwidth** command.

See also lpwidth

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Case 11: Auxiliary Output Width

Get and set the auxiliary output width.

Format oldwidth = sysstate(11,width);

Input width scalar, new output width.

Output *oldwidth* scalar, the original output width.

Remarks If *width* is 0 then the output width will not be changed.

This may also be set with the **outwidth** command.

See also outwidth

Case 12: Precision

Get and set precision for positive definite matrix routines.

Format oldprec = sysstate(12,prec);

Input *prec* scalar, 64 or 80.

Output *oldprec* scalar, the original value.

Portability Windows, UNIX

This function has no effect under Windows or UNIX. All computations are done in 64-bit precision (except for operations done entirely within

the 80×87 on Intel machines).

Remarks The precision will be changed if *prec* is either 64 or 80. Any other number

will leave the precision unchanged.

See also prcsn

Case 13: LU Tolerance

Get and set singularity tolerance for LU decomposition.

Format oldtol = sysstate(13,tol);

Input *tol* scalar, new tolerance.

Output *oldtol* scalar, the original tolerance.

h

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h

k

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S

u

V

W

x y z

Remarks The tolerance must be >= 0. If *tol* is negative, the tolerance is returned

and left unchanged.

See also croutp, inv

Case 14: Cholesky Tolerance

Get and set singularity tolerance for Cholesky decomposition.

Format oldtol = sysstate(14,tol);

Input *tol* scalar, new tolerance.

Output *oldtol* scalar, the original tolerance.

Remarks The tolerance must be ≥ 0 . If *tol* is negative, the tolerance is returned

and left unchanged.

See also chol, invpd, solpd

Case 15: Screen State

Get and set window state as controlled by **screen** command.

Format oldstate = sysstate(15,state);

Input *state* scalar, new window state.

Output *oldstate* scalar, the original window state.

Remarks If *state* = 1, window output is turned on. If *state* = 0, window output is

turned off. If state = -1, the state is returned unchanged.

See also screen

Case 16: Automatic print Mode

Get and set automatic **print** mode.

Format oldmode = sysstate(16,mode);

Input *mode* scalar, mode.

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x y z

Output oldmode scalar, original mode. Remarks If mode = 1, automatic **print** mode is turned on. If mode = 0, it is turned off. If mode = -1, the mode is returned unchanged. See also print on/off **Case 17:** Automatic lprint Mode Get and set automatic lprint mode. **Format** oldmode = sysstate(17,mode); Input mode scalar, mode. Output oldmode scalar, original mode. Remarks If mode = 1, automatic **lprint** mode is turned on. If mode = 0, it is turned off. If mode = -1, the mode is returned unchanged. See also lprint on/off **Case 18: Auxiliary Output** Get auxiliary output parameters. **Format** { state, name } = sysstate(18, dummy); Input dummy scalar, a dummy argument. **Output** state scalar, auxiliary output state, 1 - on, 0 - off. string, auxiliary output filename. name See also output **Case 19: Get/Set Format** Get and set format parameters. **Format** oldfmt = sysstate(19,fmt);

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Input	fmt	scalar or 11x1 column vector containing the new format parameters. Usually this will have come from a previous sysstate(19,0) call. See Output for description of matrix.		
Output	oldfmt	11x1 v [1] [2] [3] [4] [5] [6] [7] [8] [9] [10] [11]	ector containing the current format parameters: format type. justification. sign. leading zero. trailing character. row delimiter. carriage line feed position. automatic line feed for row vectors. field. precision. formatted flag	
Remarks	If <i>fmt</i> is scalar 0, then the format parameters will be left unchanged.			
See also	format			
Case 21:	Imaginary Tolerance Get and set the imaginary tolerance.			
Format Input Output	tol oldtol	sysstate(21,tol); scalar, the new tolerance. scalar, the original tolerance.		
Remarks	The imaginary tolerance is used to test whether the imaginary part of a complex matrix can be treated as zero or not. Functions that are not defined for complex matrices check the imaginary part to see if it can be ignored. The default tolerance is 2.23e–16, or machine epsilon. If <i>tol</i> < 0, the current tolerance is returned.			
See also	hasimag			

Case 22: Source Path

Get and set the path the compiler will search for source files.

Format oldpath = sysstate(22,path);

Input path scalar 0 to get path, or string containing the new path.

Output *oldpath* string, original path.

Remarks If *path* is a matrix, the current source path is returned.

This resets the **src_path** configuration variable. **src_path** is initially defined in the GAUSS configuration file, gauss.cfq.

path can list a sequence of directories, separated by semicolons.

Resetting **src_path** affects the path used for subsequent **run** and **compile** statements.

Case 24: Dynamic Library Directory

Get and set the path for the default dynamic library directory.

Format oldpath = sysstate(24,path);

Input path scalar 0 to get path, or string containing the new path.

Output *oldpath* string, original path.

Remarks If *path* is a matrix, the current path is returned.

path should list a single directory, not a sequence of directories.

Changing the dynamic library path does not affect the state of any DLL's currently linked to GAUSS. Rather, it determines the directory that will be searched the next time **dlibrary** is called.

UNIX

Changing the path has no effect on GAUSS's default DLL, libgauss.so. libgauss.so must always be located in the GAUSSHOME directory.

Windows

Changing the path has no effect on GAUSS' default DLL, gauss.dll. gauss.dll must always be located in the GAUSSHOME directory.

OS/2

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W

Changing the path has no effect on GAUSS's default DLL, gauss.dll. gauss.dll must always be located in the same directory as the GAUSS executable, gauss.exe.

DOS

sysstate 24 has no effect, as dlibrary and dllcall are not supported.

See also dlibrary, dllcall

Case 25: Temporary File Path

Get or set the path GAUSS will use for temporary files...

Input path scalar 0 to get path, or string containing the new path.

Output oldpath string, original path.

Remarks If path is of type matrix, the path will be returned but not modified.

Case 26: Interface Mode

Returns the current interface mode.

Format mode = sysstate(26,0);

Output scalar, interface mode flag mode

non-X mode

1 terminal (-v) mode

2 X Windows mode

Remarks A mode of 0 indicates that you're running a non-X version of GAUSS;

i.e., a version that has no X Windows capabilities. A mode of 1 indicates that you're running an X Windows version of GAUSS, but in terminal mode; i.e., you started GAUSS with the -v flag. A mode of 2 indicates

that you're running GAUSS in X Windows mode.

Case 28: Random Number Generator Parameters

Get and set the random number generator (RNG) parameters.

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Input	prms	scalar 0 to get parameters, or 3x1 matrix of new parameters.

[1] seed, $0 < \text{seed} < 2^32$

[2] multiplier, $0 < \text{mult} < 2^32$ [3] constant, $0 < \text{const} < 2^32$

Output *oldprms* 3x1 matrix, current parameters.

oldprms = sysstate(28,prms);

Portability Not supported for DOS.

Format

Remarks If *prms* is a scalar 0, the current parameters will be returned without being changed.

The modulus of the RNG cannot be changed; it is fixed at 2³².

See also rndcon, rndmult, rndseed, rndns, rndus, rndn, rndu

Case 30: Base Year Toggle

Specifies whether year value returned by date is to include base year (1900) or not.

Format oldstate = sysstate(30,state);

Input state scalar, 1, 0, or missing value.

Output *oldstate* scalar, the original state.

Portability DOS

sysstate 30 has no effect. It always returns a 4-digit year.

Remarks Internally, date acquires the number of years since 1900. sysstate 30 specifies whether date should add the base year to that value or not. If state = 1, date adds 1900, returning a fully-qualified 4-digit year.

If **state** = 0, date returns the number of years since 1900. If *state* is a missing value, the current state is returned. The default state is 1.

system

a

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system

Purpose Quits GAUSS and returns to the operating system.

Format system;
system c;

Input c scalar, an optional exit code that can be recovered by the program that invoked GAUSS. The default is 0. Valid arguments

are 0-255.

Remarks The system command always returns an exit code to the operating system

or invoking program. If you don't supply one, it returns 0. This is usually

interpreted as indicating success.

See also exec

tab

tab

Purpose Tabs the cursor to a specified text column.

Format tab(col);

print expr1 expr2 tab(col1) expr3 tab(col2) expr4
...;

Input *col* scalar, the column position to tab to.

Remarks *col* specifies an absolute column position. If *col* is not an integer, it will be truncated.

tab can be called alone or embedded in a print statement. You cannot embed it within a parenthesized expression in a print statement, though. For example:

```
print (tab(20) c + d * e);
```

will not give the results you expect. If you have to use parenthesized expressions, write it like this instead:

```
print tab(20) (c + d * e);
```

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x y z

Purpose Returns the tangent of its argument.

Format $y = \tan(x)$;

Input x NxK matrix.

Output y NxK matrix.

Remarks For real matrices, *x* should contain angles measured in radians.

To convert degrees to radians, multiply the degrees by $\frac{\pi}{180}$.

Example let x = 0 .5 1 1.5; y = tan(x); 0.000000000y = 0.54630249

 $y = \begin{array}{r} 0.54630249 \\ 1.55740772 \\ 14.10141995 \end{array}$

See also atan, pi

tanh

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V

хух

tanh

```
Purpose
              Computes the hyperbolic tangent.
 Format
              y = tanh(x);
    Input
                   NxK matrix.
  Output
                   NxK matrix containing the hyperbolic tangents of the elements of
                   х.
Example
              let x = -0.5 - 0.25 \ 0 \ 0.25 \ 0.5 \ 1;
              x = x * pi;
              y = tanh(x);
                   -1.570796
                   -0.785398
                    0.000000
              x =
                    0.785398
                     1.570796
                    3.141593
                   -0.917152
                   -0.655794
                    0.000000
              y =
                    0.655794
                    0.917152
                    0.996272
              trig.src
  Source
```

tempname

tempname

Purpose Creates a temporary file with a unique name.

Format tname = tempname(path, pre, suf);

Input path string, path where the file will reside.

string, a prefix to begin the file name with.sufstring, a suffix to end the file name with.

Output *tname* string, unique temporary file name of the form

path/preXXXXnnnnnsuf, where XXXX are 4 letters, and nnnnn

is the process id of the calling process.

Remarks Any or all of the inputs may be a null string or 0. If *path* is not specified, the current working directory is used.

If unable to create a unique file name of the form requested, **tempname** returns a null string.

WARNING: GAUSS does not remove temporary files created by **tempname**. It is left to the user to remove them when they are no longer needed.

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time

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time

Returns the current system time. Purpose **Format** y = time;Output 4x1 numeric vector, the current time in the order: hours, minutes, seconds, and hundredths of a second. **Example** print time; 7.000000 31.000000 46.000000 33.000000 See also date, datestr, datestring, datestrymd, hsec, timestr

timedt

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timedt

Purpose Returns system date and time in DT scalar format.

Format dt = timedt;

Output dt scalar, system date and time in DT scalar format.

Remarks The DT scalar format is a double precision representation of the date and

time. In the DT scalar format, the number

20010421183207

represents 18:32:07 or 6:32:07 PM on April 21, 2001.

Source time.src

See also todaydt, timeutc, dtdate

timestr

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хух

timestr

```
Purpose
             Formats a time in a vector to a string.
 Format
             ts = timestr(t);
    Input
                   4x1 vector from the time function, or a zero. If the input is 0, the
                   time function will be called to return the current system time.
  Output
                   8 character string containing current time in the format hr:mn:sc
             t = \{ 7, 31, 46, 33 \};
Example
              ts = timestr(t);
             print ts;
              Produces:
                 7:31:46
 Source
              time.src
See also
              date, datestr, datestring, datestrymd, ethsec,
              etstr, time
```

timeutc

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timeutc

Purpose Returns the number of seconds since January 1, 1970 Greenwich Mean Time. Format tc = timeutc; Output scalar, number of seconds since January 1, 1970 Greenwich Mean Time. **Example** tc = timeutc; utv = utctodtv(tc); tc = 939235033utv = 1999 10 6 11 37 13 3 278 See also dtvnormal, utctodtv

x y z

title

title

Purpose Sets the title for the graph. Library pgraph Format title(str); Input string, the title to display above the graph. Global ptitle **Output** Remarks Up to three lines of title may be produced by embedding a line feed character ("\L") in the title string. **Example** title("First title line\L Second title line\L Third title line"); Fonts may be specified in the title string. For instructions on using fonts, see "Publication Quality Graphics" in the User Guide. Source pgraph.src See also xlabel, ylabel, fonts

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tkf2eps

tkf2eps

Purpose Converts a .tkf file to an Encapsulated PostScript file.

Library pgraph

Format ret = tkf2eps(tekfile, epsfile);

Input *tekfile* string, name of .tkf file

epsfile string, name of Encapsulated PostScript file

Output ret scalar, 0 if successful

Remarks The conversion is done using the global parameters in peps.dec. You can modify these globally by editing the .dec file, or locally by setting

them in your program before calling tkf2eps.

See the header of the output Encapsulated PostScript file and a PostScript

manual if you want to modify these parameters.

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x y z

tkf2ps

tkf2ps

Purpose Converts a .tkf file to a PostScript file.

Library pgraph

Format ret = tkf2ps(tekfile, psfile);

Input *tekfile* string, name of .tkf file

epsfile string, name of Encapsulated PostScript file

Output ret scalar, 0 if successful

Remarks The conversion is done using the global parameters in peps.dec. You can modify these globally by editing the .dec file, or locally by setting

them in your program before calling tkf2ps.

See the header of the output Encapsulated PostScript file and a PostScript

manual if you want to modify these parameters.

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tocart

Purpose Converts from polar to cartesian coordinates.

Format xy = tocart(r,theta);

Input *r* NxK real matrix, radius.

theta LxM real matrix, ExE conformable with r, angle in radians.

Output $xy = \max(N,L)$ by $\max(K,M)$ complex matrix containing the X

coordinate in the real part and the Y coordinate in the imaginary

part.

Source coord.src

todaydt

todaydt

Purpose Returns system date in DT scalar format. The time returned is always

midnight (00:00:00), the beginning of the returned day.

Format dt = todaydt;

Output dt scalar, system date in DT scalar format.

Remarks The DT scalar format is a double precision representation of the date and

time. In the DT scalar format, the number

20010421183207

represents 18:32:07 or 6:32:07 PM on April 21, 2001.

Source time.src

See also timedt, timeutc, dtdate

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toeplitz

toeplitz

a Creates a Toeplitz matrix from a column vector. **Purpose** b **Format** t = toeplitz(x); Input Kx1 vector. d е Output KxK Toeplitz matrix. Example g h 1 2 x =3 4 k 5 1 2 3 4 5 m 2 1 2 3 4 n 3 2 1 2 3 4 3 2 1 2 0 5 4 3 2 1 p

```
x = seqa(1,1,5);
         y = toeplitz(x);
Source
         toeplitz.src
```

q

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token

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x y z

token

```
Extracts the leading token from a string.
Purpose
  Format
               { token, str_left } = token(str);
    Input
               str
                        string, the string to parse.
  Output
               token
                            string, the first token in str.
               str left
                            string, str minus token.
Remarks
               str can be delimited with commas or spaces.
               The advantage of token over parse is that parse is limited to tokens
               of 8 characters or less; token can extract tokens of any length.
Example
               Here is a keyword that uses token to parse its string parameter.
                  keyword add(s);
                      local tok, sum;
                      sum = 0;
                      do until s $== "";
                          \{ \text{ tok, s } \} = \text{token(s);}
                          sum = sum + stof(tok);
                      endo;
                      format /rd 1,2;
                      print "Sum is: " sum;
                   endp;
               If you type:
                   add 1 2 3 4 5 6;
               add will respond:
                   Sum is: 21.00
  Source
               token.src
```

token

See also parse

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topolar

topolar

Purpose Converts from cartesian to polar coordinates.

Format { r,theta } = topolar(xy);

Input xy NxK complex matrix containing the X coordinate in the real part

and the Y coordinate in the imaginary part.

Output r NxK real matrix, radius.

theta NxK real matrix, angle in radians.

Source coord.src

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trace

trace

Purpose

h

Allows the user to trace program execution for debugging purposes.

Format

trace new;

trace new, mask;

Input

new scalar, new value for trace flag.

mask scalar, optional mask to allow leaving some bits of the trace flag

unchanged.

Remarks

The **trace** command has no effect unless you are running your program under GAUSS's source level debugger. Setting the **trace** flag will not generate any debugging output during normal execution of a program.

The argument is converted to a binary integer with the following meanings:

decimal	meaning
1	trace calls/returns
2	trace line numbers
4	verbose trace
8	output to window
16	output to print
32	output to auxiliary output
64	output to error log
	1 2 4 8 16 32

You must set one or more of the output bits to get any output from **trace**. If you set **trace** to 4, you'll be doing a verbose trace of your program, but the output won't be displayed anywhere.

The trace output as a program executes will be as follows:

(+GRAD) calling function or procedure GRAD(-GRAD) returning from GRAD

[47] executing line 47

Note that the line number trace will only produce output if the program was compiled with line number records.

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trace

To set a single bit use two arguments:

trace 16,16; turn on output to printer trace 0,16; turn off output to printer

Example

trace 1+8; trace fn/proc calls/returns to standard output trace line numbers to standard trace 2+8; output trace 1+2+8; trace line numbers and fn/proc calls/returns to standard output trace 1+16; trace fn/proc calls/returns to printer trace 2+16; trace line numbers to printer trace 1+2+16; trace line numbers and fn/proc calls/returns to printer trace 4+8; verbose trace to screen

See also #lineson

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Purpose Sets the trap flag to enable or disable trapping of numerical errors.

Format trap new;

trap new, mask;

Input

new scalar, new trap value.

mask

scalar, optional mask to allow leaving some bits of the trap flag

unchanged.

Remarks

The trap flag is examined by some functions to control error handling. There are 16 bits in the trap flag, but most GAUSS functions will examine only the lowest order bit:

trap 1; turn trapping on
trap 0; turn trapping off

If we extend the use of the trap flag, we will use the lower order bits of the trap flag. It would be wise for you to use the highest 8 bits of the trap flag if you create some sort of user-defined trap mechanism for use in your programs. (See the function **trapchk** for detailed instructions on testing the state of the trap flag; see **error** for generating user-defined error codes.)

To set only one bit and leave the others unchanged use two arguments:

trap 1,1; set the ones bit
trap 0,1; clear the ones bit

Example

```
x = eye(3);
oldval = trapchk(1);
trap 1,1;
y = inv(x);
trap oldval,1;
```

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```
trap
if scalerr(y);
   errorlog "WARNING: x is singular";
else;
   print "y" y;
endif;
In this example the result of inv is trapped in case x is singular. The trap
state is reset to the original value after the call to inv.
Run the example
x = eye(3);
It is inverted.
Now try
ones(3,3);
It isn't.
scalerr, trapchk, error
```

See also

trapchk

trapchk

Purpose Tests the value of the trap flag.

Format y = trapchk(m);

Input m scalar mask value.

Output y scalar which is the result of the bitwise logical AND of the trap flag and the mask.

Remarks To check the various bits in the trap flag, add the decimal values for the bits you wish to check according to the chart below and pass the sum in as the argument to the **trapchk** function:

bit	decimal value	
0	1	
1	2	
2	4	
3	8	
4	16	
5	32	
6	64	
7	128	
8	256	
9	512	
10	1024	
11	2048	
12	4096	
13	8192	
14	16384	
15	32768	

If you want to test if either bit 0 or bit 8 is set, then pass an argument of 1+256 or 257 to **trapchk**. The following table demonstrates values that will be returned for:

y=trapchk(257);

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trapchk

	0	1	value of bit 0 in trap flag
0	0	1	
1	256	257	
value of bit 8 in trap flag			

GAUSS functions that test the trap flag currently test only bits 0 and 1.

See also scalerr, trap, error

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trigamma

trigamma

Purpose Computes trigamma function.

Format y = trigamma(x);

Input x MxN matrix.

Output y MxN matrix, trigamma.

Remarks The trigamma function is the second derivative of the log of the gamma function with respect to argument.

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trimr

trimr

Trims rows from the top and/or bottom of a matrix. **Purpose Format** y = trimr(x,t,b);Input NXK matrix from which rows are to be trimmed. scalar containing the number of rows which are to be removed t from the top of x. b scalar containing the number of rows which are to be removed from the bottom of x. **Output** RxK matrix where R=N-(t+b) containing the rows left after the trim. Remarks If either t or b is zero, then no rows will be trimmed from that end of the matrix. Example x = rndu(5,3);y = trimr(x, 2, 1);0.76042751 0.33841579 0.01844780 0.05334503 0.38939785 0.65029973 x =0.93077511 0.06961078 0.04207563 0.53640701 0.06640062 0.07222560 0.14084669 0.06033813 0.69449247 $0.93077511 \ 0.06961078 \ 0.04207563$ y = 0.53640701 0.06640062 0.07222560 See also submat, rotater, shiftr

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trunc

trunc

Purpose Converts numbers to integers by truncating the fractional portion.

Format y = trunc(x);

Input x NxK matrix.

Output y NxK matrix containing the truncated elements of x.

Example x = 100*rndn(2,2);

$$x = \begin{array}{ccc} 77.68 & -14.10 \\ 4.73 & -158.88 \end{array}$$

y = trunc(x);

$$y = \begin{array}{r} 77.00 & -14.00 \\ 4.00 & -158.00 \end{array}$$

See also ceil, floor, round

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type

Purpose Returns the symbol table type of the argument.

Format t = type(x);

Input x matrix or string, can be an expression.

Output t scalar.

6 matrix 13 string 15 string array

Remarks

type returns the type of a single symbol. The related function typecv will take a character vector of symbol names and return a vector of either their types or the missing value code for any that are undefined. type works for matrices, strings, and string arrays; typecv works for user-defined procedures, keywords and functions as well. type works for global or local symbols; typecv works only for global symbols.

Example

```
print k;
if type(k) == 6;
    k = ""$+k;
endif;
print k;
produces
+DEN
CHARS
```

 $k = \{\text{"CHARS"}\};$

See also

typecv, typef

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Purpose Returns the symbol table type of objects whose names are given as a string or as elements of a character vector.

Format y = typecv(x);

Input x string or Nx1 character vector which contains the names of variables whose type is to be determined.

Output y scalar or Nx1 vector containing the types of the respective symbols in x.

Remarks The values returned by **typecv** for the various variable types are as follows:

6 Matrix (Numeric, Character, or Mixed)

8 Procedure (proc)

9 Function (**fn**)

5 Keyword (keyword)

13 String

15 String Array

It will return the GAUSS missing value code if the symbol is not found, so **typecv** may be used to determine if a symbol is defined or not.

Example xvar = sqrt(5);

```
yvar = "Montana";
fn area(r) = pi*r*r;
let names = xvar yvar area;
y = typecv(names);
```

 $names = \begin{cases} XVAR \\ YVAR \\ AREA \end{cases}$

typecv

$$y = \begin{array}{c} 6 \\ 13 \\ 9 \end{array}$$

See also type, typef, varput, varget

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```
Purpose Returns the type of data (the number of bytes per element) in a GAUSS data set.
```

Format y = typef(fp);

Input fp scalar, file handle of an open file.

Output y scalar, type of data in GAUSS data set.

Remarks If fp is a valid GAUSS file handle, then y will be set to the type of the data in the file as follows:

2 2-byte signed integer

4 4-byte IEEE floating point

8 8-byte IEEE floating point

```
infile = "dat1";
outfile = "dat2";
open fin = ^infile;
names = getname(infile);
```

create fout = ^outfile with ^names,0,typef(fin);

In this example a file dat2.dat is created which has the same variables and variable type as the input file, dat1.dat. typef is used to return the type of the input file for the create statement.

See also colsf, rowsf

union

union

Returns the union of two vectors with duplicates removed. **Purpose Format** y = union(v1, v2, flag);Input v1Nx1 vector. v2Mx1 vector. flag scalar, 1 if numeric data, 0 if character. **Output** Lx1 vector containing all unique values that are in v1 and v2, sorted in ascending order. The combined elements of v1 and v2 must fit into a single vector. Remarks **Example** let v1 = mary jane linda john; let v2 = mary sally; x = union(v1, v2, 0);**JANE JOHN** x =LINDA MARY SALLY

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uniqindx

uniqindx

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Purpose Computes the sorted index of x, leaving out duplicate elements.

Format

index = uniqindx(x,flag);

Input

x Nx1 or 1xN vector.

flag

scalar, 1 if numeric data, 0 if character.

Output

index Mx1 vector, indices corresponding to the elements of x sorted in

ascending order with duplicates removed.

Remarks

Among sets of duplicates it is unpredictable which elements will be

indexed.

Example

let x = 5 4 4 3 3 2 1;

ind = uniqindx(x,1);

y = x[ind];

7

ind = 5

2

2

1

y =

3

4 5

1

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unique

unique

Purpose Sorts and removes duplicate elements from a vector.

Format y = unique(x,flag);

Input x Nx1 or 1xN vector.

flag scalar, 1 if numeric data, 0 if character.

Output y Mx1 vector, sorted x with the duplicates removed.

Example let x = 5 4 4 3 3 2 1;

y = unique(x,1);

1 2

 $y = \frac{3}{3}$

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upmat, upmat1

upmat, upmat1

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Purpose

Returns the upper portion of a matrix. **upmat** returns the main diagonal and every element above. **upmat1** is the same except it replaces the main diagonal with ones.

Format

$$u = \text{upmat}(x);$$

 $u = \text{upmat1}(x);$

Input

x NxK matrix.

Output

NxK matrix containing the upper elements of the matrix. The lower elements are replaced with zeros. upmat returns the main diagonal intact. upmat1 replaces the main diagonal with ones.

Example

The resulting matrices are

$$u = \begin{array}{c} 1 & 2 & -1 \\ 0 & 3 & -2 \\ 0 & 0 & 1 \end{array}$$

$$u1 = \begin{array}{r} 1 & 2 & -1 \\ 0 & 1 & -2 \\ 0 & 0 & 1 \end{array}$$

Source

diag.src

v See also

lowmat, lowmat1, diag, diagrv, crout

W

u

upper

upper

Purpose Converts a string or matrix of character data to uppercase.

Format y = upper(x);

Input x string or NxK matrix containing the character data to be converted to uppercase.

Output y string or NxK matrix containing the uppercase equivalent of data in x.

Remarks If x is a numeric matrix, y will contain garbage. No error message will be generated since GAUSS does not distinguish between numeric and character data in matrices.

Example x = "uppercase";

y = upper(x);

y = UPPERCASE

See also lower

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use

use

a b

Purpose

Loads a compiled file at the beginning of the compilation of a source program.

Format

use fname;

Input

fname literal or 'string, the name of a compiled file created using the **compile** or the **saveall** command.

Remarks

The **use** command can be used ONCE at the TOP of a program to load in a compiled file which the rest of the program will be added to. In other words, if xy. e had the following lines:

```
library pgraph;
external proc xy;
x = seqa(0.1,0.1,100);
```

It could be compiled to xy.gcg. Then the following program could be run:

```
use xy;
xy(x,sin(x));
```

Which would be equivalent to:

```
new;
x = seqa(0.1,0.1,100);
xy(x,sin(x));
```

The **use** command can be used at the top of files that are to be compiled with the **compile** command. This can greatly shorten compile time for a set of closely related programs. For example:

```
library pgraph;
external proc xy,logx,logy,loglog,hist;
saveall pgraph;
```

This would create a file called pgraph.gcg containing all the procedures, strings and matrices needed to run PQG programs. Other

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use

programs could be compiled very quickly with the following statement at the top of each:

use pgraph;

or the same statement could be executed once, for instance from the command prompt, to instantly load all the procedures for PQG.

When the compiled file is loaded with **use**, all previous symbols and procedures are deleted before the program is loaded. It is therefore unnecessary to execute a **new** before **use**'ing a compiled file.

use can appear only ONCE at the TOP of a program.

See also compile, run, saveall

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utctodt

utctodt

Purpose Converts UTC scalar format to DT scalar format. b **Format** dt = utctodt(utc);Input Nx1 vector, UTC scalar format. d utc **Output** dtNx1 matrix. DT scalar format. Remarks A UTC scalar gives the number of seconds since or before January 1, 1970 Greenwich Mean Time. In DT scalar format, 11:06:47 on March 15, 2001 is 20010315110647. h Example tc = 985633642;print "tc = " tc; dt = utctodt(tc); print "dt = " dt; produces: m tc = 985633642n dt = 200103261107220 Source time.src p See also dtvnormal, timeutc, utctodtv, dttodtv, dtvtodt, q dttoutc, dtvtodt, strtodt, dttostr

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utctodtv

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x y z

utctodtv

Purpose Converts UTC scalar format to DTV vector format. **Format** dtv = utctodtv(utc);Input Nx1 vector, UTC scalar format. utc Output dtv Nx8 matrix. DTV vector format. Remarks A UTC scalar gives the number of seconds since or before January 1, 1970 Greenwich Mean Time. Each row of dtv, in DTV vector format, contains: [N,1]Year [N,2]Month in Year, 1-12 [N,3]Day of month, 1-31 [N,4]Hours since midnight, 0-23 [N,5]Minutes, 0-59 [N,6]Seconds, 0-59 [N,7]Day of week, 0-6, 0 = Sunday[N,8] Days since Jan 1 of current year, 0-365 Example tc = timeutc; print "tc = " tc; dtv = utctodtv(tc); print "dtv = " dtv; produces: tc = 985633642dtv = 2001 3 26 11 7 22 1 84 Source time.src See also dtvnormal, timeutc, utctodt, dttodtv, dttoutc, dtvtodt, dtvtoutc, strtodt, dttostr

utrisol

utrisol

Purpose Computes the solution of Ux = b where U is an upper triangular matrix.

Format x = utrisol(b,U);

Input b PxK matrix.

U PxP upper triangular matrix.

Output x PxK matrix.

Remarks utrisol applies a back solve to Ux = b to solve for x. If b has more than one column, each column is solved for separately, i.e., utrisol

applies a back solve to Ux[.,i] = b[.,i].

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vals

vals

Converts a string into a matrix of its ASCII values. **Purpose Format** y = vals(s);Input string of length N where N > 0. **Output** y Nx1 matrix containing the ASCII values of the characters in the string s. Remarks If the string is null, the function will fail and an error message will be given. **Example** k0: k = key;if not k; goto k0; endif; if k == vals("Y") or k == vals("y");goto doit; else; end; endif; doit: In this example the **key** function is used to read the keyboard. When **key**

returns a nonzero value, meaning a key has been pressed, the ASCII value it returns is tested to see if it is an uppercase or lowercase "Y". If it is, the program will jump to the label **doit**, otherwise the program will end.

See also chrs, ftos, stof

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varget

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Purpose

Accesses a global variable whose name is given as a string argument.

Format

y = varget(s);

Input

string containing the name of the global symbol you wish to access.

Output

contents of the matrix or string whose name is in s.

Remarks

This function searches the global symbol table for the symbol whose name is in *s* and returns the contents of the variable if it exists. If the symbol does not exist, the function will terminate with an **Undefined symbol** error message. If you want to check to see if a variable exists before using this function, use **typecv**.

Example

See also

typecv, varput

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x y z

vargetl

vargetl

Purpose Accesses a local variable whose name is given as a string argument.

Format y = vargetl(s);

Input s string containing the name of the local symbol you wish to access.

Output y contents of the matrix or string whose name is in s.

Remarks This function searches the local symbol list for the symbol whose name is

in *s* and returns the contents of the variable if it exists. If the symbol does not exist, the function will terminate with an **Undefined symbol**

error message.

```
Example proc dog;
```

```
local x,y;
x = rndn(2,2);
y = vargetl("x");
print "x" x;
print "y" y;
retp(y);
endp;
z = dog;
print "z" z;
```

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	-0.108873	0.0648738
z		
	-0.543851	-0.181701
	-0.108873	0.0648738

See also varputl

varmall

varmall

Purpose Computes log-likelihood of a Vector ARMA model.

Format res = varmall(w, phi, theta, vc);

Input w NxK matrix, time series.

phi K*PxK matrix, AR coefficient matrices.theta K*QxK matrix, MA coefficient matrices.

vc KxK matrix, covariance matrix.

Output *ll* scalar, log-likelihood. If the calculation fails *res* is set to missing value with error code:

Error Code	Reason for Failure
1	M < 1
2	N < 1
3	P < 0
4	Q < 0
5	P = 0 and $Q = 0$
7	floating point work space too small
8	integer work space too small
9	qq is not positive definite
10	AR parameters too close to stationarity boundary
11	model not stationary
12	model not invertible
13	I+M'H'HM not positive definite

Remarks

varmall is adapted from code developed by Jose Alberto Mauricio of the Universidad Complutense de Madrid. It was published as Algorithm AS311 in Applied Statistics. Also described in "Exact Maximum Likelihood Estimation of Stationary Vector ARMA Models," JASA, 90:282-264.

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varmares

varmares

Purpose Computes residuals of a Vector ARMA model.

Format res = varmares(w, phi, theta);

Input W NxK matrix, time series.

phi K*PxK matrix, AR coefficient matrices.theta K*QxK matrix, MA coefficient matrices.

Output res NxK matrix, residuals. If the calculation fails res is set to missing value with error code:

Error Code	Reason for Failure
1	M < 1
2	N < 1
3	P < 0
4	Q < 0
5	P = 0 and $Q = 0$
7	floating point work space too small
8	integer work space too small
9	qq is not positive definite
10	AR parameters too close to stationarity boundary
11	model not stationary
12	model not invertible
13	I+M'H'HM not positive definite

Remarks

varmares is adapted from code developed by Jose Alberto Mauricio of the Universidad Complutense de Madrid. It was published as Algorithm AS311 in Applied Statistics. Also described in "Exact Maximum Likelihood Estimation of Stationary Vector ARMA Models," JASA, 90:282-264.

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varput

varput

Purpose

Allows a matrix or string to be assigned to a global symbol whose name is given as a string argument.

Format

```
y = varput(x,n);
```

Input

- x NxK matrix or string which is to be assigned to the target variable.
- n string containing the name of the global symbol which will be the target variable.

Output

y scalar, 1 if the operation is successful and 0 if the operation fails.

Remarks

x and n may be global or local. The variable, whose name is in n, that x is assigned to is always a global.

If the function fails, it will be because the global symbol table is full.

This function is useful for returning values generated in local variables within a procedure to the global symbol table.

Example

```
source = rndn(2,2);
targname = "target";
if not varput(source,targname);
  print "Symbol table full";
  end;
endif;

source = -0.93519984  0.40642598
  -0.36867581  2.57623519

target = -0.93519984  0.40642598
  -0.36867581  2.57623519
```

See also

varget, typecv

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varputl

varputl

Purpose b

Format y = varputl(x,n);

given as a string argument.

Input

x NxK matrix or string which is to be assigned to the target variable.

n string containing the name of the local symbol which will be the target variable.

Allows a matrix or string to be assigned to a local symbol whose name is

Output

scalar, 1 if the operation is successful and 0 if the operation fails.

Remarks

x and n may be global or local. The variable, whose name is in n, that x is assigned to is always a local.

Example

```
proc dog(x);
  local a,b,c,d,e,vars,putvar;
  a=1;b=2;c=3;d=5;e=7;
  vars = { a b c d e };
  putvar = 0;
  do while putvar $/= vars;
    print "Assign x (" $vars "): " ;;
    putvar = upper(cons);
    print;
  endo;
  call varputl(x,putvar);
  retp(a+b*c-d/e);
endp;
format /rds 2,1;
i = 0;
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varputl

```
do until i >= 5;
  z = dog(17);
  print " z is " z;
  i = i + 1;
endo;
Produces:
Assign x ( A B C D E ): a
  z is 22.3
Assign x ( A B C D E ): b
   z is 51.3
Assign x ( A B C D E ): c
   z is 34.3
Assign x ( A B C D E ): d
   z is 4.6
Assign x ( A B C D E ): e
  z is 6.7
```

See also varget1

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vartype

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x y z

vartype

Purpose Returns a vector of ones and zeros that indicate whether variables in a data set are character or numeric. Format y = vartype(names);Input names Nx1 character vector of variable names retrieved from a data set header file with the **getname** function. **Output** Nx1 vector of ones and zeros, 1 if variable is numeric, 0 if character. Remarks This function is being obsoleted. See vartypef. If a variable name in *names* is lowercase, a 0 will be returned in the corresponding element of the returned vector. **Example** names = getname("freq"); y = vartype(names); print \$names; print y; **AGE** PAY sex WT 1.0000000 1.0000000 0.0000000 1.0000000 Source vartype.src

vartypef

vartypef

Purpose Returns a vector of ones and zeros that indicate whether variables in a data set are character or numeric.

Format y = vartypef(f);

format.

Input f file handle of an open file.

Output y Nx1 vector of ones and zeros, 1 if variable is numeric, 0 if character.

Remarks This function should be used in place of older functions that are based on the case of the variable names. You should also use the **v96** data set

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vcm, vcx

Purpose Computes a variance-covariance matrix.

Format vc = vcm(m);

 $vc = \mathbf{vcx}(x);$

Input m KxK moment (x'x) matrix. A constant term MUST have been the

first variable when the moment matrix was computed.

x NxK matrix of data.

Output *vc* KxK variance-covariance matrix.

Source corr.src

See also momentd

vec, vecr

Purpose Creates a column vector by appending the columns/rows of a matrix to each other.

Format yc = vec(x);yr = vecr(x);

Input x NxK matrix.

Output yc (N*K)x1 vector, the columns of x appended to each other.

yr (N*K)x1 vector, the rows of x appended to each other and the result transposed.

Remarks vecr is much faster.

Example x

$$x = 1.000000 2.000000$$

 $3.000000 4.000000$

 $yc = \begin{cases} 1.000000 \\ 3.000000 \\ 2.000000 \\ 4.000000 \end{cases}$

 $yr = \begin{cases} 1.000000 \\ 2.000000 \\ 3.000000 \\ 4.000000 \end{cases}$

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Purpose Vectorizes a symmetric matrix by retaining only the lower triangular portion of the matrix.

Format

v = vech(x);

Input

x NxN symmetric matrix.

Output

(N*(N+1)/2)x1 vector, the lower triangular portion of the matrix x.

Remarks

As you can see from the example below, **vech** will not check to see if *x* is symmetric. It just packs the lower triangular portion of the matrix into a column vector in row-wise order.

Example

$$x = \begin{array}{r} 11 & 12 & 13 \\ 21 & 22 & 23 \\ 31 & 32 & 33 \end{array}$$

 $v = \begin{cases} 11 \\ 21 \\ 22 \\ 31 \end{cases}$

32 33

 $sx = \begin{cases} 11 & 21 & 31 \\ 21 & 22 & 32 \end{cases}$

31 32 33

See also x

xpnd

vector (dataloop)

vector (dataloop)

Purpose Specifies the creation of a new variable within a data loop.

Format vector [#] numvar = numeric_expression;

vector \$ charvar = character_expression;

Remarks A *numeric_expression* is any valid expression returning a numeric value.

A *character_expression* is any valid expression returning a character value. If neither '\$' nor '#' is specified, '#' is assumed.

vector is used in place of make when the expression returns a scalar

rather than a vector. **vector** forces the result of such an expression to a vector of the correct length. **vector** could actually be used anywhere that **make** is used, but would generate slower code for expressions that

already return vectors.

Any variables referenced must already exist, either as elements of the source data set, as **externs**, or as the result of a previous **make**,

vector, or code statement.

Example vector const = 1;

See also make

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vget

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vget

Extracts a matrix or string from a data buffer constructed with **vput**. **Purpose Format** { x,dbufnew } = vget(dbuf,name); Input Nx1 vector, a data buffer containing various strings and dbuf matrices. string, the name of the string or matrix to extract from dbuf. name Output LxM matrix or string, the item extracted from dbuf. dbufnew Kx1 vector, the remainder of dbuf after x has been extracted. Source vpack.src See also vlist, vput, vread

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view

view

Purpose Sets the position of the observer in workbox units for 3-D plots.

Library pgraph

Format view(x,y,z);

Input x scalar, the X position in workbox units.

y scalar, the Y position in workbox units.

z scalar, the Z position in workbox units.

Remarks The size of the workbox is set with **volume**. The viewer must be outside

of the workbox. The closer the position of the observer, the more perspective distortion there will be. If x = y = z, the projection will be

isometric.

If **view** is not called, a default position will be calculated.

Use **viewxyz** to locate the observer in plot coordinates.

Source pgraph.src

See also volume, viewxyz

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viewxyz

viewxyz

Purpose Sets the position of the observer in plot coordinates for 3-D plots.

Library pgraph

Format viewxyz(x,y,z);

Input x scalar, the X position in plot coordinates.

y scalar, the Y position in plot coordinates.

z scalar, the Z position in plot coordinates.

Remarks The viewer must be outside of the workbox. The closer the observer, the

more perspective distortion there will be.

If **viewxyz** is not called, a default position will be calculated.

Use **view** to locate the observer in workbox units.

Source pgraph.src

See also volume, view

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x y z

vlist

vlist

Purpose Lists the contents of a data buffer constructed with **vput**.

Format vlist(dbuf);

Input *dbuf* Nx1 vector, a data buffer containing various strings and matrices.

Remarks vlist lists the names of all the strings and matrices stored in *dbuf*.

Source vpack.src

See also vget, vput, vread

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vnamecv

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vnamecv

Purpose	Returns the names of the elements of a data buffer constructed with
	vout.

Format	cv	=	vnamecv	(dbuf)	;

Output	cv	Kx1 character vector containing the names of the elements of
		dbuf.

See also vget, vput, vread, vtypecv

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volume

volume

Purpose Sets the length, width, and height ratios of the 3-D workbox.

Library pgraph

Format volume (x, y, z);

Input x scalar, the X length of the 3-D workbox.

y scalar, the Y length of the 3-D workbox.

z scalar, the Z length of the 3-D workbox.

Remarks The ratio between these values is what is important. If **volume** is not

called, a default workbox will be calculated.

Source pgraph.src

See also view

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vput

Purpose Inserts a matrix or string into a data buffer. **Format** dbufnew = vput(dbuf,x,xname); Input Nx1 vector, a data buffer containing various strings and dbuf matrices. If dbuf is a scalar 0, a new data buffer will be created. LxM matrix or string, item to be inserted into dbuf. х string, the name of x, will be inserted with x into dbuf. xname **Output** dbufnew Kx1 vector, the data buffer after x and xname have been inserted. Remarks If *dbuf* already contains *x*, the new value of *x* will replace the old one. Source vpack.src See also vget, vlist, vread

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vread

vread

Purpose Reads a string or matrix from a data buffer constructed with vput.

Format x = vread(dbuf,xname);

Input dbuf Nx1 vector, a data buffer containing various strings and

matrices.

xname string, the name of the matrix or string to read from *dbuf*.

Output *x* LxM matrix or string, the item read from *dbuf*.

Remarks vread, unlike vget, does not change the contents of *dbuf*. Reading *x*

from dbuf does not remove it from dbuf.

Source vpack.src

See also vget, vlist, vput

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vtypecv

vtypecv

Purpose Returns the types of the elements of a data buffer constructed with **vput**.

Format cv = vtypecv(dbuf);

Input *dbuf* Nx1 vector, a data buffer containing various strings and matrices.

Output *cv* Kx1 character vector containing the types of the elements of *dbuf*.

See also vget, vput, vread, vnamecv

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wait, waitc

wait, waitc

Purpose Waits until any key is pressed.

Format wait;

waitc;

Remarks If you are working in terminal mode, they don't "see" any keystrokes

until ENTER is pressed. waitc clears any pending keystrokes before

waiting until another key is pressed.

Source wait.src, waitc.src

See also pause

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Purpose Walks the index of an array forward or backward through a specified dimension.

Format

ni = walkindex(i,o,dim);

Input

i Mx1 vector of indices into an array, where $M \le N$.

o Nx1 vector of orders of an N-dimensional array.

dim scalar [1-to-M], index into the vector of indices *i*, corresponding to the dimension to walk through, positive to walk the index forward, or negative to walk backward.

Output

ni Mx1 vector of indices, the new index.

Remarks

walkindex will return a scalar error code if the index cannot walk further in the specified dimension and direction.

Example

orders = (3,4,5,6,7);
a = arrayinit(orders,1);
ind = { 2,3,3 };
ind = walkindex(ind,orders,-2);

2
ind = 2
3

This example decrements the second value of the index vector *ind*.

ind = walkindex(ind,orders,3);

ind = 2 4

Using the orders from the example above and the *ind* that was returned, this example increments the third value of the index vector *ind*.

See also

nextindex, previousindex, loopnextindex

window

window

Purpose Partitions the window into tiled regions of equal size.

Library pgraph

Format window(row,col,typ);

Input *row* scalar, number of rows of graphic panels.

col scalar, number of columns of graphic panels.

typ scalar, graphic panel attribute type. If 1, the graphic panels will

be transparent, if 0, the graphic panels will be nontransparent

(blanked).

Remarks The graphic panels will be numbered from 1 to (row)x(col) starting from

the left topmost graphic panel and moving right.

See **makewind** for creating graphic panels of a specific size and position. (For more information, see "Publication Quality Graphics" in

the *User Guide*.

Source pwindow.src

See also endwind, begwind, setwind, nextwind, getwind,

makewind

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Purpose Writes a matrix to a GAUSS data set.

Format y = writer(fh,x);

Input *fh* handle of the file that data is to be written to.

x NxK matrix.

Output y scalar specifying the number of rows of data actually written to the data set.

Remarks The file must have been opened with create, open for append, or open for update.

The data in x will be written to the data set whose handle is fh starting at the current pointer position in the file. The pointer position in the file will be updated so the next call to **writer** will put the next block of data after the first block. (See **open** and **create** for the initial pointer positions in the file for reading and writing.)

x must have the same number of columns as the data set. **colsf** returns the number of columns in a data set.

writer returns the number of rows actually written to the data set. If y does not equal **rows** (x), the disk is probably full.

If the data set is not double precision, the data will be rounded to nearest as it is written out.

If the data contain character elements, the file must be double precision or the character information will be lost.

If the file being written to is the 2-byte integer data type, then missing values will be written out as -32768. These will not automatically be converted to missings on input. They can be converted with the **miss** function:

x = miss(x, -32768);

Trying to write complex data to a data set that was originally created to store real data will cause a program to abort with an error message. (See **create** for details on creating a complex data set.)

writer

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С

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x y z

```
Example
            create fp = data with x,10,8;
             if fp == -1;
                errorlog "Can't create output file";
                end;
             endif;
             c = 0;
             do until c >= 10000;
                y = rndn(100,10);
                k = writer(fp,y);
                if k /= rows(y);
                   errorlog "Disk Full";
                   fp = close(fp);
                   end;
                endif;
                c = c+k;
             endo;
             fp = close(fp);
             In this example, a 10000x10 data set of Normal random numbers is
             written to a data set called data. dat. The variable names are X01-X10.
```

See also open, close, create, readr, saved, seekr

3-967

xlabel

xlabel

Purpose Sets a label for the X axis.

Library pgraph

Format xlabel(str);

Input str string, the label for the X axis.

Source pgraph.src

See also title, ylabel, zlabel

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xlsreadm

xlsreadm

Purpose Reads from an Excel spreadsheet, into a GAUSS matrix.

Format mat = xlsreadm(file, range, sheet, vls);

Input *file* string, name of .xls file.

range string, range to read, e.g. "a2:b20" or the starting point of the

read, e.g. "a2".

sheet scalar, sheet number.

vls null string or 9x1 matrix, specifies the conversion of Excel empty cells and special types into GAUSS (see Remarks). A null string results in all empty cells and empty types being converted

to GAUSS missing values.

Output *mat* matrix or a Microsoft error code.

Remarks The *vls* argument lets users control the import of Excel empty cells and special types, according to the following table:

Row Number	Excel Cell	
1	empty cell	
2	#N/A	
3	#VALUE!	
4	#DIV/0!	
5	#NAME?	
6	#REF!	
7	#NUM!	
8	#NULL!	
9	#ERR	

To convert all occurrences of #DIV/0! to 9999.99, and all other empty cells and special types to GAUSS missing values:

```
vls = reshape(error(0),9,1);
vls[4] = 9999.99;
```

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xlsreadsa

xlsreadsa

Purpose Reads from an Excel spreadsheet, into a GAUSS string array or string.

Format s = xlsreadsa(file, range, sheet, vls);

Input *file* string, name of .xls file.

range string, range to read, e.g. "a2:b20" or the starting point of the

read, e.g. "a2".

sheet scalar, sheet number.

vls null string or 9x1 string array, specifies the conversion of Excel

empty cells and special types into GAUSS (see Remarks). A null string results in all empty cells and empty types being converted

to GAUSS missing values.

Output s string array or string or a Microsoft error code.

Remarks The *vls* argument lets users control the import of Excel empty cells and special types, according to the following table:

Row Number	Excel Cell
1	empty cell
2	#N/A
3	#VALUE!
4	#DIV/0!
5	#NAME?
6	#REF!
7	#NUM!
8	#NULL!
9	#ERR

To convert all occurrences of #DIV/0! to "Division by Zero", and all other empty cells and special types to GAUSS missing values:

```
vls = reshape("",9,1);
vls[4] = "Division by Zero";
```

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xlswritem

xlswritem

Purpose Writes a GAUSS matrix to an Excel spreadsheet.

Format ret = xlswritem(data, file, range, sheet, vls);

Input data

file string, name of .xls file.

range string, the starting point of the write, e.g. "a2".

sheet scalar, sheet number.

matrix.

vls null string or 9x1 matrix, specifies the conversion from GAUSS into Excel empty cells and special types (see Remarks). A null string results in all GAUSS missing values being converted to empty cells in Excel.

Output

ret scalar, 0 if success or a Microsoft error code.

Remarks

The *vls* argument converts values in GAUSS to Excel empty cells and special types according to the following table:

Row Number	Excel Cell		
1	. 11		
1	empty cell		
2	#N/A		
3	#VALUE!		
4	#DIV/0!		
5	#NAME?		
6	#REF!		
7	#NUM!		
8	#NULL!		
9	#ERR		

To convert all occurrences of 9999.99 in GAUSS to #DIV/0! in Excel and convert all GAUSS missing values to empty cells in Excel:

```
vls = reshape(error(0), 9,1);
vls[4] = 9999.99;
```

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xlswritesa

xlswritesa a **Purpose** Writes a GAUSS string or string array to an Excel spreadsheet. h ret = xlswritesa(data, file, range, sheet, vls); Format Input string or string array. d data file string, name of .xls file. string, the starting point of the write, e.g. "a2". sheet scalar, sheet number. null string or 9x1 string array, specifies the conversion from vls GAUSS into Excel empty cells and special types (see Remarks). A null string results in all GAUSS missing values being converted to empty cells in Excel. Output scalar, 0 if success or a Microsoft error code. ret Remarks The vls argument converts values in GAUSS to Excel empty cells and k special types according to the following table: Row Number Excel Cell

Row Number	Excel Cell
1	empty cell
2	#N/A
3	#VALUE!
4	#DIV/0!
5	#NAME?
6	#REF!
7	#NUM!
8	#NULL!
9	#ERR

To convert all occurrences of "Division by Zero" in GAUSS to #DIV/0! in Excel and convert all GAUSS missing values to empty cells in Excel:

```
vls = reshape("", 9,1);
vls[4] = "Division by Zero";
```

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xpnd

xpnd

Purpose Expands a column vector into a symmetric matrix.

Format x = xpnd(v);

Input v Kx1 vector, to be expanded into a symmetric matrix.

Output x MxM matrix, the results of taking v and filling in a symmetric matrix with its elements.

$$M = ((-1 + sqrt(1+8*K))/2)$$

Remarks If v does not contain the right number of elements, (that is, if sqrt(1 + 8*K) is not integral), then an error message is generated.

This function is particularly useful for hard-coding symmetric matrices, because only about half of the matrix needs to be entered.

Example

$$x = \{ 1, \\ 2, 3, \\ 4, 5, 6, \\ 7, 8, 9, 10 \};$$
 $y = xpnd(x);$

 $\begin{array}{r}
1\\2\\3\\4\\x=5\end{array}$

6 7 8

> 9 10

a

b

c d

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3-973

xpnd

 $y = \begin{array}{c} 1 & 2 & 4 & 7 \\ 2 & 3 & 5 & 8 \\ 4 & 5 & 6 & 9 \\ 7 & 8 & 9 & 10 \end{array}$

See also vech

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xtics

xtics

Purpose Sets and fixes scaling, axes numbering and tick marks for the X axis.

Library pgraph

Format xtics(min,max,step,minordiv);

Input *min* scalar, the minimum value.

max scalar, the maximum value.

step scalar, the value between major tick marks.

minordiv scalar, the number of minor subdivisions.

Remarks This routine fixes the scaling for all subsequent graphs until graphset

is called.

This gives you direct control over the axes endpoints and tick marks. If **xtics** is called after a call to **scale**, it will override **scale**.

X and Y axes numbering may be reversed for **xy**, **logx**, **logy**, and **loglog** graphs. This may be accomplished by using a negative step value in the **xtics** and **ytics** functions.

Source pscale.src

See also scale, ytics, ztics

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x y z

Purpose Graphs X vs. Y using Cartesian coordinates.

Library pgraph

Format xy(x,y);

Input x Nx1 or NxM matrix. Each column contains the X values for a particular line.

y Nx1 or NxM matrix. Each column contains the Y values for a particular line.

Remarks Missing values are ignored when plotting symbols. If missing values are

encountered while plotting a curve, the curve will end and a new curve will begin plotting at the next non-missing value.

Source pxy.src

See also xyz, logx, logy, loglog

3-976

XYZ

Purpose Graphs X vs. Y vs. Z using Cartesian coordinates.

Library pgraph

Format xyz(x,y,z);

Input x Nx1 or NxK matrix. Each column contains the X values for a particular line.

y Nx1 or NxK matrix. Each column contains the Y values for a particular line.

z Nx1 or NxK matrix. Each column contains the Z values for a particular line.

Remarks Missing values are ignored when plotting symbols. If missing values are

encountered while plotting a curve, the curve will end and a new curve

will begin plotting at the next non-missing value.

Source pxyz.src

See also xy, surface, volume, view

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ylabel

ylabel

Purpose Sets a label for the Y axis.

Library pgraph

Format ylabel(str);

e Input str string, the label for the Y axis.

Source pgraph.src

See also title, xlabel, zlabel

v w **x y z**

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vtics

ytics

Purpose Sets and fixes scaling, axes numbering and tick marks for the Y axis.

Library pgraph

Format ytics(min,max,step,minordiv);

Input *min* scalar, the minimum value.

max scalar, the maximum value.

step scalar, the value between major tick marks.

minordiv scalar, the number of minor subdivisions.

Remarks This routine fixes the scaling for all subsequent graphs until graphset

is called.

This gives you direct control over the axes endpoints and tick marks. If **ytics** is called after a call to **scale**, it will override **scale**.

X and Y axes numbering may be reversed for **xy**, **logx**, **logy** and **loglog** graphs. This may be accomplished by using a negative step value in the **xtics** and **ytics** functions.

Source pscale.src

See also scale, xtics, ztics

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zeros

zeros a Creates a matrix of zeros. Purpose b **Format** y = zeros(r,c);Input scalar, the number of rows. d scalar, the number of columns. ce **Output** RxC matrix of zeros. Remarks This is faster than ones. Noninteger arguments will be truncated to an integer. h **Example** y = zeros(3,2); $0.000000 \ 0.000000$ k $0.000000 \ 0.000000$ 0.000000 0.00000 m See also ones, eye n 0 p q u V

W

zlabel

zlabel

Purpose Sets a label for the Z axis.

Library pgraph

Format zlabel(str);

Input *str* string, the label for the Z axis.

Source pgraph.src

See also title, xlabel, ylabel

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ztics

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ztics

Purpose Sets and fixes scaling, axes numbering and tick marks for the Z axis.

Library pgraph

Format ztics(min,max,step,minordiv);

Input *min* scalar, the minimum value.

max scalar, the maximum value.

step scalar, the value between major tick marks.

minordiv scalar, the number of minor subdivisions. If this function is

used with contour, contour labels will be placed every

minordiv levels. If 0, there will be no labels.

Remarks This routine fixes the scaling for all subsequent graphs until graphset

is called.

This gives you direct control over the axes endpoints and tick marks. If **ztics** is called after a call to **scale3d**, it will override **scale3d**.

Source pscale.src

See also scale3d, xtics, ytics, contour

хух

Obsolete Commands A

The following commands will no longer be supported, therefore should not be used when creating new programs.

disable plotsym

editm print on/off

enable rndmod

export rndns/rndus
exportf setvmode

files WinClear

font WinClearArea

FontLoad WinClearTTYlog

FontUnload WinClose

FontUnloadAll WinCloseAll

graph WinGetActive

import WinGetAttributes

importf WinGetColorCells

line WinGetCursor

lprint on/off WinMove

medit WinOpenPQG ndpchk WinOpenText ndpclex WinOpenTTY

ndpcntrl WinPan plot WinPrint

Obsolete Commands

WinPrintPQG

WinRefresh

WinRefreshArea

WinResize

WinSetActive

WinSetBackground

WinSetColorCells

Win Set Color map

WinSetCursor

WinSetForeground

WinSetRefresh

WinSetTextWrap

WinZoomPQG

Colors Appendix

Blue
 Green
 Light Blue
 Light Green
 Cyan
 Light Cyan
 Red
 Light Red
 Magenta
 Light Magenta

0 Black

8 Dark Grey

6 Brown 14 Yellow 7 Grey 15 White